ENCYCLOPEDIA OF EXPLOSIVES
AND
RELATED ITEMS

PATR 2700
VOLUME 7

BY
BASIL T. FEDOROFF & OLIVER E. SHEFFIELD

ASSISTED BY
SEYMOUR M. KAYE & MANAGEMENT SCIENCE ASSOCIATES

U.S. ARMY RESEARCH AND DEVELOPMENT COMMAND
TACOM, ARDEC
WARHEADS, ENERGETICS AND COMBAT SUPPORT CENTER
PICATINNY ARSENAL
NEW JERSEY, USA
1975
Copies of the “Encyclopedia of Explosives And Related Items” can be obtained by requesting CD ROM from the:

National Technical Information Service
(formerly Clearinghouse)
US Department of Commerce
Springfield, Virginia 22151

1-800-553-6847 USA only
703-605-6000
www.ntis.gov/FCPC

The contents of these volumes are UNCLASSIFIED.
The distribution of these volumes is UNLIMITED.

Neither the US Government nor any person acting on behalf of the US Government assumes any liability resulting from the use or publication of the information contained in this document or warrants that such use or publication will be free from privately owned rights.

All rights reserved. This document, or parts thereof, may not be reproduced in any form without written permission of the Energetics and Warhead Division, WECAC, TACOM, ARDEC, Picatinny Arsenal.

Library of Congress Catalogue Card Number: 61-61759
PREFACE

This volume represents a continuing effort to cover comprehensively the unclassified information on explosives and related subjects in the same manner and format as in previous volumes. The reader is urged to obtain the previous volumes and to read both the PREFACE and INTRODUCTION in Volume 1 in order to understand the authors' way of presenting the subject matter.

In preparation for and during the writing of this Encyclopedia, the authors have consulted freely with and have had the cooperation of many individuals who contributed their expert knowledge and advice. This fact is acknowledged throughout the text at the end of the subject item. A listing of many others who have helped in various ways would be impractical.

Dr Julius Roth, principal scientist at Management Science Associates, Los Altos, California, contributed significantly in the literature searching and writing of many of the articles for this volume under Contract DAAA21-73-C-0725. Mr Henry Heman of the Explosives Division assisted with proofreading and literature searching. Others who contributed to the manuscript, by invitation are indicated at the end of the articles.

Dr Raymond F. Walker, Explosives Division Chief, provided the financial support and encouragement to continue this work. Further financial support is expected from the sale of copies to non-governmental agencies and individuals by the National Technical Information Service, US Department of Commerce, Springfield, Virginia 22151.

Although considerable effort has been made to present this information as accurately as possible, mistakes and errors in transcription and translation do occur. Therefore, the authors encourage the readers to consult original sources, when possible, and to feel free to point out errors and omissions of important works so that corrections and additions can be listed in the next volume. The interpretations of data and opinions expressed are often those of the authors and are not necessarily those nor the responsibility of officials of Picatinny Arsenal or the Department of the Army.

This report has been prepared for information purposes only and neither Picatinny Arsenal nor the Department of the Army shall be responsible for any events or decisions arising from the use of any information contained herein.
ADDENDUM

Heptyl, listed in Vol 7 of Encycl, p H64-L and described in Vol 1, pp A441-42 as N-(2,4,6-TRINITRO-N-nitramino)-trimethylene Trinitrate is claimed now by Dr Paolo Anmat di San Filippo in a letter of June 25, 1975 addressed to the Feltman Research Laboratory Director, to be identical with Epitrile described by him and Dr Michele Giua (who died in 1966) in Annali di Chimica, Vol 50, 1381-88 (1960). This claim is not correct because Heptyl has the formula

![Chemical structure of Heptyl](attachment://heptyl.png)

while Epitrile is

![Chemical structure of Epitrile](attachment://epitrile.png)

its chemical name is 2,4,6-TRINITROPHENYL-1,3-DINITRAMINOETHANOL DINITRATE. It was prepd by Di San Filippo by nitrating either 4,6-DINITROPHENYL-1,3-DIAMINOETHANOL,

![Chemical structure of intermediate](attachment://intermediate.png)

or 2,4,6-TRINITRO-1,3-DIAMINOETHANOL,

![Chemical structure of product](attachment://product.png)

, crysts, mp 127°. In both cases concd mixed nitric-sulfuric acid was used.
The procedure is described on pp 1382 & 1386 of Ann di Chimica 50. Epitriole separated as a greenish-yellow oily-solid which crystallized from ethanol in the form of tiny crys
tals melting at 123°. Its mw is 511.24 and N content 24.55% (calc 24.66%). Its explosive
properties were not determined.

Unfortunately, we did not have the paper of Drs Giua and Di San Filippo at the time
1,3-Di(ethylolamino)-benzene and Derivatives were described in Vol 5, p D1242-R.
This compd \( \text{NH}_2 \text{C}_2 \text{H}_4 \text{OH} \)

is identical with Phenyl-1,3-diaminethanol which may be considered as the parent compd of 2,4,6-Trinitrophenyl-1,3-dinitroaminoethanol Dinitrate designated as Epitriole. The compd prepd by K.F. Waldköttner by nitrination of
4,6-Dinitro-1,3-di(ethylolamino)-benzene, described in Recueil de Travaux Chimiques
57 1307 (1938) and in Vol 5 of Encycl is

\[ \text{N} \text{C}_2 \text{H}_4 \text{OH} \]

\[ \text{O}_2 \text{N} \]

; mw 421.27, N 23.28%; a thick sticky mass, which softened at 30°, decompd violently
at 98° and ignited at 230°. It is not identical with Epitriole and was not obtd in crystal-
line form. No attempt was made by Waldköttner to further nitate this compd called
2,4,6-Trinitro-1,3-bis(N-nitro-β-ethanol-
amino)-benzene
TABLE OF CONTENTS

Preface I
Addendum III
List of Figures and Illustrations VIII
Supplement to Abbreviations for Books and Periodicals X
Supplement to List of Books on Explosives, Propellants, Pyrotechnics and Ordnance Items XI

Descriptive Text of Encyclopedic Items

<table>
<thead>
<tr>
<th>H2 to &quot;Hyros&quot; Explosive</th>
<th>H 1 to H 263</th>
</tr>
</thead>
<tbody>
<tr>
<td>IASP to IZZO</td>
<td>I 1 to I 184</td>
</tr>
<tr>
<td>J (Poudre de chasse) to Jute</td>
<td>J 1 to J 87</td>
</tr>
<tr>
<td>K (Pulver) to KZ</td>
<td>K 1 to K 20</td>
</tr>
<tr>
<td>L-Alloy to Lysol</td>
<td>L 1 to L 63</td>
</tr>
</tbody>
</table>

Tables

<table>
<thead>
<tr>
<th>Handling Characteristics of Explosives</th>
<th>H 11</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thermodynamic Calculations of Compounds and their NFPA</td>
<td>H 18</td>
</tr>
<tr>
<td>Ratings of Potential Hazard</td>
<td></td>
</tr>
<tr>
<td>Comparison of NFPA Reactivity Rating with Reaction Hazard Index</td>
<td>H 20</td>
</tr>
<tr>
<td>Data Source, Reaction Hazard Index, and National Fire Protection</td>
<td>H 21</td>
</tr>
<tr>
<td>Association Reactivity Rating</td>
<td></td>
</tr>
<tr>
<td>Properties of HBX's</td>
<td>H 24</td>
</tr>
<tr>
<td>US Specification Requirements for Grades A and B Compositions of HBX</td>
<td>H 25</td>
</tr>
<tr>
<td>HBX Composition (Percent by Weight)</td>
<td>H 26</td>
</tr>
<tr>
<td>Standard Heats of Formation and Heats of Detonation of Common Explosives</td>
<td>H 41</td>
</tr>
<tr>
<td>Standard Heats of Formation and Heats of Detonation of Explosive Mixtures</td>
<td>H 43</td>
</tr>
<tr>
<td>Melting Points, Heats of Fusion, Heats of Vaporization and Heats of Sublimation of Common High Explosives</td>
<td>H 44</td>
</tr>
<tr>
<td>Heat Conductivity Coefficients and Specific Heats of Common Explosives</td>
<td>H 45</td>
</tr>
<tr>
<td>Some Examples of the Use of Group Additivity to Check Experimentally Determined Heats of Formation</td>
<td>H 47</td>
</tr>
<tr>
<td>Comparison of Estimated Heat of Formation with Some Measured Since 1969</td>
<td>H 49</td>
</tr>
<tr>
<td>Group Values Now Available for State and Class of Compound</td>
<td>H 51</td>
</tr>
<tr>
<td>Tables (Cont’d)</td>
<td>Page</td>
</tr>
<tr>
<td>--------------------------------------------------------------------------------</td>
<td>------</td>
</tr>
<tr>
<td>Measured and Estimated Heats of Formation of Various Compounds at 298°K in kcal/mole</td>
<td>H 52</td>
</tr>
<tr>
<td>Heats of Formation of Nitroalkanes</td>
<td>H 54</td>
</tr>
<tr>
<td>Group Values for Estimating Heats of Formation of Nitroalkanes</td>
<td>H 55</td>
</tr>
<tr>
<td>Heats of Formation of Azo Compounds Measured by Engel and Wood and Estimated by Group Activity</td>
<td>H 56</td>
</tr>
<tr>
<td>Group Values for Calculating Heats of Formation of Trans Azoalkanes and Azo Radicals</td>
<td>H 57</td>
</tr>
<tr>
<td>Compositions and Properties of HEX-24 and HEX-48</td>
<td>H 73</td>
</tr>
<tr>
<td>Solubility of Hexamethylenetetramine in Solvents</td>
<td>H 79</td>
</tr>
<tr>
<td>Rf (Relative Movements of Test Sample Thru a Chromatographic Column) for Various Explosives</td>
<td>H 99</td>
</tr>
<tr>
<td>Image Dissection Camera Data</td>
<td>H 106</td>
</tr>
<tr>
<td>Characteristics of Streak Cameras</td>
<td>H 108</td>
</tr>
<tr>
<td>Specifications for Raw Materials Used in HTA-3 Explosive</td>
<td>H 177</td>
</tr>
<tr>
<td>Composition of Type I and Type II HT-3 Explosives</td>
<td>H 177</td>
</tr>
<tr>
<td>Unreacted Hugoniot Data for Condensed Explosives</td>
<td>H 180</td>
</tr>
<tr>
<td>Critical Relative Humidities for Some Explosives and for Ingredients of Explosives and Pyrotechnic Compositions</td>
<td>H 186</td>
</tr>
<tr>
<td>Chemical and Physical Properties of Hydrazine Propellant</td>
<td>H 205</td>
</tr>
<tr>
<td>Hydrolysis of Plasticizers for NC</td>
<td>H 223</td>
</tr>
<tr>
<td>Explosive Nature of Peroxides</td>
<td>H 225</td>
</tr>
<tr>
<td>Peroxy Derivatives of Aldehydes and Ketones</td>
<td>H 227</td>
</tr>
<tr>
<td>Strength of Sulfuric Acid Required to Give Different Relative Humidities</td>
<td>H 251</td>
</tr>
<tr>
<td>Performance Data for Chlorine Trifluoride-Fuel Hypergolic Propellant Combination</td>
<td>H 256</td>
</tr>
<tr>
<td>Performance Data for Bromine Pentafluoride-Fuel Hypergolic Propellant Combination</td>
<td>H 256</td>
</tr>
<tr>
<td>Performance Data for Hydrogen Peroxide-Fuel Hypergolic Propellant Combination</td>
<td>H 257</td>
</tr>
<tr>
<td>Performance Data for Fluorine-Fuel Hypergolic Propellant Combination</td>
<td>H 257</td>
</tr>
<tr>
<td>Performance Data for Nitrogen Tetroxide-Fuel Hypergolic Propellant Combination</td>
<td>H 257</td>
</tr>
<tr>
<td>Performance Data for Red Fuming Nitric Acid-Fuel Hypergolic Propellant Combination</td>
<td>H 257</td>
</tr>
<tr>
<td>Percent Firing of Different Types of Lead Azide versus</td>
<td>I 26</td>
</tr>
<tr>
<td>Distance from the Fuse</td>
<td>I 27</td>
</tr>
<tr>
<td>Ignition Temperature Tests of Initiating Explosives by Methods a (Pérez Ara) and b (Datta &amp; Chatterje)</td>
<td>I 28</td>
</tr>
<tr>
<td>Ignition Temperature of Nitroglycerin as Determined by Method of Snelling &amp; Storm</td>
<td>I 28</td>
</tr>
<tr>
<td>Ignition Temperature of Various Explosives as Determined by Method of Micewicz &amp; Majkowski</td>
<td>I 28</td>
</tr>
<tr>
<td>Ignition Temperature of Various Explosives as Determined by Method of Taylor &amp; Rinkenbach</td>
<td>I 28</td>
</tr>
<tr>
<td>Table Title</td>
<td>Page</td>
</tr>
<tr>
<td>---------------------------------------------------------------------------</td>
<td>------</td>
</tr>
<tr>
<td>Comparison of Impact Sensitiveness Tests</td>
<td>141</td>
</tr>
<tr>
<td>Impact Sensitivity Data Obtained with the Rotter</td>
<td>141</td>
</tr>
<tr>
<td>Apparatus at ARDE</td>
<td></td>
</tr>
<tr>
<td>Comparison of Impact Sensitivity Tests on the Basis of Figures of Insensitivity</td>
<td>142</td>
</tr>
<tr>
<td>Qualitative Ranking of the Impact Sensitivity of Common Explosives</td>
<td>151</td>
</tr>
<tr>
<td>Critical Stresses &amp; Thicknesses in Instrument No 2 for Various Explosives</td>
<td></td>
</tr>
<tr>
<td>Charge Thickness for Transition from Fracture to Explosion</td>
<td>152</td>
</tr>
<tr>
<td>Critical Stresses Obtained in Different Pressures</td>
<td>154</td>
</tr>
<tr>
<td>Impact Characteristics of Some Common Explosives</td>
<td>154</td>
</tr>
<tr>
<td>Comparison of Experimental &amp; Theoretical Impulse Constants</td>
<td>160</td>
</tr>
<tr>
<td>Comparison of Measured and Theoretical Impulse Constants ((l_0)) for Gas Detonations</td>
<td></td>
</tr>
<tr>
<td>Characteristics of a Typical IMR Propellant</td>
<td>164</td>
</tr>
<tr>
<td>Materials Used for the Core of an Italian Shell</td>
<td>172</td>
</tr>
<tr>
<td>Modified Gap Tests of Various Explosives</td>
<td>190</td>
</tr>
<tr>
<td>Comparison of Approximate and Exact Calculations for Shocked Argon</td>
<td>131</td>
</tr>
<tr>
<td>Shock Temperatures of Noble Gases</td>
<td>131</td>
</tr>
<tr>
<td>Important Data Concerning the Most Used Italian Ammunition</td>
<td>143</td>
</tr>
<tr>
<td>Italian Antitank &amp; Seacoast Guns</td>
<td>144</td>
</tr>
<tr>
<td>Italian Bombs &amp; Grenades</td>
<td>146</td>
</tr>
<tr>
<td>Explosive Charges &amp; Fuzes Used in Italian Bombs of WWI</td>
<td>147</td>
</tr>
<tr>
<td>Italian Mining Explosives with Nitroglycerol</td>
<td>158</td>
</tr>
<tr>
<td>Italian Pulverulent Mining Explosives with Nitroglycerol</td>
<td>159</td>
</tr>
<tr>
<td>Italian Pulverulent Mining Explosives without Nitroglycerol</td>
<td>159</td>
</tr>
<tr>
<td>Italian Mining Explosives Containing Surplus Propellants</td>
<td>160</td>
</tr>
<tr>
<td>Chemical and Physical Characteristics and Test Methods for Jet Fuels</td>
<td>72</td>
</tr>
<tr>
<td>Thermal Stability Test Conditions for Jet Fuels</td>
<td>75</td>
</tr>
<tr>
<td>Russian Blasting Cap Compositions</td>
<td>2</td>
</tr>
<tr>
<td>Isothermal Decomposition of Explosives at CJ Conditions</td>
<td>9</td>
</tr>
<tr>
<td>Influence of Kinetics on Adiabatic Explosion Times of PETN</td>
<td>9</td>
</tr>
<tr>
<td>Kinetic and Thermochemical Parameters for the Decomposition of Molten and Liquid Explosives</td>
<td>11</td>
</tr>
<tr>
<td>Arrhenius Parameters Based on the Wenograd Test</td>
<td>12</td>
</tr>
<tr>
<td>Old and Kraftzahl (KZ) Trauzl Block Expansion Test Values of Common Explosives</td>
<td>18</td>
</tr>
<tr>
<td>US Military Specification Requirements for Lead Dioxide</td>
<td>9</td>
</tr>
<tr>
<td>US Military Specification Requirements for Lead Oxide</td>
<td>10</td>
</tr>
<tr>
<td>Sales (Millions of Pounds) of Industrial Explosives</td>
<td>20</td>
</tr>
<tr>
<td>LDX Compositions and Properties</td>
<td>21</td>
</tr>
<tr>
<td>Detonation Rate of Liquid Methane and Liquid Oxygen Mixtures</td>
<td>23</td>
</tr>
<tr>
<td>Experimental Detonation Properties of Some Liquid Explosives</td>
<td>29</td>
</tr>
<tr>
<td>Common Liquid Bipropellant Systems</td>
<td>37</td>
</tr>
<tr>
<td>Properties of “Exotic” Liquid Bipropellant Systems</td>
<td>38</td>
</tr>
<tr>
<td>Current Liquid Bipropellant Applications</td>
<td>40</td>
</tr>
<tr>
<td>Typical Performance Characteristics of LP Rockets</td>
<td>42</td>
</tr>
<tr>
<td>Loading Density of Various Explosives</td>
<td>51</td>
</tr>
<tr>
<td>Typical LX Explosive Formulations</td>
<td>62</td>
</tr>
<tr>
<td>LIST OF FIGURES AND ILLUSTRATIONS</td>
<td>Page</td>
</tr>
<tr>
<td>----------------------------------</td>
<td>------</td>
</tr>
<tr>
<td>Haftohlladung (German Adhering or Stick Hollow Charge)</td>
<td>H 2</td>
</tr>
<tr>
<td>Panzerhandmine 3 (German Magnetic Antitank Charge)</td>
<td>H 3</td>
</tr>
<tr>
<td>Haftmine (German Adhering Mine)</td>
<td>H 3</td>
</tr>
<tr>
<td>Extent of Potential Hazard Rating</td>
<td>H 16</td>
</tr>
<tr>
<td>Nomograph Linking Decomposition Temperature with Arrhenius Activation Energy</td>
<td>H 19</td>
</tr>
<tr>
<td>Example of Use of Reaction Hazard Index with Methane, Ethylene and Acetylene</td>
<td>H 20</td>
</tr>
<tr>
<td>Correlation of NFPA Reactivity Rating with Reaction Hazard Index</td>
<td>H 23</td>
</tr>
<tr>
<td>Moisture Content Apparatus</td>
<td>H 33</td>
</tr>
<tr>
<td>Apparatus for 100°C Vacuum Stability Test</td>
<td>H 33</td>
</tr>
<tr>
<td>Kammerminensprengungen in Balsalgetstein (Chamber Mine Blessing in Basalt)</td>
<td>H 64</td>
</tr>
<tr>
<td>Mechanism of Rotating Prism Photoc Camera</td>
<td>H 105</td>
</tr>
<tr>
<td>Diagram of a Simple Drum Camera</td>
<td>H 108</td>
</tr>
<tr>
<td>Rudimentary Gun Shaped Like a Bottle</td>
<td>H 122</td>
</tr>
<tr>
<td>Hochdruckpumpe (HDP) Super Gun</td>
<td>H 163</td>
</tr>
<tr>
<td>German Hollow (Shaped) Charges</td>
<td>H 165</td>
</tr>
<tr>
<td>Hollow Charge Nose Attachment for AP Bombs</td>
<td>H 166</td>
</tr>
<tr>
<td>Apparatus for Determining the Temperature of Ignition</td>
<td>H 169</td>
</tr>
<tr>
<td>Steady Plane Shock Front Propagating into Undisturbed Materials</td>
<td>H 179</td>
</tr>
<tr>
<td>Alternative Hugoniot Curves</td>
<td>H 179</td>
</tr>
<tr>
<td>Transmission of a Plane Shock Wave into a Material of Lower Impedance</td>
<td>H 182</td>
</tr>
<tr>
<td>Transmission of a Plane Shock Wave into a Material of Higher Impedance</td>
<td>H 182</td>
</tr>
<tr>
<td>Shock Waves Induced in Various Materials by Normally Incident Plane Detonation Waves</td>
<td>H 183</td>
</tr>
<tr>
<td>The Rankine-Hugoniot Curve</td>
<td>H 183</td>
</tr>
<tr>
<td>Pressure-Volume Hugoniot for 1g/cc TNT</td>
<td>H 184</td>
</tr>
<tr>
<td>Definition of Pressure Values Measured on Film Strips</td>
<td>I 1</td>
</tr>
<tr>
<td>Diagrammatic Representation of Ignition</td>
<td>I 11</td>
</tr>
<tr>
<td>Apparatus for Determining Minimum Energies for Electric-Spark Ignition by Blanc et al</td>
<td>I 11</td>
</tr>
<tr>
<td>Lewis &amp; Von Elbe Model of a Minimal Flame in Ignition Process</td>
<td>I 12</td>
</tr>
<tr>
<td>Schematic Illustration of Source Energies and Critical Source Temperatures versus Critical Heating Period</td>
<td>I 13</td>
</tr>
<tr>
<td>Relation between Ignition Energy, Mean Firing Current and Critical Heating Period for Electric Fuseheads</td>
<td>I 14</td>
</tr>
<tr>
<td>Plot of Energy versus Wire Length for Electric Fuseheads</td>
<td>I 14</td>
</tr>
<tr>
<td>Relation between Cross-Sectional Area of Wire and Minimum Critical Energy per Unit Length</td>
<td>I 14</td>
</tr>
<tr>
<td>Burning Rate for Various Models of Surface Burning Rates</td>
<td>I 15</td>
</tr>
<tr>
<td>Anvil Striker Arrangement, ERL Machine</td>
<td>I 37</td>
</tr>
<tr>
<td>Drop Weight Impact Machine, ERL Model, Type 12 Tools</td>
<td>I 37</td>
</tr>
<tr>
<td>Rotter Machine Impact Chamber and Anvil-Striker Arrangement</td>
<td>I 38</td>
</tr>
<tr>
<td>Bureau of Mines Impact Apparatus Showing Anvil-Striking Pin Assembly</td>
<td>I 38</td>
</tr>
<tr>
<td>Picatinny Arsenal Impact Test Apparatus</td>
<td>I 39</td>
</tr>
<tr>
<td>Parts of Picatinny Arsenal Impact Test Apparatus</td>
<td>I 40</td>
</tr>
</tbody>
</table>
Figures (Cont'd)
Impact on a Layer Density Close to Maximum I 46
Impact on a Layer of Low Density I 46
Design of Apparatus and Connection of the Pick-up to the Oscillograph I 49
Interpretation of Oscillograph Traces of Pressure Pulses I 49
Average Pressures upon Fracture and upon Explosion vs Charge Thickness of HMX, RDX, Tetryl and TNT H 50
Average Pressures vs Charge Thickness for PETN and Trinitrophenol I 50
Spread of the Measured Pressures upon Fracture and upon Explosion for RDX I 50
Explosion Frequency vs Impact Energy for Tetryl and RDX and HMX I 51
Explosion Frequency vs charge Thickness in the Case of HMX for Various Impact Energies I 52
Time Delay Between Impact and Explosion I 53
Impedance, Acoustic and Shock Diagram I 56
Constants for EL 506D as Functions of the Ratio of Foam Neoprene to Target Weights I 60
Laser Interferometer System I 112
Relation between the Light Energy Required for Ignition and Temperature for Nitrogen Iodide I 119
Italian Cartridge with Igniter Tube I 151
Italian Cartridge Bag I 151
Idealized Projectile Types J 1
Cross Section of Open Hearth Furnace Tap Hole Showing Jet Tapper in Position for Firing J 76
Types of Jet Propulsion Engines J 78
Comparison of Turbojet & Turboprop Engines J 79
Schematic of Ramjet Engine J 80
Schematic of Pulse Jet Fitted with Flow-Check Valves J 81
Comparison of Performance Parameters of Jet Engines J 82
Line Wave Generator of the Manifold Type L 17
Line Wave Generator of Sheet Explosive L 17
Line Wave Generator of Warped Sheet of Explosive L 18
Space-Time Representation of Initiation Behavior of Nitromethane L 31
Scoop Loading of Explosive Charges L 48
Charging Plate Loading of Explosive Charges L 49
Detonator Loading Tool L 49
Tool for Direct Loading of Component L 50
Nomograph of Loading Pressure and Density L 51
Vacuum Pressing Apparatus L 53
Hydrostatic Pressing Principle L 54
Isostatic Pressing Principle L 54
SUPPLEMENT TO THE
LIST OF ABBREVIATIONS FOR BOOKS AND PERIODICALS
GIVEN IN VOL 1, pp Abbr 66 to Abbr 76; VOL 2, pp XI to XII;
VOL 3, pp XII to XIII, and VOL 4, pp XLVII to L

Blomquist OSRD 4134 (1944)  A.T. Blomquist, "Certain Aliphatic Nitramines and Related Compounds", OSRD 4134 (Nov 1944)

Bowden & Yoffe (1952)  F.B. Bowden & A.D. Yoffe, "Initiation and Growth of Explosion in Liquids and Solids", Cambridge Univ Press, Cambridge, Gt Britain (1952)


SUPPLEMENT TO THE
LIST OF BOOKS ON EXPLOSIVES AND PROPELLANTS
GIVEN IN VOL 1, p A676; VOL 2, pp C215 to C216; VOL 3, pp XIV to XV
VOL 4, pp Li to LV; VOL 5, pp XIV to XV; and VOL 6, p X

WE Inc, Old Greenwich, Conn, Vol 1 (1968)

WE Inc, Old Greenwich, Conn, Vol 2 (1968)

Anon, “Principles of Explosive Behavior”, AMCP 706-180, Engineering Design Handbook,

Brodie, Bernard & Fawn, “From Crossbow to H-Bomb”, Indiana University Press,
Bloomington (1973), 320 pp

Collective, “The Problem of Chemical and Biological Warfare”. A study of the Historical,
Technical, Military, Legal and Political Aspects of CBW and Possible Disarmament Measures.
Stockholm International Peace Research Institute, Almqvist & Wiksell, Stockholm and
Humanities Press, NY (1973)
  Vol 2. “CB Weapons Today”, 420 pp


“Blaster’ Handbook”, Explosives Division, Canadian Industries Limited, 630 Dorchester
Bld W, Montreal, Quebec (1973), 545 pp

Materiel Command, 5001 Eisenhower Ave, Alexandria, Va 22304 (Jan 1974)

Anon, “Design of Ammunition for Pyrotechnic Effects”, Engineering Design Handbook,


M.A. Cook, “The Science of Industrial Explosives”, Graphic Service & Supply, IRECO
Chemicals, Salt Lake City, Utah (1974), 449 pp

363 pp

H₂. Designation of N-Acetylaminomethylhexamethyleneaminetetramine mononitrate described in Vol 1, p A54-L

H2 Kongo or Type 98 Explosive. Japanese WWII explosive consisting of Trinitroanisole 70 & Hexanitrodiphenylamine 30%. It was used in bomb auxiliary boosters, sea mines & depth charges

H-4. The composition of this solventless Ballistite in the form of 0.869 x 0.252 x 16 inch sticks is as follows: NC/NG/DNT/Ethyl Centralite/potassium sulfate 58.00/30.00/2.50/8.00/1.50%

This propellant was used in US Ordnance rockets
Ref: Cal Tech Lab Program 209, "Investigation of the Dry Extrusion of Solvent Prepared H4 Propellant", Status Rept 1-6 (12 Jan 1945 – 16 June 1945)

H-5. An American rocket propellant consisting of: NC/NG/Et: Cent/DNT/K₂SO₄/triacetin/Pb stearate (added) 58/20/8.2.5/1.5/10%/0.5% NC (13%). Its burning rate is 0.3 to 0.8 inches per second for 1000 to 4000 psi at 122°F. The burning rate of a Russian powder examined at the same time was the same at 1000 psi, but 10-15% faster at 4000 psi

H-6 Explosive. A grey castable expl compn contg: RDX 45, TNT 30, Al 20, D-2 Comp 5 and 0.5% CaCl₂ added; mw ca 93, OB to CO₂ –66%; density 1.71 to 1.74; Ballistic Mortar Test for Power 135% TNT; Blast Effects, % TNT: In Air 127% (Shock) & 138% (Impulse); Under Water: 118% (Shock) and 154% (Bubble): Brisance by Sand Test 49.5g sand crushed vs 48.0g for TNT; Detonation Rate 7191m/sec at d 1.71, vs 6825 for TNT pressed at 1.56; Explosion Temp 610°F (min) in 5 sec; Fragmentation Test in 90mm HE, M71 Proj, No of fragments 714 vs 703 for TNT; Friction Sensitivity – unaffected; Gas Volume 733cc/g, vs 730 for TNT; Heat of Combustion 3972cal/g; Heat of Explosiv 923cal/g; Heat Test at 100° – % loss in 1st 48 hrs 0.78cc, no loss in 2nd 48 hrs and no expl in 100 hrs; Hygroscopicity d higher than for TNT; Impact Sensitivity – same as for TNT; Rifle Bullet Sensitivity – more sensitive than TNT; Sensitivity to Initiation – min priming chge less than for TNT; Storage – dry
Usage – primarily in HE charges (Ref 1, 4 & 6)
Preparation – same method as described under HBX. US Military Specification requirements and tests are described under HBX. Its Spec is MIL-E-22267A (31 May, 1963)

H-16. Code name for 1,5-Diacetoxy-2,4,6,8-tetranitro-2,4,6,8-tetr azanonorane or 1,5-Diacetoxy-1,3,5,7-pentamethylen-2,4,6,8-tetranitramine.
See Vol 5 of Encycl, p D1118-R

H19. A designation for 4-N-Methyl-2,6-N-dinitro (Bicyclo)pentamethylenetetramine

H21 or MSX. Designations for 1-Methyl-6-acetoxy-trimethylene-1,3,5-trinitramine

Haber, Fritz (1868–1934). German chemist and Nobel prize winner. Discoverer of synthetic method for catalytically producing ammonia from nitrogen and hydrogen. This discovery
enabled Germany to manufacture explosives in WWI despite the naval blockade. Did considerable work in gas warfare

Refs:  1) Hackh's Chem Dict, Blackiston, Phila (1944), p 394  2) R. Hunsilan, SS 29, 59 (1934)

Hackh & McLeod Explosive. A preliminary investigation of the effect on the explosion props of 50/50 Amatol of incorporating with it Nitronaphthalene & benzaldehyde, in accordance with a process set forth in the patent application of Messrs Hackh & McLeod, showed conclusively that the addn of 10% alpha Nitronaphthalene ("catalyst") & 2% benzaldehyde ("promoter") renders the Amatol less sensitive to friction, less brisant, and undesirably insensitive to initiation to detonation.

The results fail to substantiate the claim of the inventors and in no way bear out the theory that the efficiency of nitroexplosives can be increased by the addn of nonexplosive materials having specified intra-atomic distances

Ref:  J.D. Hopper, PATR 731 (April 1936) & PATR 743 (July 1936)

HADN. See Hexamethylenetetraminedinitrate or Hexaminedinitrate in this VOl

Haensson-bakuyaku or Type 88 Explosive. Japanese WWII expl consisting of Ammonium Perchlorate 75, ferrosilicon 16, wood meal 6 & petroleum 3%. It was used in demolition chges, mines & depth chges.


NOTE:  Compare with Carlit or Karlito described in Vol 2 of Encycl. p C68-L

Haeussermann, Carl (1853-1918). German scientist specializing in explosives. In 1891 he developed an industrial method of preparing TNT and recommended its use as a military explosive.

Ref:  C. von Hell, SS 13, 325-27 (1918)

Hafenegger Powder. A mixture patented in 1868 in England consisting of chlorates, sulfur, charcoal and potassium ferrocyanide


**Haftohladung.** (Adhering or Sticking Hollow Charge). One of the devices consisted of a conical metallic container (filled with 3 lb 5 oz of a HE) to which was attached an elongated apex, serving as a hand grip and contg the exploder pellet (PETN/Wax) and a pull (friction) delay igniter (4½ or 7 seconds). Attached to the base of the conical section was a plywood

frame-work carrying three powerful horseshoe magnets. A brass chain with a hook was attached to the framework. Total weight 3 kg.

The device could be used either as a hand grenade or as a land mine. In the first case the cord of the friction igniter was pulled off and the grenade thrown against the approaching vehicle. In the second case, the device was buried in the ground, close to the surface, with the magnets up and with the igniter cord attached to the ground. At the approach of a vehicle the magnetic attraction caused the grenade to jump towards some iron or steel part and attach itself to it. Simultaneously the cord was pulled, thus setting off the explosive train consisting of delay igniter, exploder and main charge. (Ref 2) It was claimed that this
charge could penetrate as much as 110mm of armor (Ref 1, pp 323-4).

Another magnetic antitank charge is described in Ref 1, pp 262-3 under the name of Panzer-handmine 3. It consisted of a bottle-shaped cardboard container with 2 1/3 lb of hollow charge (TNT or RDX/TNT). Three pairs of magnets were mounted at the bottom of the bottle, and a 7/4 sec friction igniter was located in the neck of the bottle. Total weight of the device was 8 lb.

The device was apparently designed to be placed by hand on the tank and the igniter pulled after it has been positioned. If the target was of non-magnetic material such as wood, the charge could be attached by means of 3 spikes located at the bottom of the device.

In another type of adhering (sticking) antitank hollow charge there were no magnets but a sticky pad (located at the wide part of the conical body) served for attaching the charge to a tank (Ref 1, p 324).


**HAFTMINE**

**Detonator Release**

- Main Charge (TNT)
- Handle

- Thermite Charge
- Low Melting Plastic
- Protective Cloth
- Protective Cover

HAFTMINE (Adhering Mine). An antitank, hollow charge device consisting of a conical container (filled with HE), provided with a flat top and a handle. The wide portion of the cone was covered with a layer of a low melting colophony-oil plastic-resin (m.p ca 50°) retained on the surface by means of an open mesh cloth. In back of the flat top, which consisted of sheet metal, was placed a thermite-type charge (Mg + Al + KClO₃) and in back of the latter a time fuse. The operator hid in a hole and, at the approach of the tank, ignited the fuse which, in turn, ignited the thermite. Just as soon as the heat of the thermite melted the resin, the device was struck (by the operator) to the bottom armor plate of the tank. At the same time the heat of the thermite set off the detonator and this in turn initiated the main charge.

This device was in an experimental stage when the war terminated.


Hahn's Explosive. A cheap explosive mixture prepared as follows: Burnt lime was dissolved in raw, odorless nitric acid so as to produce a clear transparent fluid resembling petroleum jelly. This fluid was diluted with 1 to 5 parts of water according to the desired explosive power, and then mixed with a powdered cellulosic material such as peat, sawdust, etc., until a consistant mass was obtained. After giving this mass any suitable form, it was dried and used as a
commercial explosive, e.g. in rock blasting. It was found to be a powerful explosive insensitive to shocks, friction, or pressure
Ref: M. Hahn, USP 1751326 (1930) & CA 24, 2605 (1930)

Hahn’s Powder. One of the chlorate explosives in which sensitivity of the powder due to the presence of KClO₃ was reduced by the addition of spermaceti: KClO₃ 61/8S₂S₃ 28/Charcoal 3 & spermaceti 8%. It was patented in England in 1867 and used in primers
Refs: 1) Cundill (1889) in MP 6, 8 (1893) 2) Daniel (1902), 366 3) Gody (1907), 263 4) Pérez Ara (1945), 206

Haid, August (1886-1963). German scientist, formerly director of Chemisch-Technischen Reichsanstalt. Author of numerous papers on explosives, some of them in collaboration with H. Kast, H. Selle, A. Schmidt or H. Koenen
Ref: H. Koenen & P. Dittmar, Explosivst 1963, 54-6 (Obituary, portrait and list of publications)

Haid, Becker and Dittmar Stability Test (Designed for high explosives). 42g of powdered explosive, previously dried over phosphorous pentoxide, is introduced through a side-tube into a glass vessel connected with a manometer which is in the form of a U-tube and contains mercury covered with a layer of paraffin oil. The glass vessel is also fitted with a side tap. The ensemble is heated at 75° for several hours and pressures are recorded in mm of Hg as a function of time. The steeper the pressure-time curve the less stable is the explosive
Ref 2 gives curves for principle explosives as well as for smokeless powders

Haishokuyaku. Japanese WWII explosive consisting of Ammonium Perchlorate 77, RDX 17, silicon carbide 1.5 & paraffin 4.5%. It was used in demolition blocks

Hake Effect. Some industrial explosives, even when made of normal consistency, are subject under certain conditions to a change in storage, by which they become very insensitive. Quinan (Ref) called this the “Hake Effect” in honor of C. Napier Hake, the late Inspector of Explosives for the State of Victoria, Australia
Mr. Hake was the first man of science outside the ranks of Dynamite makers to make a study of this curious & interesting observation. If a small percentage of Blasting Gelatin that has become insensitive be reworked & mixed with a large percentage of a fresh lot, it will infect the mass, and the remade Blasting Gelatin will soon undo the change. Something more than a mechanical effect is involved; in fact, an obscure form of catalytic action seems to be indicated
There are so many unknown factors involved in the “Hake Effect” that it is impossible to take them all into account. One of these is the condition of storage, especially the temp. If a Gelatin of normal consistency & sensitiveness has been stored in a comparatively high temp and is subjected for a few months to a low temp, it seems to develop the “Hake Effect” very promptly. On the other hand, a Gelatin which threatens to develop this effect may show normal sensitiveness at least for a time if transferred to a warmer climate
Ref: W.R. Quinan, “High Explosives,” Critchley Parker, Melbourne, Australia (1912), pp 27-29. See also Aging of Dynamites in Vol. 1, pp A110-R to A112-L

Halakite. An explosive submitted during WWI to the Fr & Brit authorities, consisting of Cordite mixed with Pb nitrate, Ba nitrate & Pb chromate. It was claimed to be equally effective as an HE and as a propellant
Ref: Marshall, Dict (1920), 48

Hale, G.C. (1891-1948). American scientist who served as Chief of the Chemical Department, Picatinny Arsenal (1929-48). After military service Dr Hale was assigned in 1920 to a task group which was sent to Germany for an exhaustive study of the Ger expl industry of WWI. He made numerous inventions, held many patents, and authored extensive publications in the field of military
propellants & explosives. His name has been
given to an explosive called Haleite because of the important part he played in its develop-
ment as a military expl.

See Ethylenedinitramine in Vol 6, p E238-R & Cyclonite or RDX in Vol 3 of Encycl, p C611

Haleite. See Ethylenedinitramine under Ethy-
lenediamine and Derivatives in Vol 6 of Encycl, p E238-R

Halford-Kistiatowsky-Wilson Equation of State.
See Vol 4, p D608, Eqs 23 & 24

Hall (or Will) Powder. Patented in 1863 in Eng-
land, it contained potassium chlorate 47, po-
tassium ferrocyanide 38 & sulfur 15%
Refs: 1) Cundill (1889) in MP 6, 8.(1893)
2) Daniel (1902), 366 3) Giua, Trattato 6
(1959), 391

Hall (John) & Sons. Employed since 1875
charges of compressed. Nitrocellulose in
blasting of sandstone and clays and found that
such charges were very effective
Ref: Daniel (1902), p 366

Note: Later, compressed Nitrocellulose was
used extensively for bursting charges in shells,
bombs & mines. As late as WWI nearly all
small demolition (called pyroxylinovyhshashki)
blocks used in Russia were of compressed
pyroxyline. Bursting charges in older models
of Whitehead torpedoes also used compressed
Nitrocellulose, contg ca 18% water
Ref: Col M.M. Kostechh, Private Communication,
(1955)

HALOBIPHENYLS

2-Chloro-4',5-dinitrophenyl has been prep-
by the mixed acid nitration of 2-chlorobiphenyl,
mp 159°, further nitration may yield small
amt of a tetranitro deriv melting at 127°
Ref: G.H. Beauveau et al., JChemSoc (1961)
2749 & CA 56, 3377 (1962)

Haloclastite or Haloklastit. Same as Petroklasti-
tie (Ger): One of the pre-WWI expls used in
potash mines and stone quarries: Na-nitrate 69,
K-nitrate 5, sulfur 10, coal tar pitch 15,
K-dichromate 1%
Refs: 1) Marshall 1 (1917), 89 2) Stettbacher
(1933), 111 3) PATR 2510 (1958), Ger 130-L

Halodinitrophenylnitraminooethyl Nitrates. See
under Phenylaminoethanol & its derivatives

Halogenated Acetylens

Mono and di-halogen-substituted acetylenes and
their metal salts are very unstable, highly flammable,
and many of them are highly explosive:

Acetylene Bromide, Bromoacetylene, Bromo-
ethylyne or Ethynylbromide (called Bromathin
or Bromacetylen in Ger). BrC::CH; mw 104.93;
colorless gas, bp -4.7°, Qvap 58cal/g, d, 4.68g/l
at 0° & 1 atm; sl sol in w; sol in eth & di
HNO3; toxic. Prep'd from alc KOH & bromo-
minated hydrocarbons, also Br2C::CHBr & HgCl2
& KCN to give the mercuric salt (Ref 1).

Addnl thermodynamic data (see Acet Chl for
definitions): (Hg-Cl2)/T 9.702, (Fe-Cl2)/T 49.52,
cp 13.31 cal/°mole & S° 59.22 eu (entropy
units) (Ref 10)

It is pyrophoric and w sohs give off O2
and phosphoresce. Hg(CBr2)2 darkens and
explodes above 153°. It also explodes with shock

Acetylene Chloride, Chloroacetylene, Chloroethylyme
or Ethynyl Chloride, (called Chloroathin or

Chlor-acetylen in Ger), CIC::CH; mw 60.48;
color-
less gas, bp-29.6° Qvap 89 cal/g (at bp); d 2.0
(Ref 4).
Soln in w & alc (water soln gives off
O2 and glows in dark), nauseating odor and
probably anesthetist if inhaled. Prep'd from
Cl2C::CHCl & alc KOH (Ref 6); Cl2C::CHOCH
& Ba(OH)2; CHCl::CHCl & KOH, HgCl2 & NaCN
to give Hg(C::CCl2)2 which gives acetylene
chloride on heating with NaCN and KOH under
N and drying with H2SO4-P2O5 (Ref 2).

Addnl thermodynamic data: H2-E°/T 9.442,
\[ F_2\text{E}^0/T \text{G} 46.96 \text{ c}_p^0 \ \text{13.31} \text{ (all in cal/deg mol)} \]

\[ S^0 \text{56.40 eu, } H_2, F_2, \text{c}_p^0 \text{ & } S^0 \text{ are respectively standard heat of formation, stand free energy of form; heat capacity & std entropy). } E_\text{p}^0 \text{ is heat of formation of a perfect gas at abs zero (Ref 10)} \]

Acetylene chloride behaves very unpredictably. It is spontaneously flammable in air and it explodes when heated in air or shocked. On pyrolysis it gives off the very toxic phosgene (Ref 8). An attempt at defining its explosion limits is made in Ref 2.

The following metal salts of acetylene chloride are known (Ref 2): 1) Hg (C\text{CCl})_2$. See above for prep; mp 185°, explodes 195° 2) AgC\text{CCl}, prep by treating acetylene chloride with an ammoniacal Ag soln to obtain a very unstable white pptate which darkens in air. It explodes on shocking even under water 3) Cu C\text{CCl}, prep by treating acetylene chloride with ammoniacal CuCl; shock sensitive.

Dibromoacetylene or Dibromoethylene (called Dibromäthin or Dibromacetylen in Ger) Br\text{C\text{Br}}; mw 183.82; colorless liq. mp $\sim 24^\circ$, bp $\sim 76^\circ$, d $\sim 2$, ins in w; sol in org solns; toxic; vap inhalation produces violent headaches & dizziness. Prep by action of alc KOH on Br\text{C\text{Br}} or aq KOBr on acetylene (Ref 3). It is very unstable and will explode in N\text{2} or CO\text{2} containing traces of O\text{2}; also explodes on heating.

Dichloroacetylene or Dichloroethylene (called Dichloräthin or Dichloracetylen in Ger), C\text{Cl\text{Cl}}; mw 94.92; colorless liq. mp $\sim 66^\circ$, bp 32 to 33° (748mm); sol in Et\text{2}O & CCl\text{4}; ins in w. Prep by passing hot Cl\text{2}C\text{HCl} over KOH at 130° (Ref 4). Explosive on exposure to air and on heating to over 130°.

Diiodoacetylene or Diiodoethylene (called Diiodäthin or Diiodacetylen in Ger). ICl\text{Cl}; mw 277.82; yellowish rhombic crts (from ligno); mp 81°; bp 80 to 100° (decomp); sl sol in cold ligno; v toxic—attacks mucous membranes. Prep by action of alc KOH or NaOEt on C\text{2I4}, Cl\text{Cl\text{Cl}}; by decomposing C\text{2I4} in sunlight; from C\text{2H2} + K\text{I\text{Cl}} or KOBr in w; from C\text{2H2} or Na\text{2H2} or CaC\text{2}+ I\text{2} in liq amm (Ref 5). Addnl thermodyn data (at 373.15 K)—see Acetylene Chloride for definitions—H\text{2}, E\text{2} 13.90, F\text{c}, E\text{c} 64.75, cal/mole c\text{P} 17.44 cal/deg mole, S\text{0} 78.65 eu (Ref 9). Explos on grinding, explodes on heating above 120°, decomp at lower temps when heated in light.

Iodoacetylene, Acetylene Iodide, Ethynyliodide, or Iodoethyne (called Iodäthin or Iodoacetylen in Ger). IC\text{H}; mw 151.93, bp 32°, one of the products of the reactn of BrMgC\text{HCl} with I\text{2} (Ref 6).


**HALOGENATED AMINES**

**HALOGENATED AMIDES**

Organic compounds containing halogen atoms attached to the nitrogen. A large number of such compounds were prep by treating amines with hypochlorites. Some of these compounds are explosive:

**N,N-Dichloroformamide, HCONCl2**, mw 113.93, N 12.30%, yellowish-red liquid which may be prep by either the action of hypochloric acid or chlorine on formamide dissolved in water and cooled with ice-salt mixture. It is an explosive which already starts to decompose slowly at 0°. When brought to room temperature, it explodes spontaneously within a few seconds (Refs 1 & 4).

**N,N-Ethylidichloroformamide, N,N-Dichloropropionimid**, mw C\text{2H5}CONCl2, mw 141.99, N 9.87% bp 88-89° at 762mm. It was prepared by Riedal (Ref 3) and claimed to be a powerful explosive suitable as a replacement for NG.

The following explosive derivatives of aliphatic diamines were prep by Chattaway (Ref 2):

**Ethylentetrachloramidodiamine, or N,N,N,N’-Tetrachloroethylenediamine; Cl\text{4NCH2CH2NCI2**, mw 197.88, N 14.16%, yellow liquid, mp 4-4.5°**
bp. 76-78° at 10 mm (Ref 5); bp 116° at 50 mm (Ref 2), d 1.544 at 20° (Ref 5). Explodes with great violence on heating (Ref 2). Can also be prpd in H₂O/Cl₂/NaICO₃ soln (Ref 5).

**Ethyleneetetrabromodiamine**, Br₂NCH₂CH₂NBr₂; mw 375.68, N 7.46%; orange-colored crts, mp about 62° with decomposition, followed almost immediately by a violent explosion (Ref 2).

**s-Diacetylethylene dibromodiamine**, CH₂CONBrCH₂CH₂NBrCOCH₃; mw 301.98, N 9.28%; pale yellow prisms, mp 150-155°, giving off bubbles of gas and almost immediately exploding (Ref 2).

**s-Dipropionylethylene dibromodiamine**, CH₃CH₂CONCICOCH₂CH₂; mw 241.14, N 11.62%; yellow liquid, decomposes almost explosively on heating (Ref 2).

**s-Dipropionylethylene dibromodiamine or N,N’-Dipropionyl-N,N’-dichloroethylenediamine, CH₃CH₂CONBrCH₂CH₂NBrCOCH₂CH₃; mw 330.04, N 8.49%; flattened pale yellow needles, mp 112°; explodes at 160° (Ref 2).**

**s-Diphenylacetylene dibromodiamine, or N,N’-Dichloro-N,N’-ethylene-bis-phenylacetamide, C₆H₅CH₂CONBrCH₂CH₂NBrCOCH₂C₆H₅; mw 454.18, N 6.17%; pale yellow plates; mp 128°.**

Explodes at about 150° (Ref 2).

**Di-m-nitrobenzoyl ethylene dichlorodiamine, or N,N’-Dichloro-N,N’-ethylene-bis-3-nitrobenzamide, O₂NCCH₂CONCICH₂CH₂NClOC₂H₄NO₂; mw 427.22, N 13.12%; nearly colorless plates with slight yellowish tint, mp 173°. Stable up to about 220°, but explodes when heated in a flame (Ref 2).**

**Di-p-nitrobenzoyl ethylene dichlorodiamine, or N,N’-Dichloro-N,N’-ethylene-bis-[4-nitrobenzamide, O₂NCCH₂CONCICH₂CH₂NClOC₂H₄NO₂; mw 427.22, N 13.12%; nearly colorless plates with slight yellow tint, mp 207°; decomposes at about 215°.**

Explodes when heated in a flame (Ref 2).

**s-Dibenzenesulfonylethylene dichlorodiamine, or N,N’-Dichloro-N,N’-ethylene-bis-benzenesulfonamide, C₆H₄SO₂NCICH₂CH₂NClOC₂H₄; mw 413.32, N 6.78%; colorless prisms, mp 113°; starts to decompose at 200°.** When heated strongly in a flame it decomposes explosively (Ref 2).

**s-Dibenzenesulfonylethylene dibromodiamine, or N,N’-Dibromo-N,N’-ethylene-bis-benzenesulfonamide, C₆H₄SO₂NBrCH₂CH₂NBrCOCH₂C₆H₄; mw 502.22, N 5.58%; pale yellow prisms, mp 134°; decomposes at 165-175°. When heated rapidly in flame, it explodes (Ref 2).**

**Di-p-toluene sulfonyl ethylene dibromodiamine, or N,N’-Dibromo-N,N’-ethylene-bis-toluene sulfonamide, H₂C₆H₄SO₂NBrCH₂CH₂NBrCOCH₂C₆H₄; mw 530.28, N 5.28%; pale yellow plates, mp 165°; decomposes beginning at 170° and explodes at about 180° when heated rapidly (Ref 2).**

**Di-m-nitrobenzene sulfonyl ethylene dichlorodiamine, or N,N’-Dichloro-N,N’-propylene-bis-benzenamide, C₆H₄CONCICH₂CH₂NClOC₂H₄; mw 337.22, N 8.31%; colorless plates, mp 85°. Explodes when heated above 160° (Ref 2).**

**Refs:** 1) Bellstein, 2 (22) 2) F.D. Chattaway, JCS, 87, 381 (1905) 3) J.D. Ridge, Ger 301, 799 & CA 15, 1966 (1921) 4) P.L. Magil, IEC 26, 611 (1934) 5) L.K. Jackson et al, JACS 69, 1539 (1947) & CA 41, 5852 (1947).

### HALOGENATED DERIVATIVES OF SULFONAMIDES

The sulfonamides are distinguished by the great readiness with which they form well-crystallized derivatives in which all the amine hydrogen is replaced by halogens, when acted upon by hypochlorous acid at ordinary temperatures. The sulfonyl chloramides so produced exhibit great stability when compared with other chloramides. Those containing two chlorine atoms attached to nitrogen, melt without decomposition and can frequently be heated considerably above their melting points without undergoing any change, but they explode when heated by direct flame. The tetrachloramides derived from disulfonamides resemble nitrogen chloride itself in the violence with which they detonate. When similarly heated, the sulfonalkylchloramides do not detonate, but decompose rapidly with evolution of gas. They are used as external antiseptics — "sulfa drugs.”

The following explosive sulfonylchloramides and their salts were prepd by Chattaway (Ref 12):

**I. Sulfonylchloramides and Sulfonylalkylchloramides**

Benzenesulfonylchloramide or N,N-Dichlorobenzensulfonamide (called N,N-Dichlorobenzol-sulfamid in Ger), C₆H₄SO₂NCl₂; mw 226.08, N
6.20%; slightly yellowish crystals, mp 76°. When heated rapidly in a flame, explodes feebly (Ref 12). Its anhydrous Potassium salt, \( \text{C}_6\text{H}_5\text{SO}_3\text{K-NCl} \); mw 229.7, N 6.1%; colorless prisms, explodes when heated rapidly at 140-145° and the Sodium salt, also called Chloramine B, \( \text{C}_6\text{H}_5\text{SO}_3\text{Na-NCl} \); mw 213.6, N 6.6%; explodes at 180° to 185° (Refs 1 & 12).

Toluene-o-sulfondichloramide or \( \text{N,N-Dichloro-o-toluolsulfonanide} \), (called Dichloro-o-toluolsulfanid in Ger), \( \text{C}_6\text{H}_4(\text{CH}_3)\text{SO}_2\text{Cl}_2 \); mw 240.11, N 5.83%; mp 33°. This compound is not listed as an explosive, but its anhydrous Potassium salt explodes feebly at 145°, while its Sodium salt explodes with some violence when heated to 170-175° (Ref 2 & 12).

Toluene-p-sulfondichloramide or \( \text{N,N-Dichloro-p-toluolsulfonanide} \) (called \( \text{N,N-Dichloro-p-tolüolsulfanid} \) in Ger), \( \text{C}_6\text{H}_4(\text{CH}_3)\text{SO}_2\text{Cl}_2 \); mw 240.11, N 5.83%; mp 33°, is not listed as an explosive, but its anhydrous Potassium salt explodes with violence at 160-165° and the Sodium salt (also called Chloramine T) at 175-180°. Large quantities of the Na salt are toxic (Refs 3 & 12).

Nitrobenzene-m-sulfonchloramidine, Potassium and Sodium Salts. \( \text{C}_6\text{H}_4\text{SO}_2\text{K-NCl} \); mw 274.72, N 10.20%; and \( \text{O}_2\text{NC}_6\text{H}_4\text{SO}_2\text{Na-NCl} \); mw 258.61, N 10.83%. Both anhydrous salts are explosive. The first explodes at 155°; the second at 175° (Ref 12).

Nitrotoluene-p-sulfondichloramide, \( \text{H}_3\text{CC}_6\text{H}_4(\text{NO}_2)\text{SO}_2\text{Cl}_2 \); mw 285.11, N 9.83%; mp 101°, is not listed as an explosive, but its anhydrous Potassium salt explodes at 160°. No sodium salt is described (Ref 4 & 12).

Benzene-m-disulfonfetachloramide, \( \text{C}_6\text{H}_4(\text{SO}_2\text{NCl})_2 \); mw 374.04, N 7.49%; mp 128-30°. Colorless, transparent rhombic crystals (from pet eth). When strongly heated the melted substance explodes with a violence recalling the explosion of nitrogen chloride itself (Refs 5 & 12).

Naphthalene-2-sulfondichloramide, \( \text{C}_6\text{H}_4\text{C}_6\text{H}_4\text{SO}_2\text{NCl}_2 \); mw 276.14, N 5.07%; is not listed as an explosive, but its anhydrous Potassium salt explodes feebly at 170° and the Sodium salt at 180° (Ref 12).

Naphthalene-2,7-disulfonfetachloramide, \( \text{C}_6\text{H}_4\text{SO}_2\text{C}_6\text{H}_4\text{SO}_2\text{NCl}_2 \); mw 456.10, N 6.14%; colorless, transparent pyramids, mp 165° (Ref 6). The melted substance explodes with great violence when strongly heated. Its anhydrous Potassium, \( \text{C}_6\text{H}_4\text{SO}_2\text{K-NCl} \); mw 431.40, N 6.49% and Sodium, \( \text{C}_6\text{H}_4(\text{SO}_2\text{Na-NCl})_2 \); mw 399.18, N 7.02%; salts are in the form of colorless hair-like crystals, which decompose explosively at 145-150° and 165-170° respectively (Ref 12).

II. Arylsulfonalkylamides.

Compounds of this nature are readily produced by the action of an aqueous solution of hypochlorous acid on the sulfonalkylamides. These compounds, when rapidly heated, decompose with the evolution of gas but without explosion. Two examples are:

Benzesulfonmethylchloramide, \( \text{C}_6\text{H}_5\text{SO}_2\text{NCICH}_3 \)

Naphthalenesulfonmethylamide, \( \text{C}_6\text{H}_5\text{SO}_2\text{NHCH}_3 \)

III. Sulfondibromamides and Sulfonalkylbromamides.

Toluene-p-sulfondibromamide or \( \text{N,N-Dibromo-p-toluolsulfonanide} \), \( \text{H}_3\text{CC}_6\text{H}_4\text{SO}_2\text{NBr} \); mw 329.01, N 4.26%; orange-colored plates, mp 104°, partially sol in chl (Ref 7). When heated rapidly it decomposes explosively. Its anhydrous Potassium and Sodium salts explode mildly at 145-150° (Ref 12).

Toluene-o-sulfondibromamide, \( \text{H}_3\text{CC}_6\text{H}_4\text{SO}_2\text{NBr} \); mw 329.01, N 4.26%; orange-colored rhombic plates, sl sol in pet eth, mp 80° (Ref 2). When heated rapidly above its mp it decomposes explosively. Its anhydrous Sodium salt decomposes explosively at 135-140° (Ref 12).

Nitrobenzene-m-sulfondibromamide or \( \text{N,N-Dibromo-3-nitrobenzenesulfonanide} \), \( \text{C}_6\text{H}_4(\text{NO}_2)\text{SO}_2\text{NBr} \); mw 359.98, N 7.78%; orange rhomboids, mp 157° (Ref 8). When heated in flame it decomposes explosively. Its anhydrous Potassium and Sodium salts decompose explosively when heated.

2-Nitrotoluene-p-sulfondibromamide, \( \text{H}_3\text{CC}_6\text{H}_4(\text{NO}_2)\text{SO}_2\text{NBr} \); mw 374.01, N 7.49%; transparent prisms, sl sol chl, mp 142-143° (Ref 9). When heated quickly to a high temperature it decomposes explosively (Ref 12).

The rest of the compounds described by Chattaway (Ref 12) do not explode but merely decompose with the evolution of gas.

IV. \textit{N-Halogen Derivatives of P-Halogen substituted Benzenesulfonamides}.

Baxter and Chattaway (Ref 13) describe several compounds of this class, some of which are explosive:

p-Bromobenzensulfondichloramide or \( \text{N,N-Dichloro-3-bromobenzenesulfonanide} \),
C₆H₄Br₂SO₂NCl₂, mw 304.97, N 4.59%; mp 106°C (Ref 10). It is not listed as an explosive but its anhydrous Potassium salt explodes, without melting, at 165°C and its Sodium salt at 178°C (Ref 13).

p-Bromobenzenesulfonfildigromamide, C₆H₄BrSO₂-NBr₂, mw 393.87, N 3.56%; mp 132°C dec. It is not listed as an explosive but its anhydrous Potassium and Sodium salts explode without melting at 193°C and 211°C respectively (Ref 12). p-Iodobenzenesulfonfildigromamide, C₆H₄I₂SO₂NCl₂, mw 351.97, N 3.98%; mp 147°C dec, is not listed as an explosive but its Potassium and Sodium salts explode without melting at 150°C and 185°C respectively (Ref 12).


Halogenation. Incorporation of one of the halogen elements into an organic compound, e.g. C₂H₄ + Br₂ = C₂H₄Br₂. Review articles and books on halogenation are listed below


Halogen Azides are described in Vol 1 under Azides: BrN₃, p A525; CIN₃, p A529; FN₃, p A536; & IN₃, p A542. A recent review (Ref 1) gives the preps, chem & phys props of halogen azides, and reactions of metal halogens to give metallohalogenazides. Also discussed are reactions of metal carbynls & organometallic comps with halogen azides, & stability relations of azides (Ref 1). The heat of decomposition and max flame temp of CIN₃ have been measured recently (Ref 2). They are, respectively, -93.2 kcal/mole and 3380°K.


Halogen-Metal Interconversions with Halogenated Anilines. A highly explosive compound which appears to be p-N,N-Trilithioaniline is formed as a by-product in the reaction of BuLi and p-Bra-aniline but not o-Bra-aniline (Ref 1). More recently this reaction was repeated but no by-product is mentioned (Ref 2).


Halotetrazole Salts. The hazards heretofore encountered in the prep of halotetrazoles have been eliminated, and a safe method for diazotizing 5-aminotetrazole has been devised. Several salts of halotetrazoles were prepd, and aside from results of practical importance such as the production of comds of interest as initiator materials, this work may give rise to some important theoretical consideration as to the fundamental chemistry of initiators.

Preps of the following halotetrazoles are described by Gaughran & Kaufman (Ref): Copper Chlorotetrazole in 74% yield, Chlorotetrazole in 55% yield, Silver & Mercury Chlorotetrazole in quantitative yield, Copper Bromotetrazole in 75% yield, Bromotetrazole in 70% yield, Silver & Mercury Bromotetrazole, Iodotetrazole in about 10% yield.

It was recommended that some of the salts be thoroughly investigated for possible application as initiator materials.


1-Halo-2-vinylacetylenes. Chlorine, bromine & iodine vinylacetylenes can be prpd by the reaction, at 0°C, of the halogen in aq KOH with monovinylacetylene (Ref 1). All polymerize in air to form explosive polymers, with the 1-compds being the most explosive of the three. Polymerization is also accelerated by UV, ozonides,
benzoyl peroxides, etc. Additional methods of preparation and props of the individual comps are given below:

1-Chloro-2-vinylacetylene, 4-Chloro-1-buten-3-yne, CIC:CCH:CH₂; mw 86.52; color liq (darkens on air exposure), bp 55-57°, d 1.003 at 20°, Rf 1.4663 at 20° (Ref 2); insol in w; sol in chlf. Can be prpd by treating Me₂NCH₂CH₂:CHCl with Mel followed by treatment with base (Ref 5). Can be distilled under nitrogen but residue tends to explode, as does polymer formed in air. Hydroquinone appears to act as a preservative (Refs 2 and 3a)

1-Bromo-2-vinylacetylene or 4-Bromo-1-buten-3-yne, BrC:CCH:CH₂; mw 130.97; color liq (darkens on air exposure), bp 52-53° at 217mm, d 1.480 at 20°, Rf 1.5182 at 20°; insol in w; sol in chlf. Can be prpd by reacting Br with (CH₂:CHC:Cl)₂Hg (Ref 3). The bromopolymer is more explosive than chloropolymer. Bromopolymer formed in the absence of air is not explosive

1-Iodo-2-vinylacetylene or 4-Iodo-1-buten-3-yne, IC:CCH:CH₂; mw 177.97; colorless liq (darkens on air exposure) bp 71.5° at 102 mm (Refs 2 & 3), 78° at 125 mm, d 1.887 at 20°, Rf 1.5948 at 20° (Ref 2); insol in w; sol in chlf & Et₂O (Ref 4). Can be prpd by reaction of I and vinylacetylene in liq ammonia (Ref 4); the reaction of I and vinylacetylene-MgBr in ether; or I and (CH₂:CHC:Cl)₂Hg in chlf (Ref 2)


Halsey and Savage Explosives. Several smokeless powders, containing Ammonium Picrate, were patented in the US in 1896. They were prpd by adding finely pulverized Ammonium Picrate to an aqueous solution of potassium dichromate in such a manner as to form a plastic mass and then slowly adding a concd aqueous solution of potassium permanganate.

No 1 Cannon Powder: Amm Picrate 68, K₂Cr₂O₇ 25, KMnO₄ 7%. No 2-Sporting Powder: Amm Picrate 73, K₂Cr₂O₇ 20, KMnO₄ 7%. No 3 Rifle Powder: Amm Picrate 50, K₂Cr₂O₇ 20, KMnO₄ 7, Ba or Sr nitrate 23%

Ref: Daniel (1902) 367

Halstead Arsenal. An Armament Research Center, Kent, England

Halved Cartridge Gap Method. See under Phys Tests in Vol 1, p XIV

Hamilton patented in England in 1896 the addition of up to 15% of urea oxalate in order to lower the temperature of explosion of dynamites and smokeless powders

Ref: Daniel (1902), 367

Hancock's Explosive. A blasting composition patented in 1911 consisting of: KCIO₃ 16, KNO₃ 12, brown sugar 3, K₂Cr₂O₇ 3, S 6, charcoal 1, lamp black 1 part

Ref: W.P. Hancock, USP 995134 (1911) & CA 5, 2724 (1911)

Handhabungssichere Sprengstoffe. Ger for Explosives Safe to Handle and Transport

Ref: Davis (1943), 347

Handgrenade. A grenade to be thrown by hand. See Grenades

Handling Bombs. See Bombs in Vol 2, p B238
**Handling Explosives.** The general subject of handling explosives is discussed in Ref 2. Specific info on the handling and storing of common military explosives (taken from Ref 1) is tabulated below.

Graphite-filled PVC provides resilient coatings which can leak off dangerous electrostatic charges from explosives & propellants (Ref 3). Remote control sites for handling explosives capable of withstanding detonations of up to 10 lbs TNT are described (Ref 4).


### Handling Characteristics of Explosives

<table>
<thead>
<tr>
<th>Explosive</th>
<th>Storage</th>
<th>Precaution</th>
<th>Destroying Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mercury Fulminate</td>
<td>Wet</td>
<td>Sensitive to flame and sparks</td>
<td>Saturated sodium thiosulfate</td>
</tr>
<tr>
<td>Lead Azide</td>
<td>do</td>
<td>Sensitive to flame and sparks. Forms sensitive copper azide</td>
<td>Ammonium acetate</td>
</tr>
<tr>
<td>Lead Styphnate</td>
<td>do</td>
<td>Sensitive to flame and to static electricity</td>
<td>Sodium carbonate</td>
</tr>
<tr>
<td>DDNP</td>
<td>do</td>
<td>Sensitive to flame</td>
<td>Cold sodium hydroxide</td>
</tr>
<tr>
<td>Tetracene</td>
<td>do</td>
<td>do</td>
<td>Hydrolyzed slowly by alkalies</td>
</tr>
<tr>
<td>Nitromannite</td>
<td>do</td>
<td>Sensitive mixtures with iron and copper oxides, poison when taken orally</td>
<td>Hot sodium hydroxide</td>
</tr>
<tr>
<td>PETN</td>
<td>do</td>
<td></td>
<td></td>
</tr>
<tr>
<td>RDX</td>
<td>do</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pentolite 50/50</td>
<td>Dry</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PTX-2</td>
<td>do</td>
<td>Forms compounds with metals</td>
<td>Boiling sodium carbonate</td>
</tr>
<tr>
<td>EDNA</td>
<td>do</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tetryl</td>
<td>do</td>
<td>Dust explosion hazard (22); contact with skin may cause dermatitis.</td>
<td></td>
</tr>
<tr>
<td>Torpex-2</td>
<td>do</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Composition B</td>
<td>do</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Composition A-3</td>
<td>do</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ednatol</td>
<td>do</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Picric Acid</td>
<td>do</td>
<td>Forms sensitive lead and copper salts</td>
<td></td>
</tr>
<tr>
<td>HBX</td>
<td>do</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tritonal 80/20</td>
<td>do</td>
<td>Slightly toxic; reacts with alkalies and ammonia to form sensitive compounds</td>
<td></td>
</tr>
<tr>
<td>TNT</td>
<td>do</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Picratol</td>
<td>do</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Explosive D</td>
<td>Dry-wooden containers</td>
<td>When wet reacts slowly with lead and copper to form sensitive salts</td>
<td></td>
</tr>
<tr>
<td>Black Powder</td>
<td>Dry</td>
<td>Particularly sensitive to flame and sparks</td>
<td></td>
</tr>
</tbody>
</table>
Handy-Andy Riot-Control Cartridge. See E24 (Cartridge) in Vol 5, p E1-L.

Hangfire. A brief undesired delay in the functioning of ammunition or blasting charges after initiation. It usually refers to delay in ignition of a propelling charge. This should not be confused with "misfire", when the shot or round does not fire at all. Hangfire is of no great importance when blasting charges are fired individually or when weapons such as cannons are discharged individually. But it could be disastrous if hangfire were to take place in an automatic weapon firing at high speed, for example, in machine guns or anti-aircraft guns (Refs 1, 2 & 4).

Twelve reported hangfires since 1932 are stated to be explainable if it is assumed that the explosives involved delagated at a burning rate not exceeding 300 m/sec (Ref 3).


Hangfire Primers. A hangfire in small arms ammunition (initiated by a blow from a firing pin) can be defined as that condition which exists when initiation of the usual chain of events following the release of the trigger in a loaded weapon, occurs at a rate slower than normal; the definition of hangfire for electrically initiated ammo is similar.

A hangfire may be of very short duration or may be several seconds in duration. Hangfire in general can be caused by: weapon defects, slow burning of primer compn and/or a slow rate of ignition or combustion of the propellant charge.

In order to measure hangfire of primers, it is necessary to start a chronograph when the primer is initiated and to stop it at the first appearance of the primer flash at the flash hole. This time interval can be considered to be a measure of "primer time", and any significant increase above the average can be considered, to be a primer hangfire.


Hangfire Test. Due to the fact that delay in firing of individual shots (hangfire) in automatic weapons (especially in the case of machine guns firing synchronously with a propeller of an airplane) would bring disastrous results, it is necessary to subject the weapons and ammunition to a special test. This test usually consists of firing 600 rounds, in bursts of 75 rounds each, through a disc rotating at 1800 rpm, the groupings being measured in degrees from the leading edge of the shot nearest the zero mark. For aircraft use, the groupings must not exceed 15°; for other uses 27°.

The test may also be conducted in a special "electrostatic hangfire machine", which should give the time between the shots of 0.0014 seconds for aircraft weapons and 0.0025 sec for other weapons.


Hannan Explosives. F. Hannan in 1882 developed a chlorate expl consisting of K Chlorate 50, K nitrate 24, K ferroanide 12, charcoal 10 & paraffin 4%.


Hansen or Hydrogen Ion Concentration Stability Test. Hansen (Ref 2) gives the following test applicable to propnts, provided they contain not more than 0.5% of CaCO₃: a) Dry ca 50g of ground powder in a vacuum oven at 40° for 8 hrs b) Put 5g samples into each of 8 tubes 16 mm diam & 360 mm long, graduated to 50 ml and provided with glass stoppers c) Place the tubes into a constant temp bath maintained at 110° d) After one hr of heating, remove No 1 tube, cool 1/2 hr in air, and add distilled water of pH 7.0 up to the 50 ml mark. Shake well and determine pH by the quinhydrone method e) Remove the other tubes, one at a time, at intervals of one hr (ie after 2, 3, 4, 5, 6, 7 & 8 hrs of heating) and determine pH of each f) Plot the pH values on graph paper vs time and draw a curve
Stability ($K$) is determined from the formula:

$$K = 15 - (H) \times 10^4,$$

where ($H$) is absolute hydrogen ion concentration. For example, if $pH$ is 3.0, then $H$ is $10^{-3}$ and $K = 15 - 10^{-3} \times 10^4 = +5$, and the powder is satisfactory. If $pH$ is 2, then $H$ is $10^{-2}$ and $K = 15 - 10^{-2} \times 10^4 = -85$ and the powder does not pass the test.

In a modified method, the powder is heated under water at $100^\circ$ to induce hydrolytic decompn

Refs: 1) L. Metz, SS 21, 186-88 (1926) & CA 21, 1013 (1927) 2) N.L. Hansen, Förh III Nord Kemistmóttet (Finland) 1926, 227-30 (1928) & CA 23, 4074 (1929)

Hansen-Metz Potentiometric Test at $110^\circ$. The following modified Hansen-Metz potentiometric test at $110^\circ$ is described by da Rocha (Refs): Samples of propint are cut to pass a 2 mm sieve and retained on a 0.5 mm sieve. They are dried for 12 hrs under vacuum at 25-30$^\circ$ and placed in tubes used in the Bergmann-Junk-Mayerhofer Test (18 + 1 mm internal diam & 360 mm long). After heating up to 8 hrs, the $pH$ is determined. For an absolute hydrogen ion concentration, $H$, after 8 hrs of exposure at $110^\circ$, the stability is expressed as $K = 15 - H \times 10^4$. Positive $K$ means good stability, while negative $K$ means poor stability.

NOTE: Da Rocha proposed the terms static and dynamic stability determinations. The first term refers to a single determination of the degree of denitrination, while dynamic stability is a trend expressing the tendency to denitrination of a powder with the time of exposure to $110^\circ$ from 0 to 8 hrs.


Hardingham proposed in 1884 in France, for mining use, cartridges containing dynamite combined either with liquifide NH$_3$ or CO$_2$ or with flammable liquids such as alcohol, benzene or ether.

Ref: Daniel (1902), 368

Hardy's Powders. Several powders were known under this name, for example NaNO$_3$ 60.1, KNO$_3$ 14.1, sulfur 8.1, charcoal 11.6, sugar 4.9, moisture 1.2%.

Ref: Daniel (1902), 368

Hargreaves Theory of Aging of Blasting Gelatins. See Ageing of Dynamites in Vol 1 of Encycl, p A110-R

Harlé (Cordeau détonant de mine à la tolite). A French detonating cord used in mining. It consists of a TNT (core) enclosed in a lead tubing of 5.9 mm external diam. Its detonation velocity is 5055 to 5165 m/sec.

Ref: M. Dutour, MAF 24, 578 & 583 (1950)

Harpa or Harpago. See under Catapult in Vol 2, p C91

Harpoon Antiship Missile (US Navy). THE NAVY'S HARPOON ANTISHIP MISSILE (McDonnell Douglas) has completed six of twenty scheduled full-scale flight tests following many launchings to test techniques and components (CD, October 1972). Harpoon has been fired successfully in its basic form from aircraft; with a 300-pound, 30-inch booster from surface ships; and from submarine torpedo tubes, with folded wings and booster enclosed in a buoyant capsule.

THE MISSILE - propelled by a Teledyne turbojet engine - is 17.5 feet long, weighs 1100 pounds, and carries a conventional warhead. It flies at wave height under midcourse inertial guidance until the radar seeker locks onto the target. The missile climbs for a brief instant near the target, then dives on it.

An outstanding feature of the weapon is its adaptability to many launchers and its compatibility with ship and aircraft targeting equipment. These include the Arios ASW weapon, late-model Terrier and Tartar launchers, and A-7 attack, S-3 ASW, and P-3 patrol aircraft.
After the twentieth flight in 1974, a third Defense Council review will make a go-ahead decision on pilot production – an extra step because the Navy, to save costs, was allowed to bypass a prototype competition.  
Ref: The Common Defense Bull No 392, June 15, 1973

Harries Method of Ozonization of Organic Compounds. See under Ozonization of Organic Compounds

Harritsite. See Composition C-4 in Vol 3, p C485

Harrison Powders (1860). A mining expl consisting of potassium chlorate 70%, starch 10%, carbon 10% & sulfur 10%. An army incendiary mixt was composed of KClO₃ 65%, starch 6%, carbon 6%, sulfur 12%, licopoldio 2%, tar oil & charcoal dust 3%. A patent of 1862 reported an expl consisting of KClO₃ or NaClO₃ 56%, K ferrocyanide 28%, starch 4%, sulfur 7% & charcoal 5%.
Ref: 1) Cundill (1889) in MP 6, 9-10 (1893)  
2) Daniel (1902), 369  
3) Giua. Trattato, 6, (1959), 391

Ref: Cundill (1889) in MP 6, 10 (1892)  
2) Daniel (1902), 369

Harry Diamond Labs. The Army has classified the Harry Diamond Laboratories at Adelphi, Maryland, as a permanent US Army installation. The facility is a corporate laboratory of the US Army Materiel Command, specializing in fuzes, fluidics, special purpose radars, nuclear weapons effects, and other research, development, and engineering for weapons systems.

Harvey's Explosive. An initiating composition which was detonated by the addition of a drop of concd sulfuric acid. It contains KNO₃ 74.3%, sugar 19.2%, gallnut 6.5%.
Ref: 1) Daniel (1902), 369  
2) Giua. Trattato 6, (1959), 400

Hasethrot. A medical name for PETN, which is used as a depressor of blood pressure

Hassia — Chlorat or Spreng — Chlorat. Ger WWI expl consist of K chlorate 65% & combustibles 35%. It was claimed to be comparatively insensitive.
Ref: Marshall, Dict (1920), 48

HAST. Abbrev for High-Altitude Supersonic Target.
Ref: Maj-Gen J.G. Zierdt, Ordn 58, May-June 1973, 480-82

Haswelite. A British permitted explosive which passed the Buxton Test (See Vol 2, p B 394). It contains Am nitrate 59%, Ba nitrate 3.5%, TNT 12%, Na chloride 25.5%.
Ref: Marshall 3, (1932), 119

Haubitze. Ger for Howitzer

Hawk. See under Missiles and Glossary of Ord (1959), 146-R

Hawkins. Patented in England, in 1894, a liquid explosive which was used as a fuel in internal combustion engines. A mixture of KNO₃ 50%, sugar syrup 33.4%, KClO₃ 9.3%, and K₂Cr₂O₇ 8.3% was dissolved in 300 parts of water and was fed by means of an injector into a combustion chamber of a motor.
Ref: Daniel (1902), 369

Hawkins Brothers. Patented in 1896 in England, a mixture of the following approximate composition KClO₃ 60 to 64%, sugar 30 to 32%, wood flour or soot 2 to 4%, potassium bichromate 2 to 8%.
Ref: 1) Daniel (1902), 370  
2) Giua. Trattato 6, (1959), 391

Hawkite. A mining explosive of the Favier type: NH₄NO₃ 59 to 63%, Ba(NO₃)₂ 4 to 2%, TNT 15 to 17%, NaCl 23 to 18%. It has 76-81% of the power of standard 60% Gelnite.
Ref: Marshall 3, 119

Hay (Foin in French). Cured grass usually used as animal fodder and in:

Nitrohay (Nitrofoin in French). An explosive, which may be prpd by the nitration of dried and powdered hay. It was patented in England
in 1873 by Trench, Faure and Mackie for use in composite explosives such as Nitrohay + resin + ozocerite, collodium cotton dissolved in a volatile solvent, glycerin and charcoal or soot.

Ref: Daniel (1902), pp 522 under Nitrofoin and 773 under Trench

Haylite. British explosive of the carbonite type: NG 25 to 27, NC 0.5 to 1.5, KNO₃ 19 to 21, Ba(NO₃)₂ 19 to 21, wood meal 12 to 14, gelatinized silicic acid 6 to 8, ammonium oxalate 10 to 12%

Ref: Naoum, NG (1928), 402

Hazards of Detonations (and Explosions). See Detonation (and Explosion), Hazards of in Vol 3, p D5-L


Hazards of Detonations (and Explosions). See Detonation (and Explosion), Hazards of in Vol 3, p DS-L

Hazards, Prediction of. D.R. Stull has developed methods of linking thermodynamics and kinetics to the prediction of potential and real hazards arising from exothermic chemical reactions (including explosions). We quote extensively from his publications.

"Modern technology has been quite successful in developing tailor-made chemicals to cope with specific problems. However, this effort has also introduced some additional problems since manufacturing and handling experience is frequently inadequate to properly answer questions of hazard. Some way is needed to distinguish hazardous from nonhazardous chemicals, and to establish a measure of the extent of hazard.

As technology solves the questions it encounters, new compositions of matter are formulated. These formulations are compounded of two or more chemicals, each of which may present no special hazard but, once mixed, may be quite hazardous. Here too, some way is needed to select the composition of hazard. Although there are many sorts of hazard, we are concerned here with the risk, danger, or peril resulting from handling or processing a given chemical or a composition of matter.

The manufacture and handling of chemicals has generated a body of experience that is useful, as far as it goes. The National Fire Codes segregates hazardous materials into five levels of reactivity ranging from zero, indicating no special reaction hazard, to four, indicating the most severe reactivity hazard. This code lists a degree of reactivity for nearly 1000 chemical materials.

For many years the explosive industry has used the drop weight test to gauge reactivity by activating a material placed on an anvil with the energy imparted by a falling weight. Drop weight tests are difficult to interpret since the physical properties and characteristics vary from one sample to the next, but the comparative order of explosives by this test is known. If a sample gives a positive drop weight test, it is hazardous, but a negative test does not necessarily prove lack of hazard. Some of the less sensitive materials require more activation than the drop weight gives. For example, tests are often run in which a No. 8 blasting cap, or a 50 g charge of Tetryl are used to provide increasing steps of activation energy.

Energy Relationships Within a Molecule

The intrinsic energy of a substance determines if decomposition will be exothermic, and its ability to react exothermically with one or more additional substances. Once initiated, a hazardous compound generates energy. The energy generated may be derived from (1) the release of stored energy in the compound, or (2) a chemical reaction forming more stable products from the atoms in the compound or mixture.

An example of (1) above is acetylene that absorbs 54.2 kcal/mole (stored within the molecule) when it is made from its elements: graphite and hydrogen gas. Once decomposition of acetylene starts, the 54.2 kcal of heat is evolved which raises the temperature of the products (graphite and hydrogen gas) to 2898 K with an accompanying pressure increase if the volume is held constant. Self decomposition is regarded as the primary hazard.

An example of (2) above is ammonium nitrate that evolves 87.27 kcal/mole when syn-
thesized from its elements: hydrogen, oxygen, and nitrogen gases. Thus, ammonium nitrate is lower on an absolute energy scale than acetylene. However, once decomposition of ammonium nitrate starts, the decomposition products, nitrogen, water, and oxygen gases, are lower on the absolute energy scale than ammonium nitrate, and -28.2 kcal/mole are released heating the product gases to 1242 °K with an accompanying pressure increase if the volume is held constant.

In addition to the energy released by decomposition of a compound, there is the possibility of additional energy release by a secondary reaction with one or more other substances present. In the acetylene example above, the initial decomposition produced graphite and hydrogen gases at a decomposition temperature $T_d$ of 2898 °K. In the presence of oxygen gas, an additional reaction to carbon monoxide and water takes place with further release of energy causing a further increase of temperature and pressure. We have arbitrarily elected to call the latter temperature (with oxygen) the “flame temperature,” $T_f$. As we shall see, these two conditions, self decomposition, and oxidation to the most stable natural products at the flame temperature, are important conditions. We shall use the symbols d for decomposition and o for oxidation” (Ref 1)

After presenting the methods of calculating the requisite thermodynamic quantities Dr Stull suggests the correlations between them and potential hazard ratings shown in Figure 1 (taken from Ref 1)

The large numbers (1,2,3 & 4) in this Figure refer to NFPA Chemical Reactivity Ratings which are described below:

0....Materials which are normally stable even under fire exposure conditions and are not reactive with water

1....Normally stable materials that may become unstable at elevated temperatures and pressures, or which may react with water with some release of energy, but not violently

2....Materials which are normally unstable and readily undergo violent chemical change, but do not detonate. Includes materials which can undergo chemical change with rapid release of energy at normal temperatures and pressures or which can undergo violent chemical change at elevated temperatures and pressures. Also includes materials which may react violently with water or which may form potentially explosive mixtures with water

3....Materials which are capable of detonation or of explosive decomposition or of explosive reaction but which require strong initiating source or which must be heated under confinement before initiation. Includes materials which are sensitive to thermal or mechanical shock at elevated temperatures and pressures or which react explosively with water without requiring heat or confinement

4....Materials which are readily capable of detonation or explosive decomposition or explosive reaction at normal temperatures and pressures. Includes materials which are sensitive to mechanical or localized thermal shock

According to Dr Stull examination of Fig 1 reveals the following:

"1. Materials that achieve zero oxygen balance, can detonate, and present the greatest hazard are along the base line

2. The lower third of the chart (up to $T_o$-$T_d$ = 1,000 °K) contains hazardous explosives or materials with an NFPA rating of 3 or 4 (shock sensitive explosive materials). The single exception is hydrogen cyanide with an NFPA rating of 2 (unstable, capable of violent chemical change, but does not detonate)

3. The middle third of the chart (from 1000 to 2000 °K) contains materials that are explosive and have NFPA ratings that range
from 4 (shock sensitive explosives), such as TNT at 1,084°K, through ratings of 2 (unstable materials that do not detonate), to a rating of 1 (normally stable materials that become unstable at elevated temperatures and pressures), such as 4-nitroaniline at 2013°K. Nitroethane with an NFPA rating of 3 is a glaring exception at 1943°K, only 70°K under 4-nitroaniline, whose NFPA rating is 1

4. The upper third of the chart (above 2000°K) contains materials that are almost entirely normally stable with an NFPA rating of 0. The two exceptions are 1,3-butadiene and styrene with NFPA ratings of 2 (normally unstable and can undergo violent reaction, but do not detonate)

Thus, there is a more or less gradual progression from the most hazardous materials in the lower right corner to the “safe”, or normally stable materials, in the upper left corner. Of the 50 compounds studied, 44 were rated, and 6 were unrated. The latter, (acetamide, benzamide, hydroquinone, cellubiose, 1-(p-nitrophenoxy) ethanol, and phenylacetylene) seem to be fairly placed in the chart. The four exceptions mentioned, (hydrogen cyanide, nitroethane, 1,3-butadiene, and styrene) demonstrate that individual peculiarities will cause certain materials not to follow the general pattern exactly. A line was drawn by eye from $T_d = 3000°K$ on the baseline to $T_d = 500°K$ at $(T_0 - T_d) = 2600°K$ fits the points fairly well. This line was divided into five equal segments, then four lines were drawn from the origin through boundaries of each segment and each segmental area was given an “extent of potential hazard rating” similar to the NFPA Chemical Reactivity Rating”

The results of the thermodynamic calcns and the compounds for which they were made, together with the NFPA ratings are shown in the following Table (taken from Ref 1):
<table>
<thead>
<tr>
<th>No.</th>
<th>Compound</th>
<th>Formula</th>
<th>T&lt;sub&gt;d&lt;/sub&gt; (°C)</th>
<th>T&lt;sub&gt;n&lt;/sub&gt; (°C)</th>
<th>T&lt;sub&gt;g&lt;/sub&gt; (°C)</th>
<th>Heat in kcal/100 lbs</th>
<th>Pressure in Atmospheres</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Nitroglycerin</td>
<td>C₃N₅O₇</td>
<td>174.9</td>
<td>221.1</td>
<td>280.0</td>
<td>64.9</td>
<td>25.2</td>
</tr>
<tr>
<td>2</td>
<td>Nitrocellulose 14.24</td>
<td>C₅H₁₁N₂O₃</td>
<td>125.5</td>
<td>132.6</td>
<td>220.0</td>
<td>58.4</td>
<td>35.4</td>
</tr>
<tr>
<td>3</td>
<td>Nitrocellulose 14.7</td>
<td>C₅H₁₂N₂O₃</td>
<td>125.5</td>
<td>132.6</td>
<td>220.0</td>
<td>58.4</td>
<td>35.4</td>
</tr>
<tr>
<td>4</td>
<td>Nitrocellulose 14.1</td>
<td>C₅H₁₃N₂O₃</td>
<td>125.5</td>
<td>132.6</td>
<td>220.0</td>
<td>58.4</td>
<td>35.4</td>
</tr>
<tr>
<td>5</td>
<td>Nitrocellulose 14.0</td>
<td>C₅H₁₄N₂O₃</td>
<td>125.5</td>
<td>132.6</td>
<td>220.0</td>
<td>58.4</td>
<td>35.4</td>
</tr>
<tr>
<td>6</td>
<td>Nitromethane (9)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>7</td>
<td>Nitrous acid (20)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>8</td>
<td>Nitric acid (20)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>9</td>
<td>Nitric oxide (30)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>10</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>11</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>12</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>13</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>14</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>15</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>16</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>17</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>18</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>19</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>20</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>21</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>22</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>23</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>24</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>25</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>26</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>27</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>28</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>29</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
<tr>
<td>30</td>
<td>Nitrile (33)</td>
<td>C₂H₅NO₂</td>
<td>173.1</td>
<td>218.0</td>
<td>278.0</td>
<td>63.4</td>
<td>24.7</td>
</tr>
</tbody>
</table>

(Informal Fire Protection Association classification indicated by number in a column, thus O.)

The parenthetic number is the ord. number required to activate 3 in the U.S. Army. When 2 or 3 for mg. low weight test. These underlined are converted unit notars. envelope.] (2 is a known explosive.)

- T<sub>d</sub> = flame temp, T<sub>n</sub> = decomposition temp
- **T<sub>n</sub>** = heat of combustion; **ΔH<sub>n</sub>** = heat of explosion
- ***P<sub>n</sub>*** = combustion pressure; **P<sub>d</sub>** = explosive pressure; both at const vol
The foregoing dealt with the prediction of potential hazards arising from exothermic chemical reactions. To predict the real hazard of a reaction one must also consider how fast the reaction takes place. Thus it is necessary to link thermodynamics with chemical kinetics because a potentially dangerous reaction may be so slow under a given set of conditions that it constitutes no real danger. The classic example of this situation is the \( \text{H}_2 + \text{O}_2 = \text{H}_2\text{O} \) reaction which is highly exothermic, and consequently potentially dangerous, but exceedingly slow at ambient conditions in the absence of external stimuli (sparks, flame, etc).

Dr. Stull presents the problem very well and we quote (Ref 2):

“In the processing and handling of a chemical or a combination of chemicals, there are two main questions to be answered. First, will there be a reaction? Equilibrium chemical thermodynamics can provide an unequivocal answer to this question if the thermodynamic data for the system is at hand. The previous paper (Ref 1) dealt with this aspect of the problem and was able to identify those systems where energy releases were possible, and related the degree of “potential hazard” to the magnitude of the energy release. However, we do not live in a static, equilibrium world of potential hazards. It is necessary to couple the potential hazard evaluation with the rate of energy release by the reaction, to evaluate the real extent of hazard. The chemical transformation and its associated energy release will range from well behaved to violent depending upon the rate of the reaction.

The second main question that must be considered is: what is the time rate of energy release? Once activated, potentially hazardous materials undergo a nonequilibrium chemical reaction forming the most stable products under the prevailing circumstances. The potentially hazardous systems are those capable of generating heat. They may require different levels of activation, but the smaller the energy of activation, the more readily activated. Some materials (those capable of polymerization for example) may be thermally activated by the ambient temperature of the system. Catalysis also plays a vital role in promoting low level activations. Regardless of the mode of activation, if the heat generated by the reaction can be continuously transferred to the surroundings without creating an increase in temperature, the reaction will proceed quietly in a well behaved manner.

If the heat from the reaction is not all continuously transferred to the surroundings, the temperature of the reaction will increase, slowly at first, but will finally reach a temperature where the reaction is catastrophic. Such thermal run-away reactions are referred to as “thermal explosions”. These thermal explosion reaction types convert “potential” hazardous systems into “real” hazardous systems. Thus, it is necessary to answer both of these two main questions to evaluate the real hazard of a system.”

The actual linking of chemical (Arrhenius) kinetics and thermodynamics was done as follows (Ref 2):

“A square with \( T_d \) increasing upward from 0 to 3000°K was plotted on the left side, while \( E_a \) was plotted on the right side with values from 0 to 100 kcal/mole increasing in the opposite direction (downward). The diagonal line connecting the zeros of the two scales was named the Reaction Hazard Index, or RHI for short. The end of the RHI line at 0°K would unquestionably be at minimum hazard (or zero index), while the end of the RHI line at 0 kcal/mole (zero activation energy) must surely

\[
\text{Reaction Hazard Index} \quad \text{RHI} = \frac{10 T_d}{2 + 30 E_a}
\]

Fig 2 Nomograph Linking the Decomposition Temperature \( \text{T}_d \) °K with the Arrhenius Activation Energy \( E_a \) in kcal/mole (Ref 2)
correspond to maximum hazard (assigned an index value of 10). Thus, RHI values from 0 to 10 cover the whole range from minimum to maximum hazard. The intercept on the RHI line is given by the simple relationship RHI = 10 T_d / T_d + 30 E_a). A single value of the Reaction Hazard Index between 0 and 10 provides a numerical index for a given material.

Figure 3 shows an example of use comparing the reaction hazard of three hydrocarbon gases, methane, ethylene and acetylene, with the values of T_d, E_a, and RHI given in Table 2. The RHI figures correctly place methane as least reactive, acetylene as most reactive, and ethylene roughly midway between the two."

![Fig 3 Example of Use of Reaction Hazard Index with Methane, Ethylene, and Acetylene (Ref 2)](image)

TABLE 2
Comparison of NFPA Reactivity Rating with the Reaction Hazard Index (Ref 2)

<table>
<thead>
<tr>
<th></th>
<th>T_d °K</th>
<th>E_a kcal/mole</th>
<th>NFPA Reactivity Rating</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH₄ Methane</td>
<td>298</td>
<td>103</td>
<td>3.9</td>
</tr>
<tr>
<td>C₂H₄ Ethylene</td>
<td>1005</td>
<td>46.5</td>
<td>4.2</td>
</tr>
<tr>
<td>C₂H₂ Acetylene</td>
<td>2898</td>
<td>40.5</td>
<td>7.1</td>
</tr>
</tbody>
</table>

Again we quote Dr Stull:

"An extended effort was made to find data for as many chemical compounds as possible where the NFPA Reactivity Rating could be compared directly with the Reaction Hazard Index. Table 3 lists 80 compounds where the direct comparison could be made, and includes all of the basic data necessary to calculate the Reaction Hazard Index. The 80 compounds are grouped into the five NFPA Reactivity Ratings as follows: 38 compounds with 0 rating (the largest and best established group), 13 compounds in rating 1 and in rating 2, 6 compounds in rating 3, and 10 compounds in rating 4. A cross plot of the NFPA Reactivity Rating versus the Reaction Hazard Index is presented in Figure 4. The average RHI for each NFPA rating is given in Table 3, and is plotted as a + in Figure 4. The best dotted straight line through the average indices shows a remarkable accord between these two methods of reactivity hazard rating.

The NFPA Reactivity Ratings represent the consensus of a committee of experts, and because of the number of factors taken into consideration the values agreed upon might in some instances border on the next higher (or lower) value, and thus be indeterminate by one unit. Hence, acetylene (No 72) rightfully carries the maximum hazardous NFPA Reactivity Rating of 4, and based on the evidence in Table 3, it is not unreasonable that vinylcyclohexane (No 68) should also.

The Reaction Hazard Index method presented here is based on experimentally observed quantities representing the Arrhenius energy of activation for the decomposition process and the energy generated by the decomposition as indicated by the adiabatic decomposition temperature reached by the decomposition products. The method used develops a single numerical value that is characteristic of the reaction hazard for the compound measured and represents the real hazard. Minimum definite measurements can be prescribed for compounds where handling experience is lacking. This objective method can be used to evaluate the real degree of hazard for a single compound or for a mixture of chemicals, and can be used to develop a Reaction Hazard Index system that will effectively rate all chemical reaction hazards with respect to each other."
<table>
<thead>
<tr>
<th>No.</th>
<th>Formula</th>
<th>Gaseous Compound</th>
<th>$\Delta H^0_{f,298}$ kcal/mole Ref.(24)</th>
<th>$T_d^{0,K}$</th>
<th>Activation Energy for Decomposition kcal/mole Ref.**</th>
<th>Reaction Hazard Index RHI</th>
<th>NFPA Reactivity Rating</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>CHCl₃</td>
<td>Chloroform</td>
<td>24.2</td>
<td>683</td>
<td>47.0</td>
<td>(1)</td>
<td>3.26</td>
</tr>
<tr>
<td>2</td>
<td>CH₂O₂</td>
<td>Formic Acid</td>
<td>90.49</td>
<td>800</td>
<td>30.6</td>
<td>(1)</td>
<td>4.66</td>
</tr>
<tr>
<td>3</td>
<td>CH₃Cl</td>
<td>Methyl Chloride</td>
<td>20.63</td>
<td>744</td>
<td>74.0</td>
<td>(4)</td>
<td>2.55</td>
</tr>
<tr>
<td>4</td>
<td>CH₄</td>
<td>Methane</td>
<td>17.89</td>
<td>298</td>
<td>103.0</td>
<td>(1)</td>
<td>0.88</td>
</tr>
<tr>
<td>5</td>
<td>CH₃N</td>
<td>Methylamine</td>
<td>5.50</td>
<td>767</td>
<td>58.0</td>
<td>(21)</td>
<td>3.06</td>
</tr>
<tr>
<td>6</td>
<td>C₂H₄Cl₂</td>
<td>1,1-Dichloroethane</td>
<td>31.05</td>
<td>847</td>
<td>53.5</td>
<td>(10)</td>
<td>3.45</td>
</tr>
<tr>
<td>7</td>
<td>C₂H₄Cl₂</td>
<td>1,2-Dichloroethane</td>
<td>31.00</td>
<td>849</td>
<td>47.0</td>
<td>(21)</td>
<td>3.76</td>
</tr>
<tr>
<td>8</td>
<td>C₂H₅Br</td>
<td>Ethylbromide</td>
<td>15.30</td>
<td>670</td>
<td>53.2</td>
<td>(21)</td>
<td>2.96</td>
</tr>
<tr>
<td>9</td>
<td>C₂H₅Cl</td>
<td>Ethylchloride</td>
<td>26.70</td>
<td>701</td>
<td>56.5</td>
<td>(1)</td>
<td>2.93</td>
</tr>
<tr>
<td>10</td>
<td>C₂H₆</td>
<td>Ethane</td>
<td>20.24</td>
<td>597</td>
<td>89.5</td>
<td>(11)</td>
<td>1.82</td>
</tr>
<tr>
<td>11</td>
<td>C₂H₇N</td>
<td>Dimethylamine</td>
<td>4.50</td>
<td>792</td>
<td>43.4</td>
<td>(21)</td>
<td>3.78</td>
</tr>
<tr>
<td>12</td>
<td>C₂H₇N</td>
<td>Ethylamine</td>
<td>11.00</td>
<td>740</td>
<td>43.4</td>
<td>(21)</td>
<td>3.62</td>
</tr>
<tr>
<td>13</td>
<td>C₃H₆</td>
<td>Cyclopropane</td>
<td>12.74</td>
<td>936</td>
<td>65.5</td>
<td>(11)</td>
<td>3.22</td>
</tr>
<tr>
<td>14</td>
<td>C₃H₆O</td>
<td>Acetone</td>
<td>51.99</td>
<td>774</td>
<td>68.0</td>
<td>(21)</td>
<td>2.75</td>
</tr>
<tr>
<td>15</td>
<td>C₃H₇Cl</td>
<td>1-Chloropropane</td>
<td>31.10</td>
<td>699</td>
<td>55.0</td>
<td>(21)</td>
<td>2.98</td>
</tr>
<tr>
<td>16</td>
<td>C₃H₈</td>
<td>Propane</td>
<td>24.82</td>
<td>626</td>
<td>63.3</td>
<td>(21)</td>
<td>2.48</td>
</tr>
<tr>
<td>17</td>
<td>C₄H₈</td>
<td>1-Butene</td>
<td>0.03</td>
<td>825</td>
<td>59.1</td>
<td>(1)</td>
<td>3.18</td>
</tr>
<tr>
<td>18</td>
<td>C₄H₈</td>
<td>Cyclobutane</td>
<td>6.37</td>
<td>865</td>
<td>62.5</td>
<td>(7)</td>
<td>3.16</td>
</tr>
<tr>
<td>19</td>
<td>C₄H₈O</td>
<td>2-Butanone</td>
<td>56.97</td>
<td>755</td>
<td>67.2</td>
<td>(21)</td>
<td>2.72</td>
</tr>
<tr>
<td>20</td>
<td>C₄H₈O₂</td>
<td>Ethylacetate</td>
<td>105.86</td>
<td>735</td>
<td>48.0</td>
<td>(11)</td>
<td>3.38</td>
</tr>
<tr>
<td>21</td>
<td>C₄H₉Br</td>
<td>1-Bromobutane</td>
<td>25.65</td>
<td>668</td>
<td>50.9</td>
<td>(21)</td>
<td>3.04</td>
</tr>
<tr>
<td>22</td>
<td>C₄H₉Cl</td>
<td>1-Chlorobutane</td>
<td>35.20</td>
<td>701</td>
<td>57.0</td>
<td>(21)</td>
<td>2.91</td>
</tr>
<tr>
<td>23</td>
<td>C₄H₁₀</td>
<td>Butane</td>
<td>30.15</td>
<td>633</td>
<td>86.3</td>
<td>(11)</td>
<td>1.96</td>
</tr>
<tr>
<td>24</td>
<td>C₄H₁₀O</td>
<td>2-Methylpropane</td>
<td>32.15</td>
<td>611</td>
<td>53.5</td>
<td>(21)</td>
<td>2.76</td>
</tr>
<tr>
<td>25</td>
<td>C₄H₁₀O</td>
<td>tert-Butyl Alcohol</td>
<td>77.87</td>
<td>628</td>
<td>61.6</td>
<td>(1)</td>
<td>2.54</td>
</tr>
<tr>
<td>26</td>
<td>C₄H₁₀O</td>
<td>Ethyl Ether</td>
<td>61.88</td>
<td>761</td>
<td>78.0</td>
<td>(1)</td>
<td>2.46</td>
</tr>
<tr>
<td>27</td>
<td>C₅H₁₀O₂</td>
<td>Iso-propylacetate</td>
<td>115.40</td>
<td>696</td>
<td>45.0</td>
<td>(1)</td>
<td>3.40</td>
</tr>
<tr>
<td>28</td>
<td>C₅H₁₂</td>
<td>2, 2-Dimethylpropane</td>
<td>39.67</td>
<td>597</td>
<td>80.4</td>
<td>(1)</td>
<td>1.98</td>
</tr>
<tr>
<td>29</td>
<td>C₅H₁₂</td>
<td>2-Methylbutane</td>
<td>36.92</td>
<td>626</td>
<td>58.6</td>
<td>(21)</td>
<td>2.62</td>
</tr>
<tr>
<td>30</td>
<td>C₅H₁₂</td>
<td>Pentane</td>
<td>35.00</td>
<td>645</td>
<td>61.2</td>
<td>(21)</td>
<td>2.60</td>
</tr>
<tr>
<td>31</td>
<td>C₅H₁₂O</td>
<td>2, 2-Dimethylpropanol</td>
<td>70.00</td>
<td>725</td>
<td>60.0</td>
<td>(1)</td>
<td>2.87</td>
</tr>
<tr>
<td>32</td>
<td>C₅H₁₂</td>
<td>Cyclohexane</td>
<td>29.43</td>
<td>677</td>
<td>64.1</td>
<td>(19)</td>
<td>2.60</td>
</tr>
<tr>
<td>33</td>
<td>C₆H₁₂O₂</td>
<td>Butyric acid</td>
<td>116.7</td>
<td>715</td>
<td>46.0</td>
<td>(1)</td>
<td>3.41</td>
</tr>
<tr>
<td>34</td>
<td>C₇H₈</td>
<td>Toluene</td>
<td>11.95</td>
<td>859</td>
<td>85.0</td>
<td>(1)</td>
<td>2.52</td>
</tr>
<tr>
<td>35</td>
<td>C₇H₄Cl</td>
<td>Methylcyclohexane</td>
<td>36.99</td>
<td>660</td>
<td>57.9</td>
<td>(19)</td>
<td>2.75</td>
</tr>
<tr>
<td>36</td>
<td>C₇H₇O₂</td>
<td>tert-Amylacetate</td>
<td>122.7</td>
<td>705</td>
<td>40.3</td>
<td>(1)</td>
<td>3.68</td>
</tr>
<tr>
<td>37</td>
<td>C₈H₁₀</td>
<td>Ethylbenzene</td>
<td>7.12</td>
<td>830</td>
<td>73.0</td>
<td>(1)</td>
<td>2.75</td>
</tr>
<tr>
<td>38</td>
<td>C₈H₁₀</td>
<td>4-Xylene</td>
<td>4.29</td>
<td>817</td>
<td>79.5</td>
<td>(1)</td>
<td>2.55</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Average of 38 compounds: 2.89

<p>| 39  | C₉H₈N₂ | Methylhydrazine     | 20.40                                   | 1022        | 51.9                                                | (1)                      | 3.96                   | 0                      |
| 40  | C₉H₄O₂ | Acetic Acid         | 104.93                                  | 634         | 67.5                                                | (1)                      | 2.38                   | 0                      |
| 41  | C₉H₈N₂ | 1, 1-Dimethylethydrazine | 20.30                           | 953         | 49.6                                                | (1)                      | 3.90                   | 0                      |
| 42  | C₃H₅Br | Allylbromide        | 11.80                                   | 988         | 45.5                                                | (21)                     | 4.20                   | 0                      |</p>
<table>
<thead>
<tr>
<th>No.</th>
<th>Formula</th>
<th>Gaseous Compound</th>
<th>ΔH°298 kcal/mole</th>
<th>Tg⁰K</th>
<th>Activation Energy for Decomposition kcal/mole</th>
<th>Reaction Hazard Index RHI</th>
<th>NFPA Reactivity Rating</th>
</tr>
</thead>
<tbody>
<tr>
<td>43</td>
<td>C₃H₅N</td>
<td>Propionitrile</td>
<td>+ 12.10</td>
<td>903</td>
<td>72.7</td>
<td>2.93</td>
<td>1</td>
</tr>
<tr>
<td>44</td>
<td>C₃H₆</td>
<td>Propylene</td>
<td>+ 4.88</td>
<td>866</td>
<td>78.0</td>
<td>2.70</td>
<td>1</td>
</tr>
<tr>
<td>45</td>
<td>C₃H₆O</td>
<td>Propionaldehyde</td>
<td>- 45.90</td>
<td>819</td>
<td>50.3</td>
<td>3.52</td>
<td>1</td>
</tr>
<tr>
<td>46</td>
<td>C₄H₆O₃</td>
<td>Acetic Anhydride</td>
<td>- 137.60</td>
<td>793</td>
<td>34.5</td>
<td>4.34</td>
<td>1</td>
</tr>
<tr>
<td>47</td>
<td>C₅H₁₀O₃</td>
<td>Diethyl Carbonate</td>
<td>- 151.60</td>
<td>753</td>
<td>46.0</td>
<td>3.53</td>
<td>1</td>
</tr>
<tr>
<td>48</td>
<td>C₆H₁₄O</td>
<td>Iso-propyl Ether</td>
<td>- 76.20</td>
<td>712</td>
<td>63.5</td>
<td>2.72</td>
<td>1</td>
</tr>
<tr>
<td>49</td>
<td>C₇H₁₄Cl</td>
<td>Benzyl Chloride</td>
<td>- 7.30</td>
<td>831</td>
<td>68.0</td>
<td>2.89</td>
<td>1</td>
</tr>
<tr>
<td>50</td>
<td>C₁₀H₁₂</td>
<td>Dicyclopentadiene-endo</td>
<td>+ 46.60</td>
<td>990</td>
<td>34.0</td>
<td>4.93</td>
<td>1</td>
</tr>
<tr>
<td>51</td>
<td>C₁₀H₁₂</td>
<td>Dicyclopentadiene-exo</td>
<td>+ 46.60</td>
<td>990</td>
<td>38.5</td>
<td>4.61</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Average of 13 compounds</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>3.58</td>
</tr>
<tr>
<td>52</td>
<td>C₂H₄</td>
<td>Ethylene</td>
<td>+ 12.50</td>
<td>1005</td>
<td>46.5</td>
<td>4.19</td>
<td>2</td>
</tr>
<tr>
<td>53</td>
<td>C₂H₄</td>
<td>Ethylene</td>
<td>+ 12.50</td>
<td>1005</td>
<td>37.7</td>
<td>4.71</td>
<td>2</td>
</tr>
<tr>
<td>54</td>
<td>C₂H₄O</td>
<td>Acetaldehyde</td>
<td>- 39.76</td>
<td>866</td>
<td>48.0</td>
<td>3.76</td>
<td>2</td>
</tr>
<tr>
<td>55</td>
<td>C₂H₆O</td>
<td>Propylene Oxide</td>
<td>- 22.17</td>
<td>948</td>
<td>58.0</td>
<td>3.53</td>
<td>2</td>
</tr>
<tr>
<td>56</td>
<td>C₂H₆O₂</td>
<td>Dioxolane</td>
<td>- 70.0</td>
<td>911</td>
<td>53.6</td>
<td>3.64</td>
<td>2</td>
</tr>
<tr>
<td>57</td>
<td>C₄H₆</td>
<td>1,3-Butadiene</td>
<td>+ 20.40</td>
<td>991</td>
<td>79.4</td>
<td>2.94</td>
<td>2</td>
</tr>
<tr>
<td>58</td>
<td>C₄H₆</td>
<td>1,3-Butadiene</td>
<td>+ 20.40</td>
<td>991</td>
<td>24.7</td>
<td>5.72</td>
<td>2</td>
</tr>
<tr>
<td>59</td>
<td>C₄H₈O</td>
<td>Glycol Toluol</td>
<td>- 37.40</td>
<td>863</td>
<td>41.0</td>
<td>4.12</td>
<td>2</td>
</tr>
<tr>
<td>60</td>
<td>C₄H₈O</td>
<td>Vinyl Ethyl Ether</td>
<td>- 33.50</td>
<td>880</td>
<td>44.4</td>
<td>3.98</td>
<td>2</td>
</tr>
<tr>
<td>61</td>
<td>C₅H₈O</td>
<td>Vinyl Allyl Ether</td>
<td>- 12.8</td>
<td>959</td>
<td>30.6</td>
<td>5.11</td>
<td>2</td>
</tr>
<tr>
<td>62</td>
<td>C₆H₈</td>
<td>Styrene</td>
<td>+ 35.22</td>
<td>993</td>
<td>19.2*</td>
<td>6.33</td>
<td>2</td>
</tr>
</tbody>
</table>

*For the polymerization process.

| 63  | C₈H₁₂   | Vinyl Cyclohexane                       | + 16.8          | 876  | 62.2                                          | 3.18                      | 2                      |
| 64  | N₂H₄    | Hydrazine                               | + 22.75         | 1338 | 60.5                                          | 4.24                      | 2                      |
|     |         | Average of 13 compounds                 |                 |      |                                               |                          | 4.27                   | 2                      |
| 65  | C₂H₆O   | Ethylene Oxide                          | - 12.58         | 1062 | 57.4                                          | 3.81                      | 3                      |
| 66  | C₂H₅NO₂ | Nitroethane                             | - 24.20         | 1161 | 45.0                                          | 4.62                      | 3                      |
| 67  | C₂H₅NO   | 1-Nitropropane                          | - 29.80         | 1046 | 47.7                                          | 4.22                      | 3                      |
| 68  | C₂H₄     | Vinylacetylene                          | + 72.80         | 2317 | 28.3                                          | 3.73                      | 3                      |
| 69  | C₅H₁₀O₂  | tert-Butyl Peroxide                     | - 81.50         | 850  | 37.0                                          | 4.31                      | 3                      |
| 70  | C₁₂H₁₆N₄O₁₈ | Cellulose Nitrate               | - 229.80        | 2213 | 46.7                                          | 6.12                      | 3                      |
|     |         | Average of 6 compounds                  |                 |      |                                               |                          | 5.07                   | 3                      |
| 71  | CH₃NO₂  | Nitromethane                            | - 17.86         | 2621 | 59.0                                          | 5.97                      | 4                      |
| 72  | C₂H₂    | Acetylene                               | + 54.19         | 2898 | 40.5*                                         | 7.05                      | 4                      |
| 73  | C₂H₄O₃  | Peracetic Acid                          | - 97.73         | 976  | 32.0                                          | 5.04                      | 4                      |
| 74  | C₂H₅NO₃ | Ethyl Nitrate                           | - 36.8          | 2094 | 39.9                                          | 6.36                      | 4                      |
| 75  | C₃H₅N₃O₉ | Nitroglycerine                         | - 73.20         | 2895 | 40.3                                          | 7.05                      | 4                      |
| 76  | C₄H₆O₄  | Acetyl Peroxide                         | - 116.1         | 983  | 29.5                                          | 5.26                      | 4                      |
| 77  | C₄H₉O₂  | tert-Butyl Hydroperoxide                | - 62.9          | 919  | 37.8                                          | 4.48                      | 4                      |
| 78  | C₅H₉O₂  | Diethyl Peroxide                        | - 46.1          | 968  | 37.3                                          | 4.64                      | 4                      |
| 79  | C₆H₁₈O₂  | Di-tert-butyl Peroxide                  | - 81.5          | 850  | 37.5                                          | 4.30                      | 4                      |
| 80  | C₉H₁₂O₂  | Cumene Hydroperoxide                    | - 21.9          | 989  | 29.0                                          | 5.32                      | 4                      |

*For the polymerization process.

**See Ref. 2.
Fig 4 Correlation of the NFPA Reactivity Rating with the Reaction Hazard Index (Ref 2)

Refs: 1) D.R. Stull, Chem Eng Prog, Loss Prevention, 4, 16 (1970)
2) D.R. Stull, Ibid, 7, 67 (1973)

Hazards—Prevention of Industrial Gas Explosions.
In the prevention of industrial gas explosion disasters, the most useful terms for evaluating the hazards of various fuel/oxidant systems are believed to be limits of flammability and min spontaneous ignition temp (Ref 3).

The addition of a flammable diluent is suggested as a method of preventing explosions in processes involving flammable gas mixts; e.g., the addition of sufficient methane to a potentially explosive mixt of H + O in a hydrogen peroxide plant suppresses explosive reactions without excessive loss of plant throughput capacity which would have resulted if an inert suppressant (diluent) such as nitrogen were added to the mixture. Butane or pentane are even more effective than methane (Ref 1).

Spraying with 10-20 psi steam is suggested before introducing an air stream (carrying wood preservatives) into an organic vapor atm (Ref 2)

Ref: 1) E. Jones, Chem Eng 59 (6), 185 (1952) & CA 46, 8374 (1952) 2) M. Hudson, USP 2633429 (1953) & CA 48, 5592 (1954)

HBX (High Blast Explosive). High explosive compositions containing RDX, TNT, Al and a wax desensitizer. HBX-1 consists of RDX 40, TNT 38, Al 17, D-2 wax 5% & 0.5% added CaCl₂. It was developed during WWII as a desensitized modification of Torpex (Ref 1). A further modification, HBX-3, consists of RDX 31, TNT 29, Al 35, D-2 wax 5% & -5% added CaCl₂. HBX-1 and HBX-3 are prepared by adding water-wet RDX to molten TNT and heating until the water is evaporated. The D-2 wax (84% paraffin, 14% NC, 2% lecithin) and CaCl₂ are then added. Both HBX-1 and HBX-3 are powerful and brisant explosives whose physical and explosive properties are tabulated here after description of prep

HBX expl comps are prep by melting TNT in a steam jacketed kettle equipped with a mechanical stirrer. Water-wet RDX is added slowly with stirring and heating until all the water is evapd. Powdered Al is added and the mixture is continued to be stirred until uniform. D-2 wax and Ca chloride are then added and the mixture is cooled from temp 95-100° to a temp considered suitable for casting (the lowest practicable pour temp)

HBX can also be prep by adding the calcd amt of molten TNT to molten Comp B to obtain the desired proportion of RDX/TNT and then the appropriate weights of other ingredients are added to complete the compn (Ref 6, p 163)

NOTE: The desensitizer wax, also known as Composition D-2, consists of 84% paraffin wax, 14% NC and 2% lecithin. Its Military Specification Requirements and Tests are in MIL-C-1864
### TABLE
Properties of HBX's

<table>
<thead>
<tr>
<th>Properties</th>
<th>HBX-1</th>
<th>HBX-3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mol Wt's</td>
<td>102</td>
<td>64</td>
</tr>
<tr>
<td>OB to CO₂, %</td>
<td>−68</td>
<td>−75</td>
</tr>
<tr>
<td>OB to CO, %</td>
<td>−35</td>
<td>−49</td>
</tr>
<tr>
<td><strong>Ballistic Mortar:</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>% TNT in Air</td>
<td>121 (Shock)</td>
<td>116 (Shock)</td>
</tr>
<tr>
<td>% TNT in Air</td>
<td>121 (Impulse)</td>
<td>125 (Impulse)</td>
</tr>
<tr>
<td><strong>Blast Effects:</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>% TNT under water</td>
<td>111 (Shock)</td>
<td>101 (Shock)</td>
</tr>
<tr>
<td>% TNT under water</td>
<td>145 (Bubble)</td>
<td>191 (Bubble)</td>
</tr>
<tr>
<td>Brisance by Sand Test, % TNT</td>
<td>102 (123)</td>
<td>93.5</td>
</tr>
<tr>
<td>Density, g/cc</td>
<td>1.69−1.70</td>
<td>1.81−1.84</td>
</tr>
<tr>
<td>Detonation Rate, m/sec</td>
<td>7224</td>
<td>6927</td>
</tr>
<tr>
<td>Expln Temp (5 sec)</td>
<td>480°</td>
<td>500°</td>
</tr>
<tr>
<td>Fragmentation Test in 90mm, HE, M71 Shell</td>
<td></td>
<td></td>
</tr>
<tr>
<td>No of Fragments vs TNT</td>
<td>910</td>
<td>703</td>
</tr>
<tr>
<td>Friction Sensitivity</td>
<td>Unaffected</td>
<td>Not given</td>
</tr>
<tr>
<td>Gas Volume</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Heat of Combustn, cal/g</td>
<td>3882</td>
<td>4495</td>
</tr>
<tr>
<td>Heat of Expln, cal/g</td>
<td>919</td>
<td>877</td>
</tr>
<tr>
<td>Heat of Formation, cal/g</td>
<td>758</td>
<td>491</td>
</tr>
<tr>
<td>Heat Test at 100°</td>
<td></td>
<td></td>
</tr>
<tr>
<td>% Loss in 1st 48 hrs</td>
<td>0.058</td>
<td>0.70</td>
</tr>
<tr>
<td>% Loss in 2nd 48 hrs</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Expln in 100 hrs</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Hygroscopicity, % Loss (in 7 days at 30° and 95% RH)</td>
<td>2.98</td>
<td>2.01</td>
</tr>
</tbody>
</table>

**Impact Sensitivity**
- comparable to TNT
- more sensitive than TNT
- Require slightly lower minimum priming charge

**Rifle Bullet Test**
- Principally in HE charges, such as in GP bombs

**Sensitivity to Initiation**
- Dry
- Principally in HE charges, such as in GP bombs

**Storage Usage**
- Dry
- Principally in HE charges, such as in GP bombs

---

*Note: HBX-1 was recommended for use in booster charges and in depth charges (Refs 4, 5 & 6)*

---

HBX-1, HBX-3 and H-6 Explosive Compositions
US Military Specification Requirements and Tests, as described in MIL-E-22267A (31 May 1963)

Requirements:

3.2 Form. Unless otherwise specified the compns HBX-1, HBX-2 & H-6 shall be supplied in the form of buds or as strips 1.5 inches wide, 1 inch deep & 3 inches long

3.4 Components. The components used in the preparation of the Grade A HBX and H-6 composition shall comply with the specifications listed in 2.1 and shall be of the following grade or class as applicable:

Composition B  Grade A
TNT          Grade 1
Al Powder    Type III, Grade F, Class 7

3.5 Grade A Compositions. The composition of the Grade A HBX composition shall conform to the nearest tenth percent as required in Table I when tested as specified in 4.4.3.2 or 4.4.3.3

Table 1. Grade A Compositions

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>HBX-1 Percent by Weight</th>
<th>HBX-3 Percent by Weight</th>
<th>H-6 Percent by Weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>RDX plus Nitrocellulose plus **Calcium Chloride plus ***Calcium Silicate</td>
<td>40.4 ± 3.0%</td>
<td>31.3 ± 3.0%</td>
<td>45.1 ± 3.0%</td>
</tr>
<tr>
<td>TNT</td>
<td>37.8 ± 3.0%</td>
<td>29.0 ± 3.0%</td>
<td>29.2 ± 3.0%</td>
</tr>
<tr>
<td>Aluminum</td>
<td>17.1 ± 3.0%</td>
<td>34.8 ± 3.0%</td>
<td>21.0 ± 3.0%</td>
</tr>
<tr>
<td>**** Wax plus lecithin</td>
<td>4.7 ± 1.0%</td>
<td>4.9 ± 1.0%</td>
<td>4.7 ± 1.0%</td>
</tr>
</tbody>
</table>

* Note. All of the RDX component and portions of the TNT and wax are added as Composition B.
** A separate calcium chloride determination need only be done if required in the contract or order (See 6.2). Percentage requirement of Calcium Chloride is 0.5% ± 0.1%
*** Calcium Silicate shall be determined as specified in 4.4.1.2 when used in the formulation of HBX compounds. The Calcium Silicate content shall be a minimum of 1.25 weight percent of the TNT content of the mixture and shall only be used in the HBX compositions when specified by the procuring agency (See 6.2). Calcium Silicate is designated for use only in Army formulations of HBX type explosive compositions.
**** The major portion of the wax and all of the Nitrocellulose and lecithin are added as Composition D-2

3.6 Grade B Compositions. The composition of the Grade B HBX & H-6 Composition shall conform to the nearest tenth percent as required in Table II when determined as specified in 4.4.3.4

Table II. Grade B Compositions

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>HBX-1 Percent by Weight</th>
<th>HBX-3 Percent by Weight</th>
<th>H-6 Percent by Weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>RDX plus Nitrocellulose plus Calcium Chloride plus *Calcium Silicate</td>
<td>40.4 ± 3.0%</td>
<td>31.3 ± 3.0%</td>
<td>45.1 ± 3.0%</td>
</tr>
<tr>
<td>TNT</td>
<td>37.8 ± 3.0%</td>
<td>29.0 ± 3.0%</td>
<td>29.2 ± 3.0%</td>
</tr>
<tr>
<td>Aluminum</td>
<td>17.1 ± 3.0%</td>
<td>34.8 ± 3.0%</td>
<td>21.0 ± 3.0%</td>
</tr>
<tr>
<td>Wax plus lecithin</td>
<td>4.7 ± 1.0%</td>
<td>4.9 ± 1.0%</td>
<td>4.7 ± 1.0%</td>
</tr>
</tbody>
</table>

Hot melt (asphaltic lining material) — maximum allowable 0.75%
A separate Calcium Chloride detn need only be done if required in the contract or order. An additional one-half percent of Calcium Chloride shall be added whenever preparing the reclaimed explosive for re-use. Percentage requirement is 0.7% ± 0.3%
* The Calcium Silicate content shall be a minimum of 1.25 weight percent of the TNT content of the mixture and shall only be used in the HBX compositions when specified by the procuring agency (See 6.2). Calcium Silicate shall be determined as specified in 4.4.1.2
3.7 Moisture Content, when tested as specified in
4.4.4 shall be 0.20% max for Grade A and 0.50% for Grade B
3.8 Vacuum Stability. The max volume of gas
liberated when tested as specified in 4.4.5 shall
be 2cc/g/48 hrs at 100° for Grade A and B
4.2 Lot Size. For the purpose of sampling a lot
shall be limited to 1300 lbs max and to one batch
from a single vessel. Tests shall be conducted on
each lot
4.3.1 Preproduction Samples. After award of
contract but prior to entering quantity production,
a preproduction sample shall be prepd for in-
spection and acceptance tests to determine con-
formance of the sample with the requirements of
the specification (See 6.2)
4.3.1.1 Preproduction Sample for Subsequent
Contracts. The necessity for such a sample will be
detd by the procuring activity when production
under a new contract by the same contractor at
the same location follows the prep of any HBX
or H-6 compn covered by this spec
4.3.2.1 Sample for Chemical Composition Analysis
is taken by catching a portion of the molten expl
compn, from the approx center of the batch, and
pouring into shallow Al dish 4 inches in diam in
order to obtn on cooling a wafer approx ¼ inch thick
4.3.2.2 Sample for Moisture Analysis consists of
approx 50g reserved from 4.3.2.1
4.4.1 Process Verification. Unless otherwise speci-
fied, provss verification used as the inspection
shall be subject to Government verification at the
time the first batch is prepared and at random in-
tervals during the production, but not less than
once during each week of continuous operation.
Verification will consist of surveillance of the
process and related equipment to determine that
practices, methods and procedures are being pro-
perly applied, and that the products are produced
under the requirements of this specification. A
record shall be made of each batch of explosive
prepared to ensure that the following requirements
have been met:
(a) Explosive components – Components shall
meet the requirements of 3.4
(b) Composition – The quantities of each compo-
nent required shall be calculated and weighed to
give the correct composition. Component weights
may be calculated from Table III
(c) Order of addition – The order of addition shall
be recorded. Order of addition shall be mandatory
as specified in 3.3
(d) Temperature – The temperature shall be re-
corded, prior to, during, and after each phase of
batching
(e) Agitation – Agitation shall be maintained from
the time of the addition of the first component
until the kettle has been drained
(f) Mixing time – The mixing time following the
addition of the last component

Table III. HBX Compositions
(Percent by Weight)

<table>
<thead>
<tr>
<th></th>
<th>HBX-1</th>
<th>HBX-3</th>
<th>H-6</th>
</tr>
</thead>
<tbody>
<tr>
<td>B</td>
<td>66.08</td>
<td>51.33</td>
<td>74.20</td>
</tr>
<tr>
<td>D-2</td>
<td>4.66</td>
<td>4.73</td>
<td>4.69</td>
</tr>
<tr>
<td>Aluminum</td>
<td>17.10</td>
<td>34.75</td>
<td>20.61</td>
</tr>
<tr>
<td>Additional TNT</td>
<td>11.66</td>
<td>8.69</td>
<td></td>
</tr>
<tr>
<td>Calcium Chloride</td>
<td>0.50</td>
<td>0.50</td>
<td>0.50</td>
</tr>
<tr>
<td>Calcium Silicate</td>
<td>0.47</td>
<td>0.36</td>
<td>0.37</td>
</tr>
</tbody>
</table>

Each component shall be based on the percent
by weight limitation listed in Table I or II for each
HBX formulation

a) When specified by the procuring activity,
Calcium Silicate used in the formulation of HBX
components shall be a minimum of 1.25 weight
percent of the TNT content
4.4.1.2 Determination of Calcium Silicate. When
specified by the procuring activity (See 6.2), Cal-
cium Silicate used in the HBX & H-6 compositions
shall be determined by visual verification of the
weight of calcium silicate added to the batch.
Quantities added shall be calculated in terms of
weight percent of TNT content
4.4.2 Lot Acceptance Tests. Unless otherwise
specified in the contract or order, samples selected
from each inspection lot shall be subjected to all
the requirements and tests of this specification.
Failure of the HBX compositions to meet any of
the requirements or tests of this specification shall
be considered cause for rejection of the lot
4.4.3 Composition Analysis
4.4.3.1 Sample Preparation. Reduce sample thru
a 20 mesh screen (US Standard Sieve Series)
using a wooden mortar and pestle
4.4.3.2 Method of Analysis. Grade A:
4.4.3.2.1 Determination of Aluminum. Weigh
accurately a sample calculated to contain 0.4g of
TNT into a tared sintered glass filtering crucible,
medium porosity, 30-ml capacity. Extract on a
Fisher filtrator with ethylene chloride (purified
1,2 dichloroethane) at approx 20°C, 5 equal
portions totaling 45ml, contact time of 1 min each.
Collect filtrate in a 50ml volumetric flask. Reserve filtrate for TNT determination.

The residue remaining on the filtering crucible is extracted with hot cyclohexanone (highest purity grade) at approx 100°C. Five consecutive 3-second extractions of 3ml each with continuous vacuum filtration and no stirring are made. This is followed by four consecutive 30-sec extractions of 10ml each with stirring. After the cyclohexanone extraction, wash down the sides of the crucible with approx 2ml of reagent grade acetone. Repeat for a total of 3 washings. Dry in an oven at 90°C ±10°C for 30 mins, cool in a desiccator and weigh. The residue is aluminum.

Percent Al = [(A−B) x 100] / W

where: A = Wt of crucible with Al
B = Wt of empty crucible
W = Wt of sample

4.4.3.2.2 TNT Determination. Dilute the TNT filtrate obtained in 4.4.3.2.1 to 50ml with ethylene chloride. Compare its spectral absorbance at 20° with that of a soln, 0.400g of TNT in 50ml ethylene chloride at 367 millimicrons. Det TNT by reference to a graph prepd in advance from known soln. The prepn of the solns and the graph is described in 4.4.3.5.1. A Beckman DK-2 or equivalent spectrophotometer may be used to det the spectral absorbance.

When using the Beckman DK-2 recording spectrophotometer, one proceeds under the following conditions:

a) 10mm cell with 9mm spacer
b) Wavelength setting of 367 millimicrons
c) Time constant of 0.6
d) Scanning time of 1
e) Scale expansion 2X
f) Sensitivity 11.5
g) Adsorption scale of −0.3 ±0.7
h) Hydrogen lamp
i) Photomultiplier position 1X

Percent TNT = [(A x 100)] / W

where: A = Wt of TNT from graph
W = Wt of sample

4.4.3.2.3 Determination of Wax + Lecithin. Weigh a 3-g sample and transfer it to a small, accurately tared, test tube. Add exactly 3ml of tetrachloroethylene and reweigh the tube with contents. Heat it in a steam bath to 90° while stirring vigorously and transfer to a water bath used to maintain the prisms of an Abbé refractometer at 50±0.5°. After cooling, reweigh the tube with contents to det exact amt of tetrachloroethylene lost by volatilization. Replace the exact amt of loss and insert the tube in the water bath. Set the refractometer to read 1.4910 as the RI of tetrachloroethylene at 50±0.5° and pour several drops of the supernatant liquid from the test tube into the prisms of the refractometer. Quickly close the prisms and at the end of 30 secs det the RI to sodium D light. The wax + lecithin concn is detd by reference to a graph previously prepd from known solns. The prepns of the solns and the graph are described in 4.4.3.5.2.

Percent Wax + Lecithin = (C x 100) / W

where: C = Wt of wax + lecithin from the graph
W = Wt of sample

4.4.3.2.4 Determination of Calcium Chloride. Weigh accurately a 2-g sample into a tared sintered glass filtering crucible, medium porosity, 30-ml capacity. Using a Fisher filtrator with water vacuum, extract sample with 100ml of distilled w, 10 extractions, 10ml each, of 30 sec duration. Collect filtrate in a 250 Erlenmeyer flask. Cool to RT and add 2ml of 0.1M K chromate indicator to the filtrate. (For the prepn of 0.1M K chromate see 4.4.3.5.3). Titrate to the end point with 0.1N Ag nitrate. (For the prepn of 0.1N Ag nitrate soln, see 4.4.3.5.4). The end point is noted by the first of a permanent colored red ppt of Ag chromate. The end point should be detd by using a white background or by transferring filtrate plus a minimum of distilled w washings to a white casserole.

Percent Ca Chloride = [V x N x (0.0555)] / W

where: V = ml of Ag nitrate
N = Normality of Ag nitrate
0.0555 = Millequivalent weight of Ca chloride
W = Weight sample

4.4.3.2.5 Determination of RDX + Nitrocellulose. The percentages of TNT, Ca chloride, aluminum, and wax + lecithin are added and their sum subtracted from 100 percent. The remainder is taken to be the percentage of RDX plus Nitrocellulose.

4.4.3.3 Alternate Method of Analysis. Grade A

4.4.3.3.1 Determination of RDX + Nitrocellulose. Weigh accurately a sample calculated to contain between 0.35 and 0.40g of TNT into a tared 100-ml beaker. Add 20ml of RDX saturated benzene to the beaker. The prepn of the RDX saturated benzene is described in 4.4.3.5.5. Cover beaker with a watch glass and place on a steam bath for 30 mins, swirl soln frequently. (An oscillating hot
plate may be used if available). Do not boil the benzene! After removal from the steam bath, cool sample to RT. Allow a minimum of 1 hour cooling

Make a quantitative transfer of the sample to the original tared filtering crucible. Filter using a Fisher filtrator with water suction. Use a small polyethylene wash bottle which contains benzene saturated with RDX to make the transfer. Make a total of 3 washings of the beaker and residue on the crucible, using between 10 and 15 ml of benzene saturated RDX per rinse. Make a final rinse of the crucible contents with 5 ml of benzene saturated with RDX. Collect the filtrate which contains TNT and wax in a 100-ml volumetric flask. Reserve the filtrate to det the TNT content. Aspirate the crucible for 1 min after the final rinse.

The crucible plus its contents are placed on a Fisher filtrator, and extracted with hot distd water, 3 portions of 5 ml each, 30 secs contact time for each washing. This is necessary to remove any Ca chloride that may be remaining in the residue. Dry crucible and its contents in an oven at 90°C ± 10°C for 1 hour. Cool the sample in a desiccator and weigh. The crucible and its contents are placed on a Fisher filtrator, attached to a water aspirator. Extract the sample with four 20-ml portions of hot, reagent grade, acetone. Allow 30 secs contact time between solvent and sample before applying suction for each extraction. Wash down the sides of the crucible with two additional portions of 10 ml each of hot acetone. Place crucible and its residue in an oven to dry at 90°C ± 10°C for 30 mins, cool in a desiccator and weigh. RDX and Nitrocellulose are removed in the filtrate. Aluminum remains as the residue on the crucible.

Percent RDX + Nitrocellulose = 

\[(\frac{(A-B) \times 100}{W})\]

where: 
A = Wt of sample plus crucible before acetone extraction
B = Wt of residue plus crucible after acetone extraction
W = Wt of sample

4.4.3.2 Determination of Aluminum. The aluminum is the residue remaining on the crucible after the acetone extraction.

Percent aluminum = 

\[(\frac{(B-D) \times 100}{W})\]

where: 
B = Wt of residue plus crucible after acetone extraction
D = Wt of original crucible
W = Wt of sample

4.4.3.3 Determination of TNT. For method of prepn and standardization of soln used in the TNT detn see 4.4.3.5.6 thru 4.4.3.5.13

Dilute the TNT and wax filtrate obtnd in 4.4.3.3.1 to 100 ml with benzene satd with RDX. Transfer a 10-ml aliquot of the TNT soln which conts 35-40 mg of TNT to a 300-ml reduction flask (Scientific Glass Apparatus Co JD 2776 or equivalent is satisfactory). Evaporate the benzene to dryness with a slow stream of dry air (connect a drying tube between the supply and inlet tube of flask).

Dissolve the residue in 25 ml of reagent grade glacial acetic acid (measure in a cylinder). Sweep the flask with N₂ or CO₂ gas for 5 mins. Pipette accurately 25 ml of 0.2 N titrns chloride soln into the flask. Add 25 ml of 6N HCl. A current of CO₂ or N₂ should be passed thru the reaction flask during the refluxing, cooling and titration periods to prevent air oxidation of the titanous ion. Reflux for 15 mins, using glass beads to reduce bumping. (Ground glass joints on the flask and condenser are to be preferred. A Glass-Col heating mantle is the most convenient source of heat, though a hot plate may be used). Cool the flask to RT without disconnecting the reflux condenser. (Lift the flask and condenser and substitute a pan of cold water for the heater). Titrate the sample in the flask with standard ferric ammonium sulfate (0.15 N) soln, using a magnetic stirrer if available. As the end point is approached [the Ti(III) color gets light] add 5 ml of 20% ammonium thiocyanate and continue the titration to the appearance of the red color.

At least 4 ml of 0.15 N Fe(III) soln should be required in back titration. If less, repeat adding more Ti(III) soln in excess to the TNT–wax filtrate. [If less than 4 ml of Fe(III) soln is used in the back titration in either standardization of Ti(III) soln or in the detn of TNT, low values may be obtd. If the excess Ti(III) at the end of the refluxing is too small, reduction of the nitro groups may be incomplete. On the other hand, if more than 10 ml of Fe(III) soln is used, an unnecessarily large excess of Ti(III) soln is being added]. Run a blank on a volume of RDX satd benzene equal to the volume of the aliquot of sample soln used in the preceding titration.

Percent TNT =

\[(\frac{0.1261 \times (AN-BF) - (CN-DF) \times 100}{W})\]
where: \( A = \text{ml of Ti(III) soln} \)
\( N = \text{Normality of Ti(III) soln} \)
\( B = \text{ml of Fe(III) soln} \)
\( F = \text{Normality of Fe(III) soln} \)
\( C = \text{ml Ti(III) added to blank} \)
\( D = \text{ml Fe(III) added to blank 0.0126l} \)
\( \text{TNT} = 1 \text{ meq of TNT} \)
\( W = \text{Wt of sample represented by aliquot AN–BF = Meq Ti(III) used by TNT & RDX} \)
\( \text{CN–DF = Meq Ti(III) used by RDX} \)

4.4.3.3.4 Determination of Calcium Chloride.
Calcium chloride is determined as in 4.4.3.2.4

4.4.3.3.5 Determination of Wax + Lecithin. The percentages of TNT, RDX plus Nitrocellulose, Ca chloride, and aluminum are added and their sum subtracted from 100 percent. The remainder is taken to be the percentage of wax plus lecithin

4.4.3.4.1 Method of Analysis. Grade B

4.4.3.4.1 Determination of Hot Melt – Visually. Weigh accurately a 1 g sample of HBX into a tared sintered glass filtering crucible, medium porosity, 30-ml capacity. Using approx 5ml of ethylene chloride, divided into 5 equal portions, 1 min contact time each, extract sample on a Fisher filtrator with water vacuum. Collect filtrate in a 50-ml volumetric flask and dilute to the mark with ethylene chloride. Compare the color of the filtrate visually with previously prep'd standards to det the weight of hot melt present. The prep of the hot melt standard solns are described in 4.4.3.5.14

Percentage of hot melt = \( (A \times 100) / W \)

where: \( A = \text{Wt of hot melt from comparison with standards} \)
\( W = \text{Wt of sample} \)

4.4.3.4.2 Determination of Hot Melt – Spectrophotometer. Compare the spectral absorbance at 20°C of the ethylene chloride filtrate obtd in 4.4.3.2.1 or 4.4.3.4.1 with that of a standard soln (0.4g of TNT/50ml of ethylene chloride) at 430 millimicrons. The weight of hot melt is detd by reference to a graph prep'd in advance from standard hot melt solns. The prep of the solns and graph are described in 4.4.3.5.14. The TNT + wax + hot melt filtrate collected in 4.4.3.3.1, diluted to 50ml with RDX satd benzene may be substituted for the ethylene chloride–TNT soln obtd in 4.4.3.4.1. All graphs and standard solns needed will substitute RDX satd benzene for ethylene chloride

Percentage of hot melt = \( (A \times 100) / W \)

where: \( A = \text{Wt of hot melt obtd from the graph} \)
\( W = \text{Wt of sample} \)

4.4.3.4.3 Determination of TNT – Spectrophotometer. Hot melt interferes with the method for the absorbance of TNT as described in 4.4.3.2.2. Corrected TNT values can be obtd by comparing the TNT soln at 430 millimicrons in addition to 367 millicrons. At 430 mu, the absorption is due to the hot melt alone. Correct the weight for TNT obtd from TNT spectral absorbance graph by subtracting from it the value obtd from a correction graph prep'd in advance as described in 4.4.3.5.15. The TNT + wax + hot melt filtrate collected in 4.4.3.3.1, diluted to 50ml with benzene satd RDX may be substituted for the ethylene chloride soln in 4.4.3.2.2. All graphs and standard solns needed will substitute RDX satd benzene for ethylene chloride

Percent TNT = \( [(A–B) \times 100] / W \)

where: \( A = \text{Wt of TNT from TNT spectral calibration graph} \)
\( B = \text{TNT correction value from the TNT correction graph} \)
\( W = \text{Wt of the sample} \)

4.4.3.4.4 Determination of TNT – Titanous Chloride Method. Hot melt is present in the TNT wax filtrate obtd in 4.4.3.3.1. This filtrate is used in the TNT detn by the titanous chloride method. Hot melt does not interfere with the TNT detn

4.4.3.4.5 Determination of Aluminum. The method described in 4.4.3.2.1 or 4.4.3.3.2 may be used to detn the aluminum content

4.4.3.4.6 Determination of RDX + Nitrocellulose. The method described in 4.4.3.2.5 or 4.4.3.3.1 may be used to detn the RDX + Nitrocellulose content

4.4.3.4.7 Determination of Calcium Chloride. The method described in 4.4.3.2.4 is used to detn the Ca chloride content

4.4.3.4.8 Determination of Wax + Lecithin. The percentages of TNT, RDX plus Nitrocellulose, Ca chloride, aluminum, and hot melt are added and their sum subtracted from 100%. The remainder is taken to be the percentage of wax plus lecithin

Where the method of analysis described in 4.4.3.3 is not used in its entirety, the wax plus lecithin content can be detd as follows: The crucible plus residue obtd after the RDX satd benzene extraction as described in 4.4.3.3.1 is extracted with an additional 50ml of RDX satd benzene (5–10ml portions, 30 secs each). The sample is dried at 100°C for 30 mins. The crucible
plus residue are cooled in a desiccator and weighed. The loss in weight of the sample is equal to the weight of the TNT + wax + hot melt extracted.

Percent wax + lecithin =
\[
\frac{[A-(B+C)] \times 100}{W}
\]

where:  
- A = Loss in wt of sample after RDX satd benzene extraction
- W = Wt of sample
- B = Wt of TNT detd in 4.4.3.2.2 or 4.4.3.3.3
- C = Wt hot melt detd in 4.4.3.4.1 or 4.4.3.4.2

The refractive index method for the detn of wax plus lecithin described in 4.4.3.2.3 cannot be used when hot melt is present.

4.4.3.5 Preparations of Solutions, Graphs and Standardization of Solutions:

4.4.3.5.1 Preparation of TNT Standard Solutions and TNT Spectral Calibration Graph. Prepare a stock TNT soln. Accurately weigh and transfer 8g of TNT into a 500-ml volumetric flask. Dilute to mark with ethylene chloride.

Into each of seven 50-ml volumetric flasks, pipette the following quantities of the TNT stock soln respectively: 22ml, 23ml, 24ml, 25ml, 26ml, 27ml, 28ml. Dilute with ethylene chloride to the mark.

Weight of TNT in each standard solution =
\[
A \times 0.016
\]

where:  
- A = Volume of TNT stock soln added to 50-ml volumetric flask
- 0.016 = Weight of TNT (gms)/ml of TNT stock soln

Prepare the TNT spectral calibration graph by detg the absorbance at 367 mu of the standard TNT solns at 20°C compared to a soln of 0.40g of TNT/50ml of ethylene chloride using 1 mm path cells. Plot absorbance against weight of TNT.

4.4.3.5.2 Preparation of Wax Standard Solutions and Wax Refractive Index Calibration Graph. Prepare 3-g HBX samples with different weights of wax and lecithin. The samples must be prepared with the same materials used to prep the HBX being analyzed. The weight of wax + lecithin in the sample is based on the wax from the D-2 and Comp B (if used). The standard wax samples are treated as described in 4.4.3.2.3 to obtn the wax in soln.

Prepare the wax refractive index calibration graph by detg the refractive index of the various solns of known wax concentration under the same conditions described in 4.4.3.2.3. Plot the refractive index readings from the samples against weight of wax + lecithin.

4.4.3.5.3 Preparation of 0.1M Potassium Chromate Solution. Accurately weigh 1.942g of reagent grade K chromate into a 100-ml volumetric flask. Dilute to mark with distd w. Mix well.

4.4.3.5.4 Preparation of 0.1N Silver Nitrate Solution. Accurately weigh 16.989g of reagent grade Ag nitrate into a liter volumetric flask. Dilute to mark with distd w. It is not necessary to standardize the Ag nitrate soln if care is taken in the prep of the soln. Store the Ag nitrate soln in a dark place.

4.4.3.5.5 Preparation of RDX Saturated Benzene. To a gallon of reagent grade benzene, add RDX in excess of solubility, and stir for several hours. Let stand overnight. The soln should be prepd and kept at the same temp as will prevail at the time of filtering the extracted sample. Filter soln just before use.

4.4.3.5.6 Preparation of 20 Percent Solution of Ammonium Thiocyanate. Dissolve 60g of reagent grade ammonium thiocyanate (NH₄SCN) in 240ml of distd w. Filter until clear and store.

4.4.3.5.7 Preparation of 6N Hydrochloric Acid. Add 250ml of reagent grade 38% HCl to 250ml of distd w. Mix well.

4.4.3.5.8 Preparation of 0.2N Titious Chloride Solution. Mix 150ml of 20% titious chloride soln with 100ml of 38% HCl soln. (As the concn of the 20% titious chloride varies from bottle to bottle, one may use an adjusted volume for the prep of additional 0.2N Ti(III) soln rather than the 150ml of 20% soln). Dilute quantitatively to 1 liter with distd w. Mix by bubbling a stream of nitrogen or carbon dioxide gas thru the soln, filter thru Whatman No 41 (fluted) or equivalent paper, store in a system arranged so that only carbon dioxide or nitrogen gas will be drawn into the stock bottle as the soln is used. (Scientific Glass Apparatus Co JB-7670, burette, automatic, for titious chloride solution, improved form; or JB-7615, burette, automatic, or equivalent is satisfactory. Teflon stopcocks are to be preferred).

4.4.3.5.9 Preparation of 0.15N Ferric Ammonium Sulfate Solution. Dissolve 75g of hydrated ferric ammonium sulfate (FeNH₄(SO₄)₂·12H₂O), reagent grade, in 600ml distd w. Add to this 25ml of 95% reagent grade sulfuric acid. When dissolved, dilute with distd w to 1 liter; filter, mix thoroughly, and store.
4.4.3.5.10 Preparation of 0.200N p-Nitroaniline Solution. Use p-nitroaniline with a mp of 147–148°C, and recrystallize once from ethanol. Dry in a desiccator. For a 0.200N soln weigh exactly 1.151g of recrystallized p-nitroaniline, dissolve in reagent grade glacial acetic acid, transfer to a 250-ml volumetric flask and dilute to the mark with glacial acetic acid. Mix well.

4.4.3.5.11 Comparison of Titanous Chloride and Ferric Ammonium Sulfate Solutions (Determination of R). This value is necessary in the determination of the normality of the Ti(III) and the normality of the Fe(III). R equals the ml of Ti(III) reacting with 1.00ml of Fe(III) soln. Sweep the air from a 300-ml reduction flask with a current of nitrogen or carbon dioxide gas for 5 mins. Continue to pass the current of nitrogen or carbon dioxide gas thru the flask until the titration is completed.

Pipette 50.00ml of approx 0.15N Fe(III) soln into the air free reduction flask. Add 25ml of 6N HCl (use a cylinder to measure the HCl). Titrate with approx 0.2N Ti(III) soln until near the end point (the Ti(III) color gets light). Add 5ml of 20% ammonium thiocyanate soln (by cylinder), and continue the titration until the red color just disappears. Repeat the procedure until two successive values for R agree to within 1 part per thousand of their mean.

\[ R = \frac{A}{B} \]

where: \( A = \text{ml of Ti(III) soln} \)
\( B = \text{ml of Fe(III) soln} \)

4.4.3.5.12 Standardization of 0.2N Titanous Chloride Solution. Sweep the air from a 300-ml reduction flask with a current of nitrogen or carbon dioxide gas for 5 mins. Continue to pass the current of gas thru the flask until the titration is completed.

Pipette 20.00ml of the 0.200N p-Nitroaniline soln into the reduction flask. Pipette 25ml of titanous chloride approx 0.2N into the same flask. Add 25ml of 6N HCl (by cylinder), and a few glass beads. Connect the reduction flask to a reflux condenser and boil for 15 mins. Cool the flask to RT without disconnecting the condenser (lift the flask and condenser, remove the heater, and let the flask down into a pan of cold water).

Titrate the excess Ti(III) with Fe(III) soln. As the end point is approached, Ti(III) color gets light, add 5ml of 20% ammonium thiocyanate (by cylinder), as an indicator and titrate to the appearance of the red color. Run a blank, substituting 20ml of glacial acetic acid for the p-nitroaniline soln.

\[ \text{Normality of Ti(III) solution} = \frac{4.000}{[A \text{ml} - B \text{ml} - (C \text{ml} - D \text{ml})]} \]

where: \( 4.000 = \text{Millequivalents of p-nitroaniline in 20.00ml of 0.200N soln} \)
\( A = \text{ml of Ti(III) soln} \)
\( B = \text{ml of Fe(III) soln} \)
\( C = \text{ml of Ti(III) in blank} \)
\( D = \text{ml of Fe(III) in blank} \)
\( R = \text{ml of Ti(III) reacting with 1.00ml of Fe(III) soln} \)

The term \( (C - D) \) should be zero. If the value lies outside the range +0.10 and -0.10, either R is incorrect, or the acetic acid is contaminated. Repeat the detn of R. Continued high or low values for \( (C - D) \) probably mean impure acetic acid.

If normality of Ti(III) soln falls outside of the range 0.19–0.22, add 20% titanous chloride soln or water as required to bring normality to approx 0.2N. Then repeat the detn of R and repeat the detn of the normality.

4.4.3.5.13 Determination of the Normality of the Ferric Ammonium Sulfate Solution. Using the value of R and the value of the normality of titanous chloride detd in 4.4.3.5.11 and 4.4.3.5.12, calculate the normality of the ferric ammonium sulfate soln.

\[ F = \frac{\text{Normality of Fe(III) soln} \times (R)}{N} \]

where: \( R = \text{ml of Ti(III) soln reacting with 1.00ml of Fe(III) soln} \)
\( N = \text{Normality of Ti(III) soln} \)

4.4.3.5.14 Preparation of Hot Melt Standard Solutions and Hot Melt Spectral Calibration Graph. Prepare a stock TNT soln by accurately weighing and transferring 8g of TNT into a 500-ml volumetric flask. Dilute to mark with ethylene chloride. Prepare a stock hot melt soln using material conforming to Specification MIL-C-3301. Accurately weigh and transfer 0.45g of hot melt into a beaker and dissolve in a 100ml of ethylene chloride. Filter soln thru a sintered glass filtering crucible, medium porosity, 30-ml capacity until filtrate shows no sign of sedimentation.

To det weight of hot melt per ml of filtrate, accurately pipette 15ml of filtrate into a tared evaporating dish with cover. Evaporate to dryness on a steam bath or hot plate below 75°C. Replace cover, cool in a desiccator to RT and weigh.

\[ \text{Weight of hot melt per ml of solution} = \frac{A - B}{V} \]
where: \( A = \text{Wt of evaporating dish plus cover plus residue} \)
\( B = \text{Wt of evaporating dish plus cover} \)
\( V = \text{Volume of hot melt filtrate pipetted into evaporating dish} \)

Into each of six 50-ml volumetric flasks, pipette accurately 25ml of the TNT stock soln, add to each of the flasks respectively, by pipetting accurately, 0.5ml, 1.0ml, 1.5ml, 2.0ml, 2.5ml, 3.0ml of the hot melt filtrate. Dilute with ethylene chloride to the mark.

Weight of hot melt in standard = WD

\( D = \text{ml of hot melt added to the standard} \)

Prepare the hot melt spectral calibration graph by detg the absorbance at 430mu of the standard hot melt and TNT smlns at 20°C compared to a soln of 0.400g of TNT/50ml of ethylene chloride using 1 mm path cels. Plot absorbance against weight of hot melt.

4.4.3.5.15 Preparation of the TNT Correction Graph for the Presence of Hot Melt. Determine the spectral absorbance of the hot melt and TNT standard solns at 367mu, at 20°C, using 1 mm path cels. The prep of the solns is described in 4.4.3.5.13. From this data determine the apparent weight of the TNT from the TNT spectral calibration graph described in 4.4.3.5.1. Subtract the actual weight of TNT (0.4g) in the standard hot melt soln from the apparent weight of TNT. This gives the TNT correction value.

Determine the spectral absorbance of the hot melt standard solns at 430mu as described in 4.4.3.4.14.

For the TNT correction graph for the presence of hot melt plot the absorbance of the standard hot melt solns at 430mu against the TNT correction values obtd at 367mu.

4.4.4 Moisture Determination. Remove sample of HBX from closed sample container. Sample size shall be approx 100g for Grade A material and 50g for Grade B material. Break up quickly and transfer the sample to a tared liter Erlenmeyer flask with a ground glass neck. Stopper flask and record weight of flask plus sample. (When sample is broken up, it should be exposed to the air as little as possible as the Ca chloride in the HBX quickly takes up moisture).

To the flask containing the sample add 200ml of toluene and a magnetic stirring bar. Attach flask to a Bidwell-Sterling trap (graduated 5ml in 0.1ml). To the top of the trap is attached a cold water condenser. (See Figure 1). (The use of Teflon sleeves on the ground glass joints in place of a lubricant will facilitate the cleaning of the glassware).

Heat the assembly on a magnetic stirrer hot plate, with agitation provided by the magnetic stirrer. Heat to boiling. (The toluene-water azeotrope condenses and falls back into the trap. Water being heavier, collects in the trap and the excess toluene returns to the boiling flask). Continue boiling until no change in the water volume has been observed for 15 mins. Boil a minimum of 1 hour.

CAUTION: Keep soln well agitated at all times. When soln begins distilling, position flask so that an air space is between the flask and hot palate. This is to prevent a hot spot from building up in the chunk explosive. Sample should not be left unattended during the moisture detn. Care must be taken that the cold water condenser is operating properly at all times. If the condenser should fail to trap the toluene and the flask were to boil dry, there could be a possibility of overheating the expl.

Care should be taken to prevent condensation of atmospheric moisture in the condenser. Five minutes before heating is discontinued wash down the condenser with toluene to remove any water clinging to the tip of the condenser.

After the moisture has been collected, transfer the trap to a constant temp bath at 40°C ± 2°C. Allow about 30 mins for trap contents to reach the temp of the bath. By means of a wire loop work any droplets of water trapped along the side to the bottom of the trap. Read the volume of the water collected.

Run a blank on the toluene to det water present in the toluene. Calculate the percentage of water in the sample.

\[
\text{Percentage of water} = \frac{(A-B) \times C \times 100}{W}
\]

where: \( A = \text{ml of water} \)
\( B = \text{ml of water in toluene} \)
\( C = \text{Density of water at temp of the water bath} \)
\( W = \text{Wt of the sample} \)

4.4.5 Vacuum Stability Test, 100°C

4.4.5.1 Calibration of Glass Tube. Determine the volume in mls of the 15.5cm heating tube by running in mercury from a buret until the tube is filled to the level at which the ground glass
joint of the capillary tube will make contact with the mercury. Subtract from the indicated buret reading, the volume of expl used in the test. The difference shall be represented by the symbol A. Transfer 7.0 ml of mercury to the cup at the lower end of the capillary tube. Clamp the tube in an upright vertical position, and measure the height in mm of the mercury column in the capillary tube (approx 25mm). Measure the length in mm of each of the 3 parts of the capillary tube and add these values to obtain total length. From the total length subtract the height of the mercury column in the capillary tube as previously obtained. Represent this difference by the symbol B1. From the total length subtract the height of the column of mercury in the capillary tube measured at the end of the test described in 4.4.5.1.

Represent this difference by the symbol B. Determine the capacity of the capillary tube per unit of length as follows: Transfer an accurately weighed sample of approx 10g of mercury to the cup at the lower end of the capillary tube. Manipulate the tube so that when it is horizontal, mercury is contd in a continuous section of the longest part of the tube and measure the length of the mercury column. Repeat this twice with the mercury in 2 other parts of the long section of the tube. Calculate the average of the 3 measured lengths of the mercury column. Represent the unit capacity in ml per mm of the capillary tubing by the symbol C. This can be obtained from the formula:

\[ C = \frac{W}{d \bar{x}} \]

where: 
C = Unit capacity of capillary tubing in ml per mm
W = Grams of mercury
d = Density of mercury at temp of determination
\( \bar{x} \) = Average measured lengths of mercury column in mm

4.4.5.2 Test Procedure. Transfer a 1 g sample, dried at 65°C for 2 hours, to the heating tube of the apparatus shown in Figure 2. Connect the capillary tube to the heating tube. Clamp the apparatus so that the long section of the capillary tube is in a nearly vertical position. Transfer 7.0 ml of mercury to the cup at the lower end of the capillary tube. Connect a vacuum pump to the lower end of the capillary tube and evacuate the system until the pressure is reduced to approx 5mm of mercury. (Evaluation of the capillary
tube is facilitated by placing the cup of the tube in a horizontal position so that mercury does not block the capillary opening). After evacuation, disconnect the pump. Seal the connection between the capillary tube and the heating tube with 1 ml of mercury. Measure the total vertical height of the column of mercury in the capillary tube. Measure and subtract the vertical height of the mercury in the cup. The difference shall be represented by the symbol \( H_1 \). Note the RT \( t_1 \) and the barometric pressure. Subtract the value \( H_1 \) from the barometric pressure in mm. Represent this difference by the symbol \( P_1 \). Insert the heating tube in a constant temp bath consisting of a soln of glycerin and water (sp gr 1.05). Maintain at a temp of 100.00°C ± 0.5°C for 48 hours. Remove the heating tube from the bath and allow to cool to RT. Measure the total vertical height of the column of mercury in the capillary tube and subtract the vertical height of the mercury in the cup. This difference shall be represented by the symbol \( H \). Note the RT \( t \) and the barometric pressure in mm. Represent this difference by the symbol \( P \).

4.4.5.3 Calculation of Liberated Gas Volume.
Calculate the gas vol in ml (at standard conditions) liberated in the test described in 4.4.5.2, using the values described by the symbols in 4.4.5.2 and 4.4.5.1 in the following formula:

\[
V = \frac{[A+C(B-H)] \times 273P}{760(273+t)} - \frac{[A+C(B_1-H_1)] \times 273P_1}{760(273+t_1)}
\]

6.3 General Safety Precautions. The preparation and handling of the items covered by this specification, and the subassemblies thereof, involve hazardous operations and therefore require explosives safety precautions. Use of this specification will not be construed as to relieve the contractor or manufacturer of responsibility for the safety of his operations. Listed below are certain minimum provisions which a contractor or manufacturer (who prepares the item covered) should observe in order to fulfill his responsibility for safety. At Bureau of Naval Weapons, Navy Department, and other government plants, these provisions are mandatory. Such other warnings and precautions, pertinent to the operational effectiveness or safety during preparation of the specified items, are included in detailed technical requirements of the specification.

6.3.1 All handling and bathing operations should be conducted in a neat and orderly manner.
6.3.2 Safe equipment and methods should be utilized for transporting and handling explosives components and mixtures. Where required, remote controlled barricaded handling equipment shall be used for explosives operations, such as mixing, pouring, weighing, charging, sifting, drying, casting, etc.
6.3.3 The exposure of explosive materials and related parts should be so controlled as to minimize the absorption of moisture from the atmosphere or other sources during handling and batching operations.
6.3.4 All explosive components and mixtures should be stored in suitable storage magazines located in accordance with American Table of Distances (ATD) or other applicable safety standards; and while in process, in safety lockers and chests if in loading rooms, or in adequate ready or service magazines located in accordance with Intraplant distances when outside of loading rooms. For Navy managed explosive loading plants, the provisions of the Armed Services Explosives Safety Board covering quantity-distance relations for explosives will apply.
6.3.5 Proper care must be exercised at all times to protect personnel from accidents, fires, or explosions, and to limit damage to equipment and loading areas. In this connection, the precautionary measures in the following paragraphs should be observed.

6.3.5.1 Employ properly proportioned and properly located protective barricades, screens or shields at all required points.
6.3.5.2 Keep only minimum limited quantities of explosives components and expl mixtures at each stage of operations.
6.3.5.3 Keep explosives and expl components in approved covered receptacles with covers in place when material is not being taken out of or put into the receptacles. Where necessary, receptacles should be conductive to ground electrostatic charges.
6.3.5.4 Protect operations from electrostatic charges by effectively grounding all machinery, equipment and fixtures; and, where necessary, employ suitable grounded conductive coverings for floors, 2 ork benches and tables, and workers' conductive shoes. Workers' clothing of a type to minimize the accumulation of static charges.
should be employed. Fabrics such as silk and nylon, which promote static generation should be avoided. Additional precautions should include mechanical shielding to contain fragments and blast, also electrical shielding from induced electric currents generated by sources such as lightning, static, radiations from communications apparatus, radar, or high frequency heat apparatus, etc. Additional grounding devices such as grounding bracelets for workers should be employed where operations are conducted with items which are unusually sensitive to initiation by static electricity. Where necessary for safety, humidity of work rooms should be appropriately increased, as required, to lessen electrostatistic effects but without inducing excessive moisture absorption by any of the components.

6.3.5.5 Protect all exp loaded operations from effects of electric current originating from equipment such as soldering irons, heaters, switches, wiring, motors, lights, test instruments, etc, by suitable insulation, grounding, separation or shielding. Such electric sources may initiate exps by heat, sparks, arcs. Circuits may be inadvertently completed, for example: from a defective soldering iron thru a ground contact.

6.3.5.6 Enforce, where necessary, the wearing of suitable safety footwear, gloves, goggles, respirators, and impregnated garments to protect personnel against burns, poisoning, and associated industrial hazards.

6.3.5.7 Allow no fires or exposed electrical or other sparking equipment, and little or no flammable material to be present in loading, handling and storage spaces. Enforce proper “Match” and “No Smoking” rules where necessary.

6.3.5.8 Enforce good housekeeping and maintain effective policing, inspection and supervisory methods throughout the loading area and surroundings. Employ effective cleaning methods periodically to minimize the accumulation of explosives and explosives dust and other contaminants upon, and assure its removal from floors, walls, ceilings, ledges, tables, benches, piping and equipment or the item loaded; also, clean up any spilled material immediately.

6.4 Manufacture by Government Activities. When the HBX type explosive compns are to be prpd in accordance with this specification by government activities, the requirements given herein for bidders and contractors shall apply to such government activities.

HBX-1 Analytical Procedure. Pristera (Ref) briefly describes its analysis as follows: Extract TNT wax and lecithin with carbon tetrachloride; then determine TNT by titration reduction as described on p 466 of Ref; extract with water the Ca chloride from carbon tetrachloride insoluble residue and extract RDX, with hot toluene, from water-insol residue using Wiley extractor. Extract NC from toluene-insol residue with cyclohexanone, acetone, or tetrahydrofuran; the remainder is Al. 


HBX-3 Analytical Procedure. Accdg to Pristera, the procedure is the same as for HBX-1. The procedure was also described by S. Semel et al in PATR 2459 (1957), entitled “A Chemical Method for the Composition Analysis of Cast HBX-3”.

H-6 Analytical Procedure. Accdg to Pristera, the procedure is the same as for HBX-1. 


HC Mixture. A German smoke screen mixture which partly replaced the less stable BM Mixture during WWI. The HC mixture contains Zn dust and hexachloroethane (solid) along with relatively small percentages of NH₄Cl, NaClO₃, and Mg CO₃. It is stable, safe and easily handled, but its cost is comparatively high. It was used in candles, the 4-inch Stokes mortar-shell (burning type) and HC grenades. 

Refs: 1) H.W. Walker, IEC 17, 1064 (1925) 
2) Anon, Field Artillery Journal 33, 352 (1943)

HDP Supergun. See Hochdruckpumpe or V-3 (Vergeltungswaffe Drei).

HE. Abbr: for High Explosives

Headaches Arising from Contact with NG or Dynamites. Test exposure (oral and skin contact) to a 20% NG Dynamite produced severe headaches, nausea and vomiting. These effects are ascribed to intracranial vasodilatation produced by the NG vapors or liquid. Headaches were alleviated by intramuscular injection of caffeine-Na-benzoate followed by oral administration of...
amph.etamine sulfate. Headaches were prevented by daily oral administration of amphetamine sulfate for 2 or 3 days prior to NG exposure (Ref 1).

Reaction to NG among pharmaceutical workers is described. Included in the discussion are physiological actions, susceptibility and habituation, toxic effects and methods of control (Ref 2).

NG headaches are ascribed to distention of intracranial veins or reduction of venous tone or insufficient tone resulting from enhanced arterial blood flow (Ref 3).


Health Hazards of Explosives and Propellants. Health hazards may be encountered in the manufacture, handling and use of explosives and propellants. These hazards vary with materials and degree of exposure and can range from mild dermatitis to severe poisoning.

More serious than dermatoses caused by skin contact with Tetryl, TNT, DNT, Hg-Fulminate, solvents etc, during explosives and ammunition production, are exposures to toxic dusts, fumes and vapors. Among these are TNT, DNT, oxides of N, Pb-dusts and vapors, and solvent vapors. Special skin cleansing agents and solns for detecting these harmful materials on the skin are discussed in Ref 1.

In propellant and missile production hazards may arise from contact with: ammonia, aniline, MeOH, furfuryl alc, hydrazine, JP-4, hydrogen peroxide and red fuming nitric acid (Ref 2), as well as other propellant ingredients such as NG, epoxy comps, polyurethanes, Amm Perchlorate etc (Ref 5).

In actual field use explosion product fumes can be hazardous, eg Ammonium Nitrate-fuel oil mixtures (ANFO) are not recommended for underground blasting because they produce toxic fumes (Ref 3). Highly toxic NO is often found in explosion fumes. The oxidation of NO to the less dangerous NO2 was found to be less rapid under mining conditions than was previously believed (Ref 4).


HEAT. Abbr for High Explosive Antitank

Heat (Definitions and Selected General References) A form of energy. The mean energy transferred from one system to another system as a result of purely thermal interactions (temperature gradients) is called heat.


Heat of Adsorption (see also Adsorption). The heat released when a gas is adsorbed on a surface. The differential heat of adsorption is given by:

\[ Q_{\text{diff}} = \Delta H_{\text{g}} - \Delta H_{\text{ads}} \]

where \( \Delta H_{\text{g}} \) and \( \Delta H_{\text{ads}} \) are heat contents or enthalpies (see below) of the gas and adsorbed phases.

Heat Capacity is the ratio of the heat absorbed to the rise in temperature for an infinitely small increase in temperature. For a process occurring at constant volume the heat capacity \( C_V = \frac{\delta E}{\delta T} \) where \( \delta E \) is the internal energy change of the process over a temperature change \( \delta T \).

Similarly for a constant pressure process \( C_P = \frac{\delta H}{\delta T} \) where \( \delta H \) is the change in heat content (enthalpy) of the process. When dealing with a unit weight of a substance, a small (c) is employed and this ratio is also known as specific heat. In cgs units c is given in calories per gram per degree. For many organic explosive compounds \( c \approx 0.3 \text{ cal/g}\circ C \) at rm temp


Heat of Combustion. See Vol 4, p D369 (Tables D380-1) of Encycl under DETONATION (AND EXPLOSION), DEFLAGRATION (AND COMBUSTION), AND FORMATION, HEATS OF

Heat of Condensation is the reverse of Heat of Evaporation or Vaporization (See under Heat, Latent)

Heat Conductivity, Specific or Coefficient of Thermal Conductivity (\( \lambda \)) is the quantity of heat in gram-calories transmitted per second through a plate of material one centimeter thick and one square centimeter in area, when the temperature differential between the two sides is 1\(^\circ\)C. When it is desired to express it in Btu per inch/second/degree F, the above value has to be multiplied by 0.00560

If two opposite faces of a cube, made from the substance to be examined, are maintained at temperatures \( (T_1) \) and \( (T_2) \), the heat conductivities across the section of the cube \( (A) \) cm\(^2\) and \( (D) \) cm thick, the specific heat conductivity:

\[ \lambda = \frac{K(T_2 - T_1)}{D} \]

where \((K)\) is a constant depending on the nature of the substance and \((i)\) is the time in seconds. See Table 4 of this Vol for heat conductivities of common explosives


Heat Content or Enthalpy. A thermodynamic property closely related to energy. It is defined by \( H = E + PV \) where \( E \) is the internal energy of the system, \( P \) is the pressure on the system and \( V \) is the volume of the system. Often it is used in differential form as in \( \Delta H = \Delta E + P\Delta V \) for a constant pressure process

Heat of Crystallization. The number of calories liberated or absorbed per mol, or gram when a substance passes into the crystalline state. In cases of solidification of crystalline compounds, the heat liberated on freezing may be considered as the heat of crystallization

Heat of Decomposition. The heat evolved or absorbed in a particular decomposition process. Thus the heat of decomposition of \( A \rightarrow B + C \) is usually different from \( A \rightarrow B + D \). However, \( A \rightarrow E + F \rightarrow B + C \) has the same net heat effect as \( A \rightarrow B + C \). The heat of decomposition is generally expressed in kg calories per gram molecule or in calories per gram

Heat of Deflagration. See Vol 4 of Encycl under DETONATION (EXPLOSION, DEFLAGRATION, COMBUSTION AND FORMATION) HEATS OF, p 374-R

Heat of Detonation. See Vol 4 of Encycl pp D 370 to 375

Heat of Dilution. Is the quantity of heat consumed or liberated when a solution is diluted
Heat of Dissociation is the heat involved in the disruption of a chemical bond.

Heat of Dissolution or of Solution is the quantity of heat liberated or absorbed when a solid is dissolved or two miscible liquids are mixed.

Heat Effects — Data for Common Explosives. The data required for computing various heat effects involving explosives and explosions are: standard heats (also called enthalpies) of formation, heats of detonation (or explosion), heats of fusion, vaporization and/or sublimation, heat conductivity, and specific heat.

Tables of these data for common explosives are given below. Examples of the use of these data are shown in the next section. In conformance with modern usage exothermic heat effects have a negative sign.


1) Heat of nitrification of glycerin. Suppose we wish to estimate the heat evolved in nitrating 1 mole of glycerin with mixed acid. To simplify this illustrative example we will assume: initial mixture — 1 mole glycerin/3 moles 100% nitric acid/6 moles 100% sulfuric acid; final mixture — 1 mole nitroglycerin completely separated from the spent acid which is now diluted by 3 moles of water; the entire heat of dilution is due to 3 moles water dissolving in 6 moles of 100% sulfuric acid. Thus

\[ \Delta H_f^\circ = -159.8 \text{ kcal/mole} \]

\[ \Delta H_a = -9.4 \text{ kcal/mole} \]

b) 6H\(_2\)SO\(_4\)(100%) = 6H\(_2\)SO\(_4\) (aq) (½ H\(_2\)O)

\[ \Delta H_f^\circ = -6x193.9 = -6x197.67 \]

\[ \Delta H_b = -22.6 \text{ kcal/mole} \]

The net heat effect is \( \Delta H_a + \Delta H_b = -32 \) kcal/mole glycerin, most of which comes from the heat of dilution of H\(_2\)SO\(_4\). The \( \Delta H_f^\circ \)'s for these calculations were taken from Table 1 and Refs 1 & 7.

2) Heat of Detonation. Price (Ref 3) has shown that the heat of detonation of an explosive correlates well with the blast effects (shock, impulse and bubble energy) that the explosive produces in air and underwater. This correlation is confirmed in Ref 9, where it is also pointed out that explosive power, as measured in the ballistic mortar or lead block, can be related to the heat of detonation.

Furthermore, heat of detonation can be an important parameter in establishing the equation of state of detonation products or in thermal explosion calculations. Consequently, the heat of detonation is an important property of an explosive and methods of its accurate evaluation can contribute greatly to explosive technology.

To obtain the true value of the heat of detonation one must know the composition of the detonation products at C-J conditions. At present such compositions can be obtained only by theoretical calculations. These calculations depend strongly on the choice of the equation of state of the detonation products. For military CHNO explosives the main equilibria that determine product composition are (Ref 9):

\[ \text{CO}_2 + \text{H}_2 = \text{CO} + \text{H}_2\text{O} \]
\[ 2\text{CO} = \text{CO}_2 + \text{C} \]
\[ \text{CO} + \text{H}_2 = \text{H}_2\text{O} + \text{C} \]
\[ \text{N}_2 + 3\text{H}_2 = 2\text{NH}_3 \]
\[ \text{C} + 2\text{H}_2 = \text{CH}_4 \]

if the equil mixture of (a), (b) and (c) contains appreciable amounts of H\(_2\). Of these equilibria only (a) is relatively independent of pressure and only moderately dependent on temperature in the expected C-J range. For oxygen-balanced explosives equil (a) is controlling. Thus, the composition of detonation products of oxygen-balanced explosives should be nearly independent of the choice of eqn of state.

For military explosives that are deficient in oxygen (negative OB), equil (b) and (c) must be considered and the choice of eqn of state becomes important. Currently the BKW eqn of state (see Vol 4, p D284) is used for most calculations, eg Ref 4. From Mader's calcs (Ref 4) combined with the \( \Delta H_f^\circ \)'s of Table 1, the heats of detonation of NG, PETN, HMX, RDX, PBX 9404, LX-04, TNT and NM are
respectively: 1.48, 1.51, 1.475, 1.48, 1.41, 1.31 (Ref 10), 1.24 and 1.30 kcal/g. These values are to be compared with experimental $\Delta H_d$'s (for heavily confined samples) (Tables 1 and 2) of: $-1.49, 1.48, 1.42, 1.38, 1.31, 1.09$ and $1.23$ kcal/g. This excellent agreement between theory and experiment (except for TNT) is really quite amazing. Theory considers $H_2O$ as gaseous, whereas it is liquid in the calorimeter. Furthermore, the detonation product composition in the calorimeter is some frozen equilibrium composition (probably corresponding to a freeze-out temperature of $\approx 1600^\circ$K), while CJ temperatures are of the order of 2000-4000$^\circ$K. What appears to be happening in the confined calorimeter samples is a series compensating heat effects namely: $H_2O(g) + H_2O(l)$ and $N_2 + 3H_2 = NH_3$ exothermic & the partial reversal of equil (b) & (c) (decrease of $C_2$) endothermic. Also at $T \approx 1600^\circ$K equil (a) is shifted towards $CO_2$ as compared to $T_{CJ}$ which tends to counteract to some extent the reversal of equilibria (b) and (c).

For a quick approximate computation of the heat of detonation of military CHNO explosives, the following is suggested (Ref 9) for obtaining an approximate product composition:

All $H \rightarrow H_2O(g); all remaining O is used to form CO$_2$; any remaining C $\rightarrow C(s); all N \rightarrow N_2$

As an example of the use of this approximate method, let us estimate $\Delta H_{det}$ for Haleite (EDNA):

$$C_2H_6N_4O_4 + 3H_2O(g) + \frac{1}{2}CO_2 + 3/2C_2 + 2N_2$$

$$\Delta H_f^0's = -24.6 \quad 3(-57.8) \quad \frac{1}{2}(-94.05) \quad 0 \quad 0$$

$$\Delta H_{det} = 3(-57.8) + \frac{1}{2}(-94.05) + 0 + 0 + (-24.6)$$

$$= -195.8 \text{ kcal/mole}$$

$$\Delta H_{det} = 195.8 \text{ kcal/mole} = 1.30 \text{ kcal/g}$$

which compares amazingly well with a reported measurement (Ref 6) of 1.3 kcal/g.

3) Heat of decomposition. Since the products of a relatively slow ("low" temp) decomposition of an explosive will often be quite different from deton products, is not surprising that $\Delta H_{decomp} \neq \Delta H_{det}$. Hall (Ref 8) measured the decomposition of solid nitramines in the temp range of 365-540$^\circ$K. In the following tabulations some of his data are compared with $\Delta H_{det}$ taken from Table 1.

<table>
<thead>
<tr>
<th>Expl</th>
<th>$\Delta H_{decomp}$ (kcal/g)</th>
<th>$\Delta H_{det}$ (kcal/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HMX</td>
<td>0.56 at $\sim 540^\circ$K</td>
<td>1.48</td>
</tr>
<tr>
<td>RDX</td>
<td>0.61 at $\sim 479^\circ$K</td>
<td>1.42</td>
</tr>
<tr>
<td>Tetryl</td>
<td>0.33 at $\sim 445^\circ$K</td>
<td>1.17</td>
</tr>
</tbody>
</table>

4) Heat Balance in Estimating Critical Dimensions for Thermal Explosion. For thermal explosion to occur the rate of heat generation must exceed the rate of heat loss. The general equations for this situation can be solved only by numerical means. Specialized cases are, however, amenable to analytical solutions. An interesting case is that of a semi-infinite slab of explosive for which Friedman (Ref 5) obtained a series of approximate analytical solutions. These enable one to estimate the critical thickness of the slab and the time to explosion as functions of the initial slab temperature and the kinetic and thermal parameters of the explosive.

The approximate equations in dimensionless form are:

$$a_{cr}^2 = 4\theta_0 \exp(1/\theta_0) \ln[0.608(\theta_0 - \theta_0')/\theta_1^2]$$

(1)

$$\tau_0 = \theta_0^2 \exp(1/\theta_0) \alpha_{cr}^2$$

(2)

and

$$\theta_0 = \frac{RT_0}{E} \quad \theta_1 = \frac{RT_1}{E}$$

(3)

$$d^2 = a_{cr}^2 \lambda E / \rho RQZ$$

(4)

$$\tau_0 = \frac{\tau_0 c \rho d^2}{\lambda}$$

(5)

where $T_0$ is the slab temp, $T_1$ = ambient temp, $E$ & $\zeta$ are the activation energy and frequency factor of the explosive, and $\lambda$, $\rho$, $c$ and $Q$ are its thermal conductivity, density, specific heat and heat of reaction. The quantities $d$ and $\tau_0$ are respectively the critical $1/2$ thickness of the slab and the time to explosion at $T_0$ if the process is adiabatic.

To illustrate the use of the heat data in the preceding section for estimating thermal explosion parameters, let us determine the critical thickness of a semi-infinite slab of Tetryl kept at 445$^\circ$K. From Ref (2) we take $E \equiv 35000$ cal/mole and $Z \equiv 10^{18}$sec$^{-1}$. Substituting these values in eqn (3) and then into eqns (1) and (2) gives:

$$a_{cr}^2 = 3.24 \times 10^{15} \quad \& \quad \tau_0 \equiv 0.032$$
Now taking $Q = 330 \text{ cal/g}$ (Ref. 8) and the
other thermal data from Table 4, solutions of
eqns (4) and (5) lead to:

$$d = 5.4 \text{ cm and } T_0 = 132 \text{ sec at}$$

$$T_0 = 445^\circ \text{K and } \rho = 1.6 \text{ g/cc}$$

In qualitative terms this means that quite a
thick slab of Tetryl must be heated to around
$170^\circ \text{C}$ for over 2 minutes before explosion
occurs

*Written by J. ROTH*

*Refs:* 1) F. Rossini et al, NBS Circ 500 (1952)
59, 801 (1959) 4) C.L. Mader, LASL, LA
2900, July 1963 5) M.H. Friedman, Combust
& Flame 11, 239 (1967) 6) See Ref. 12 of
Table 1 7) See Ref. 14 of Table 1 8) P.G.
Hall, Trans Farad Soc 67, 556 (1971) & CA 74,
103822 (1971) 9) Anon, “Principles of Ex-
plosive Behavior”, AMCP 706-180 (April 1972)
10) UCRL-S1319 “Props of Chem Expls and
Exp1 Simulants”, Dec 1972
Table 1
Standard Heats of Formation and Heats of Detonation of Common Explosives

<table>
<thead>
<tr>
<th>EXPLOSIVE</th>
<th>Chemical Name</th>
<th>Code or Common Name</th>
<th>$\Delta H_f^0$ kcal/mole</th>
<th>Ref</th>
<th>$\Delta H_{det}$ kcal/g</th>
<th>Ref</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bis (trinitroethyl) nitrazine</td>
<td>BTNEN</td>
<td>-76.9</td>
<td>12</td>
<td>1.50</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>Bis (N,N-trinitroethylurea)</td>
<td>BTNEU</td>
<td>-32.7</td>
<td>12</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>1,3-Diamino-2,4,6-trinitro-benzene</td>
<td>DATB</td>
<td>-29.8</td>
<td>15</td>
<td>1.01</td>
<td>9</td>
<td></td>
</tr>
<tr>
<td>4,6-Dinitro-2-diazophenol</td>
<td>DDNP</td>
<td>-27.8</td>
<td>8</td>
<td>$\approx 0.8(d)$</td>
<td>14</td>
<td></td>
</tr>
<tr>
<td>Diethyleneglycolinitrate</td>
<td>DEGDN</td>
<td>-102(a)</td>
<td>14</td>
<td>$\approx 0.8(d)$</td>
<td>14</td>
<td></td>
</tr>
<tr>
<td>Diethanolnitraminedinitrate</td>
<td>DIINA</td>
<td>-78(a)</td>
<td>12</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>2,4-Dinitrotoluene</td>
<td>DNT</td>
<td>-17.1</td>
<td>13</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Ethylenedinitramine</td>
<td>EDNA or Haleite</td>
<td>-24.6</td>
<td>13</td>
<td>$\approx 1.3(d)$</td>
<td>14</td>
<td></td>
</tr>
<tr>
<td>Ammonium Picrate</td>
<td>ExplD</td>
<td>-92.7</td>
<td>13</td>
<td>$\approx 0.8(d)$</td>
<td>14</td>
<td></td>
</tr>
<tr>
<td>Bis (2,2-dinitro-2-fluoroethyl) formal</td>
<td>-</td>
<td>-178.8</td>
<td>11</td>
<td>1.28</td>
<td>11</td>
<td></td>
</tr>
<tr>
<td>Cyclotetramethylene tetramine</td>
<td>HMX</td>
<td>+17.93</td>
<td>12</td>
<td>1.48</td>
<td>11</td>
<td></td>
</tr>
<tr>
<td>Cyclotrimethylenetrimine</td>
<td>RDX</td>
<td>+14.71</td>
<td>12</td>
<td>1.42</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1.38(c)</td>
<td>9</td>
</tr>
<tr>
<td>Hydrazine Nitrate</td>
<td>HN</td>
<td>-60.5</td>
<td>4</td>
<td>0.88(e)</td>
<td>10</td>
<td></td>
</tr>
<tr>
<td>2,2',4,4',6,6'-Hexanitroazobenzene</td>
<td>HNAB</td>
<td>+57.8</td>
<td>17</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Hexanitrostilbene</td>
<td>HNS</td>
<td>+13.8</td>
<td>13</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Lead Azide</td>
<td>LA</td>
<td>+115.8</td>
<td>7</td>
<td>0.40</td>
<td>7</td>
<td></td>
</tr>
<tr>
<td>Lead Trinitroresorcinate</td>
<td>Lead</td>
<td>+204(a)</td>
<td>8</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Styphnate (LSt)</td>
<td>+107(a)</td>
<td>5</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Mercury Fulminate</td>
<td>MF</td>
<td>+65</td>
<td>14</td>
<td>$\approx 0.4(d)$</td>
<td>14</td>
<td></td>
</tr>
<tr>
<td>Nitrocellulose</td>
<td>NC</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>dinitro</td>
<td></td>
<td>-180(b)</td>
<td>1</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>trinitro</td>
<td></td>
<td>-156(b)</td>
<td>1</td>
<td>$\approx 1.1(d)$</td>
<td>14</td>
<td></td>
</tr>
<tr>
<td>Nitroglycerine (2)</td>
<td>NG</td>
<td>-88.6</td>
<td>13</td>
<td>1.48(e)</td>
<td>10</td>
<td></td>
</tr>
<tr>
<td>Nitroguanidine</td>
<td>NGu</td>
<td>-22.1</td>
<td>13</td>
<td>$\approx 0.7(d)$</td>
<td>14</td>
<td></td>
</tr>
<tr>
<td>Nitromethane (3)</td>
<td>NM</td>
<td>-27.03</td>
<td>13</td>
<td>1.23</td>
<td>11</td>
<td></td>
</tr>
<tr>
<td>Mannitol Hexanitrate</td>
<td>Nitromannite</td>
<td>-154(a)</td>
<td>12</td>
<td>$\approx 1.5(d)$</td>
<td>14</td>
<td></td>
</tr>
<tr>
<td>Pentaerythritotetranitrate</td>
<td>PETN</td>
<td>-128.7</td>
<td>13</td>
<td>1.49</td>
<td>11</td>
<td></td>
</tr>
<tr>
<td>Pentaerythritotrintrate</td>
<td>Petrin</td>
<td>-128(a)</td>
<td>13</td>
<td>$\approx 1.2(d)$</td>
<td>14</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>-133(a)</td>
<td>6</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>2,4,6-Trinitrophenol</td>
<td>Picric Acid (PA)</td>
<td>-51.3</td>
<td>13</td>
<td>1.00(c)</td>
<td>4</td>
<td></td>
</tr>
</tbody>
</table>
**Table 1 (Continued)**

**Standard Heats of Formation and Heats of Detonation of Common Explosives**

<table>
<thead>
<tr>
<th>EXPLOSIVE</th>
<th>Chemical Name</th>
<th>Code or Common Name</th>
<th>$\Delta H_f^\circ$ kcal/mole</th>
<th>Ref</th>
<th>$\Delta H_{det}$ kcal/g</th>
<th>Ref</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tetranitro-1,2,5,6-tetrazadibenzo-cyclooctatetrene</td>
<td>TACOT</td>
<td>+128</td>
<td>17</td>
<td>0.98(c)</td>
<td>17</td>
<td></td>
</tr>
<tr>
<td>1,3,5-Triamino-2,4,6-trinitrobenzene</td>
<td>TATB</td>
<td>-36.9</td>
<td>14</td>
<td>—</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td>Triethylene glycolinitrate</td>
<td>TEGDN</td>
<td>-151(a)</td>
<td>2</td>
<td>—</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td>2,4,6-Trinitrophenylmethylnitramine</td>
<td>Tetryl</td>
<td>+6(a)</td>
<td>13</td>
<td>1.14(c)</td>
<td>9</td>
<td></td>
</tr>
<tr>
<td>1,3,5-Trinitrobenzene</td>
<td>TNB</td>
<td>-8.9</td>
<td>15</td>
<td>—</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td>Tetranitromethane (C)</td>
<td>TKN</td>
<td>+8.9(a)</td>
<td>13</td>
<td>—</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td>2,2,2-Trinitroethyl-4,4,4-trinitrobutyrate</td>
<td>TNE</td>
<td>-118.6</td>
<td>4</td>
<td>1.47</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>2,4,6-Trinitrotoluene</td>
<td>TNT</td>
<td>-16.0</td>
<td>13</td>
<td>1.09</td>
<td>11</td>
<td></td>
</tr>
</tbody>
</table>

(a) Uncertain  
(b) Per monomer unit  
(c) Unconfined or lightly confined — these are really $\Delta H_{expl}$ and not $\Delta H_{det}$  
(d) Conditions of measurement are unknown  
(e) Computed; for these oxygen-balanced materials the computed $H_d$ should be nearly independent of the eqn of state of the deton products

Tables of heat data are presented below:

Table 2
Standard Heat of Formation and Heats of Detonation of Explosive Mixtures

<table>
<thead>
<tr>
<th>Compound</th>
<th>$\Delta H_f^\circ$ cal/g</th>
<th>Ref</th>
<th>$\Delta H_{det}$ kcal/g</th>
<th>Ref</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baratol</td>
<td>-708</td>
<td>1</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Comp B-3</td>
<td>+11</td>
<td>2</td>
<td>1.20(a)</td>
<td>1</td>
</tr>
<tr>
<td>Comp C-4</td>
<td>+33</td>
<td></td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>LX-04(b)</td>
<td>-215</td>
<td>1</td>
<td>1.31(a)</td>
<td>1</td>
</tr>
<tr>
<td>LX-11-0(c)</td>
<td>-307.3</td>
<td>1</td>
<td>1.23(d)</td>
<td>1</td>
</tr>
<tr>
<td>Octol</td>
<td>+26</td>
<td>1</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>PB9404(e)</td>
<td>+0.8</td>
<td>1</td>
<td>1.38(a)</td>
<td>1</td>
</tr>
<tr>
<td>Pentolite 50/50</td>
<td>-239</td>
<td>2</td>
<td>1.23(d)</td>
<td>1</td>
</tr>
<tr>
<td>XTX-8003(f)</td>
<td>-444</td>
<td>1</td>
<td>1.16(d)</td>
<td>1</td>
</tr>
</tbody>
</table>

(a) Moderate confinement
(b) 85/15 HMX/Viton
(c) 80/20 HMX/Viton
(d) Heavy confinement
(e) 94/3/3 HMX/NC/binder
(f) 80/20 PETN/Silicone rubber

Refs: 1) UCRL-51319 "Props of Chem Expls and Expl Simulants," Dec 1972 2) Algebraic addn of $\Delta H_f^\circ$'s of individual explosive compositions taken from Table 1
Table 3

Melting Points, Heats of Fusion, Heats of Vaporization and Heats of Sublimation of Common HE

<table>
<thead>
<tr>
<th>Explosive</th>
<th>mp(a) °C</th>
<th>ΔH\text{fus}(a) \text{kcal/mole}</th>
<th>ΔH\text{vap}(a) \text{kcal/mole}</th>
<th>ΔH\text{subl}(a) \text{kcal/mole}</th>
</tr>
</thead>
<tbody>
<tr>
<td>DATB</td>
<td>290(12)</td>
<td>–</td>
<td>–</td>
<td>33.5(10)</td>
</tr>
<tr>
<td>β HMX</td>
<td>286(16)</td>
<td>–</td>
<td>–</td>
<td>41.9(10)</td>
</tr>
<tr>
<td>HNS</td>
<td>318(15)</td>
<td>–</td>
<td>–</td>
<td>43.0(10)</td>
</tr>
<tr>
<td>NG</td>
<td>12.8(4)</td>
<td>6.0(5)</td>
<td>20.0(8)</td>
<td>–</td>
</tr>
<tr>
<td>Nitroform</td>
<td>15</td>
<td>3.6(9)</td>
<td>7.8(9)</td>
<td>11.1(9)</td>
</tr>
<tr>
<td>NMethane</td>
<td>-29</td>
<td>–</td>
<td>9.2(11)</td>
<td>8.4(8)</td>
</tr>
<tr>
<td>PETN</td>
<td>141(12)</td>
<td>–</td>
<td>23.0(1)</td>
<td>36.3(4)</td>
</tr>
<tr>
<td>Picric Acid</td>
<td>122(12)</td>
<td>4.7(12)</td>
<td>21.0(1)</td>
<td>–</td>
</tr>
<tr>
<td>RDX</td>
<td>205.5(13)</td>
<td>8.5(13)</td>
<td>26.0(1)</td>
<td>31.1(10)</td>
</tr>
<tr>
<td>TATB</td>
<td>330(12)</td>
<td>–</td>
<td>–</td>
<td>40.2(10)</td>
</tr>
<tr>
<td>Tetryl</td>
<td>126.5(13)</td>
<td>6.2(13)</td>
<td>26.0(1)</td>
<td>–</td>
</tr>
<tr>
<td>TNA(b)</td>
<td>188</td>
<td>–</td>
<td>–</td>
<td>27.7(10)</td>
</tr>
<tr>
<td>TNB(c)</td>
<td>121</td>
<td>4.0(14)</td>
<td>17.5(7)</td>
<td>–</td>
</tr>
<tr>
<td>TNT</td>
<td>81(12)</td>
<td>5.4(2)</td>
<td>17(1)</td>
<td>28.3(2)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5.1(12)</td>
<td>18.4(8)</td>
<td>24.4(3)</td>
</tr>
</tbody>
</table>

(a) Numbers in ( ) are Refs; underlined values are preferred
(b) 2, 4, 6-Trinitroaniline
(c) Sym-Trinitrobenzene

<table>
<thead>
<tr>
<th>Explosive</th>
<th>Coeff of Heat Conduction (cgs) x 10^4</th>
<th>temp (°C)</th>
<th>Specific Heat c_p (cal/deg g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baratol</td>
<td>11.8 at 2.5 g/cc (3)</td>
<td>75</td>
<td>0.280 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0</td>
<td>0.213 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>25</td>
<td>0.201 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>100</td>
<td>0.171 (6)</td>
</tr>
<tr>
<td>Comp B</td>
<td>6.3 at 1.71 g/cc (6)</td>
<td>75</td>
<td>0.235 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0</td>
<td>0.220 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>25</td>
<td>0.254 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>50</td>
<td>0.305 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>75</td>
<td>0.376 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>85</td>
<td>0.354 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>100</td>
<td>0.312 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>18-40</td>
<td>0.324 (7)</td>
</tr>
<tr>
<td>DNT</td>
<td>6.2 at 1.32 g/cc (5)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HMX</td>
<td></td>
<td>75</td>
<td>0.153 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0</td>
<td>0.228 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>25</td>
<td>0.248 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>50</td>
<td>0.266 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>100</td>
<td>0.295 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>150</td>
<td>0.315 (6)</td>
</tr>
<tr>
<td>β HMX</td>
<td></td>
<td>-70 to 250°C</td>
<td>0.0935 + 5x10^-4T (5)</td>
</tr>
<tr>
<td>Lead Azide</td>
<td>1.6 (5)</td>
<td>50 to 50</td>
<td>0.110 (6)</td>
</tr>
<tr>
<td></td>
<td>4 at 3.62 g/cc (2)</td>
<td>100</td>
<td>0.100 (2)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>200</td>
<td>0.117 (2)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>250</td>
<td>0.116 (2)</td>
</tr>
<tr>
<td>Lead Styphnate</td>
<td></td>
<td>-50</td>
<td>0.141 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0</td>
<td>0.158 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>25</td>
<td>0.164 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>50</td>
<td>0.167 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>100</td>
<td>0.164 (2)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>200</td>
<td>0.191 (2)</td>
</tr>
<tr>
<td>Mercury Fulminate</td>
<td>1 (5)</td>
<td>110 to 125</td>
<td>0.120 (2)</td>
</tr>
<tr>
<td></td>
<td>2.9 (1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NMethane</td>
<td>liq</td>
<td></td>
<td>0.414 (4)</td>
</tr>
<tr>
<td>NG</td>
<td>solid</td>
<td></td>
<td>0.315 (6)</td>
</tr>
<tr>
<td></td>
<td>liq</td>
<td></td>
<td>0.356 (6)</td>
</tr>
<tr>
<td>LX-04</td>
<td>9.3 at 1.86 g/cc (6)</td>
<td>20</td>
<td>≈0.28 (7)</td>
</tr>
<tr>
<td>PBX9404</td>
<td>10.1 at 1.83 g/cc (6)</td>
<td>-55</td>
<td>0.18 (3)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>-10</td>
<td>0.26 (3)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>22</td>
<td>0.24 (3)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>55</td>
<td>0.25 (3)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>85</td>
<td>0.28 (3)</td>
</tr>
<tr>
<td>PETN</td>
<td>6 at 1.46 g/cc (2)</td>
<td>115</td>
<td>0.272 (1)</td>
</tr>
</tbody>
</table>
Table 4 (Continued)
Heat Conductivity Coefficients and Specific Heats of Common Explosives

<table>
<thead>
<tr>
<th>Explosive</th>
<th>Coeff of Heat Conduction $\chi(\text{cal}) \times 10^4$</th>
<th>temp ($^\circ$C)</th>
<th>Specific Heat $c_p$ (cal/deg g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Picric Acid (PA)</td>
<td>6.2 at 1.41 g/cc (5)</td>
<td>0</td>
<td>0.235 (6)</td>
</tr>
<tr>
<td></td>
<td>2.4 at 1.7 (1)</td>
<td>30</td>
<td>0.258 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>60</td>
<td>0.282 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>120</td>
<td>0.337 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>?</td>
<td>0.26 (1)</td>
</tr>
<tr>
<td>RDX</td>
<td>6.9 at 1.26 g/cc (5)</td>
<td>20</td>
<td>0.298 (6)</td>
</tr>
<tr>
<td></td>
<td>7.0 at 1.53 g/cc (5)</td>
<td>60</td>
<td>0.330 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>100</td>
<td>0.406 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>140</td>
<td>0.446 (6)</td>
</tr>
<tr>
<td>Tetryl</td>
<td>5.8 at 1.39 g/cc (5)</td>
<td>-100</td>
<td>0.182 (6)</td>
</tr>
<tr>
<td></td>
<td>6.8 at 1.53 g/cc (5)</td>
<td>-50</td>
<td>0.200 (6)</td>
</tr>
<tr>
<td></td>
<td>2.3 at ? (1)</td>
<td>0</td>
<td>0.212 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>50</td>
<td>0.223 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>100</td>
<td>0.236 (6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>?</td>
<td>0.225 (1)</td>
</tr>
<tr>
<td>TNT</td>
<td>5.3 at 1.19 g/cc (5)</td>
<td>0</td>
<td>0.309 (6)</td>
</tr>
<tr>
<td></td>
<td>7.1 at 1.51 g/cc (5)</td>
<td>20</td>
<td>0.328 (6)</td>
</tr>
<tr>
<td></td>
<td>6.2 at 1.6 g/cc (6)</td>
<td>50</td>
<td>0.353 (6)</td>
</tr>
<tr>
<td></td>
<td>$\text{at } c^\circ$</td>
<td>80</td>
<td>0.374 (6)</td>
</tr>
<tr>
<td></td>
<td>4.8 at 1.6 $\text{liq}$</td>
<td>93 liq</td>
<td>0.376 (4)</td>
</tr>
<tr>
<td></td>
<td>3.5 at 0.8 $\text{liq}$</td>
<td>?</td>
<td>0.35 (1)</td>
</tr>
</tbody>
</table>

Refs: 1) A.F. Belyaev & N. Matyushko, Compt Rend URSS 30, 629 (1941) & CA 37, 531 (1943)
Heat Effects — Methods for Estimating $\Delta_H^o_f$. In the preceding sections we have emphasized the importance of knowing the heats of formation of explosive materials in order to estimate $\Delta_H^o_d$, detonation product compositions, fire & explosion hazards of potentially dangerous materials, and critical diameters for thermal explosions. This is by no means a complete list of the uses of heats of formation. Thus the $\Delta_H^o_f$ of a compound is indeed one of its most useful properties. However, how does one proceed if the $\Delta_H^o_f$ for a material of interest has not been determined experimentally? The obvious answer to this is by estimating $\Delta_H^o_f$ on the basis of theoretical or semi-empirical methods. Many such methods exist. The following article contributed by Dr. Robert Shaw of SRI presents a method that is particularly applicable to estimating the heats of formation of explosive compounds.

Estimation of Heats of Formation of Organic Chemical Explosives by Group Additivity. Introduction. During the past decade there have been significant developments of empirical methods for estimating the thermodynamic properties of organic compounds based on group additivity methods.

Advances in scientific and technological aspects of chemistry are dependent on advances in basic information. thermochemistry is important in our understanding of molecular structure including stabilities, strain energies, and interaction with radiation. It is also important in predicting reaction equilibria and yields, in process design, in thermal engineering applications, and in evaluating fire and explosion hazards. So far, the ideal gas state has received a large portion of thermochemists’ efforts. For example, the first page of the filing order for JANAF Thermochemical Data (Ref 6) contains 58 gases, 10 liquids, and 28 solids. This emphasis on gas phase stems partly from its more advanced theoretical status and partly from its importance in military problems. However, in the incoming requests for data at Stanford Research Institute from chemical engineers, we have had many more inquiries for condensed phases than for gases.

Estimated data have many advantages. Measurements are time-consuming and costly, and in many cases estimates are all that are needed for screening purposes. On the other hand, a minimum of good experimental data is required as a basis for estimates. Estimates are especially important in explosives thermochemistry, because so many of the experimental measurements are buried in difficult-to-reach technical reports and in the classified literature.

An important use of accurate estimates, such as those provided by Group Additivity, is that they provide a check on experimental measurements. In all cases where discrepancies have existed between measured data and those estimated by Group Additivity, repeating or checking the reported experimental data revealed errors. Some examples are given in Table 1.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Original Measurement</th>
<th>Estimated by Group Additivity</th>
<th>$\Delta$ (Meas.-Est.)</th>
<th>Revised Measurement</th>
<th>$\Delta$ (Meas.-Est.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-Methylcyclopentene</td>
<td>-1.5$^a$</td>
<td>-0.3$^a$</td>
<td>-1.2</td>
<td>0.6$^b$</td>
<td>0.2</td>
</tr>
<tr>
<td>Tetralin</td>
<td>2.6$^a$</td>
<td>6.6$^a$</td>
<td>-4.0</td>
<td>7.3$^b$</td>
<td>0.7</td>
</tr>
<tr>
<td>t-Butyl hydroperoxide</td>
<td>-52.1$^a$</td>
<td>-57.1$^a$</td>
<td>5.0</td>
<td>-58.8$^b$</td>
<td>-1.7</td>
</tr>
<tr>
<td>Hexanitrostilbene</td>
<td>29.2$^c$</td>
<td>58.6$^d$</td>
<td>29.4</td>
<td>57.0$^e$</td>
<td>1.6</td>
</tr>
</tbody>
</table>

$^a$Ref 14
$^b$Ref 15
$^c$Ref 8
$^d$Ref 20
$^e$Ref 9.
Estimating techniques point out where experimental data are lacking and where sufficient experimental data already exist. For example, Table II shows heats of formation that have been reported since the 1969 paper on Group Additivity was published (Ref 14). The agreement between experimental data and that predicted by Group Additivity is very gratifying, but it raises questions of cost/benefit regarding the selection of these compounds for experimental study.

Having discussed the need for estimated data, we turn now to the question, “why Group Additivity?” The most cogent answer is, user acceptance. In the present state of the art, there is a bewildering variety of methods of estimation, most of which are complicated, time-consuming, and of limited usefulness. The Group Additivity method is simple, fast, and accurate, and is quickly gaining wide acceptance as the best and most general method for estimation of gas-phase data. For example, ASTM Committee E-27 on the Hazard Potential of Chemicals has adopted Group Additivity as the method of choice for estimating thermochemical data. CIBA-GEIGY Inc has recently sent its Director of Explosion Hazard Research to SRI for a year to study estimating techniques with Dr. Benson, E.S. Domalski of the National Bureau of Standards has used Group Additivity methods to estimate the thermochemical properties of the air pollutants PAN (peroxycynoxtetralate/nitrate) and PBN (peroxynbenzylnitrate) (Ref 17). Dr. Benson has been invited by the new ACS journal, Chemical Technology, to write a paper entitled, “The Use of Empirical Methods for the Prediction of Thermochemical Kinetic Rate Data and the Uses of Limitations of Such Data in Laboratory Practice and in Plant Design.” The Environmental Protection Agency has recently sponsored a project at SRI to apply computer techniques to the estimation of thermochemical and kinetic data using Group Additivity. The McDonnell-Douglas Laboratories, sponsored by a contract with the Ballistic Research Laboratory, is developing a computer program to use Group Additivity for gases. Two papers at the 65th Annual Meeting of the American Institute of Chemical Engineers (“Thermophysical Properties of Pure Chemical Compounds,” by P.L. Chueh and C.H. Deal, and “Implementation of Combined Second Order Additivity and Group Equations Methods for the Estimation of Chemical Thermodynamic Data,” by W.H. Seaton and E. Freedman) dealt with applications of Group Additivity.

A powerful argument in favor of Group Additivity is that a start has already been made in the application of Group Additivity to condensed phases. In 1969, groups were derived (Ref 11) for the heat capacities of liquids at 298°K that improved the precision of estimation from ± 4 to better than ± 1.5 cal/(mole-K). Since then, groups have been derived for heats of formation of solid nitroaromatics (Ref 20) and for solid and liquid nitroalkanes (Ref 22) Group Additivity. The basic idea behind Group Additivity is that chemical thermodynamic properties of molecules consist of contributions from the individual groups that make up the molecule. Group Additivity is therefore an extension of the series atom additivity, bond additivity, . . . , and turns out to be an excellent compromise between simplicity and accuracy.

A useful way to look at it is to consider an example of Group Additivity applied to the ΔH_f° of gases at 25°C. The n-hexane molecule is made up of two methyls bonded to carbon atoms, indicated by 2[C-(C)H_3], and four methylenes bonded to two carbon atoms, indicated by 4[C-(C)_2H_4]. Similarly, n-octane is composed of 2[C-(C)H_3] + 6[C-(C)_2H_4]. Thus

\[ \Delta H_f°(\text{cal/mole}) \]

\[ 2[C-(C)H_3] + 4[C-(C)_2H_4] = -40.0 \]

\[ 2[C-(C)H_3] + 6[C-(C)_2H_4] = -49.8 \]

and \[ [C-(C)_2H_4] = -4.9 \] and \[ [C-(C)H_3] = -10.2 \]

The ΔH_f° of n-decane may then be estimated as 2[C-(C)H_3] + 8[C-(C)_2H_4] = -59.6 (cf -59.7 observed). The other groups were obtained similarly. In practice, when there is a large amount of measured data, the groups are derived by computer, using a least squares regression program, giving \[ [C-(C)H_3] = -10.1 \] and \[ [C-(C)_2H_4] = -5.0 \] kcal/mole.
Table II
Comparison of Estimated Heats of Formation with Some Measured Since 1969\(^a\)
(The units are kcal/mole for the ideal gas state.)

<table>
<thead>
<tr>
<th>Compound</th>
<th>Measured Since 1969</th>
<th>Estimated Using Group Values</th>
<th>(\Delta) (Meas.-Est.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>n-Pentane</td>
<td>-35.08</td>
<td>-35.10</td>
<td>0.02</td>
</tr>
<tr>
<td>2-Methylbutane</td>
<td>-36.73</td>
<td>-36.29</td>
<td>-0.44</td>
</tr>
<tr>
<td>2,2-Dimethylpropane</td>
<td>-40.14</td>
<td>-39.82</td>
<td>-0.32</td>
</tr>
<tr>
<td>n-Propylcyclohexane</td>
<td>-46.0</td>
<td>-46.3</td>
<td>-0.3</td>
</tr>
<tr>
<td>1-Methyl-1-ethylcyclohexane</td>
<td>-46.6</td>
<td>-45.8</td>
<td>-0.8</td>
</tr>
<tr>
<td>1-Methyl-cis-2-ethylcyclohexane</td>
<td>-45.7</td>
<td>-44.9</td>
<td>-0.8</td>
</tr>
<tr>
<td>1-Methyl-trans-2-ethylcyclohexane</td>
<td>-46.6</td>
<td>-46.5</td>
<td>-0.1</td>
</tr>
<tr>
<td>1-Methyl-cis-3-ethylcyclohexane</td>
<td>-48.2</td>
<td>-48.1</td>
<td>0.1</td>
</tr>
<tr>
<td>1-Methyl-cis-4-ethylcyclohexane</td>
<td>-46.3</td>
<td>-46.5</td>
<td>0.2</td>
</tr>
<tr>
<td>1-Methyl-trans-4-ethylcyclohexane</td>
<td>-49.1</td>
<td>-49.1</td>
<td>0.0</td>
</tr>
<tr>
<td>n-Propylcyclopentane</td>
<td>-35.3</td>
<td>-34.6</td>
<td>-0.7</td>
</tr>
<tr>
<td>3-Methylcyclopentene</td>
<td>2.0</td>
<td>1.8</td>
<td>0.2</td>
</tr>
<tr>
<td>Acenaphthene</td>
<td>37.4</td>
<td>36.8</td>
<td>0.6</td>
</tr>
<tr>
<td>Acenaphthalene</td>
<td>61.6</td>
<td>60.8</td>
<td>0.8</td>
</tr>
<tr>
<td>Cyclopropylacene</td>
<td>18.4</td>
<td>17.3</td>
<td>1.1</td>
</tr>
<tr>
<td>2-Methyl-1-butanethiol</td>
<td>-27.42</td>
<td>-27.24</td>
<td>-0.18</td>
</tr>
<tr>
<td>3-Methyl-1-butanethiol</td>
<td>-27.40</td>
<td>-27.24</td>
<td>-0.16</td>
</tr>
<tr>
<td>3-Methyl-2-butanethiol</td>
<td>-28.91</td>
<td>-29.36</td>
<td>0.45</td>
</tr>
<tr>
<td>2,2-Dimethyl-1-propanethiol</td>
<td>-30.76</td>
<td>-30.77</td>
<td>0.01</td>
</tr>
<tr>
<td>2,3-Dimethyl-2-butanethiol</td>
<td>-35.22</td>
<td>-36.55</td>
<td>1.33</td>
</tr>
<tr>
<td>2-Methyl-2-pentanethiol</td>
<td>-35.37</td>
<td>-35.17</td>
<td>-0.20</td>
</tr>
<tr>
<td>o-Toluene thiol</td>
<td>21.90</td>
<td>21.80</td>
<td>0.00</td>
</tr>
<tr>
<td>Diethyl ketone</td>
<td>-61.7</td>
<td>-62.0</td>
<td>0.3</td>
</tr>
<tr>
<td>Methyl-n-propyl ketone</td>
<td>-61.9</td>
<td>-61.7</td>
<td>0.2</td>
</tr>
<tr>
<td>Methyl-isopropyl ketone</td>
<td>-62.8</td>
<td>-63.3</td>
<td>0.5</td>
</tr>
<tr>
<td>Methyl-n-butyl ketone</td>
<td>-66.9</td>
<td>-66.7</td>
<td>-0.2</td>
</tr>
<tr>
<td>Methyl-t-butyl ketone</td>
<td>-69.5</td>
<td>-70.3</td>
<td>0.8</td>
</tr>
<tr>
<td>Ethyl-n-propyl ketone</td>
<td>-66.5</td>
<td>-66.9</td>
<td>0.4</td>
</tr>
<tr>
<td>Compound</td>
<td>Measured Since 1969</td>
<td>Estimated Using Group Values</td>
<td>Δ (Meas.-Est.)</td>
</tr>
<tr>
<td>--------------------------------</td>
<td>---------------------</td>
<td>------------------------------</td>
<td>---------------</td>
</tr>
<tr>
<td>Ethyl-isopropyl ketone</td>
<td>-68.4</td>
<td>-68.5</td>
<td>0.1</td>
</tr>
<tr>
<td>Ethyl-t-butyl ketone</td>
<td>-75.0</td>
<td>-75.5</td>
<td>0.5</td>
</tr>
<tr>
<td>Diisopropyl ketone</td>
<td>-74.4</td>
<td>-75.1</td>
<td>0.7</td>
</tr>
<tr>
<td>Isopropyl-t-butyl ketone</td>
<td>-80.8</td>
<td>-82.1</td>
<td>1.3</td>
</tr>
<tr>
<td>Di-n-butyl ketone</td>
<td>-82.4</td>
<td>-81.8</td>
<td>-0.6</td>
</tr>
<tr>
<td>Di-t-butyl ketone</td>
<td>-82.6</td>
<td>-89.1</td>
<td>6.5</td>
</tr>
<tr>
<td>Methyl-hexyl ketone</td>
<td>-82.5</td>
<td>-76.6</td>
<td>-5.9</td>
</tr>
<tr>
<td>Ethyl-pentyl ketone</td>
<td>-80.9</td>
<td>-76.8</td>
<td>-4.1</td>
</tr>
<tr>
<td>Propylbutyl ketone</td>
<td>-83.5</td>
<td>-76.8</td>
<td>-6.7</td>
</tr>
<tr>
<td>Di-isobutyl ketone</td>
<td>-85.5</td>
<td>-84.3</td>
<td>-1.2</td>
</tr>
<tr>
<td>t-Butyl-neopentyl ketone</td>
<td>-94.2</td>
<td>-94.0</td>
<td>-0.2</td>
</tr>
<tr>
<td>Di-n-pentyl ketone</td>
<td>-92.6</td>
<td>-91.7</td>
<td>-0.9</td>
</tr>
<tr>
<td>Acrolein</td>
<td>-17.8</td>
<td>-17.8</td>
<td>0.0</td>
</tr>
<tr>
<td>But-2-enal</td>
<td>-25.5</td>
<td>-25.6</td>
<td>0.1</td>
</tr>
<tr>
<td>Benzaldehyde</td>
<td>-8.9</td>
<td>-8.9</td>
<td>0.0</td>
</tr>
<tr>
<td>3,5-Dioxaheptane</td>
<td>-90.1</td>
<td>-98.8</td>
<td>-0.3</td>
</tr>
<tr>
<td>1,1-Diethoxyethane</td>
<td>-108.4</td>
<td>-109.2</td>
<td>0.8</td>
</tr>
<tr>
<td>3,6-Dioxaoctane</td>
<td>-97.6</td>
<td>-99.0</td>
<td>1.4</td>
</tr>
<tr>
<td>2,4,6-Trimethyl-3,5-dioxaheptane</td>
<td>-125.7</td>
<td>-126.4</td>
<td>0.7</td>
</tr>
<tr>
<td>6-Ethyl-5,7-dioxaundecane</td>
<td>-132.1</td>
<td>-133.9</td>
<td>1.8</td>
</tr>
<tr>
<td>3,5,7-Trioxanonane</td>
<td>-138.9</td>
<td>-138.2</td>
<td>-0.7</td>
</tr>
<tr>
<td>3,5,7,9-Tetraoxaundecane</td>
<td>-177.1</td>
<td>-177.5</td>
<td>0.4</td>
</tr>
<tr>
<td>3,5,7,9,11-Pentaoxastridecane</td>
<td>-216.5</td>
<td>-216.8</td>
<td>0.3</td>
</tr>
<tr>
<td>Propionaldehyde</td>
<td>-44.5</td>
<td>-44.4</td>
<td>-0.1</td>
</tr>
<tr>
<td>Dimethoxyethane</td>
<td>-83.2</td>
<td>-82.7</td>
<td>-0.5</td>
</tr>
<tr>
<td>1,1-Dimethoxyethane</td>
<td>-93.1</td>
<td>-92.9</td>
<td>-0.2</td>
</tr>
</tbody>
</table>

*The hydrocarbon and amine data are from Ref 19, the sulfur data are from Ref 25, and the data for oxygenated compounds are from Ref 21.*
For a second example, from Stull, Westrum, and Sinke, (Ref 12) for n-pentane $\Delta H_f^o$(liquid) = -41.4 kcal/mole, and for n-hexane, $\Delta H_f^o$(liquid) = -47.5 kcal/mole. n-Hexane has one more

$[\text{C}(\text{C})_2(\text{H})_4]^g = -47.5 - (-41.4) = -6.1$ (where the subscript g denotes liquid phase). Substituting the above value of -6.1 we obtain the methyl group $[\text{C}(\text{C})(\text{H})_3]^g = [-47.5 - 4(-6.1)]/2 = [-47.5 + 24.4]/2 = -23.1/2 = -11.6$

Check: $\Delta H_f^n$-heptane $^g = 2[\text{C}(\text{C})(\text{H})_3]^g + 5[\text{C}(\text{C})_2(\text{H})_4]^g = 2(-11.6) + 5(-6.1) = -23.2 - 30.5 = -53.7$ kcal/mole, cf -53.6 measured (Ref 12). As an added bonus, we obtain groups for estimating heats of vaporization. From previous work (Ref 14) we know that for the ideal gas value for the group $[\text{C}(\text{C})(\text{H})_3]^g$ is -10.1 kcal/mole (where the subscript g denotes ideal gas). Therefore, $[\text{C}(\text{C})(\text{H})_3]^vap = [\text{C}(\text{C})(\text{H})_3]^g - [\text{C}(\text{C})(\text{H})_3]^g = -10.1 - (-11.6) = 1.5$ kcal/mole.

The heat of vaporization of n-heptane at 298 K = 2$[\text{C}(\text{C})(\text{H})_3]^vap + 5[\text{C}(\text{C})_2(\text{H})_4]^vap = 2(1.5) + 5(1.1) = 3.0 + 5.5 = 8.5$ kcal/mole, cf. 8.7 measured (Ref 12).

To date, group values have been derived for heats of formation of the states and classes of compounds shown in Table III.

<table>
<thead>
<tr>
<th>State</th>
<th>Class of Compound</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ideal gas</td>
<td>Organics in general</td>
<td>14</td>
</tr>
<tr>
<td>Solids</td>
<td>Nitroaromatics</td>
<td>20</td>
</tr>
<tr>
<td>Solids, liquids,</td>
<td>Nitroalkanes</td>
<td>22</td>
</tr>
<tr>
<td>Ideal gas</td>
<td>Oxygen-containing</td>
<td></td>
</tr>
<tr>
<td></td>
<td>organics</td>
<td>21</td>
</tr>
<tr>
<td>Ideal gas</td>
<td>Azo. compounds</td>
<td>26</td>
</tr>
</tbody>
</table>

Heats of Formation of Solid Nitroaromatics. In principle, there are two methods of using group additivity to determine heats of formation of solids. If the groups are available, then the heat of formation of the ideal gas can be calculated. If the heat of sublimation is known, then the heat of formation of the solid follows. The second method is to develop group values for solids and to use these directly to calculate the required heat of formation. The heats of formation of some solid nitroaromatic compounds present an interesting example of the two approaches. The results suggest that the direct approach of using group values for solids is the better method at present. The Ideal Gas Method. The heat of formation of a nitroaromatic compound in the ideal gas state requires that the C6H5NO2 group be known (where C6 represents an aromatic, i.e., benzene carbon atom). This group was not given in the recent ideal gas group additivity review (Ref 14). The group may be obtained from the heat of formation of any nitroaromatic compound in the ideal gas state if the other groups are known. For example, for nitrobenzene $\Delta H_f^o = 5$C6H5 + 1(C6H4NO2) = 16.9 kcal/mole (Table IV).

From the known value of (C6H5) = 3.3 kcal/mole, (C6H4NO2) = 16.9 - 16.5 = 0.4 kcal/mole.

In Table IV the heats of formation of several nitroaromatic compounds in the ideal gas state are calculated from the measured heats of formation of the solid and the measured heats of sublimation. In cases where the heat of sublimation is not measured at 298 K there should be a correction for the differences in heat capacities of the solid and ideal gas. The data required to make these corrections are not available but in general it is expected that the corrections will be small and can be neglected. From the heats of formation of each compound in the ideal gas state, a value for the group C6H5NO2 (ideal gas) has been derived (Table IV). A weighted-average value (C6H5NO2(ideal gas)) = 3.0 kcal/mole was used, and a heat of formation was estimated for each compound. In Table IV the difference between observed and estimated heat of formation in the ideal gas state is less than +3.2 kcal/mole in all cases.
## Measured and Estimated Heats of Formation of Various Compounds at 298 K in Kcal/mole

**Table 1V**

<table>
<thead>
<tr>
<th>Compound</th>
<th>Measured</th>
<th>Estimated</th>
<th>Heat of</th>
<th>Molar</th>
<th>Mass</th>
<th>Formula</th>
<th>Heat of</th>
<th>Estimated</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Formation</td>
<td></td>
<td></td>
<td></td>
<td>Formation</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

- "H N" denotes "Heat of"
The Group Additivity for Solids Method. The large number of compounds for which the heat of formation of the solid has been measured makes the group additivity for solids an attractive method. From the measured data (Table IV), groups were derived by inspection. The values for the groups are (C\(_6\)-H) = 0, (C\(_6\)-NO\(_2\)) = -3, (C\(_6\)-NH\(_3\)) = -9, and (C\(_6\)-CH\(_3\)) = -6 kcal/mole. If the groups had been derived statistically, there would probably be a slight improvement, but the convenience of having (C\(_6\)-H) = 0 and integral values for the other groups would be lost in the process. Using these groups, the heats of formation in the solid state for a number of compounds have been calculated. The differences between observed and estimated values are small and in the range expected for group additivity

Conclusion. In general, the fit between observed and estimated values suggests that the principle of group additivity can be successfully applied to the heats of formation of solids. At present this approach is more convenient than using ideal gas groups together with heats of sublimation

Heats of Formation of Nitroalkanes. The heats of formation of several liquid and solid poly-nitroalkanes have recently been measured by Lebedeva and Ryadnenko (Ref 10). Their results support and extend some earlier work by Holcomb and Dorsey (Ref 2) and by Ménard and Thomas (Ref 5). Data for the liquid and gaseous mononitroalkanes have been evaluated by Stull, Westrum, and Sinke (Ref 12). JANAF (Ref 6) have reported some unpublished results by Shomate. All the above results are listed in Table V with estimates based on group additivity using the group values derived by us and listed in Table VI. The agreement between observed and estimated values is very good with a few exceptions. The most spectacular exception is 1,1,3,5,5-Heptanitropentane where, even allowing for 6 alkyl gauche interactions, the compound is made unstable by about 14 kcal/mole, perhaps because of NO\(_2\) crowding

Another anomaly is found for the 1,1,1-fluorodinitroalkanes. The experimental ΔH\(_f\) for 1,1,1-DNE and 1,1,1-DNP both give values for the group C-(C)(NO\(_2\))\(_2\)(F) which differ by 9 kcal/mole. For lack of definite information, the difference is split equally in setting the group value. However, the following comparison suggests that the experimental ΔH\(_f\) for 1,1,1-DNE is suspect:

\[
\begin{align*}
\text{EtH} + \text{F} + \text{EtF} + \text{H} &+ 9.1 \text{ kcal/mole} \\
n\text{-PrH} + \text{F} + n\text{-PrF} + \text{H} &+ 9.2 \text{ kcal/mole} \\
1,1\text{-DNE} + \text{F} &+ 1,1,1\text{-FDNE} + \text{H} - 1.3 \text{ kcal/mole} \\
1,1\text{-DNP} + \text{F} &+ 1,1,1\text{-FDNP} + \text{H} + 7.3 \text{ kcal/mole}
\end{align*}
\]

More work is needed on the experimental values of ΔH\(_f\) for the 1,1,1-fluorodinitroalkanes
<table>
<thead>
<tr>
<th>Table V</th>
<th>Heats of Formation of Nitroalkanes</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>[SOLID]</td>
</tr>
<tr>
<td></td>
<td>gauche corrections</td>
</tr>
<tr>
<td>1-NITROALKANES</td>
<td></td>
</tr>
<tr>
<td>Nitroethane</td>
<td>33.912</td>
</tr>
<tr>
<td>1-Nitropropane</td>
<td>40.312</td>
</tr>
<tr>
<td>1-Nitrobutane</td>
<td>46.012</td>
</tr>
<tr>
<td>2-NITROALKANES</td>
<td></td>
</tr>
<tr>
<td>2-Nitropropane</td>
<td>12</td>
</tr>
<tr>
<td>2-Nitrobutane</td>
<td>43.312</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>1,1-DINITROALKANES</td>
<td></td>
</tr>
<tr>
<td>1,1-Dinitroethane</td>
<td>1-alk-NO₂</td>
</tr>
<tr>
<td>1,1-Dinitropropane</td>
<td>1-alk-NO₂</td>
</tr>
<tr>
<td>1,1-Dinitrobutane</td>
<td>1-alk-NO₂</td>
</tr>
<tr>
<td>1,1,1-FLUORODINITROALKANES</td>
<td></td>
</tr>
<tr>
<td>1,1,1-Fluorodinitroethane</td>
<td>1-alk-NO₂</td>
</tr>
<tr>
<td>1,1,1-Fluorodinitropropane</td>
<td>1-alk-NO₂</td>
</tr>
<tr>
<td>ALPHA-Omega DINITROALKANES</td>
<td></td>
</tr>
<tr>
<td>1,2-Dinitroethane</td>
<td>42.710</td>
</tr>
<tr>
<td>1,3-Dinitropropane</td>
<td>-51.3</td>
</tr>
<tr>
<td>1,4-Dinitrobutane</td>
<td>-59.610</td>
</tr>
<tr>
<td>PRIMARY-SECONDARY DINTrOALKANES</td>
<td></td>
</tr>
<tr>
<td>1,2-Dinitropropene</td>
<td>1-alk-NO₂</td>
</tr>
<tr>
<td>2,2-DINITROALKANES</td>
<td></td>
</tr>
<tr>
<td>2,2-Dinitropropane</td>
<td>46.010</td>
</tr>
<tr>
<td>2,3,3,3-Tetranitrobutane</td>
<td>4 alk-NO₂</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>1,1,1-TRINITROALKANES</td>
<td></td>
</tr>
<tr>
<td>1,1,1-Trinitroethane</td>
<td>6 NO₂-NO₃</td>
</tr>
<tr>
<td>Hexanitroethane</td>
<td>6 NO₂-NO₃</td>
</tr>
<tr>
<td>1,1,1-Trinitropropane</td>
<td>2 alk-NO₂</td>
</tr>
<tr>
<td>1,1,1,3,3,3-Tetranitropropane</td>
<td>2 alk-NO₂</td>
</tr>
<tr>
<td>1,1,1,4,4,4-Tetranitrobutane</td>
<td>2 alk-NO₂</td>
</tr>
<tr>
<td>1,1,1,3,3,5,5-Heptanitropropane</td>
<td>6 alk-NO₂</td>
</tr>
<tr>
<td>TERTIARY NITROALKANES</td>
<td></td>
</tr>
<tr>
<td>2,3-Dimethyl-2,3-dinitrobutane</td>
<td>6 alk-NO₂</td>
</tr>
<tr>
<td>NITROALKANES</td>
<td></td>
</tr>
<tr>
<td>Nitromethane</td>
<td></td>
</tr>
<tr>
<td>1-Nitropropylne</td>
<td></td>
</tr>
<tr>
<td>2-Nitropropionyl</td>
<td></td>
</tr>
</tbody>
</table>

[a] Calculated from data in reference 2
[b] Square brackets indicate that the compound was the sole source of this group, so that there is necessarily a fit between observed and estimated data
<table>
<thead>
<tr>
<th>Group</th>
<th>Solid</th>
<th>Liquid</th>
<th>Ideal</th>
</tr>
</thead>
<tbody>
<tr>
<td>C-(C)H$_2$(NO$_2$)</td>
<td>-22.2</td>
<td>-21.5</td>
<td>-14.4</td>
</tr>
<tr>
<td>C-(C)$_2$(H)NO$_2$</td>
<td>(-21)$^a$</td>
<td>-21.2</td>
<td>-13.6</td>
</tr>
<tr>
<td>C-(C)$_2$(NO$_2$)</td>
<td>-17</td>
<td>-18.3</td>
<td></td>
</tr>
<tr>
<td>C-(C)(H)(NO$_2$)$_2$</td>
<td>-24.0</td>
<td>-9.9</td>
<td></td>
</tr>
<tr>
<td>C-(C)(H)(NO$_2$)$_2$</td>
<td>-60.2</td>
<td>-46.9</td>
<td></td>
</tr>
<tr>
<td>C-(C)$_2$(NO$_2$)$_2$</td>
<td>-21.2</td>
<td>-21.0</td>
<td>-10.2</td>
</tr>
<tr>
<td>C-(C)(NO$_2$)$_3$</td>
<td>-13.6</td>
<td>-13.0</td>
<td></td>
</tr>
<tr>
<td>alk-NO$_2$ gauche</td>
<td>2</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>NO$_2$-NO$_2$ gauche</td>
<td>9</td>
<td>9</td>
<td></td>
</tr>
<tr>
<td>C-(C)(H)$_2$</td>
<td>-13.5$^b$</td>
<td>-11.8$^c$</td>
<td>-10.1$^d$</td>
</tr>
<tr>
<td>C-(C)$_2$(H)$_2$</td>
<td>-6.85$^b$</td>
<td>-6.1$^c$</td>
<td>-5.0$^d$</td>
</tr>
<tr>
<td>C$_d$-(H)(NO$_2$)$^a$</td>
<td></td>
<td>7.1$^e$</td>
<td></td>
</tr>
<tr>
<td>C$_d$-(C)(NO$_2$)$^a$</td>
<td></td>
<td>4.4$^e$</td>
<td></td>
</tr>
</tbody>
</table>

$^a$Based on liquid phase group
$^b$Based on nC$_{18}$H$_{38}$ and nC$_{32}$H$_{66}$ (Ref 12)
$^c$Based on nC$_3$H$_{12}$ and nC$_6$H$_{14}$ (Ref 12)
$^d$Ref 14
$^e$Estimated from kinetics (Ref 22)
### Table VII
Heats of Formation (in kcal/mole) of Azo Compounds Measured by Engel and Wood (Ref 23) and Estimated by Group Additivity

<table>
<thead>
<tr>
<th>Compound</th>
<th>$\Delta H^\circ_{\text{f liquid}}$</th>
<th>$\Delta H_{\text{vap}}$</th>
<th>$\Delta H^\circ_{\text{f gas}}$</th>
<th>$\Delta H^\circ_{\text{f gas}}$</th>
<th>Difference</th>
<th>obs</th>
<th>est</th>
<th>obs - est</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\left[\text{CH}_2\text{CH}_2\text{CH}_2\text{N}\right]_2$ trans</td>
<td>4.0</td>
<td>8.0</td>
<td>12.0</td>
<td>12.0</td>
<td>[0]</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\left[(\text{CH}_3)_2\text{CN}\right]_2$ trans</td>
<td>0.4</td>
<td>8.0</td>
<td>8.4</td>
<td>8.4</td>
<td>[0]</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>cis</td>
<td>0.4</td>
<td>10.0</td>
<td>10.4</td>
<td>9.4</td>
<td>1.0</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\left[(\text{CH}_3)_2\text{CN}\right]_2$ trans</td>
<td>-18.4</td>
<td>8.8</td>
<td>-9.6</td>
<td>-9.6</td>
<td>[0]</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

![Diagram 1](image1.png)

![Diagram 2](image2.png)

![Diagram 3](image3.png)

![Diagram 4](image4.png)
A small but mathematically significant deviation from group additivity is shown by 1,2-dinitroethane in the liquid phase. The difference of 3.3 kcal/mole between the observed and estimated values is too large and cannot reasonably be reduced by altering the group value. An experimental value of the heat of vaporization of 1,2-dinitroethane is needed. Azo Compounds in Ideal Gas State. As Benson and Walsh have commented (Ref 14), "(Thermoch- emical) information on azo compounds is particularly lacking." The only azo compounds for which experimentally determined heats of formation have been published are difluoro diazine FNNF, diimide HNNH, and azoisopropene (CH₂)₂CHNNCH(CH₂)₂. However, Engel and Wood (Ref 23) recently measured the heats of combustion and vaporization of a large group of azo compounds. Their results are summarized in Table VII.

Engel and Wood's value for the heat of formation of azoisopropene is 11 kcal/mole less stable than Coates and Sutton's 1948 value (Ref 1) of 19.4 kcal/mole. A consequence of this change is that ΔHf° of azomethane calculated from graph values (Table VIII) becomes 34.0 kcal/mole instead of the previous value (Ref 6) of 44.7 kcal/mole. A further consequence is that the heat of hydrogenation of azomethane then becomes 12.4 kcal/mole, which is very different from the heat of hydrogenation of diimide (27.4 kcal/mole). However, Engel and Wood's data are very self-consistent as shown by the reasonable agreement between their experimental results and those calculated by group additivity for the cyclic compounds (Table VIII). Therefore the more recent work by Engel and Wood is favored.

### Table VIII.

<table>
<thead>
<tr>
<th>Group</th>
<th>ΔHf° (kcal/mole)</th>
</tr>
</thead>
<tbody>
<tr>
<td>N₂A⁻¹(N₂A⁻¹)(C)</td>
<td>56.4</td>
</tr>
<tr>
<td>N₂A⁻¹(C)</td>
<td>27.0</td>
</tr>
<tr>
<td>C⁻¹(N₂A⁻¹)(H)₃</td>
<td>-10.1</td>
</tr>
<tr>
<td>C⁻¹(N₂A⁻¹)(C)(H)₂</td>
<td>-6.0</td>
</tr>
<tr>
<td>C⁻¹(N₂A⁻¹)(C)₂(H)</td>
<td>-3.4</td>
</tr>
<tr>
<td>C⁻¹(N₂A⁻¹)(C)₃</td>
<td>-3.0</td>
</tr>
</tbody>
</table>

*a*This table was compiled from data in references 23 and 18 and after considerable discussion with S.W. Benson. There are some typographical errors on p 32 of O'Neal and Benson's monograph (Ref 15). The groups [C⁻¹(N)(H)₂]₁, [C⁻¹(N)(H)(C)]₁, and [C⁻¹(N)(C)₂]₁ in the third column of Table V-8 are all missing a carbon atom. They should read [C⁻¹(C)(N)(H)₂]₁, [C⁻¹(N)(H)(C)₂]₁, and [C⁻¹(N)(C)₃]₁. Also in the text DH₁° - DH₂° should read DH₁° + DH₂°.

**Conclusions.** Group additivity, an established technique for estimating the heats of formation, has been well developed for application to organic chemicals, in general, in the ideal gas state. However, very little has been done in deriving the group values for the many classes of organic explosives in condensed phases, especially the solid state. The principles are there; it is only a case of doing the work.

Written by R. Shaw

Heat of Evaporation. See under Heat, Latent

Heat Evolution (or Liberation) in Explosives. For a detonation to be self-sustaining the detonating explosive must liberate a considerable amount of heat. If energy is lost to the surroundings the detonation characteristics are altered, and if sufficient energy is lost the detonation fails. For a steady detonation, the energy evolved in the detonation reactions must obviously be greater than the energy lost to the surroundings.

[See Vol 4 of Encycl, under Detonation (and Explosion), Heat of]

Heat of Formation. See Vol 4 of Encycl under DETONATION (AND EXPLOSION), DEFLAGRATION (AND COMBUSTION), AND FORMATION, HEATS OF & TABLES 1 & 2 this Vol

Heat of Fusion. See under Heat, Latent and Table 3 of this Vol

Heat of Hydration is the amount of heat absorbed or liberated when a substance combines with water

Heat, Latent ("Latent" means "lying hidden" or "not manifest"). It is the heat absorbed by a substance without causing a rise in temperature.

There are two kinds of latent heat:

a) Latent heat of fusion is the amount of heat necessary to transform (melt, fuse) a unit quantity of solid into a liquid at the same temperature and standard pressure. Tables usually express it in calories per gram.

b) Latent heat of vaporization (or evaporation) (L) is the quantity of heat required to convert a unit quantity of liquid to vapor at the same temperature. Tables usually express it in calories per gram. (L) can also be calculated, approximately, by the Trouton's rule: L = CT/M, where (C) is a constant equal to 21 for most liquids, (T) boiling point in absolute degrees and (M) molecular weight. Example: L for acetone = 21 x (56 + 273)/58 = 120 cal/g against 124 actually measured

Heat, Molecular is the amount of heat required to raise the temperature of one mol of a substance, 1°C, ie, the specific heat of the substance multiplied by its molecular weight.

Heat Pulse. (Also see Detonation, Flash-Across, Heat Pulse and Hypervelocity Phenomena in Vol 4, p D348-49). A concept advanced by M.A. Cook (Refs 1 & 2) to provide a theoretical mechanism for the shock initiation of explosives. Cook also used the heat pulse concept in his explanation of certain unusual luminosity effects observed primarily in the detonation of liquid explosives. Briefly stated, Cook believes that detonation is initiated when a result of rising temperature, produced by reaction in the already shocked region of an explosive, a portion of the explosive becomes thermally "super-conductive" and a "heat-pulse" flashes thru it and catches up with the shock front. Studies conducted by Kendrew & Whitbread (Ref 3) tend to discount the necessity for postulating a "heat-pulse" in a theoretical explanation of shock initiation or the above unusual luminosity effects. More recent studies of shock initiation have also failed to produce any conclusive evidence of a "heat-pulse" (Ref 4).

Refs: 1) Cook (1958) pp 83-89 & 164-166
2) M.A. Cook et al, Proc Third Symposium on Deton (1960), pp 150-183
3) E.L. Kendrew and E.G. Whitbread, Ibid, pp 202-204 & pp 574-583
4) Several papers on shock initiation in the Fifth Symp on Deton (1970)

Heat Sensitization of Explosives and Memory Effect. See under Detonation, Heat Sensitization and Memory Effect in Vol 4, p D367-R.


Heat of Solution. Same as Heat of Dissolution.

Heat of Sublimation is the number of calories required, in a constant temperature process, to convert a unit weight of solid directly into a vapor.

Heat of Swelling is the heat evolved when a colloid (eg gelatin, NC) absorbs a liquid (such as H₂O, acetone etc).

Heat of Transition is the quantity of heat liberated or absorbed when a substance changes from one allotropic crystal form to another.

Heat Tests. Under this term may be placed all the stability tests which involve heating an explosive, eg Abel Heat Test, Acidity Measurements (pH measurements), American Test at 65.5°C or 80°C, Bergmann-Junk Test, Brame's Test, Brunswig's Test, Chiaraviglio & Corbino Test, Conductivity Method (De Bruin & de Pauw), Continuous Test, Deflagration Test, Desmaroux Test, Dupre's Vacuum Test, Dutch Test, German 132°C Test, Grottanelli's Test, Guttmann Test, Haid, Becker & Dittmar Test, Hansen's Test, Hess Test, Hoitsma Test, Horn-Soefert Test, International 75°C Test, Jensen's Test, Loss of Weight Test, Marqueyrol's Test, Meerscheidt-Hüllessem Test, Mittasch's Test, Obermuller's Test, Pavlik's Test, Methyl Violet Test, Silvered Vessel Test, Simon Thomas Test, Spica Test, Sy Test, Taliani Test, Taylor's Test, Tomanari's Test, Vacuum Stability Test, Vieelle Test, Warmlagermethode 75°C (German Storage Test), Will Test.

These tests are described separately in alphabetical order.

Heat Value is the number of calories obtained by the complete combustion of a unit weight of substance.

Heat of Vaporization. See under Heat, Latent.

Heath and Frost patented in 1877 the use of a water-retaining paste such as soap 5, starch 0.5, glue 0.5 & water 94% for tamping bore-holes, loaded with dynamite, in order to prevent the ignition of firedamp.

Ref: Daniel (1902), 371.

Hebler Powder or Wellite. Smokeless powder used in Switzerland before the introduction of NC smokeless powders. It contains KNO₃ 62.3, NH₄NO₃ 15.8, charcoal 11.5, sulfur 9.5, moisture 0.9%. Use of this powder was abandoned because it is too hygroscopic.

**Heda Powder.** An American Dynamite containing 20 to 75% NG absorbed by wood pulp, sodium nitrate etc
*Ref:* Daniel (1902), 371

**Hedgehog.** A British antisubmarine device for projecting 24 spigot type projectiles equipped with contact fuzes. They contain approx 30 lbs of HE each

**HEI.** Abbreviation for High Explosive Incendiary

**Height of Burst of Ground Type Pyrotechnic Signals, Determination.** See under Position Finders

**HEIT.** Abbreviation for High Explosive Incendiary Tracer

**Heliophanite.** See under Pandacolites

**Hellhoff Explosive.** According to GerP12122 of 1880, it was prepared by the nitration of purified-tar oil, followed by washing, drying and mixing of the nitroaromatic with oxygen carriers, such as K (or Na) nitrate (or chlorate), etc. It was claimed that this explosive mixture was very powerful
*Ref:* See under Hellhoffit

**Hellhoffit (Hellhoffite).** One of the Sprengel type explosives, invented about 1870 by Hellhoff and Grüson. It consisted of 28 parts of nitrobenzene and 72 parts of fuming nitric acid. This liquid was sometimes used absorbed on kieselguhr (See Gehrrottoffit). The disadvantage of these Sprengel type explosives was their extreme corrosiveness (Ref 1)

According to Thorpe (Ref 2), Hellhoffit was tried in shells, the two ingredients being mixed during flight & exploded on impact (see also Anilite under French explosives)

Stettbacher (Refs 3 and 4) investigated Hellhoffit and its modifications and found that the glass-lined depth charges (Tiefenbomben) containing Hellhoffit, were much more effective than those loaded with Picric Acid. The mixture consisting of fuming nitric acid (d 1.52) 64.51, nitrobenzene 25.81 carbon disulfide 6.45 and aluminum bronze 3.23% was found to be one of the most effective. A mixture prepd by dissolving 66.7 parts of Dinitrobenzene in 100 parts of fuming nitric acid was also claimed to be effective

**Hemelitene and its Nitrated Compounds.** See under 1,2,3-Trimethylbenzene

**Hemicelluloses (or Semicelluloses).** A group of gummy substances intermediate in composition between cellulose and sugars. They are insoluble in hot water but readily soluble in dilute (4-5%) aqueous NaOH solutions. They are converted by warming with dilute acids (hydrolysis) at normal pressures into pentoses and hexoses (with galactose and glucose predominating), whereas cellulose itself yields only glucose (Ref 2)

According to Karre (Ref 4) hemicelluloses comprise a series of complex polysaccharides, which like lichenin, occur as constituents of the cell walls, as well as reserve substances, occasionally becoming reconverted into sugar by the plant. On hydrolysis, many of them give galactose and mannose in addition to glucose; they are therefore known as galactans, mananns etc. Whether these substances are true compounds is very doubtful. Certain enzymes found in plants and in the intestinal tract convert them into sugars

Dorée (Ref 3) describes them as a polysaccharides, which in their natural state are insoluble in boiling water, but readily soluble in dilute caustic and easily hydrolyzable on warming with dilute acids

Hemicellulose may be divided into pentosans-those yielding pentoses on hydrolysis, hexosans-those yielding hexoses and xylans-those yielding xyloses

The hemicelluloses give explosive compounds on nitrination, but the yields are small and stabilization difficult (Ref 1)
*Refs:* 1) N. Solechnik, J Applied Chem (USSR) 6, 93-8 (1933) & CA 27, 5964 (1933) 2) Marsh & Wood, Cellulose (1945), 2 & 75 3) Dorée (1947) 31 4) Karre (1950), 357
HEMP. A hydrodynamic code used for detonation product flow calculations. See Vol 4, p D152

Hemp Fiber (Chanvre in French). The bast-fiber obtained from the plant Cannabis indica or C sativa, which is a perennial herb (Ref 1). The plant is native of western and central Asia, but has long been cultivated in Brazil and tropical Africa, and is now extensively cultivated in many countries. Its fiber is used for preparation of ropes and paper, but was also proposed by Trench (Ref 2), in 1877 to be nitrated to an explosive

Hemp Nitrate (Nitrohemp) (In French Nitrochanvre). Nitrohemp resembles cotton nitrocellulose or nitroguite in its properties. It may be prepared by nitrating hemp with mixed nitric-sulfuric acid. Trench (Ref 2) proposed using it as a basic ingredient in commercial explosives. Other components were colloidion cotton, resin, ozokerite, glycerin etc
Ref:  Daniel (1902), 773

Hemp Hurds. The refuse from hemp (stems of cannabis sativa) fibers. A US patent claims their use as absorbents for liquid explosives such as NG
Ref:  A.S. Fox & F. Raper, USP 1979681 (1934) & CA 29, 346 (1935)

Hengstite (Hengstite). A smokeless powder patented by Hengst in 1888 in Germany and France. It was prep'd from nitrated "strawpulp" by treating it first with potassium permanganate and then with sodium carbonate. In order to render the mass cohesive and plastic it was mixed with glue prep'd by boiling crushed flaxseed with dextrin. Finally to this were added KNO₃, KClO₃ and ZnSO₄
When intended to use as a military propellant, the mass was either granulated or used in the form of fibers. When used as a mining explosive, it was a plastic pressed into boreholes
The same inventor proposed a propellant prep'd by nitrating of esparato grass (matweed) or of fibers covering coconut shells
Ref:  Daniel (1902), 373

Honeite. Brit 33 grain NC rifle powder manufactured since 1900
Refs:  1) Daniel (1902) 373  2) Marshall I, 326

HEO. Di(hydroxyethyl) oxamide used in prepa of Ditinitroli(nitroxyethyl) oxamide; see Diethylol-oxanide in Vol 5, p D1243-R

HEP. High Explosive, Plastic (HEP) is defined as ammunition designed to defeat armor by spalling or scabbing. Conventional HEP shells have thin walls, ogival noses, base fuzes, and large filler capacities
The expl chge is base detonated upon impinging on the target producing a high energy impulsive load. The incident shock wave is reflected from the rear surface of the target as a tension wave. When the stress exceeds the critical fracture stress of the target, part of the rear surface is "thrown off" as a spall
Development of HEP ammo based upon this principle was initiated in England during WWII after "wall buster" shell were found capable of scabbing armor. These shells were named HE/SH (High Explosive/Squash Head) since they "squash" against the target prior to deton of the filler
The US Army became interested in this method of defeating armor in 1947. Chamberlain Corp, Waterloo, Iowa prep'd in 1960 an "Annotated Bibliography of HEP Information" which provides the first centralized source of info on HEP ammunition

Hept-RDX. See 1,3,5-Trinitro-1,3,5-triazacycloheptane listed under Triazacycloheptane

HEPTANE AND DERIVATIVES
Heptane, Dipropylmethane, Heptylhydride, CH₃(CH₂)₇CH₃; mw 100.23; colorless liq., mp -90.5°, bp 98.4°, flp 25°F, d 0.684 at 20°, RI at 20°, 1.3876, insol w, sol org solvs. Obtained by fractional distillation of petroleum. Irritating to respiratory tract; narcotic in large doses. Dangerous when exposed to gt or flame
Azidoheptanes, C₇H₁₅N₃, mw 141.25, N 29.75. The following derivs. are described in Beil:

1-Azido-heptane. N₃CH₂C₆H₁₄; liq, bp 74° at 18mm, d 0.868 at 20°, RI 1.4343. Prepd by action of NaN₃ on 1-chloro or 1-iodo heptane (Ref 1).

2-Azido-heptane. H₃CCH(N₃)C₅H₁₁; liq, bp 65-66° at 20mm, d 0.862, RI 1.4323 at 20°. Prepd from 2-chloroheptane & NaN₃; NaN₃ + Hg(OAc)₂ + 1-heptane (Ref 2).

3-Azidoheptane, 3-Triazaoheptane or 3-Heptylazide. C₅H₇CH(N₃)C₄H₉; liq, bp 79-81° at 43mm (Ref 2), d 0.858 at 25°, RI 25° 1.4298 (Ref 3); ΔH° = -29 kcal/mole (Ref 4). Prepd in sealed tube reaction of 3-iodoheptane with NaN₃ in acq MeOH (Ref 2).


4-Azidoheptane. (C₅H₇)₂CH(N₃); liq, bp 64-65° at 20mm, d 0.864 at 20°, RI 1.4327 at 20°. Prepd from 4-chloroheptane + NaN₃ (Ref 2).


1,7 Diadiazoheptane (called Heptandiylidiazid in Ger). N₃(CH₂)₂N₃; mw 182.27, N 46.12%; liq, bp 116° at 5mm, RI 1.4700 at 20°. Prepd from 1,7-Dibromoheptane + NaN₃ (Ref 2).

Ref: Beil 1 [397].

Many nitro and nitroso derivatives are also known:

4-Nitroso-4-nitroheptane. H₂C(CH₃)C(NO₂)₂(NO₂)(CH₃)₂CH₃; mw 160.20, N 17.49%. Crystals, mp 72-73° with decmp, sl sol in eth. Prepd from C₅H₇CNOH + NO in cold eth. in dark. Oxidation with CrO₃ in HAC gives 4,4-dinitroheptane. No expl props mentioned for either (Ref 2).

Ref: Beil 1, 156.

Dinitroheptanes, C₇H₁₄(NO₂)₂, mw 190.23, N 14.73%. The following derivs are known:

1,1-Dinitroheptane. (NO₂)₂CH₂C₆H₁₃; liq, bp 81° at 1 mm, d 1.091 at 20°, RI 1.4432 at 20°. Prepd by reacting 2-hexylacetocetoic acid - Et ester with acq nitric acid (Ref 2).

Ref: Beil 1, <396>.

1,7-Dinitroheptane. O₂NC₅H₁₄NO₂; pale yellow liq, bp 198-200° at 10mm, RI 1.458 at 20°(Ref 2).

Prepd from 1,7-diodeheptane + AgONO in eth; the decarboxylation of 2,8-dinitrononanedicarboxylic acid (Ref 2).


3,5-Dinitroheptane. C₅H₇CH(NO₂)₂CH₂CH(NO₂)₂C₅H₇; needles (from MeOH), mp 33°, bp 115-116° at 3mm, d 1.09, RI 1.4453 at 20° (Ref 4). Prepd by reacting 1-nitropropane with 2-nitro-a-buten in ac; heating 1-nitropropane with paraformaldehyde in presence of Et₂NH; from 2-nitrobutene and K-salt of 1-nitropropane in acq soln at 10° (Ref 4).

EtCH(NO₂)CH₂OAc in MeOH, H₂O, NaN₃ soln reactd with PrNO₂ & Triton B catalyst (Ref 2).

Refs: 1) Beil 1 [434] <396> 2) G.L.


2,4-Dinitroheptane. H₃CCH(NO₂)₂CH₂CH(NO₂)₂C₅H₇; bp 98° at 3mm, d 1.10 at 20°, RI 1.4482 at 20°. From 2-nitropropane and K-salt of 1-nitrobutene in ac at 10° (Ref 4).


Heptanedinitrile, 4,4-Dinitro or 4,4-Dinitropimelitoniitrile. CN(CH₂)₇C(NO₂)₂(CH₂)₇CN; mw 240.25, N 23.33%; cryst, mp 79°. Prepd from HC(NO₂)₂:2NO₂K & CH₂:2CHCN in water at 35-45° (Ref 1); or (NO₂)₂C(CH₂OH)₂ & CH₂:2CHCN in acq KOH (Ref 2). Claimed as an ingredient in rocket propellant charges (Ref 3).


Trinitroheptanes, C₇H₁₃(NO₂)₃, mw 235.23, N 17.87%, ob to CO₂ -98.6%. The following isomers are known:

1,1,1-Trinitroheptane, (O₂N)₃C₆H₁₃; liq, bp 80°, d 1.212 at 20°, RI 1.4482 at 20°. Prepd from C₅H₇₁ + AgC(NO₂)₂ in MeCN.

3,3,5-Trinitroheptane. \( \text{C}_2\text{H}_5\text{C(NO}_2\text{)}_2\text{CH}_2\text{CH-} \) 
\( \text{(NO}_2\text{)}_2\text{C}_2\text{H}_5 \), liq, bp 108–110° at 0.005mm, RI 1.4685 at 25°. Prepd by heating 1,1-dinitropropane in a soln of 2-nitrobutylacetate in MeOH, w & NaOH at 40–45° for 2 hrs. Claimed to be a high explosive.

**Note:** Among the polynitroheptanes this is the only found reference to their being explosive.

**Refs:** 1) Beil, not found 2) K. Klager, USP 3000968 (1961) & CA 56, 3715 (1962)

2,6-Dichloro-2,4,6-trinitroheptane. \( \text{CH}_2\text{CCl(NO}_2\text{)}_2\text{-} \) 
\( \text{CH}_2\text{CH(NO}_2\text{)}_2\text{CH}_2\text{CCI(NO}_2\text{)CH}_3 \); mw 304.11, N 13.82%, cryst, mp 73–119°. Prepd by adding the product of reaction of aq base & \( \text{CICH}_2\text{CH}_2\text{NO}_2 \)

a soln of 2-nitroallylacetae in \( \text{CH}_2\text{Cl}_2 \) and heating 3 hrs at 53°. Further chlorination with \( \text{Cl}_2 \)
in basic MeOH gave the 2,4,6-trichloro-2,4,6-trinitroheptane.

**Refs:** 1) Beil, not found 2) M.B. Frankel et al, USP 3440282 (1969) & CA 71, 91105 (1969)

Tetranitro derivs were not found in the literature.

2,2,4,6,6-Pentanitroheptane. \( \text{CH}_3\text{C(NO}_2\text{)}_2\text{CH}_2\text{-} \) 
\( \text{CH(NO}_2\text{)}_2\text{CH}_2\text{C(NO}_2\text{)}_2\text{CH}_3 \); mw 325.23, N 21.54%, OB to CO\(_2\) –46.7%. Prepd by reacting 2-nitro-3-acetoxy-1-propene with 1,1-dinitroethane

**Refs:** 1) Beil, not found 2) M.B. Frankel, Tet 19, Suppl 1, 215 (1963) & CA 59, 11236 (1963)

Hexanitroheptanes, \( \text{C}_7\text{H}_{10}\text{(NO}_2\text{)}_6 \); mw 370.23, N 22.70%, OB to CO\(_2\) –30.2%. The following isomers are known:

1,1,3,3,5,5-Hexanitroheptane. \( \text{(O}_2\text{N)}_2\text{CHCH}_2\text{-} \) 
\( \text{CCH(NO}_2\text{)}_2\text{CH}_2\text{C(NO}_2\text{)}_2\text{C}_2\text{H}_5 \). Claimed to be a burning rate accelerator for rocket fuels; no prep given in CA.

**Refs:** 1) Beil, not found 2) M.B. Frankel, USP 3000970 (1961) & CA 56, 4618 (1962)

2,2,4,4,6,6-Hexanitrohexaheptane. \( \text{CH}_3\text{C(NO}_2\text{)}_2\text{CH}_2\text{-} \) 
\( \text{C(NO}_2\text{)}_2\text{CH}_2\text{C(NO}_2\text{)}_2\text{CH}_3 \); crysts, mp 132°, d 1.70. Prepd from 2,2,4,6,6-pentanitroheptane K-salt & \( \text{N}_2\text{O}_4 \) in \( \text{CCl}_4 \) at –15 to –25° for 4 hrs; or by nitric acid nitration of the pentanitroheptane (Refs 4 & 5); also from \( \text{MeNH}_2 \) & dichlorotetranitroethane (Ref 2). Its hot bar ignition temp is 338° & its sensitivity is approx that of pentolite (Refs 3 & 4)


**Heptanedioic Acid, 4,4-Dinitromalonic acid.** [\( \text{HOOC-C}_2\text{H}_4\text{(NO}_2\text{)}_2\text{C}_2\text{H}_4\text{-} \) \( \text{COOH} \). mw 354.22, N 15.82%, cryst, mp 137°. Obt’d from its di-methyl ester, by refluxing with HCl, which is formed by reaction of HC(NO\(_2\))\(_2\): \( \text{NO}_2 \)K and \( \text{CH}_2\text{CHCO}_2\text{Me} \) in water (Ref 1). It is claimed to be useful in propellant compositions (Ref 2)

**Refs:** 1) Beil, not found 2) L. Herzog et al, JACS 73, 749 (1951) & CA 45, 5609 (1951)

3) USP 2918489 (1960) & CA 54, 8638 (1960)

N,2,2,2,2',2',2'-Heptanitrodiamine. Designated as HOX, also called Bis-2,2,2-trinitroethyl-nitrazine or Bis(2,2,2-trinitroethyl)-nitrazine, abbr as BTNEN. It is listed as an expl der of Diethylenimine in Vol 5, p D1224-R

**Heptanitrohydrocellulose.** See under Hydrocellulose in this Vol.

**Heptanoic Acid and Derivatives**

Heptanoic Acid or Enanthic Acid. \( \text{CH}_3\text{(CH}_2\text{)}_5\text{COOH} \). mw 130.21, colorless, oily liq, mp –10°, bp 223.5°, d 0.913 at 25°, RI 1.4216 at 20°, sl sol in water; sol in alc & eth. Prepd by oxidizing heptan with dichromate & sulfuric acid. Low toxicity & flammability. Used in organic synth & production of special lubricants & brake fluid for aircraft.


The following nitro deriv is described in the literature:

4,6,6-Trinitroheptanoic acid, Me ester. \( \text{MeOOC-C}_2\text{H}_4\text{(NO}_2\text{)}_2\text{CH}_2\text{C(NO}_2\text{)}_2\text{CH}_3 \); mw 279.24, N 15.05%; needles (from ethn), mp 56°. Prepd from 4-nitro-4-pentanoate + Me\( \text{CH(NO}_2\text{)}_2\) + NaOMe
in MeOH at 40-45°; no explosive props are mentioned, but it must be expl
Refs: 1) Beil, not found 2) K. Klager, JOC 20, 650 (1956) & CA 50, 8576 (1956)

**Heptanone and Derivatives**

4-Heptanone, Dipropylketone or Butyronitrile
\((CH_2)\_2CO\); mw 114.21; colorless liq, mp 
-32.6°, bp 144°, fl p 120°, d 0.817 at 20°, R1 
1.4073 at 20°; insol in water, sol in alc & eth.
Toxicity undetermined; moderate fire hazard;
used as a solvent and in polymers & resins
Refs: 1) Beil, Dinitro, C\(_7\)H\(_2\)\(_2\)N\(_2\)O\(_4\), Trinitro, 
C\(_7\)H\(_7\)N\(_3\)O\(_7\), Tetranitro, C\(_7\)H\(_10\)N\(_4\)O\(_9\), and
Pentanitro, C\(_7\)H\(_9\)N\(_5\)O\(_9\), derivs were not found
2) CondChemDict, 8th Ed (1971), p 325

**Heracline.*** British explosive (1875), containing
about equal amounts of each of the following
compounds: KNO\(_3\), NaNO\(_3\), sulfur and saw-
dust, to which was added about 0.5% of Picric
Acid.
Ref: Daniel (1902), 373

**Heraklin (Ger).** An expl patented by Dickerhoff,
was prepared by soaking sawdust in a concd
aqueous soin of equal parts of Picric Acid &
Amm nitrate. The resulting product was dried
and mixed with various ams of pulverized
sulfur and K or Na nitrate
Refs: 1) Gody (1907), 551 2) PATR 2510,
(1958), Ger 88:1

**Hercoblastings.** (Called Kammerminensprengungen
in Ger and Cámaras de mina in Spat). In this
method which might be translated as Chamber
Mines Blasting, a large area consisting of the
material to be mined is selected and horizontal
tunnels are drilled under it, deep in the ground.
Then charges of explosives are placed in the
tunnel, as shown, for example in Figs 259 &
260 copied from Ref 1, p 388. The layout of
the Figs is for a basalt quarry blasting
conducted in 1931 in Pfalz, Germany. All
the shown charges were exploded simultaneously
That method also proved to be popular in
blasting old asphalt or macadam pavements,
where in order to obtain large pieces of materi-
al, expls of low detonation velocity and large
volume of gases produced on exphn should be used. When blasting hard rock, exphs with high deton velocity should be selected.

In Refs 2 & 3 it is suggested that for blasting hard rock, tunnels can be dug deep underground and the charges are uranium-plutonium atomic devices instead of conventional explosives.

Refs: 1) Stettbacher (1933), 397-88 2) Stettbacher (1948), 168 3) Stettbacher, Pólvoras (1952), 212

Hercules Incorporated. Hercules Incorporated, known as Hercules Powder Co prior to 1966, a diversified, world-wide chemical company with estimated annual sales of about $1 billion in 1973, began its corporate history on October 17, 1912. On that date, a decree of the federal court, under the Sherman Act, ordered the dissolution of the E.I. duPont de Nemours and Company, and a regrouping of the properties into the same company plus two new corporations. Hercules Powder Company and the Atlas Powder Company were the new corporations (Ref 3).

The name Hercules Powder was originally given to a dynamite manufactured by the California Powder Works as long ago as 1870 (Ref 1). This was obviously done with the object of comparing the strength of the dynamite to that of the mythological Hercules. Later, there existed a Hercules Powder Company, formed by Lammot duPont in 1882. It operated two dynamite plants whose operations were assumed by others, and its charter was cancelled in 1904. This historic name, and the brand name of the principal dynamite assigned to the new company were influential in the chosen name Hercules Powder Company. At the opening of business on January 1, 1913, Hercules Powder Company, incorporated in Delaware, began with nine black powder mills and two active dynamite plants:

**Black Blasting Powder Mills**
- Rosendale, NY
- Ringtown, PA (2 mills)
- Youngstown, Ohio
- Pleasant Prairie, Wis
- Columbus, Kansas
- Santa Cruz, Calif

**Dynamite Plants**
- Kenvil, NJ
- Hercules, Calif

"A" Blasting and Sporting Powder Mills

Hazardville, Conn
Valley Falls, NY

Black Powder mills began operating as early as 1843 and the dynamite plants began operations in about 1872, producing explosives for the iron, copper, gold, silver, and anthracite coal industries. During the intervening years between the 1870's and founding of Hercules Powder Company in 1912, many advancements in products and process technology had taken place. Low-freezing Nitroglycerin was known, gelatin dynamite was a commercial product, the use of active "dope" (sodium nitrate, coal, sulfur, rosin) was common. Ammonium Nitrate dynamite (extra dynamite) was also an important product. The importance of safety in coal mining had been recognized and use of hydrated cooling salts such as sodium carbonate, or magnesium sulfate had begun. Equipment for manufacture of dynamite such as the Talley mixer, and the Hall packing machine were operational. In principle at least, this equipment design is still in use today.

In addition to dynamite manufacture, the Kenvil plant was equipped shortly after the assignment to Hercules to manufacture about 1500 lbs/day of smokeless shot gun and rifle powder. The brands which became Hercules property included Bullseye* and Unique*. These brands still exist today.

In 1914 the Independent Powder Company of Missouri was acquired, with its dynamite plant near Joplin, Missouri.

In 1914-1915 a new dynamite and black blasting powder plant was constructed and began operation at Bacchus, Utah.

During the years of World War I the Kenvil and Hercules plants were equipped to manufacture TNT. Capacity eventually reached 7,000,000 lbs/month. In addition, Hercules operated the Nitro, West Virginia smokeless powder plant for a short time. The Union plant in Gillespie, N.J., was also acquired by Hercules for production of smokeless propellant and later, for manufacture of nitrocellulose for dynamites and lacquer as well. Total Hercules production of smokeless propellant during World War I was over 100,000,000 lbs. In addition,
A gasless delay fuse which did not require venting was developed for electric caps. This new design improved water resistance and helped prevent misfires or burning of the explosive charge. In addition, semi-gelatin, permissible dynamites containing sodium chloride as a coolant were developed. A broader range of Hercomite* brands of low-cost ammonium nitrate dynamites was introduced. The practice of seismic exploration required explosives with reliable performance after exposure to high water pressures for long periods of time, and so the Vibrogel* seismic gelatins were developed. Red Dot* smokeless shotgun powder was first used, and has been used by champions in trap and skeet shooting ever since.

In the late 30's and early 40's blasting caps for seismic use were developed which enabled the user to determine to within a few tenths of milliseconds the instant of detonation by monitoring the cap circuit. These caps were named Vibrocaps*. Dynamite packaging improvements also followed. Long lengths of thick paper cartridges were developed to enable them to be threaded, or screwed together to improve loading into the borehole. This was Hercules patented Spiralok*. In addition, the paper shell for other dynamites was improved so that it could be loaded into boreholes and then expanded without previously having being slit with a knife. This so called Tamplite* method of shooting resulted in better performance due to better coupling of the explosive in the borehole to the media to be blasted and had the additional advantage of making the loading operation safer for the miner. Currently plastic shells have replaced Spiralok in seismic use.

In World War II Hercules was called upon to conduct research programs to develop new types of ordnance, particularly propellants for guns and rockets. The first bazooka propellant grains were extruded at the Kenvil plant, and many other propellant processing and application improvements followed. Hercules designed, constructed and operated Radford Arsenal, and Sunflower and Badger Ordnance Works for the U.S. Government. These plants turned out all the rocket propellants fired by U.S. Forces during World War II, as well as a substantial portion of the conventional propellants for guns and mortars. In addition Hercules operated the New River bag loading plant, and Volunteer TNT plant. Hercules continued as the contractor for Radford and Sunflower after the war, and in addition, assumed responsibility for the operation of the Allegany Ballistics Laboratory for the Navy. Propellant charges for such operational missiles as Nike, Honest John, Little John, Talos, Terrier, and Rat were designed and developed at ABL (Ref 4).

In about 1953, ANFO (ammonium nitrate and fuel oil) figuratively exploded on the scene of the blasting industry. The market for large diameter, low cost dynamites was rapidly taken over by the even lower cost ANFO. Hercules and other explosives suppliers faced a diminishing dynamite market, and excess capacity. Hercules Black Powder plants had all been closed by this time due to lack of a market, and dynamite came under pressures of excess capacity. However, innovations in explosives had not yet ceased. In the early 50's, Hercules introduced long-length small diameter dynamite cartridges (king-size) for improved economy in loading into boreholes. Hercules patented the equipment design for packaging such long lengths and demand for these products has continued. Improvements in static resistance of blasting caps were introduced as Vibrocap SR* in the mid 50's. Vibronite B* was introduced for a high volume, but short-lived history in offshore seismic operations. It is a blasting agent packaged in metal cans to withstand the rigors of handling storage and use in offshore seismic operations.

Vibrocol*, long length (ca 65 ft or more), pre-packaged columnar charges of explosives were especially suited for some land seismic exploration. Low-priced, packaged blasting agents, Dynatex* and Tritex* were useful in wet locations where ANFO could not be used.

The Chemical Propulsion Division of Hercules was formed in 1958, and began construction of a major facility for design development and manufacture of large rockets and propellants at the Bacchus, Utah plant. In 1958 Hercules acquired the design, development, and production facilities of Young Development Laboratories of Rocky Hill, N.J. The filament-wound, glass fiber-reinforced plastic Spirally* structures

*Indicates registered trademark.
operations for manufacture of acetic acid and acetone were conducted.

In 1921 Hercules last major acquisition in the explosives business was Aetna Explosives Company. With this acquisition came the following active plants:

- Empornum, Pa — Dynamite
- Ishpeming, Mich — Dynamite
- Birmingham, Ala — Dynamite & black powder
- Goes, Ohio — Black powder
- Port Ewen, NY — Blasting caps
- Prescott, Ontario — Fulminate

Electric blasting caps as well as regular blasting caps were added to the Hercules product line with the Aetna acquisition. The plant at Port Ewen, NY had been in operation since 1913. The features of the electric caps which were manufactured at Port Ewen included the insulated wires, copper shell, a base charge (pressed fulminate) an electrical heater (bridge wire), primer (loose fulminate), plug, waterproofing, and sulfur seal. These caps were intended to fire instantaneously when electric current was applied. By 1927 delay electric caps had been developed at Port Ewen, and were commercially available (Ref.2). The design of the delay fuse required venting of the delay element.

In its first 25 years, (ie from 1912 to 1937) Hercules Powder Company introduced many additional innovations. In 1929 Hercules Powder Company introduced a patented semi-gelatin dynamite called Gelamite*. It was an advance in the art due to its ability to replace gelatin dynamites where extreme water resistance was not required, with substantial cost savings. Produced at Rocky Hill proved to be of great importance for rocket cases, motor housings, nose cones, and fuel tanks because of their strength and light weight. From this beginning, Hercules developed the Spiralloy* rocket cases into production scale rockets. Along with the cases, Hercules high energy propellants achieved a breakthrough in mass fraction for high performance space motors and upper stages of ballistic missiles. The Minute Man missile system was one of major importance to the nation and to Hercules Chemical Propulsion Division in these early days. Following the success of the Minute Man, Hercules has also done significant design, development and production work on the Polaris and Poseidon Systems.

In 1958 Hercules began producing \( \text{N}_2\text{O}_4 \), most of which has been used by the Air Force and NASA as an oxidizer in liquid propulsion systems.

In the late 1950's and early 1960's the entire explosives industry was intensively working on a new explosive system, which has since become known as water-gel or aqueous slurry explosives. Hercules was among the leaders in innovation and promotion of water gels. Packaged Flogel*, an aqueous solution of oxidizing salts sensitized with smokeless powder became a forerunner of many such products. The product was packaged in plastic bags. It found ready acceptance on the Iron Ranges of Minnesota and Michigan where jet-pierced boreholes could now be completely filled with this gelled explosive, to improve the blasting performance. The economy, water resistance, and safety of these water gel systems were a significant marketing attraction. Early in the 1960's, Hercules also began to supply water gels in bulk trucks, pumped directly into the borehole, and due to the added savings in labor costs, the use of this concept rapidly accelerated nationwide.

A highly significant technological improvement in blasting technique, called "pre-splitting", using dynamite was introduced by Hercules in 1961. Through the use of precise drilling and special blasting procedures this technique could produce very precise shearing to improve the appearance & durability of the remaining rock, and the economy of construction blasts. This has become a standard technique where overbreak is undesirable, and is widely used in highway construction.

In the late 1960's Hercules water-gel explosives were being used nationwide in bulk operations and packaged products. Sturess was a Hercules patented method of gasifying water gels to produce improved performance due to increased sensitivity and energy output.

SuperSeis* a patented explosive system for offshore seismic exploration, jointly developed by Hercules and Western Geophysical Company, became operational in 1971. It consists of small explosive charges that are ejected from the exploration boat via a long pressured tube and detonate after an appropriate delay, at
water depths of ca 40 feet. Because these charges are percussion initiated they are safer than the conventional electrically initiated seismic charges (Ref 5).

The product outlook for the future includes the Trident I, C4 overseas long-range missile, which is a joint effort of Hercules and Thiokol Chemical Corporation. The work is carried on at the Bacchus and ABL facilities. In addition work continues on Polaris and Poseidon missiles, land-based minuteman missiles and other rockets. Advanced composites are finding increasing use in structural applications requiring high strength and modulus with minimum weight. Explosives-water-gel products will probably accelerate in market acceptance and use because of increased safety in handling and shipping. Their future use in small cartridges to replace NG dynamites is likely to increase.

The third 25-year span of Hercules explosives, smokeless powder and propulsion activities is continuing with the following principal manufacturing facilities:

Bacchus, Utah - Propellants; rockets
Bessemer, Ala - Dynamite, nitric acid, PETN, water gel
Carthage, Mo - Dynamite, water gel
Donora, Pa - Ammonium nitrate, nitric acid
Hercules, Cal - N₂O₄ nitric acid, ammonium nitrate
Ishpeming, Mich - Water gel
Kenil, N J - Smokeless powder, diazo
Louisiana, Mo - Ammonium nitrate, nitric acid
Parlin, N J - Nitrocellulose
Port Ewen, N Y - Detonators, initiators
Rocket Center, W Va - Propellants
Radford, Va - Government owned, company operated
Sunflower, Kansas - Government owned, company operated

which manufacture the following products:

**BLASTING CAPS**

**No 6 Strength Initiators**: aluminum shell No. 6 strength initiators for explosives set off from the spit by the safety fuse

**ELECTRIC BLASTING CAPS**

**Coaldet**: copper-bronze alloy shell, static-resistant, waterproof, millisecond delay electric blasting caps designed especially for use in the coal mining industry. Available in eight accurately timed delays with nominal firing times from 25 through 500 milliseconds. Firing times are printed on blasting cap shell as positive and permanent identification of delay. In addition, Coaldet electric blasting caps are furnished with water-resistant tags that display the delay number in easy-to-see numerals and leg wire insulation is color-coded in a brilliant, high gloss that is color-distinguishable in an underground coal mine environment. All Coaldet blasting caps have iron leg wires for magnetic removal and all conform to the U.S. Bureau of Mines permissible recommendations

**Instader**: bronze shell No. 6 strength, static-resistant, waterproof, instantaneous initiators for explosives set off by an electric current. Available with various length leg wires of copper or iron as required

**Millider**: bronze or aluminum shell No. 8 strength, static-resistant, waterproof, millisecond delay electric blasting caps with delay elements supplying 18 controlled (0 to 17 periods) firing times from 12 to 700 milliseconds with no overlapping of delay periods. Available with various length leg wires of copper or iron as required

**Superdet**: No 6 strength bronze or aluminum shell static-resistant, waterproof, approximately ½-second delay electric blasting caps with delay elements supplying 19 controlled (0 to 15 periods plus ¼, ½, and ¾ fractional delay periods) firing times from 12 milliseconds to 15.6 seconds with no overlapping of delay periods. Available with various length leg wires of copper or iron as required

**Vibrodet SR**: static-resistant, No 8 strength E.B. caps designed to meet the requirements of the seismographic industry for a high-quality, no time lag, waterproof E.B. cap. Vibrodet SR WW is a self-disarming Vibrodet SR designed for offshore blasting work. Vibrodet types are available with copper leg wires only

**BOOSTERS**

**Titan Booster 25, 150, 225, 350, 500, and 2500**: high-strength, relatively insensitive explosive initiators designed for use with commercial explosives, slurries, and blasting agents. The Titan Booster 25XT is a specially designed, self-disarming booster for offshore seismographic work

The self-disarming Titan Booster 25XT and Vibrodet SR WW, aimed at eliminating the
possibility of primed offshore blasting charges being washed ashore, have proved very effective and, as safety devices, were quickly adopted by the offshore seismic industry

**DETONATING CORD**

Detonating Cord: core of pentaerythritol tetranitrate (PETN) contained within a waterproof covering of textile or plastic; available in several grades, each on spools of 500 or 1,000 feet. Manufactured by the Ensign-Bickford Company, Coast Fuse, Inc., and Austin Powder Company.

**SAFETY FUSE**

Safety Fuse: internal burning fuse with black powder core for conducting flame from collar of blasthole to blasting cap or to black powder charge. Hercules Incorporated markets various grades manufactured by The Ensign-Bickford Company and Coast Fuse, Inc.

**BLASTING MACHINES**


Generator-Type: manually operated electric generators for initiating electric caps. Four styles: two 10-cap, a 30-cap, and a 50-cap machine manufactured by Fidelity Electric Company, Lancaster, Pa., 17604; for Hercules Incorporated.

**SQUIBS**

Electric Squibs: means for initiating black powder electrically; supplied with leg wires of 4- and 6-foot lengths.

**BLASTING ACCESSORIES**

Blasting cap crimpers, blasting galvanometers, connecting wire, explosives magazines, fuse lighters, Ignitacord* and Quarrycord*, connectors, leading wire, detonating cord clips and connectors, Quarrycord, rheostats, tamping bags, and V A O blasting meters.

**EXPLOSIVES**

Nitroglycerin Dynamites: in cartridges ranging from 7/8 to 8-inch diameter with varying weight strengths:

- Blasting Gelatin* Hercomite*
- Ditching Dynamite* Hercor
- Extra Dynamite Hercospit
- Gas-Well Explosives High-Pressure Gelatin
- Gelamite* Nitroglycerin
- Gelatin Dynamites
- Gelatin Extra Oil-Well Explosives
- Hercol* Stumping Dynamite
- Unigel

**Permissible Dynamites**: Nitroglycerin-based explosives tested and approved by the U.S. Bureau of Mines as safe for use in blasting in gaseous and dusty coal mines, provided they are stored and used in accordance with conditions established by the Bureau of Mines:

- Collier-C* Red H C
- Hercogel*-A Red H D
- Red H*A Red H F
- Red H B Red H L

**Seismograph-Grade Explosives**: for use in seismic prospecting on land and offshore. Various grades are available in diameters from 2 to 8 inches packaged in fiber cartridges, metal cans, or plastic:

- Gelamite S Vibrogel 5
- Vibrocol* Vibronite* S Primer
- Vibrogel* 3

**BLASTING AGENTS**

Ammonium Nitrate Prill and Fuel Oil Mixtures (AN/FO): available in 50- and 80-pound multiwall paper bags, packages from 5 to 12 inches in diameter, and in bulk deliveries to field storage bins or direct to the blasthole:

- Freemix 3 Hercomix 1-4
- Freemix 33 Hercomix T
- Hercomix*

**Nitro Carbo Nitrate Blasting Agents**: a series of low-cost blasting products containing no “high explosives” ingredients. They have limited water resistance and are available in varying cartridge strengths and packages with diameters of 3 inches and up:

- Dynatex* Tritex* 2
- Dynatex WR

**Seismograph-Grade Blasting Agents**:

- Vibronite B Vibronite S
- Vibronite B High Vibronite S-1

**Energy**

Super AN/FO: ammonium nitrate and fuel oil mixtures with metal additives to obtain increased strength. Super AN/FO is available in packages or can be delivered direct to the blasthole in bulk form.
BLASTING AGENT INGREDIENTS

Herco-Prills*: ammonium nitrate prills designed for use with No 2 diesel fuel oil to produce a nitro carbo nitrate blasting agent. Herco-Prills are readily absorbtent, nonsetting, and offer excellent fuel oil absorbency. Available in bulk, and in 50- or 80-pound multiwall, moisture-resistant bags

SLURRY BLASTING MATERIALS

Slurrex: is a major technological breakthrough by Hercules in blasting with slurry explosives. Developed through the Hercules Energy Management concept, Slurrex provides a varying-density technique to any slurry formula for obtaining a precise amount of volume energy in the borehole. This is caused by the injection of a chemical reagent at the time of bulk loading or packaging. This chemical reagent provides millions of tiny gas bubbles that act as "hot spots," or initiating surfaces, to improve the efficiency of the detonation reaction. Hercules is able to design very precisely a formula for maximum energy and then vary the volume energy with no significant formula change. As an example of the value of this energy control, Hercules is able to provide any level of measured volume energy from 50 million to 110 million ft-lbs/cu ft.

The Slurrex technique is usually applied to produce the maximum measured energy at the bottom of the hole. There the gas bubbles, or "hot spots," are more highly compressed than they are at the top—hence, an optimum energy and density result where these factors are most needed. Each succeeding foot up the borehole has a different energy level, and this directly meets the blast requirements at each horizon on the face. This patented technique, exclusive with Hercules, can be applied to any of its slurry products

Nitro Carbo Nitrate (NCN) Slurries: plastic, water-resistant products that consist essentially of inorganic nitrates, fuels, and metals, in which none of the ingredients are classified as explosives; must be detonated by adequate-strength Titan Boosters. Available in various formulations in either packaged or bulk forms, both of which offer the Slurrex blasting technique

Floget*: water-resistant, highly plastic, non-nitroglycerin slurry explosives classified as a Propellant Class B for shipping purposes.

Must be detonated by adequate-strength Titan Boosters. Available in various grades in either polyethylene bags or bulk, both of which offer the Slurrex blasting technique

PROPELLANTS

GRANULATED DOUBLE-BASE TYPES
(for use in guns)

Blue Dot*: a premium powder designed especially for magnum shotgun loads

Bullseye*: for use in loading pistol and revolver cartridges

Green Dot*: designed for use in medium shotshell loads for all gauges, also some revolver and pistol loads

Herco*: coarse grain, for use in heavy and magnum shotgun loads, also some revolver and pistol loads

Hercules Red Dot*: especially suited for use in light and standard shotshell loads in all gauges, also some revolver and pistol loads

Hercules 2400*: progressive-burning, for use in small-capacity center-fire rifle cartridges, for reduced loads in larger capacity rifle cartridges, for 410-gauge shotshells, and high-velocity loads in some revolvers

HiTemp*: for use in oil-well perforator guns when exposure to high temperature is important

Military Propellants: a wide variety of compositions for standard or special applications

Unique*: an all-around propellant for use in pistol and revolver cartridges, light or gallery rifle loads, and light through heavy shotshell loads

ELECTRIC INITIATORS, EXPLOSIVE DEVICES, ASSEMBLIES, SUBSYSTEMS,

Actuators
Bellows motors
Blasting caps
Blasting supplies
Bolts, explosive
Boosters
Cartridge actuators
Destruct units
Detonators
Dimple motors
Electroexplosive devices
Explosive actuators
Flame initiators
Flash squibs
Fuze subassemblies
Gas actuators

Gas generators
Igniters
Indicators for EED'S
Piston actuators
Primers
Safe and arm systems
Separation devices
Simulators
Squibs
Switch modules
Switches
Thursters
Time-delay devices
CHEMICAL PRODUCTS

Ammonium Nitrate: special purified product for industrial chemical use, available in solution, prill, and grained form. Also high-density grained material for formulation of military explosive compositions

Composition D-2 Wax: a densitizing compound containing wax, nitrocellulose, and a wetting agent; used in military explosive formulations

Diazodinitrophenol (Dazo): yellow, crystalline compound used as priming charge in blasting caps

Mixed Nitrating Acids: a mixture of nitric and sulfuric acid in various ratios and adjusted to the desired DVS values (dehydrating value of sulfuric acid)

Mixed Oxides of Nitrogen (MON): mixtures of N₂O₅ and NO in various proportions as required up to 40% NO

Nitric Acid: basic chemical of commerce available in strengths 57 to 98%

Nitric Acid, Red Fuming: strong nitric acid containing free NO₂. Available in strengths up to 30% NO₂. Also made with and without HF corrosion inhibitor

Nitric Oxide (NO): powerful nitrosating agent in organic reactions. Available under 500 psig in cylinders and tube trailers. A conversion unit is also available for the manufacture of NO from N₂O₅ at the point of use. This is attractive to the large user because it reduces the freight cost

Nitrogen Tetroxide (N₂O₄): an energetic and versatile oxidizer, nitrat, and nitroson. Available as a liquid in steel cylinders and tank cars under moderate pressure (0 to 10 psig)

Nitrogen Trioxide (N₂O₃): especially useful for organic nitrosations and the in situ manufacture of nitrosyl sulfuric acid and sodium nitrite. Also used as an oxidizing agent.

Available in same shipping containers as nitrogen tetroxide

Pentaerythritol Tetranitrate (PETN): white, crystalline compound used as a detonating agent in blasting caps and detonating fuse

CHEMICAL PROPULSION SYSTEMS

Propulsion Systems

AND GAS GENERATORS

Complete solid-fueled rocket motors and gas generators for all applications from tiny satellite spin-control rockets to propulsion for large ballistic missiles, such as Poseidon, Polaris, and Minuteman. A wide variety of systems are in development and production using metal and Spiraloy* motor cases and both composite and double-base solid propellants

Hercapel: a unique all-epoxide cure composite solid propellant with excellent mechanical and ballistic properties. Its outstanding performance in extended environments makes it well suited for tactical missiles

Double Base Solid Propellants: a wide variety of physical and ballistic properties which can be tailored to meet specific performance requirements. Their high specific impulse and excellent reproducibility are two of the many reasons Hercules double-base propellants are found in many of our rocket motors and gas generators used for both military and space applications

ADVANCED SYSTEMS

ADVANCED COMPOSITE MATERIALS

Advanced composite materials from Hercules consist of high-strength, high-modulus graphite fibers that have been preimpregnated with polymer matrices. These materials possess strength-to-weight ratios greater than steel; uses include fabrication of aerospace and aircraft structures, industrial equipment, and other structures or systems requiring high strength and modulus with minimum weight. Hercules capabilities include production of the graphite fibers and materials as well as research, development, design and production of the structure or systems that utilize them

Written by C. W. EILO


Hercules powder. No 1: NG 75.00, KNO₃ 2.10, MgCO₃ 20.85, KClO₃ 1.05, sugar 1.00%. No 2: NG 40.00, KNO₃ 31.00, MgCO₃ 10.00, KClO₃ 3.34, sugar 15.66%
Thorpe (Ref 3) gives two formulas for Hercules powders: (1) NG 77, NaNO₃ 1, woodpulp 2, MgCO₃ 20, strength 106% of Guhr Dynamite No 2. (2) NG 42, NaNO₃ 43.5, woodpulp 11.0, MgCO₃ 3.5, strength 86% of Guhr Dynamite No 1

Herculite. A perchlorate explosive which was at one time on the British "Permitted List". KClO₄ 27/NG 33/collodion cotton 1/woodmeal 9/NH₄ oxalate 29/H₂O 1; limit charge 16 oz 
Ref: Barnett (1919), 137

Herculin. One of the old mining explosives containing KNO₃ together with sawdust and camphor. Other ingredients were also added 
Ref: Daniel (1902), 374

HERO (Hazards of Electromagnetic Radiation to Ordnance). Any Ordnance Item is defined as being "HERO UNSAFE ORDNANCE" when: 1) Its internal wiring is physically exposed; 2) Tests are being conducted on the item that result in additional electrical connections to the item; 3) EED's (Electroexplosive Devices) having exposed wire leads are present, handled or loaded; 4) The item is being assembled or disassembled; or 5) The item is in a disassembled condition (Ref 1)
If such unsafe ordnance is exposed to RF (radio frequency) fields above a certain amplitude level, sufficient RF energy can be induced into the firing circuits to explode the EED's. Therefore, for the handling of HERO UNSAFE ORDNANCE, restrictions of field intensities to certain levels, such as the o.g volts per meter maximum for the 2 to 32 MHz (mega Hertz) range, have been established to ensure safe operations. These criteria levels are specified in Ref 1, which is confidential


Hertz, E. von ( ). German scientist who specialized in explosives. He was the author of numerous publications and patents. He proposed nitro-bis-diazobenzene perchlorate as an initiator and patented the use of Lead Styphnate in initiating compositions

Hertz's Explosives. Patented in 1923 CA 18, 1573-4 (1924) BritP 207563, several compositions, such as ortho-, and para-nitrated quinone diazides of the polymeric phenols or their metallic salts either (1) as a top charge over a main (base) charge such as Tetryl, TNT, or PETN, or (2) in admixtures with other compounds. Following are the primary and nitrating compounds proposed by von Hertz:
1) Bis[Dinitro-hydroxy-azo-quinone], called by v. Hertz sym-Tetranitro-dioxy-diphenol-quinone- tetrazide, or Tetraniitro-diresorcin-diazo-anhydride
2) Dinitro-di-hydroxy-azo-quinone, called by v. Hertz Dinitro-3,5-di-azo-quinone-diazide, or Dinitro-phloroglucin-diazo-anhydride
3) Dinitro-hydroxy-azo-quinone, called by v. Hertz Dinitro-m-azo-quinone-diazide, or Dinitro- resorcin-diazo-anhydride
4) Mono- (and di)-nitro-hydroxy-azo-quinones, called by v. Hertz Mono- (and di)-nitro-para- oxogenine-diazides, or Mono- (and di)-nitro-hydroquinone-diazo-anhydride

HES 4138. One of the Hercules Powder Co solventless sheet double-base, propellants prep'd and investigated during WWII: NC (13.13% N) 49.8, NG 40.0, Et Centr 1.0, K nitrate 1.5, DNT 6.0, DBuPh 1.5 & candellia wax 0.2% 
Ref: R.B. Corey et al, OSRD Reps 1103 (1942), 1 und 1558 (1943) 2

Hesilit. A Rus pre-WWII mining expl: NG (gelatinized with CC) 30.75/DNT 5.25/AN 18/ rye meal or dextrine 39% 
Ref: Anon, SS 12, 409 (1917)

HE SHELL. Same as High Explosive Shell
HE/SH Shell. High Explosive Squash Head Shell. See under HEP in this Vol

HES, P. German scientist who designed several tests for explosives, such as Hess' Crusher Test, Deflagration Test etc. Author of several publications on physical testing of explosives:

Hess' Crusher Test (Hess' Brisance Meter). See Brisance Meter of Hess (See Vol 3 of Encycl., p C492-R, under Compression (or Crusher) Tests)

Hess' Deflagration Test. This test, which is no longer used, consisted of heating the explosive at 70° until spontaneous combustion (deflagration) took place (Ref 1). In a modified method, the explosive was heated at 75° under pressure until deflagration took place

Note: American investigator, A.P. Sy, also designed a "Deflagration Test"

Hess' Test. In this test, a sample of an explosive was heated at 70° in a current of air which was then passed into a solution of zinc iodide and starch. The time taken to impart a blue color to the solution was noted (Ref 2)

Refs: 1) Reilly (1938), 81 2) Ibid, 78

HET. Abbrev for High Explosive Tracer

Heterocyclic Compounds are ring structure carbon comps in which one or more of the atoms in the ring are not C. Frequently these other atoms are N. General refs to heterocyclic comps are:


Heuschrecke (Ger for Grasshopper). A series of weapon carriers (Waffen träger) such as for 105 mm Gun, developed by the Germans early in the WW II. They are described in vol III of the Illustrated Record of German Army Equipment 1939-1945, War Office, London (1947)

HEX (High Energy Explosive). Two types of these expl comps, HEX-24 and HEX-48 are described in Ref 2, pp 164-69

Origin: The development of "slow-burning" expl mixtures which would produce increased blast effects in enclosed or nearly enclosed spaces directed attention to their use for possible military application. In 1950 PicArsn developed a "high capacity" filler for 20mn projectiles consisting of 85/10/5—RDX/Al/desensitizer which was more powerful than Tetryl filler. However, in comparison with MOX (Metal Oxidizer Explosives) Type (described in Ref 2, pp 213-225) there was little doubt as to the superiority of MOX mixts. HEX mixts were developed at PicArsn in 1953 by Sheffield & Murray (Ref 1) as superior high blast comps suitable for use in small caliber projectiles

The HEX comps were prep'd by blending the appropriate weight of the dry ingredients in a Patterson-Kelly Twin-Shell Blender for at least 30 mins

An alternate procedure for 100 to 200g batches used a Cradle-Roll Mixer. This device consisted of a half-barrel type container constructed of wood and lined with an electrical conductive material. A plastic roll was allowed to move over the ingredients by remote control action. The roll action prevented caking of the mixt but had no adverse effect on particle size of the ingredients. The period of time to obt a intmate mixt was approx 15 mins

Compositions and Properties

<table>
<thead>
<tr>
<th>Components</th>
<th>HEX-24</th>
<th>HEX-48</th>
</tr>
</thead>
<tbody>
<tr>
<td>KClO₄ (17 microns)</td>
<td>32</td>
<td>32</td>
</tr>
<tr>
<td>Al (atomized 20 microns)</td>
<td>48</td>
<td></td>
</tr>
<tr>
<td>Al, flaked (1 micron)</td>
<td></td>
<td>48</td>
</tr>
<tr>
<td>RDX (thru 325 mesh)</td>
<td>16</td>
<td>16</td>
</tr>
<tr>
<td>Asphaltn (thru 100 mesh)</td>
<td>4</td>
<td>4</td>
</tr>
</tbody>
</table>

Properties

<table>
<thead>
<tr>
<th>Property</th>
<th>HEX-24</th>
<th>HEX-48</th>
</tr>
</thead>
<tbody>
<tr>
<td>Molecular Weight</td>
<td>47.6</td>
<td>47.6</td>
</tr>
<tr>
<td>OB to CO₂, %</td>
<td>-42</td>
<td>-42</td>
</tr>
<tr>
<td>OB to CO, %</td>
<td>-34</td>
<td>-34</td>
</tr>
<tr>
<td>Powder, color</td>
<td>gray</td>
<td>gray</td>
</tr>
<tr>
<td>Density, apparent, g/cc</td>
<td>1.39</td>
<td>0.69</td>
</tr>
<tr>
<td>Density, pressed at 20000 psi</td>
<td>2.1</td>
<td>1.62</td>
</tr>
<tr>
<td>Ballistic Mortar</td>
<td>Not Given</td>
<td>Not Given</td>
</tr>
<tr>
<td>Blast Effects</td>
<td>Not Given</td>
<td>Not Given</td>
</tr>
<tr>
<td>Brisance by Sand Test, g sand crushed</td>
<td>12.5</td>
<td>23.7</td>
</tr>
<tr>
<td>Brisance for TNT</td>
<td>48.0g</td>
<td></td>
</tr>
<tr>
<td>Detonation Rate</td>
<td>Not Given</td>
<td>Not Given</td>
</tr>
</tbody>
</table>
Properties (cont'd)

Explosion Temp, °C 520 545
(5 sec)

Flame Temp, °K 2552 2382

Fragmentation Not Given Not Given

Friction Test, steel shoe Detonates Partially deton

Friction Test, fiber shoe Unaffected Unaffected

Gas Volume, cc/g 159 200

Heat of Combustn, cal/g 4197 4119

Heat of Expln, cal/g 1858 1735

Heat Test at 100°

% loss in 1st 48 hrs 0.15 Not Detd

% loss in 2nd 48 hrs 0.00 Not Detd

Expln in 100 hrs None —

Hygroscopicity, % None Not Given

Impact Sensitivity 16 inches Not Given

PicArsn App, 2kg wt (TNT 14)

Rifle Bullet Test Not Given Not Given

Sensitivity to Initiation LA 0.20g LA 0.20

(Min priming charge) Tetryl 0.25 Tetryl 0.25

Storage Dry Dry

Trauzl Test Not Given Not Given

Vacuum Stability Test 1.25 1.52

at 100°, cc/g/48 hrs

Usage, press-loaded at HE filler for

20000 psi small caliber

shells

Note: Above expl props are listed in Ref 2

Refs: 1) O.E. Sheffield & E.J. Murray, “Development

of Explosives — Metallized Explosives — High

Blast Fillers for Small Caliber Shell”, PicArsn

Memorandum Report No MR-49, (Dec 1953)

2) US Army Materiel Command, “Explosives

Series, Properties of Explosives of Military

Interest”, AMCP 706-177 (Jan 1971), pp 164-69

Note: There are no US Military Specifications

for HEX’s

Hexa, Hexamin, Hexamit, Hexyl or Hexanitro-
diphenylamine (HNDPh). It is described in
Vol 5, pp D1434-R to D1438-R, under
DIPHENYLAMINE AND DERIVATIVES. Also
in PATR 2610 (1958), p Ger 88-R

Hexa S-22, S-26 and E4. German “Substitute
Explosives” (Ersatz-sprengstoffe) used during
WWII (See Vol 5, Table E15, p E122)

Hexachlorethane, Perchloroethane or Carbon
Hexachloride. Cl₃C-CCl₃; mw 236.72, mp 185°

(sublimes), bp 185.5° at 777 mm, d 2.091 at

20°/4°. Exists in three crystalline forms;

usually as colorless rhombic crystals with a

camphor-like odor. May be prep’d by the

pressure chlorination of tetrachloroethylen

in an enamel vessel and in presence of light

(Ref 3). Nearly insol in w (0.005 at 22°)

very sol in alc and eth. It is toxic, being

similar in action to CCl₄ and results in

lowered blood pressure, kidney and liver

injuries. Explodes in mixtures with Zn powders,

and to lesser extent Al powders (Ref 3). Used

in smoke screen compositions (Ref 2), in ex-

plosive mixtures and pyrotechnics, and as a

camphor substitute in celluloid and other NC

plastics. Tech grade MIL-H-235A (See also Ref 5)

Refs: 1) Beil 1,87, (26) [58] 2) G. Reure, MP 38, 419-21 (1956)

3) A. Lamberous & J. Meyer, MP 39, 343 (1957) & CA 52, 21107

(1958) 4) Kirk & Ortherm 3, 773 (1949); 2nd

edit 5, 166 (1964) 5) CondChemDict (1971), 440

Hexachloronaphthalene or Chloronaphthalene

Chlorinated white tar). C₁₀H₈Cl₆; mw 334.82.

Crystals; prep’d by chlorination of napththalene

(Ref 2, 185). This compd is highly toxic (Ref

3). It was proposed as an addition (4–5%) to

explosives containing large amounts of NaN₃

together with NG, sawdust, etc. It is supposed to

act as a sensitizer in these mixtures

Refs: 1) Beil, not found 2) J.W. Dawson & W.M. Dehn, USP 2255653 (1941) and CA 36,

272 (1942) 3) CondChemDict (1971), 440

2,4-Hexadiyne-1,6-dihydroperoxide. See under

Diacetylenic Dihydroperoxides, Vol 5, p D1120-R

Hexaethylidenetetramine (called Tricrotvlidenetetra-
min in Ger). C₁₂H₂₄N₄; mw 224.40, N 24.97%,

crystals (from hot water, forms numerous salts, some

of which are unstable on heating (Ref 1)

It is reported that the compd is formed by the

reaction of acetaldehyde-ammonia &

ammonia (Ref 1). This reaction yielded a max of

87% gum from which the separation of

not more than 3% of Hexaethylidenetetramine

was possible. Nitration attempt of the product

was not considered practicable. The above

reaction is analogous to the hexamethylen-

tetramine from which RDX is prep’d by
nitrated, and was considered to yield a product which might be nitrated to Cyclotriethylideneminnenitrinamine (Ref 2).


Hexahydrobenzene or Hexamethylene. See Cyclohexane and Derivatives in Vol 3, pp C595-L & R

Hexahydrocatechol. See Cyclohexanediol and Derivatives in Vol 3, p C595-R - C596-L

Hexahydro-1,3-dinitro-5-nitroso-5-triazine,

\[
\begin{array}{c}
\text{H}_2\text{C} & \text{N}^2_2 \\
\text{ONN} & \text{C} \\
\text{CH}_2 & \text{NNO}_2 \\
\text{H}_2 &
\end{array}
\]

mw 206.15, N 40.78%; mp 165° (Ref 3) & 177° (Ref 2); sol in alc. Prep'd by adding Cyclotri-methyleneaminotriazine to a soln of AN in concd sulfic acid at -25°. Mixt is kept at -15° until all solid has dissolved and then warmed and filtered at 0° (Ref 3); 1,9-Diacetoxy-2,4,6,8-tetranitro-2,4,6,8-tetrazalone is reacted with a soln of 99% nitric, hydrogen peroxide & water at -40° to ppt subject material (Ref 2). Nitration of subj mat produces RDX (Ref 3)


Hexahydro-3,6-dinitro-1,3,6-oxadiazepine or 3,6-Dinitro-1-oxa-3,6-diazacycloheptane.

\[
\begin{array}{c}
\text{H}_2\text{C} & \text{N} \\
\text{O}_2\text{NN} & \text{C} \\
\text{CH}_2 & \text{NNO}_2 \\
\text{H}_2 &
\end{array}
\]

N 29.15%; crysts, mp 148-155° (Ref 3) & 154-155° (Ref 2). Prep'd from \(\text{[CH}_2\text{HNNO}_2\]_2\) in 40% formalin reacted with acetyl hyd at 20°, followed by digestion & hydrolysis at 65° (Ref 2); N,N-bis(piperidinomethyl)-ethylendinitramine refluxed in acetylcholoride, then poured into iced water, the pptate is then sequentially poured into MeOH & EtO. It is then dried & recryst from chlf. The intermed thus formed, \([\text{CICH}_2\text{N(NO}_2\text{CCH}_2\]_2\), on standing gives the subject product.


Hexahydro-1,3-dinitro-1,3,5-triazine Nitrate, PCX,

\[
\begin{array}{c}
\text{H}_2\text{C} & \text{N} \\
\text{HNO}_3 & \text{N} \\
\text{C} & \text{NNO}_2 \\
\text{H}_2 &
\end{array}
\]

mw 239.16, N 35.15%

crys, mp 99° (decomp); insol in cold alc & eth. Prep'd by adding hexamine dinitrate to 93% nitric acid at -40°, then adding water and filtering ppt (Ref 4). PCX reacted with sulfuric acid gives RDX (Ref 4)

PCX is claimed to be an intermediate in RDX synth (Ref 2) but this claim is refuted in Ref 3


Hexahydro-1-(methoxymethyl)-3,5-dinitro-triazine or 1-Methoxymethyl-3,5-dinitro-1,3,5-triazoacyclohexane.

\[
\begin{array}{c}
\text{H}_2\text{C} & \text{O} \\
\text{O}_2\text{NN} & \text{N} \\
\text{CH}_2 & \text{NNO}_2 \\
\text{H}_2 &
\end{array}
\]

crys (acet-chlf), mp 134°. Prep'd by addn of 3,5-Dinitro-1,3,5-triazacyclohexane nitrate to a soln of MeOH and 40% qaq formaldehyde (Ref 2); HNO\textsubscript{3} in SO\textsubscript{2} (at -30°) + (CF\textsubscript{3})\textsubscript{2}O to give subject compd, mp 138° (Ref 3). Nitrat of subj comp with HNO\textsubscript{3} was studied in Ref 4

Hexahydro-1-methyl-3,5-dinitro-s-triazine or 1,6-Dinitro-3-methylhexahydro-1,3,5-triazine.

\[
\begin{align*}
&\text{Hexahydro-2-oxo-1,3-dinitropyrimidine or} \\
&N,N'-\text{Dinitro-N,N'-trimethyleneara,} \\
&\begin{array}{c}
\text{CH}_3 \\
\text{H}_2 \text{C} \\
\text{H}_2 \text{C} \\
\text{O}_2 \text{N} \\
\text{N} \\
\text{N} \\
\text{N} \\
\text{N} \\
\text{N} \\
\text{H}_2 \text{C} \\
\text{H}_2 \text{C} \\
\end{array} \\
\text{CO} \\
\text{H}_2 \text{C} \\
\text{H}_2 \text{C} \\
\text{O}_2 \text{N} \\
\text{N} \\
\text{N} \\
\text{N} \\
\text{N} \\
\end{align*}
\]

OB to CO$_2$ - 50.5%; nds (from alc), decomp in boiling water. Prepd from N,N'-trimethylene and conc nitric acid

Refs: 1) Beil, not found 2) A.P.N. Franchimont & H. Friedmann, Rec 26, 218 (1907) & CA 1, 2881 (1907)

Hexahydrophenol. See Cyclohexanol and Derivatives in Vol 3, pp C596-R & C597-L

Hexahydropyrogallol. See Cyclohexanetriol and Derivatives in Vol 3, p C496-L & R

Hexahydroterephthalic Acid Diazide; 1,4-Cyclohexanedicarbonylazide (called Trans-Hexahydroterephthalsiurediazid in Ger),

\[
\begin{align*}
\text{CON}_3 \\
\text{H}_2 \text{C} \\
\text{H}_2 \text{C} \\
\text{CON}_3 \\
\end{align*}
\]

needles,mp 63° with decmp, insol in cold w: sol in ligroin, benz & eth. Prepd by reacting the 1,4-dihyrazid with NaN$_2$ in s dl HAc

Product decmp slowly on standing. On heating, decomposition becomes rapid and almost explosive. The 1,3 isomer may have been an intermediate in a synth of diamino-cyclohexane (Ref 2)


Hexahydro-3,5,7-trinitro-1,3,5,7-tetrazocine-1(2H)-methanolacetate or 1 Acetoxyethyl-3,5,6-trinitro-1,3,5,7-tetrazacyclooctane.

\[
\begin{align*}
\text{NO}_2 \\
\text{O}_2 \text{N} = \text{CH}_2 - & \text{N} - \text{CH}_2 - \text{NNO}_2 \\
\text{H}_2 \text{C} \\
\text{H}_2 \text{C} \\
\end{align*}
\]

\[\text{H}_2 \text{C} \quad \text{N} \quad \text{CH}_2 \quad \text{H}_2 \text{COCCCH}_3 \]

mw 335.28, N 29.25%; crystals, mp 152–53°(Ref 4), no recryst sol found. Prepd by adding 98% HNO$_3$ rapidly to 1,5 methylene-3,7 dinitro
1,3,5,7-tetrazacyclooctane in acetic anhyd (Ref 2); or adding (in increments) 1 acetyl-3,7-dinitro-5-nitroso-1,3,5,7-tetrazacyclooctane to a soln of 50% H₂O₂ in 98% HNO₃ at -50° and raising temp to 0° for 12 hrs (Ref 3) Refs: 1) Beil, not found 2) W.E. Bachmann & E.L. Jennett, JACS 73, 2773 (1951) & CA 46, 2085 (1951) 3) W.E. Bachmann & N.C. Deno, JACS 73, 2777 (1951) & CA 46, 2085 (1951) 4) R.A. Marcus & C.A. Winkler, CanJChem 31, 602 (1953) & CA 47, 12219 (1956)

Hexahydro-1,3,5-trinitro-1,3,5(1H)-triazepine or 1,3,6-Trinitro-1,3,6-triazacycloheptane, Cyclonite Homolog. See Vol 3, p 630. Additional data: nmr spectra (Ref 1); differential thermal analysis (DTA) gives 281° as the exotherm max (Ref 2) Refs: 1) J.A. Bell & I. Dunstan, JChemSoc(C) (1966) 870 & CA 64, 19616 (1966) 2) Y.P. Carigan & D.R. Satriana JOrgChem 32, (2), 285 (1967) & CA 70, 10960 (1969)

Hexahydro-3,5-trinitro-1,3,5-triazine. See Cyclotrimethylenetetraminamide in Vol 3, p 611

Hexahydro-1,3,5-trinitroso-1,3,5-triazine. See Cyclotrimethylenetetraamino in Vol 3, p 630


Hexahydroxymethylbenzene Hexanitrate. See Hexahydroxymethylbenzene Hexanitrate

Hexakis (2,2,2-trinitroethyl)-melamine,
[(O₃N)₃CCH₂]₂N-C-N[N(CH₂C(NO₂)₃]₂

N(CH₂C(NO₂)₃]₂

mw 1104.51, N 30.44%; red oil, which flashes on a hot plate but cannot be detonated by impact. Two synthetic routes to its prepn were considered to be unsuccessful: a) replacement of the chlorine atoms of cyanuric chloride,

Cl-C-N-N-Cl

N
C
C
N

by bis-(trinitroethyyl)amine, [(O₃N)₃CCH₂] NH, in the presence of pyridine; b) reaction of cyanamide with 2 molar equivalents of trinitroethanol to give bis-(trinitroethyl)-cyanamide, followed by trimerization of the latter Refs: 1) Beil, not found 2) N.K. Sundholm, T.C. Richards & D.L. Schoene, Naugatuck Chem Div, US Rubber Co Progress Rept (15 Feb 1950 to 15 April 1950), NORD 10121, p 4

Hexal. A Swiss explosive consisting of RDX & Al, particularly suitable for AA ammunition. Its d is 1.8, expln temp 225 (RDX 225°), Trangl value 420 cc (RDX also 420) & deton vel 7900 m/sec (RDX 8500) (Ref 1) The Al is coated with a water-insol-wax to make it impervious to water-Al reactions that liberate hydrogen (Ref 2) Refs: 1) PATR 2510 (1958), p Ger 88-R 2) P. Aubertin, FrP 1180530 (1959) & CA 54, 20210 (1960) 3) M. Freiwald, Explosivst 6, 133 (1961)

3,3,7,7,11,11-Hexamethanol-1,5,9-trioxacyclocodecanehexanitrate or 1,5,9-Trioxacyclocodecane-3,3,7,7,11,11-hexamethanolhexanitrate.

[CH₂ONO₂][CH₂ONO₃][CH₂ONO₂]

O-CH₂-C-CH₂-O-CH₂-C-CH₂-O-CH₂-C-CH₂

CH₂ONO₂

CH₂ONO₂

CH₂ONO₃

mw 624.45, N 13.46%, OB to CO₂ -53.8%; tacky solid. Pentaoerithritol is polymerized with dil sulfuric acid and nitrated with 98% nitric acid at 0°. Used as sensitizer for AN explosives Refs: 1) Beil, not found 2) J.A. Wyler, USP 2465776 (1949) & CA 43, 6223 (1949)

2,3,5,2',4',5'-Hexamethyloxazobenzene-6-azoide or 6-Azido-2,3,5,2',4',5'-hexamethyloxazobenzene.

(CH₃)₃C₂(N₂)₃C₆H₄N·N·C₆H₄(CH₃)ₖ; mw 307.44, N 22.78%; red nds (from eth); mp 90°-91°(dec). explodes mildly on rapid heating or on contact with concd H₂SO₄. May be prepd from 6-amino-2,3,5,2',4',5'-hexamethyloxazobenzene as described in Ref 2 Refs: 1) Beil 16, 76 2) T. Zincke, H. Jaenke, Ber 21, 546 (1888)
Hexamethyldiphenylethane and Derivatives

Hexamethyldiphenylmethane also called 3,5-Dimethyphenylpropane

\[
\begin{align*}
\text{CH}_3 & \quad \text{CH}_3 \\
\text{C(CH}_3_2 \text{)} & \quad \text{CH}_3 \\
\text{CH}_2 & \quad \text{CH}_2 \\
\end{align*}
\]

252.43; crystals (from alc), mp 174.5°. Prepd by treating m-xylene with 2,2-dichloropropene in the presence of AlCl₃. Nitrination with 65% nitric acid gives the mononitro and dinitro derivatives, neither of which is explosive, although the latter decomposes at 250°. Further nitrination with 97% nitric acid produces the tetranitro derivative which decomposes at 295-300°.

Hexanitroxohexamethyldiphenylethane or 2,2-Bis[2,4,6-trinitro-3,5-dimethylphenyl] propane, [(CH₃)₂C₆(NO₂)₃]₂C(CH₃)₂; mw 522.43, N 16.09%, OB to CO₂ – 107.2%; yellowish crystals (from isomyl alc), color deepens on exposure to light; sl sol in warm isomyl alc; decomposes with a slight explosion at 205°. Prepd by nitratating 2,2-Bis[2,4-dinitro-3,5-dimethyl-phenyl] propane with fuming sulfuric acid mixed with 97% nitric acid. It is an expl

Ref.: 1) Bell 5, [527] 2) H. Goudet & F. Schenker, Helv 10, 134, 139 (1927)

3,3,6,9,9-Hexamethyl-1,2,4,5,1,2,4,5-tetraoxane. See Acetone Compounds under 2,5-Bis(hydroperoxy)-2,5-dimethyl-hexene in Vol 2, p B144-R

Hexamethylenediamine and Derivatives

Hexamethylenediamine (HMDA). H₂N(CH₃)₂NH₂; mw 116.24, N 24.10% silk-like leaves, mp 42° bp 205° (subl), sol in water; sl sol in h eth. Prepd by hydrogenation of adiponitrile over Raney Ni or Co; chlorination of butadiene then reactn with NaCN & hydrogenation. Acute local irritant-mod flame hazard. Used in high polymer synth & as a cross-linking agent (Refs 1, 3 & 4).

In alc soin HMDA reacts with TNT to give the following derivs:

\[\beta \text{TNT} = 3,3'-\text{dimethyl-2,2',6,6'-tetranirotrodi}-\]
\[\text{phenyl-N,N'-hexamethylenediamine}, \text{mp 176-177°}
\]

\[\gamma \text{TNT} = 4,4',6,6'-\text{deriv, mp 187-190°}
\]

With 2,4 (NO₂)₂C₆H₃Cl, HMDA reacts to give 2,2',4,4',6,6'-tetranirotrodi(phenyl-N,N'-hexamethylenediamine, mp 206-208° (Ref 2)

Ref.: 1) Bell 4, [597] 2) M. Giua & G. Musso,
(V.) \[ \text{NO}_2 \cdot \text{NH} \cdot \text{CH}_2 \cdot \text{N(NO}_3 \text{)} \cdot \text{CH}_2 \cdot \text{NH}_2 \cdot \text{HNO}_3 \]

(VI.) \[
\begin{array}{c}
\text{H}_2 \text{C} \\
\text{N-CH}_2 - \text{CH}_2 - \text{N-CH}_2 - \text{N-NO}_2 \\
\text{H}_2 \text{C} \\
\text{N-CH}_2 - \text{CH}_2 - \text{N-CH}_2 - \text{N-NO}_2
\end{array}
\]


Hexamethylenediamine Peroxide.

\[ (\text{CH}_3)_2 \text{NH}_2 \cdot \text{H}_2 \text{O}_2 ; \text{mw} 150.26, \text{N} 18.64\% \text{; crys} \text{ts. HMDA reacts with aq or ethereal H}_2 \text{O}_2 \text{ to give a solid adduct} \]

Refs: 1) Bell, not found 2) Ya.A. Fialkov & A.A. Shokol, UkrainKhimZhur 15, 318 (1949) & CA 48, 5091 (1954)

Hexamethylenetetramine and Derivatives. Hexamethylenetetramine (HMTA), Hexamethylenecaine, Hexin, Urotropine, Hexamine, Forminycystogen, Aminoform, Hex, Naphthamine or Uriton (Ger)

\[
\begin{array}{c}
\text{H}_2 \text{C} \\
\text{N-CH}_2 - \text{CH}_2 - \text{N-CH}_2 - \text{N-CH}_2 - \text{N} \\
\text{H}_2 \text{C}
\end{array}
\]

\( \text{mw} 140.19, \) N 39.97%; colorless, odorless and sweet-tasting rhombic crystals; d 1.27 at 25°, mp 280°; when heated in a sealed tube it decomposes with charring at a temp above 280°; when heated in air, it decomposes without melting; when heated in vacuum, it sublimes at 230-270° without appreciable decomposition, fl p 482°F

When hexamine is ignited it burns with a pale blue flame. Its std heat of formation is +29.65 kcal/mole for the solid and 47.6 for the gas (Ref 2)

Pure hexamine may be taken internally by some persons in small amounts and it is used in medicine as a urinary antiseptic, but with some persons it is toxic. It can also be a skin irritant

Hexamine serves as a parent compound for the preparation of RDX, HMX & Hexamethyleneperoxidediamine. It is also used in a mixture with sodium phenate, as an adsorbent of phosphine in gas masks. Its use as a cross-linking agent for guar gum in AN-slurry explosives has been patented (Refs 9 & 10)

An early review of the uses of hexamine in the explosive industry is given by Konrad (Ref 2). Also, see further under "Hexamethylenetetramine Explosives". Other more recent reviews are Refs 7 & 9. For the nitration of hexamine see Vol 3, pp C611-624

According to requirements of the U.S.
yel crysts; was prep'd by Hofmann (Refs 1 & 2) by the interaction of hexamine with chromic acid and 30% hydrogen peroxide.

Very sl sol in water. Explodes (but not violently) on heating.

**Hexamethylenetetramine-Dichromic Acid.**

\[
[(\text{CH}_2)_6\text{N}_4]^+ \cdot \text{H}_2\text{Cr}_2\text{O}_7^-; \text{mw} 498.46, \text{N} 22.48%;
\]

yel crysts; was prep'd by Cambier and Brochet (Refs 1, 2 & 3). Explodes violently.

**Hexamethylenetetramine-Tetrachromic Acid.**

\[
[(\text{CH}_2)_6\text{N}_4]^+ \cdot 2\text{H}_2\text{Cr}_4\text{O}_{13}^-; \text{mw} 698.46, \text{N} 22.48%;
\]

crys; was prep'd by Cambier and Brochet (Refs 1, 2 & 4). Explodes more violently than the previous compd.

Lit search in CA (1947-71) revealed no other expl hexamine Cr compds.

**References:**


**Hexamethylenetetramine-Chromium Compounds.**

There are several explosives containing hexamine combined with either chromic oxide or chromic acids:

**Hexamethylenetetramine-Chromium Tetroxide,**

\[(\text{CH}_2)_6\text{N}_4\text{CrO}_4^-; \text{mw} 256.11, \text{N} 21.88%; \text{reddish brn}\]

Armed Forces as covered by Spec MIL-H-502A (June 1968), there are two grades of Hexamine (1 & II) both of which must comply with the following requirements:

1. **Purity** — minimum 99.5%, maximum 100.2% (as determined by the method described below, under Tests)

2. **Formaldehyde** — none

3. **Ammonia** — max 0.02%

4. **Chlorides** — max 0.02%

5. **Ash** — max 0.05%

6. **Water insoluble material** — max 0.05%

7. **Solubility in glacial acetic acid** — completely soluble without turbidity.

In addition, there are specific requirements for each grade:

8. **Moisture** — max, Grade I, 0.50%, Grade II, 0.25%

9. **Granulation** — through U.S. Standard sieves, conforming to Federal Spec RR-S-366;

   **Grade I** — sieve No. 16 — 100% (min.), No. 60 — 30% (max); No. 100 — 10% (max)

   **Grade II** — sieve No. 16 — 100% (min.), No. 60 — 50% (max), No. 100 — 20% (max)

**References:**


**Hexamethylenetetramine Explosives.** A powerful solid explosive was claimed to have been prep'd by oxidizing hexamine with a solution of hydrogen peroxide, treating the resulting product with nitric acid and then reoxidizing with H_2O_2 (Ref 1).

After drying, this was mixed with AN, castor oil and turpentine (Ref 2).

**References:**

1) E.L. d'Asteck, USP 1835697 (1931) & CA 26, 1125 (1932) 2) L.E. d'Asteck Callery, Brit J415900 (1934) & CA 29, 929 (1935)

**Hexamethylenetetramine Explosives.** Several explosive compositions containing hexamine in admixtures with oxidants or as chromium, copper nitrate, perchlorate, & peroxide compds are known. Several hexamine/oxidant mixtures have been patented as explosive & propellant compositions:

91/9% AN/HMTA used dry or as slurries (Refs 3 & 5); HMTA/Al_2O_3-9H_2O/KClO_3 (Ref 1); an HMTA-AN adduct prep'd from aq solns or melts and mixed 1 part adduct to 4 parts AN (Ref 2); compositions that are detonator-sensitive consisting of 90/9/1% AN/HMTA/abietic acid; prep'd by making a homogenous melt at ca 145°C and quickly cooling it to get finely crystal AN. This compd passed thru 16 mesh screens; and loaded into 1.5 inch diam steel tubes, deto-
nates at 18,700 ft/sec at d 1.24 & 13,400 ft/sec at d 1.44 g/cc (Ref 4). The use of HMTA in AN slurry explosives is claimed in Refs 6 & 7.


Hexamethylenetetramine Nitrates. The following nitrates were reported to have been prep'd by various investigators:

Hexamethylenetetramine Mononitrate. (CH₃)₆N₄ . HNO₃; was prep'd by Delepine by the action of dilute HNO₃ on aqueous solns of hexamine at 0⁰ (Ref 4). When more concentrated acid is used, the dinitrate is obtained.

Hexamethylenetetramine Dinitrate (HDN) (Hexamime Dinitrate). (CH₃)₆N₄ . 2HNO₃; mw 266.22, N 31.57, OB to CO₂ -78.1%; white crystals; d 1.63, mp 160-165⁰, ΔH_comb 3583 cal/g, ΔH₂O -92.8 kcal/mole (Ref 9). UV spectra for both the mono and dinitrates are given in Ref 11. May be prep'd by nitration of hexamine with mixed nitric-sulfuric acid. For this, Hale (Ref 6) gradually added nitric acid (d 1.42) to a 25% aqueous solution of hexamine at 0⁰. The precipitated salt was separated from the acid by filtration through glass wool and dried after washing with cold 50/50 alcohol-ether.

It may also be prep'd by passing formicdehyde into a cold or alcoholic soln of NH₃ and treating the resulting product with nitric acid (Ref 7).

The obsd low ylds in the nitration of hexamine have been ascribed to the formation of Trinitrodiaminodimethylamine, O₂NN(CH₃NNO₂)₂ (Ref 10). Further nitration of hexamine-dinitrate with concd nitric acid gives RDX (See Vol 3, p C614).

If hexamine is treated with nitric acid, or Cyclonite Dinitrate with AC₂O in sulfuric acid, and then neutralized to a pH of 5.6, 3,7 Dinitropentamethylenetetramine is claimed to be formed (Ref 12).

HDN is readily sol in water; but decomposes on standing. It is sol in alc, eth, chl, & CCl₄.

It is an explosive compound, which is less powerful and much less brisant than TNT.

Power (by Trauzl Test) - 65% TNT; Brisance (by Sand Test) 18.1 g against 48 g for TNT, or 38% of TNT; Impact Sensitivity (with 2 kg weight), Picatinny Arsenal App 13” against 14” for TNT and 9” for Tetryl; Bur of Mines App 40 cm against 60+ for TNT and 26 cm for Tetryl; Pendulum Friction Test - negative; Minimum detonating charge for 0.4 g HDN is 0.24 g. Tetryl when init'd with 0.24 g MF; Thermal Stability, as determined by the 120⁰ Heat Test - S.P. (salmon pink) 145 min., Red Fumes 300+ mins; no explosion in 5 hrs, which might be considered satisfactory; Heat of Combustion 948.4 kcal/mole.

HDN is one of the intermediate products obtained during preparation of Cyclonite from hexamine. It has also been proposed for use in the manufacture of an explosive, tentatively considered to be 1,3,5-trinitrohexahydros-triazine (Ref 7). Its use in an AN-slurry expl has been patented (Ref 12).


Hexamethylenetetramine-Nitroform Salt.

(CH₃)₆N₄ . 3HNO₂; mw 592.77, N 33.70%, OB to CO₂ -79.6%; crystals, mp 145⁰ (dec). It was prep'd by refluxing for 1.5 hrs a mixt of hexamethyl-
ene tetramine & nitroform in abs alc until a yel
flocculant formed. The ppt was filtered,
was twice with abs alc and once with ether.
The product was not analyzed. Detonates
weakly and burns easily
Refs: 1) Beil, not found 2) J.J. Schnauffer,
US Rubber Co Quarterly Progress Repi No 5,
(1 Oct 1948-1 Feb 1949), p 47

Hexamethylenetetramine Perchlorates. An
explosive perchlorate was prep. in 1916 by
Riedel (Refs 1 & 3). It was the monoperchlo-
rate. Later, Hassel (Ref 4) prep. the mono-
bi- & triperchlorates by treating an aqueous
solution of hexamine with perchloric acid,
with or without alcohol
Following are the properties of the mono-
and diperchlorates:
Hexamethylenetetramine Monoperchlorate.
(CH₂)₆N₂·HClO₄; mw 240.65, N 23.28%,OB to
CO₂ & Cl₂ -96.4%, mp 158°. Crystals; easily
sol in w imparting an acidic reaction to it
It is an explosive about 65% as powerful as
TNT and less sensitive to impact (Ref 4)
Hexamethylenetetramine Diperchlorate.
(CH₂)₆N₂·2HClO₄; mw 341.12, N 16.43%,OB to
CO₂ & Cl₂ -28.1%. Crystals. It is an explo-
sive about 125% as powerful as TNT and
comparable in sensitivity to impact to Tetryl.
Lit search of CA (1947-71) uncovered no new
refs to explosive HMTA Perchlorates
Refs: 1) Beil 1, (308) 2) J.B. Riedel, GerP
292284 (1916) 3) O. Hassel, NorwegianP
57831 (1937) & CA 31, 6466 (1937)
4) Blutt, OSRD 2014 (1944) 5) Walker (1953),
283 & 331

Hexamethylenetetramine Peroxides. The first
compound of this type was prep. by von
Girsewald (Refs 1, 2 & 3) in 1912 by evaporat-
ing under vacuum at 40-50°, the solution
obtained by dissolving hexamine in a slight
excess of 30% H₂O₂ containing a small amount
of mineral acid, and then concentrating the
product in vacuum. The resulting Hexamethylen-
etetramine Hydroperoxide (Hexamethylene-
Wasserstoffperoxyd in German) was a colorless
crystalline compound, readily sol in w & alc.
It is an extremely explosive material which
deflagrates on rapid heating or when brought
in contact with concd H₂SO₄. It is stable only
to about 70°

Note: If instead of treating the free hexamine
with peroxide, its salts are used (such as the
citrate), or if hexamine is treated with peroxide
in the presence of substantial quantities of
acid, the resulting product is Hexamethylen-
etriperoxidediamine (cy)

Another peroxide compound was prep. by
Leulier (Ref 4) and repeated by v. Girsewald
(Ref 5) on treating hexamine first with hydra-
gen peroxide and then with nitric acid. The
resulting white crystals were presumed to be.
H₂N(CH₂)₂(CH₂-OH)₂·N₂·2HClO₄; mw 137.14,N 10.21%.
It is insol in w, alc, eth & other organic sol-
vents. When wet, it is inactive, but when dry
it is detonated violently by the blow of a
hammer or when heated to about 130°
Refs: 1) Beil 1, (308) 2) Walker (1953) 289
3) C. von Girsewald, Ber 45, 2574 (1912) &
CA 7, 346 (1913) 4) A. Leulier, JPChimChim
15, 222 (1917) & CA 11, 2277 (1917) 5) C.
von Girsewald, Ber 54B, 490 (1921) & CA
15, 2416 (1921)

Hexamethylenetetramine Styrphate or Hexa-
mine Styrphate. Probably C₁₂H₁₂N₂O₆ (struct
formula unknown); mw 385.34, N 25.45%,crysts
[from NM], mp 197-198° by fast heating
Richmond et al (Ref 2) prep. it by adding
one equivalent of hexamine to a satd aq soln
of Styrphic Acid. The pptn of hexamine mono-
styrphate was quantitatve; if the Styrphic
Acid was saturated into ethanol instead of
water, the loss by solubility in 1:1 water-
ethanol was only 0.25 g per 100 cc of soln.
The mp was 196° (6° per minute rise). This
styrphate is much more useful than picrate as
a hexamine derivative largely because Amm
Styrphate is sol in water or water-ethanol.
Thus a soln contg one mole hexamine as a 50% soln
plus 2 moles of ammonia gave a 98% yield
of Hexamine Styrphate, mp 196°
Refs: 1) Beil, not found 2) H.H. Richmond
et al, JACS 70, 3663 (1948) & CA 43, 1316
(1949)

Hexamethylenetetramino-Tetrazido-Copper.
(N₂)₃Cu(CH₂)₆N₂Cu(N₂)₃; mw 435.33, N
51.49%; is described by Cirulis & Straumanis,
JPChim 162, 321 (1943) & CA 38, 1970 (1944).
It is sol in aq ammonia, in Et diamine & in boiling
dil nitric acid. It burns quietly in a flame but
confined it explodes at 180-185°. With 1 kg
falling weight it explodes at 10cm
Hexamethylenetetramine- Trinitro-m-cresylate.  
\((\text{CH}_3)_2\text{N} \cdot \text{C}_6\text{H}(\text{CH}_3)(\text{OH})(\text{NO}_2)\text{C}_6\)  
mw 383.32, N 25.58%. crysts, mp 175°. Was first prep by Datta et al (Ref 2) by mixing hot benzene, alcohol solutions of 1 mol of hexamine and 1 mol of trinitro-m-cresylate and then allowing to cool. The resulting compound exploded at 325°  
Refs:  1) Beil, not found  2) R.L. Datta, L. Misra & J.C. Bardhan, JACS 45, 2432 (1923)  3) Walker (1953), 331

Hexamethylenetetramine Triperoxide and Hexamethylenetriazine Triperoxide. Depending on conditions HC\textsubscript{2}O + H\textsubscript{2}O\textsubscript{2} + oxalic acid (sic) give the tetramine or triazine peroxides. Their use as primary explosives has been patented  
Refs:  1) Beil, not found  2) Gévelot and Gaupilat, FrP 893942 (1944) & CA 47, 8373 (1953)

Hexamethylenetriperoxidiamine; 3,4,8,9,12,13-
Hexaaza-1,6-diazabicyclo-[4,4,4]-tetradecane  
(HMTD) (formerly called by Langler, Hexa-oxymethylenediamine). Ni (\text{CH}_3 \cdot \text{O} \cdot \text{O} \cdot \text{CH}_3) \cdot \text{N} \cdot \text{N}  
mw 208.17, N 13.46%, OB to CO\textsubscript{2} 92.22%;  
d (cryst) 1.57; loading density 1.05 at 100 atm, 1.15 at 200 and 1.30 at 800 atm/ml  
145° Colorless plates or rhombic crystals.  
IR spectra are given in Ref 19. Was originally prep by Baeyer and Villiger (Ref 2) by adding 40% formaldehyde to a solution of ammonium sulfate in 3% H\textsubscript{2}O\textsubscript{2} at 55°. Later C. von Girsewald (Refs 3, 4 & 5a) prep it by dissolving hexamine and citric acid in 30% hydrogen peroxide  
Taylor & Rinkenbach (Ref 6) modified the Girsewald procedure and prep HMTD as follows:  
a) Dissolve, with stirring, 56 g of powdered hexamine in a liter beaker containing 185 g of 30% hydrogen peroxide (or its equivalent of stronger solution up to 40%)  
b) While cooling the beaker in water, gradually add, with stirring, 84 g of powdered, crystalline citric acid, keeping the temperature in the beaker below 30°  
The following reactions taken place, as was suggested by Marotta and Alessandrin (Refs 7a and 16, p 289)
\[
\begin{align*}
\text{CH}_3 \cdot \text{N} \cdot \text{CH}_2 \\
\text{2N} \cdot \text{CH}_2 \cdot \text{N} \cdot \text{CH}_2 + 3\text{H}_2\text{O}_2 & \rightarrow \\
\text{CH}_2 \cdot \text{N} \cdot \text{CH}_2 \\
\text{CH}_3 \cdot \text{O} \cdot \text{O} \cdot \text{CH}_3 \cdot \text{N} + 2(\text{CH}_2 \cdot \text{NH}) & \rightarrow \\
\text{CH}_3 \cdot \text{O} \cdot \text{O} \cdot \text{CH}_2
\end{align*}
\]

HMTD  
Trimethylenetriamine Ammonia also forms during this reaction, but it is absorbed by acid with formation of a salt  
c) After all the acid has dissolved and the temperature starts to drop, stop the agitation and allow the beaker to stand overnight in a bath at room temperature  
d) Separate the crystals by filtering through a Büchner funnel and rinse them with distilled water, followed by 95% alcohol (to facilitate the drying) and then dry in air  
e) Allow the filtrate to stand until the next day and if there are any additional crystals, separate them out and purify as above  
This method gives yields between 68.5 and 71%. Davis (Ref 15) uses a method of prep which requires about 5 hours and cooling to 0° & below. Leullier (Ref 5) prep HMTD from hexamine, hydrogen peroxide and nitric acid. Konrad (Ref 11) used the same procedure as of Taylor & Rinkenbach and also by treating 50 g of ammonium sulfate with 50 g of 30% hydrogen peroxide and 5 g of 40% formaldehyde.  
Properties of HMTD were examined in detail by Taylor & Rinkenbach (Refs 6 & 7) and also by Konrad (Ref 11) 
Solubility at 22° (g per 100 g of solvent): water 0.01; abs alc less than 0.01; ether 0.017; 
CS\textsubscript{2} less than 0.01; CCl\textsubscript{4} 0.013; glacial acetic acid 0.14; chloroform 0.64; acetone 0.33; 
glycolacetate 0.90 :  
HMTD is much more sol in hot water, especially on long standing. Its aqueous solns are stable  
Solubilities in hot water:  
At 60° after 2 hours 0.10%; 8 hrs 0.35; 24 hrs 0.50; 48 hrs 0.50  
At 75° after 2 hours 0.25%; 8 hrs 0.60; 24 hrs 1.30; 48 hrs 2.25  
At 100° after 2 hours 3.25%; 8 hrs 39.00%; 24 hrs 68.00.
It is very hygroscopic and does not seem to be toxic. HMTD is not compatible with most metals even when it is dry. It attacks Al, Sn, Zn, brass, Cu, Pb and iron (Ref 16). Mixtures of HMTD with PA, RDX, PETN and Tetryl appeared to be stable at 50°. Mixtures of HMTD with TNT or KClO₃ were less stable (Ref 16).

HMTD is destroyed by caustic soln with liberation of ammonia (Ref 16). It can be nitrated at low temps using HNO₃ to give ca 26% RDX (Ref 18).

HMTD is an explosive of the initiating type; it is more powerful and brisant than MF, rather being comparable to LA and Cyanuric Triazide. It does not become dead-pressed even at pressures of 11000 psi (an advantage over MF).

Explosive properties:

Behavior Toward Flame: when a small quantity is ignited, it flashes like NC, giving a flame several inches in height.

Brisance by Sand Test: 0.5 g at 1000 psi pressure crushes 42.5 g compared with 44.2 g for Cyanuric Triazide and 16.5 g for MF.

Power: by Trauzl Test, about 60% of TNT or 230% of MF (Ref 11).

Explosion temperature: explodes instantly at 200° or open flame; ignites in 3 secs at 149° when thrown on a heated metallic surface; explodes at 130° when heated gradually (Ref 16).

Friction Sensitivity: extremely sensitive.

Heat of Combustion: 4295 cal/g at constant volume, H₂O liq.

Heat of Formation, 385 cal/g or 80 kg cal/mol (Ref 10).

Impact Sensitivity: it is very sensitive to impact even when wet (Ref 16).

Velocity of Detonation: 4511 m/sec at d 0.88 in column 0.22" diam S100 m/sec at d 1.1.

Initiating Action: The minimum amounts of HMTD required to detonate 0.4 g of the following HE were determined by increasing the amounts of HMTD until the amount of sand crushed was at a maximum. (Explosives and initiator were loaded in No 8 cap at 1000 psi pressure.)

TNT required 0.08 g using reinforcing cap and 0.10 g without reinforcing cap. PA required 0.05 g using reinforcing cap and 0.06 g without reinforcing cap.

Tetryl required 0.05 g using reinforcing cap and 0.06 g without reinforcing cap. When 0.05 g HMTD was pressed at 1000 psi into a No 8 capsule, it detonated a stick of ordinary 40% dynamite and a stick of Blasting Gelatin which had become insensitive through age.

HMTD is very unstable in storage. Its weight loss is 79% in 300 days at 50°, 150 days at 70° and 5 to 20 days at 90°, for material stored dry. Even under water it showed considerable decomp in 4 months (Ref 16).

Although C.v. Girsewald patented the use of HMTD as a primary explosive for detonators, its sensitivity to heat, friction and impact, its incompatibility with metals and its poor storage qualities make it unsuitable for practical use (Ref 16).


Hexamethylolbenzene and Derivatives

Hexamethylolbenzene or Benzenhexamethanol. C(CH\(_2\)OH\(_6\)); mw 198.25; crystals, mp 302-311\(^\circ\)C (Refs 2 & 3), sol in w. Prepd from C\(_6\)(CH\(_2\)OAc\(_6\) + alc KOH (Ref 2); by trimerizing H\(_2\)CH\(_2\)OH with Ni(o) bistrimethyl catalyst under N (Ref 3) or Co(CO\(_3\))NO catalyst (Ref 4).

Refs: 1) Beil, not found 2) M. Chaigneau, CR 233, 692 (1951) & CA 46, 7071 (1951) 3) N. von Kutepow & F. Meier, Ger P 1159951 (1963) & CA 60, 9198 (1964) 4) Anon, FrP 1397664 (1965) & CA 63, 8235 (1965)

Hexamethylolbenzene Hexanitrate or Benzenehexamethylhexanitrate. C\(_6\)(CH\(_2\)ONO\(_2\))\(_6\); mw 528.30, N 15.91%, OB to CO\(_2\) 36.35%; wh powd, mp 176\(^\circ\)C, decomp. D\(_\text{H}_{\text{comb}}\) 2653 cal/g. Prepd by HNO\(_3\) nitration of hexamethylolbenzene (Ref 2).

Impact sensitivity 3 cm or less with 2 kg wt on B of M machine; wt loss in 100\(^\circ\)C heat test: 0.70% first 48 hrs, 0.32% second 48 hrs, no expl in 100 hrs


Hexamine. Same as Hexamethylenetetramine, described on p H 79ff

Hexamine Derivatives are described under Hexamethylenetetramine on pp H80 to H84

Hexamine Explosives are described as Hexamethylenetetramine Explosives

Hexamit. See Novit

Hexamite (Called Hexamite or NTD2 by the U.S. Navy; Schiesswolle 8 and TSMV 1-101 by the Germans; OtsuB or Type A by the Japanese). It consists of TNT 60/HeNDPhA 24/Al 16; greenish-gray solid, d 1.72, mp 80-90\(^\circ\)C (can be cast loaded); Ballistic Strength 130% TNT (due to the addn of Al); Briscante ca 46 g sand crushed vs 43 g for TNT; Explosion Temp 200-260\(^\circ\)C, Deton Vel 6900 m/sec; Impact Sens (Pic Ars app w 2 kg wt) 10" vs 14" for TNT; Rifle Bullet Sensitivity

detonates from the impact of 30 cal bullet, fired from a distance of 90 ft. It is stable and does not attack metals, but it is very toxic

Uses:
Germany: in mines, torpedoes, depth charges and Naval demolition containers

Japan: As type A explosive it was intended to be used in underwater ammunition to replace the type 97 explosive, TNT 60, HeNDPhA 40


HEXAMINES are Ammonium or Ammoniates (described in Vol 1, pp A275-L to A286-R) contg six NH\(_3\) groups. The following explosive hexamines are described in Vol 1:

Hexamminecadmium (II) Bromate, [Cd(NH\(_3\))\(_6\)](BrO\(_3\))\(_2\) (p A277)
Hexamminecadmium (II) Chlorate [Cd(NH\(_3\))\(_6\)](ClO\(_3\))\(_2\) (p A277)
Hexamminecadmium (II) Perchlorate [Cd(NH\(_3\))\(_6\)](ClO\(_4\))\(_2\) (p A277)
Hexaminechromium (III) Azide [Cr(NH\(_3\))\(_6\)](N\(_3\))\(_3\) (p A277)
Hexaminechromium (III) Nitrate [Cr(NH\(_3\))\(_6\)](NO\(_3\))\(_3\) (p A277)
Hexaminechromium (III) Perchlorate [Cr(NH\(_3\))\(_6\)](ClO\(_4\))\(_2\) (p A277)
Hexaminecobalt (III) Chlorate, Monohydrate or "Luteocobalttriammine Chlorate," [Co(NH\(_3\))\(_6\)](ClO\(_4\))\(_2\) \(\cdot\) H\(_2\)O (p A280)
Hexaminecobalt (III) Chlorite, Monohydrate, [Co(NH\(_3\))\(_6\)](ClO\(_4\))\(_2\) \(\cdot\) H\(_2\)O (p A280)
Hexaminecobalt (III) Hexanitrocobaltate, [Co(NH\(_3\))\(_6\)]Co(NO\(_3\))\(_2\) (p A280)
Hexaminecobalt (III) Iodate, [Co(NH\(_3\))\(_6\)](IO\(_4\))\(_2\) (p A280)
Hexaminecobalt (III) Nitrate [Co(NH\(_3\))\(_6\)](NO\(_3\))\(_2\) (p A280)

Note: Addnl info is given below

Hexaminecobalt (III) Perchlorate, Monohydrate or "Luteocobalttriammine Perchlorate" [Co(NH\(_3\))\(_6\)](ClO\(_4\))\(_2\) \(\cdot\) H\(_2\)O (p A280)
Hexaminecobalt (III) Perchlorate, [Co(NH\(_3\))\(_6\)](ClO\(_4\))\(_2\) (p A280)
Hexaminecobalt (II) Perchlorate, [Co(NH\(_3\))\(_6\)](ClO\(_4\))\(_2\) (p A280)
Hexamminecopper (II) Chlorate, 
\([Cu(NH_3)_6]_2(ClO_3)_2\) (p A281)

Hexamminenickel (II) Azide, 
\([Ni(NH_3)_6]_2(N_3)_2\) (p A281)

Hexamminenickel (II) Bromate 
\([Ni(NH_3)_6]_2(\text{BrO}_3)_2\) (p A281)

Hexamminenickel (II) Chlorate, 
\([Ni(NH_3)_6]_2(ClO_3)_2\) (p A281)

Hexamminenickel (II) Iodate, 
\([Ni(NH_3)_6]_2(\text{IO}_3)_2\) (p A281)

Hexamminenickel (II) Perchlorate, 
\([Ni(NH_3)_6]_2(ClO_4)_2\) (p A281)

Hexamminezinc (II) Chlorate, 
\([Zn(NH_3)_6]_2(ClO_3)_2\) (p A282)

Hexamminezinc (II) Nitrate, 
\([Zn(NH_3)_6]_2(NO_3)_2\) listed on p A282 but not described. It refers to Ref 23, which is D.B. Donskay & M.A. Portnov, ZhObshchKhim 9, 526-31 (1939) & CA 33, 9091 (1939)

Ref: See Vol 1, pp 1, 3, 7, 8, 10, 11, 12a, 14, 15, 16, 22, 23, 24, 39, 41, 42, 43, 46, 49, 67, 68, 69, 71, 72

Note: Destruction of cobaltic ammimes, such as of hexammine by means of thiosulfate in acid soln was described by L.M. Ordlo, ZavodLab 8, 502 (1939) & CA 36, 6935 (1942), which is listed in Vol 1, p A286-R

Hexammine, Calcium. Probably Ca(NH_3)_6. The compd decomposes explosively when heated in vacuum. The temp of explosion depends on the degree of vacuum. Products of the decomp are Ca nitride & hydride and hydrogen (Ref 1)

However, later studies (Ref 2) suggest that Ca Hexammine explosions are due to impurities which catalyze its decomp


Hexammine Chromic Perchlorate. \([Cr(NH_3)_6]_2(ClO_4)_2\) mw 452.57. This inorg coordination compd was studied from the standpoint of its stab sensitivity vs loading density for possible use as an initiating compd. It was found to increase in sensitivity until an apparent max point was reached. The stab sensitivity of the compd also appeared to be affected by humidity. The compd exhibited a sensitivity of 28 inch-ounces as compared to 12 inch-ounces for material dried over Ca sulfate (Ref 4)

See also Table A in Vol 1 of Encycl, p A277 and the following Refs


Hexamminecobalt (III) Nitrate. \([Co(NH_3)_6]_2(NO_3)_3\) mw 347.20, N 36.32%; soln in hot sl acid w/ insol in alc; \(\Delta H^e = -311.5 \text{ kcal/mole}\) (Ref 2)

Prepd by reacting a cobalt nitrate hexahydrate soln with AN and concd ammonia and then air oxidizing Co (II) to Co (III) or reacting hexamine cobalt (II) chloride with concd nitric acid (Ref 1); by reacting Co(NH_3)_6 with AN in liq ammonia at 100° in an autoclave (Ref 3)

DTA and gas evolution curves are given in Ref 4. Explosive characteristics such as: output, impact sensitivity, and autoignition temp are given in Ref 6. Additional impact and friction sensitivity data are found in Ref 5 which also gives confined deton vel as 3000 m/sec & unconfined deton vel as 2700 m/sec


Hexammons. See Ghekamony in Vol 6, pG73-R
HEXANE AND DERIVATIVES

n-Hexane (called Hexan in Ger); CH₃(CH₂)₄CH₃; mw 86.17; colorless vol liq, mp -94.3°C, bp 68.9°C, bp -9°F, auto ignition temp 500°F, d of liq 0.659 at 20°C, vap press 100 mm at 16°C, ΔHf° = -39.96 kcal/mol, refr index 1.3749; sl sol in w, sol in alc, acet, eth, and chl. Mod toxic in inhalation and ingestion; pure hexane reacts explosively with O₂ at 227°C (Ref 2). Hexane occurs in crude petroleum and can be obtained by fractional distillation. It is used as a solvent, in low temp thermometers, as a paint diluent, and as a polymerization reaction medium. It is the parent comp for the following potentially exl derivs:

Refs: 1) Beil 1, 142, (51) [105] & {374}; 2) C.F. Cullis, Bull Fr 1950, 863-8 & CA 47, 2689 (1953)

Azidohexanes, C₆H₁₃N₃, mw 127.22, N 33.04%. The following isomers are known:

2-Azidoxane, Triazooxane or Hexyl Azide.

N₃CH₂C₆H₁₁; liq, bp 156-157°C, RI 1.4325 at 20°C (Ref 2), sol in eth, Qcomb 1043kcal/mole, ΔHf° = -36.2kcal/mole (Ref 6). Prepd by heating hexyl bromide, NaN₃, aq MeOH in sealed tube for 6 hours; adding dropwise hexyl iodide to aq NaN₃ in aq PROH & refluxing (Ref 4). Low toxicity less than NaN₃ (Ref 3)

Used as a foaming agent in caprolactam polymerization (Ref 5)


2-Azidoxane. CH₃CH(N₃)C₆H₅; liq, bp 96-98°C at 160mm, RI 1.4353 at 20°C, d 0.857, opt rot +27.8° at 25°C (Ref 2). Prepd by adding 2-iodohexane to NaN₃ in aq MeOH and heating in sealed tube

Refs: 1) Beil 1, (396) 2) P.A. Levene et al

Nitrohexanes, C₆H₁₃NO₂; mw 131.20, N 10.68%. The following isomers are known:

1-Nitrohexane, O₂NCH₂C₆H₁₁; colorless fluid liq, bp 193-194°C, d 0.949 at 20°C; insol in water; sol in alc. IR and UV spectra given in Ref 2. Prepd by reacting hexane with fuming nitric acid which also gives:

2-Nitrohexane, CH₃CH(NO₂)C₆H₅; liq, bp 176°C, d 0.936 at 20°C. It pyrolizes unimolecularly with an activation energy of 40-45kcal/mole (Ref 3)


Dinitrohexanes, C₆H₁₂(NO₂)₂; mw 176.20, N 15.90%. The following isomers are known:

1,1-Dinitrohexane, (NO₂)₂CH₂C₆H₁₁; yel oil, bp 80.5°C at 1 mm, d 1.091, RI 1.4432 at 25°C (Ref 2), insol in w, sol in alc & eth. Prepd by nitrating methylhexyketone or hexane


1,2-Dinitrohexane, NO₂CH₂CH(NO₂)C₆H₅; liq. Prepd by reacting 1-hexane with an equilibrium mixt of NO₂ & N₂O₄ to give 1,2-dinitrohexane & 1-nitro-2-hexanol nitrate

Refs: 1) Beil, not found 2) G.A. Bonetti et al, USP 3192248 (1965) & CA 63, 11361 (1965)

2,4-Dinitrohexane, CH₃CH(NO₂)CH₂CH(NO₂)C₂H₅; liq, bp 177-79°C at 0.5mm, d 1.100, RI 1.4482 at 25°C (Ref 2)


3,4-Dinitrohexane, C₆H₂(CH(NO₂)C(NO₂)CH₂H; pale yellow liq, bp 50-54°C at 0.3mm, RI 1.4512 at 25°C, sol in chl. Prepd by reacting 1-nitropropane in an alkaline chloroform mixt with an aq soln of Na-persulfate, Na-acetate & Ag-nitrate (Ref 2)


Tetranitrohexanes, C₆H₁₀N₄O₈; mw 266.20, N 21.05%. Two isomers are known:

1,1,1,3-Tetranitrohexane, CH₃(CH₂)₂CH(NO₂)CH₂C(NO₂)₃; crystals (from hexane), mp 37.7-38°C; was prepd by adding 1,1,1-trinitro-3-aci-nitrohexane to a vigorously stirred soln of urea in MeOH (Refs 1 & 4)
**3,3,4,4-Tetranitrohexane**, 
C\textsubscript{4}H\textsubscript{6}C(NO\textsubscript{2})\textsubscript{2}C(NO\textsubscript{2})\textsubscript{2}C\textsubscript{6}H\textsubscript{5}; wh waxy crys, mp 106–07°; was prepd by treating 3,4-dimethoxy-
hex-3-en with liq N\textsubscript{2}O\textsubscript{4} at 85° (Refs 2 & 3) 
**Refs:** 1) Beil, 1, <357>  
2) Beil, 1, <358>  
3) C.E. Gabriel, JACS 77, 1293 (1955) & CA 50, 1571 (1956)  
4) S.S. Novikov et al, Bull Acad Science(USSR) 1959, 1765 (Engl)

Hexadiol and Derivatives

1,6 Hexanediol or 1,6 hexamethylenglycol. 
CH\textsubscript{2}OH(CH\textsubscript{2})\textsubscript{4}CH\textsubscript{2}OH; mw 118.17; low melting 
needles-like solid or colorless liq, mp 42°, bp 250°; n\textsubscript{D}26° 1.3953; sol in water & alc; 
sol in hot eth; low toxicity. Prepd by reduction or hydrogenation of adipic acid 
esters. Used as solvent and resin intermediate. It is the 
parent compd for the following derivs, none of which 
appears to be explosive 
**Refs:** 1) Beil, 1, 484 (251) [551] & (220)  
2) Cond Chem Dct (8 Edit) (1971) p 443

Dinitrohexanediols, (HO)\textsubscript{2}C\textsubscript{6}H\textsubscript{10}(NO\textsubscript{2})\textsubscript{2}; mw 
208.20, N 13.46%. The following isomers are known: 

2,5-Dinitro-3,4-hexanediol. 
H\textsubscript{2}CCH(NO\textsubscript{2})\textsubscript{2}CH(OH)CH(OH)CH(NO\textsubscript{2})\textsubscript{2}CH\textsubscript{3}; 
crys, mp 111-112° (from EtOH/water), 163-
165° (from EtOH) (Refs 2, 3 & 4). Prepd by 
treating glyoxal in alk soln with EtNO\textsubscript{2}, 
neutralizing Na-salt with HOAc, & extracting 
with nitromethane (Ref 1); by adding O\textsubscript{2}\textsubscript{N} 
(CH\textsubscript{2})\textsubscript{4}NO\textsubscript{2} to qaq NaOH and when salt form 
is complete treating with qaq formal and HOAc 
(Ref 2 & 3) 
**Refs:** 1) Beil, not found 2) H. Plaut, USP 
2616923 (1952) & CA 49, P 11704 (1955)  
3) H. Feuer & R. Miller, JOrgChem 26, 1348-57 
(1961) & CA 56, 1334 (1961)  
4) H. Feuer A.T. Nielsen & C.E. Colwell, Tetrahedron, 18, 
Suppl 1, 57-64 (1963) & CA 59, 11229 (1963)  
5) E.S. Lipina, V.V. Perekalin, Ya. S. Bobovich, 
ZhObshchKim 34(11), 3653-40 (1964) & CA 
62, 9043 (1965)

5,5-Dinitro-1,2-hexanediol. 
H\textsubscript{2}C(OH)CH(OH)C\textsubscript{2}H\textsubscript{5}C(NO\textsubscript{2})\textsubscript{2}CH\textsubscript{3}, liq, RI 
1.4804 at 25°, sol in CH\textsubscript{2}Cl\textsubscript{2}. Prepd by adding 
MeOH soln of CH\textsubscript{2}C(NO\textsubscript{2})\textsubscript{2}(CH\textsubscript{2})\textsubscript{4}COCH\textsubscript{3}OH 
to cold alk soln of NaBH\textsubscript{4} & acidifying with 
H\textsubscript{2}SO\textsubscript{4} (Ref 2 & 3). Has anti-bacterial, fungicidal 
& anti-inflammatory props (Ref 4) 

**Refs:** 1) Beil, not found 2) Aerojet-General 
Corp FP 1319705 & CA 69, P9791 (1963)  
3) G.B. Linden, USP 3101378 (1956) & CA 60, 
P1589 K1964  
4) H. Gold, Nitro Compsd Proc Intern Symp, Warsaw 1963, 449-61 & CA 64, 
1184 (1966)

Tetranitrohexanediols, (HO)\textsubscript{2}C\textsubscript{6}H\textsubscript{10}(NO\textsubscript{2})\textsubscript{4}; mw 
298.20, N 18.79%, OB to CO\textsubscript{2} -37.6%. The following isomers are known: 

1,1,6,6-Tetranitro-2,5-hexanediol. 
(NO\textsubscript{2})\textsubscript{2}CHCH(OH)CH\textsubscript{2}CH\textsubscript{2}CH(OH)CH(NO\textsubscript{2})\textsubscript{2}; 
brown liq; sol in CH\textsubscript{2}Cl\textsubscript{2}. Prepd by suspending 
di-K salt of Me(NO\textsubscript{2})\textsubscript{2} in water and adding 
(CH\textsubscript{2}CHO)\textsubscript{2} & then neutralizing with sulfuric 
acid. Used as an intermediate. It is exnl 
**Refs:** 1) Beil, not found 2) H. Plaut, USP 
2544403 (1951) & CA 45, 7587 (1951)

2,2,5,5-Tetranitro-1,6-hexanediol. 
OH\textsubscript{2}CH\textsubscript{2}(NO\textsubscript{2})\textsubscript{2}CH\textsubscript{2}CH\textsubscript{2}OH; crystals, mp 
126–27° (recrystd from w); sol in Et\textsubscript{2}O & w. 
Prepd by nitrixing MeOH soln of tetranitrobutane 
with formalin and adding small amount of NaOH; 
adding 1,4-dinitrobutane toaq NaOH & then 
adding formalin & finally anaq soln of NaNO\textsubscript{2} & 
AgNO\textsubscript{3}. No expl props mentioned, but it is exp 

**Refs:** 1) Beil, not found 2) H. Feuer et al, 
JOrgChem 27, 3598 (1962) & CA 57, 14921 (1962)

2,5-Hexanedioldinitrite. CH\textsubscript{2}CH(ONO)\textsubscript{2}CH\textsubscript{2} 
(ONO)(OH)CH\textsubscript{2}; liq, bp 56° at 33mm, d 1.042, RI 
1.3978 at 25°; insol in w. Prepd by treating aq 
(CH\textsubscript{3}CHOHCH\textsubscript{2})\textsubscript{2} at 0° withaq NaNO\textsubscript{2} and then 
adding 6N HCl. The kinetics of its decompn were 
also studied 
**Refs:** 1) Beil, not found 2) L.P. Kuhn et al 
JACS 78, 2719 (1956) & CA 50, 14506 (1956)

2,5-Hexanedioldinitrate, or 2,5 Dinitroxyhexane. 
CH\textsubscript{3}CH(ONO)\textsubscript{2}C\textsubscript{2}H\textsubscript{5}CH(ONO)\textsubscript{2}CH\textsubscript{3}; mw 
208.20, N 13.46%, OB to CO\textsubscript{2} -92.2%; crys, mp 50.4°; 
vap press 0.98–1.33, 6.1–9.4 &19–31 at 20, 30 
& 40° resp, heat of fusion 13.2 kcal/mole, heat of 
vaporization 13.0 kcal/mole. Prepd by nitration of 
2,5 hexanediol. No expl props mentioned 
**Ref:** 1) Beil, not found 2) M.D. Kemp et al, 

2,3,4,5-Hexanetetronic-tetroxime Diperoxide or 
Dimethyltetranitro tetraoxime Diperoxide 

\[
\text{H}_3\text{C}_6 : \text{C} \overset{\text{O} \cdot \text{O} \cdot \text{O} \cdot \text{N}}{\text{C}} \text{C} \text{H}_3
\]

; mw 186.15, N 30.10%
OB to CO₂ −77.4%; crysts (from eth), mp 181°. Prepd by reacting (C₆N₄O₆)C(NOH)CH₂I in ether with N₂O₅; or reacting the above oxime with 50% nitric acid and aq KMnO₄. No explosive props are mentioned.

Refs: 1) Beil, not found 2) G. Ponzio & V. Bernardi, AttiAccadSciTorino 59, 693 (1924) & CA 19, 469 (1925) 3) G. Ruggieri, Gaz chim ital 54, 72 (1924) & CA 19, 2187 (1925) 4) Nothing more recent found in CA

**Hexanetriol and Derivatives**

1,2,6-Hexanetriol. OHCH₂CH(OH)(CH₂)₂CH₂OH; mw 134.20; colorless liq, mp 32.8°, bp at 5mm. 178°, flp 380°F, d 1.10, RI 1.476 at 20° (Ref 3); sol in w; low toxicity. Prepd by hydrogenation of 2-hydroxyxadic aldehyde over Raney Ni (Ref 2). Used as resin intermed, softener & solvent.


1,2,6-Hexanetritrintritate (HTTN). O₂NCH₂ CH(NO₂)₂C₃H₈CH₂NO₃; mw 269.20, N 15.61%, OB to CO₂ −50.5%; light tan liq melts below −51°, d 1.39, s ol water, v sol eth, acet, non-hygroscopic. Obtained by mixed acid nitrations of hexanetriol at 0°, separating HTTN, washing and stabilizing with NaHCO₃ soln. Proposed as a substitute for NG in dynamos.

It has an impact of 76 cm on B of M machine with 2 kg weight & will not explode in friction pendulum test. Its vacuum stability is 0.8 cc of gas in 40 hours at 100°.

Refs: 1) Beil, not found 2) J.B. Bronstein Jr, USP 2683164 (1954) & CA 48, 11062 (1954)

**Hexanexol.** See Dulcitol and Derivatives in Vol 5, p D1567-R

**Hexanites or Schiesswolke neuer Art. (in Ger)** are cast charges of 60/40 TNT/Hexanitrodiphenylamine.

Ref: Anon "Explosivstoffe" Wasag-Chemie AG, Essen, (1961) p 76

**Hexanitrate Dulcitol.** See Dulcitol Hexanitate in Vol 5, p D1567-R

**Hexanitroazobenzene.** See under Azobenzene Vol 1, p A649

**Hexanitrocabanilide or sym-Dipicrylurea.** See N,N'-Bis(2,4,6-trinitrophenyl)-urea in Vol 2, p B156-L

**Hexanitrocompounds** are described under corresponding parent substances.

**Hexanitrodiphenylamine.** See under Diphenylamine in Vol 5, p D1434-R

**Hexanitrodiphenylsulfide.** See under Diphenylsulfide in Vol 5, p D1477-R

**Hexanitrodiphenylsulfone.** See under Diphenylsulfone in Vol 5, p D1480-L

**Hexanitrodiphenyloxide.** Same as Hexanitrodiphenylether, Vol 5, p D1453-R

2,4,6,2',4',6'-Hexanitrodiphenyl-ethylene-dinitramine or N,N'-Di(2,4,6-trinitrophenyl-dinitramino)-ethane (Code named Bitetrol, Ditetrol or Octyl). See 1,2-Bis(2',4',6'-trinitro-nitranilino)-thane in Vol 2, p B131-R

**Hexanitrodiphenyl Guanidine.** See N,N'-Bis(2,4,6-trinitrophenyl)-guanidine in Vol 2, p B154-R

2,4,6,2',4',6'-Hexanitrodiphenylmethylamine. See under Diphenylmethylamine in Vol 5, p D1468-L

N,N'-Hexanitrodiphenylpropane-1,3-dinitramine. One of the names for Methylene Ditetryl described as 1,3-Bis(2',4',6'-trinitro-N-nitranilino)-propane in Vol 2, p B133-L

N,N'-Hexanitrodiphenyl-propylene-1,3-dinitramine (Code named Methylene Ditetryl). See 1,3-Bis(2',4',6'-trinitro-N-nitranilino)-propane in Vol 2, p B133-L

**Hexanitrodiresorcin or 2,4,6,2',4',6'-Hexanitro-3,5,3',5'-tetrahydroxydiphenyl.** See Hexanitro-biresorcinol under Biresorcinol and Derivatives in Vol 2, p B127-L
Hexanitrodiresorcinal. See Hexanitrobiuresorcinal under Biresorcinal and Derivatives in Vol 2, p B127-L.

Hexanitrodulcita. See under Dulcitol Hexanitrate in Vol 5, p D1567-R.

Hexanitroethane. See under ethane and derivatives in Vol 5, p E149-L.

Hexanitrohydrazonobenzene or N,N'-Dipicrylhydrazine. See Vol 5, p D1463-L.

Hexanitrohydrocellulose. See under Hydrocellulose Nitrates in this Vol.

Hexanitroinositol. See under Inositol and Derivatives in this Vol.

Hexanitromethylphenylether or Hexanitro-4-methyldiphenyl ether. \( (\text{O}_2\text{N})_3\text{C}_6\text{H}_2\text{OC}_6\text{H}(\text{CH}_3) \) \( (\text{NO}_2)_3 \) (Note: The position of the NO\(_2\) groups has not been established; mw 452.23, N 18.59%, OB to CO\(_2\) -53.1%, mp 103.5°). Crystals (from hot alcohol); may be prepd by boiling 4-nitro-4'-methylphenylether (or 4-nitro-2'-methylphenylether) with strong nitric acid for 45 minutes and then pouring the solution into a large amount of cold water. This separates a light yellow oil which turns into crystals that very slightly sol in water, sol in benzene, ethyl acetate, carbon disulfide, and hot alcohol; difficult to dissolve in cold alcohol. When ignited in the flame of a lamp it flashfugates.

Recently the 3',5'-Methyl-2,2',4,4',6,6'-hexanitro-diphenyl Ether, mp 141-42°, has been prepd by the mixed acid nitration of the corresponding pentanitro diphenyl ether (Ref 4).


3,3,5,7,7 Hexanitro-1,9-nonane Diisocyanate. mp 127 ± 1°. Prepd by heating at 80° 4,4,6,6,8,8 hexanitro-1,11-undecanediocic acid (mp 84 ± 1°) which in turn is made by treating the corresponding dibasic acid with sodium azide. The diisocyanate is claimed to be useful in the prep of high energy polyurethane propellants.

Ref: 1) Beil, not listed. 2) K. Klager USP 2978475 (1961) & CA 55, 19799 (1961)

Hexanitrooxanilide or N,N'-Dipicyroxamide. See under Oxamide.

Hexanitrophenolanthrone. The name given in JCS 100(I), 656 (1911) for the compd listed in Vol 2, p B147-L as 10,10-Bis(4-hydroxyphenyl)-x,x,x,x,x,x-hexanitro-9-anthranone.

Hexanitrosobenzene. See under Benzene and Derivatives in Vol 2, p B45-L.

Hexanitrosorbitol. See Sorbitol Hexanitrate under Sorbitol and Derivatives.

Hexanitrosulfobenzene. Same as Hexanitrodiphenylsulfone, Vol 5, p D1480-L.

Hexanols and Derivatives

Hexanols, \( \text{C}_6\text{H}_{14}\text{O} \), mw 102.17. The following isomers are known:

1-Hexanol or n-Hexyl Alcohol,
H\(_3\)C(CH\(_2\))\(_4\)CH\(_2\)OH; colorless liq., sp gr 0.822 (15/4°), bp 157.5°, RI 1.4169 (25°).

2-Hexanol or Butylmethylcarbinol,
H\(_3\)CC(=O)CH\(_2\)CH\(_2\)CH\(_3\); colorless liq., sp gr 0.818 (16.8/4°), bp 140°, RI 1.4126 (Ref).

3-Hexanol, \( \text{C}_3\text{H}_7\text{CHOHCH}_3 \); colorless liq., sp gr 0.818 (20/4°), bp 135°.

Ref: 1) Beil 1, 407-08, (202), [437-38] & (1660).

Dinitrohexanols, \( \text{C}_6\text{H}_{12}\text{N}_2\text{O}_5 \); mw 192.20, N 14.58%. The following isomers are known:

2,6-Dinitro-1-hexanol.
OH(CH\(_2\))\(_2\)(NO\(_2\))\(_2\)(CH\(_2\))\(_3\)CH\(_2\)NO\(_2\); yel liq., bp decomp, RI 1.4745 at 20°; sol in CH\(_2\)Cl\(_2\). Prepd by reacting 1,5 dinitropentane withaq NaOH then formalin (37% soln) and acetic acid.


5,5 Dinitro-2-Hexanol. \( \text{H}_2\text{CCH(OH)(CH}_3\text{)}\text{O_2} \text{N}_2 \text{O}_2 \text{C}_6\text{H}_3 \); yel liq., bp 104°at 0.6mm, RI 1.4656 at 20°; d 1.24; sol in eth. Prepd by addingaq NaBH\(_4\) to MeOH soln of CH\(_3\)CO(CH\(_2\))\(_2\)CH\(_3\) with the pH of
mixt constantly adjusted with H₂SO₄ at 3–4
Refs: 1) Beil, not found 2) H. Shechter et al, JACS 74, 3664 (1952) & CA 47, 5886 (1953)
1-Nitro-2-Hexyl Nitrate. O₅NCH₂(CHONO₂)₃(CH₂)₃CH₃; prepd. by first obtaining 1-nitro-2-
hexyloperoxynitrate by reacting 1-hexene in CCl₄ with
Oxygen & N₂O₄ and then treating with NO₂.
It is used as a fuel additive & intermediate in
preps of surfactants, insecticides, etc
Refs: 1) Beil, not found 2) D.R. Lachowicz
& K.L. Kreuz USP 3282983 (1966) & CA 66, 10577 (1967)

Hexanones and Derivatives
Hexanones, C₆H₁₂O; mw 100.16. The following
isomers are known:
2-Hexanone or n-Butylmethyl Ketone,
CH₃COC₆H₅, colorless liq, sp gr 0.816 (15/4°), fr p
–56.9°, bp 127.2°, fl p 95.8°F. Very toxic
Ref: Beil 1, 689, (354), [745] & [2826]
3-Hexanone, C₉H₆COCH₂C₂H₅; colorless liq, sp gr
0.813 (21.8/4°), bp 123–24°
Ref: Beil 1, 690, (354), [746] & [2830]

Dinitrohexanones, C₆H₁₈N₂O₄; mw 190.18, N
14.73%. The following isomers are known:
5,5-Dinitro-2-hexanone,
H₃CCO(CH₂)₂C(NO₂)₂CH₃; liq, sp gr 1.264, RI
1.4607 (20°), bp 110° at 1.3 mm; sol in CH₂Cl₂.
Prepd. by adding dioxane solns of CH₂(NO₂)₂
& PhCH₂NMMe₃OH to C₂H₅CH₂CH₂. After
65 hrs stirring mix is poured into w & acid-
ified and product extracted with CH₂Cl₂. It is expl
Refs: 1) Beil, not found 2) H. Shechter &
L. Zeldin, JACS 73, 1276 (1951) & CA 45,
9459 (1951)
4,4-Dinitro-3-hexanone,
H₂C₅COC(NO₂)₂C₂H₅; liq, bp 69–71° at 1 mm, RI
1.4548. Prepd. by reacting Etc:EtC with N₂O₄ in
dry Et₂O and distilling the resulting oil. It is expl
Refs: 1) Beil, not found 2) J.P. Freeman &
W.D. Emmons, JACS 70, 1712 (1957)

HEXAZADIENES (Hexaziden in German).
Hexazadienes are derivatives of NH₂:NNH:NNH.
N:NN Some of them are explosive, for instance:
1,6-Bis-[tetrazolimidylen-(5)]-hexazadiene or
1,6-Di-[tetrazoly-(5)-hexazadiene] (Bisdiazot-
tetrazolhydrizide old German),

\[
\begin{align*}
N,NNH< & C:N:N:N=N:N \leq NH,N
\end{align*}
\]

(3 of all known organic compounds, this is the
richest in nitrogen), mp (explodes at
about 90°). Leaflets, was prepd. by Hofmann
et al from 5-diazotetrazol hydrochloride, hydra-
zine hydrochloride and sodium acetate at a
very low temperature

According to Karrer (Ref 4), it may be
prepd. by diazotizing 5-aminotetrazole and
coupling the resulting compound with hydra-
azine

Stable at room temp; powerful explosive. If
dry crystals are scattered loosely on a
piece of cardboard and one of the crystals
 touched with a flame or heated object (rod etc),
the material will detonate with such brisanse
that each crystal will puncture the cardboard
Refs: 1) Beil 26, (123-24) 2) K.A. Hofmann
& H. Hock, Ber 44, 2953 (1911) 3) F.R.
Benson, Chem Revs 41, 9 (1947) 4) Karrer
(1950 ed) p 803 (under Tetrazole)

Hexazidocuprate-Lithium Salt. Li₄ [Cu(N₃)₆]; rw
343.49, N 73.42%; crystals with 3H₂O which is
lost above 120°. The anhydrous compound ex-
plodes at 224-5°. Prepd. by dissolving Cupric
Azide in an or alc soln of Lithium Azide
Refs: 1) Gmelin, Syst No 60, Teil B, 150 (1958)
2) M. Straumanis & A. Cirulis, ZAnorgChem 252,
9 (1943) & CA 38, 3563 (1944)

Hexazidotannate-Sodium Salt. Na₂Sn(N₃)₆,
probably Sn(N₃)₂·2NaN₃; mw 416.85, N 60.50%
white solid, sl sol in eth or benz. Prepd. by heating a
tetrahydrofuran soln of SnCl₄ & NaN₃. Hydro-
lyzes in moist air. Explodes on boiling over an
open flame
Refs: 1) Gmelin, Syst No 46, Teil 61, 145–46
(1972) 2) E. Wiberg & H. Michard, ZNatur-
forsch 9b, 500 (1954) & CA 49, 768 (1955)

Hexazobenzene. See Diazidobenzene in Vol 2,
p B42-R
Hexenes and Derivatives

Hexenes or Hexylenes, C<sub>8</sub>H<sub>12</sub>; mw 84.18. Six isomers are described in Beil. Three of them: 1-, 2-, and 3- are described below together with their nitrated derivs.

1-Hexene or Hexylene, C<sub>8</sub>H<sub>10</sub>::CH<sub>2</sub>; mw 84.16, colorless liq, fr p < -139.8°, bp 63.6°, fl p < -15°, d 0.673, RI 1.3876 at 20°; insol in w; sol in alc. Prepd by reaction of PrMgBr & allyl bromide. Highly flammable and moderately toxic

Refs: 1) Beil 1, 215, (89), [191], (800) & <828> 2) CondChemDict, 8th Edit (1971), p 444

1-Nitro-1-hexene. H(NO<sub>2</sub>)<sub>2</sub>:CH<sub>2</sub>H<sub>2</sub>; mw 129.18, N 10.84%; liq, bp 54–55° at 1.5 mm, RI at 25° 1.4581. Prepd by refluxing BuCH(OAc)CH<sub>2</sub>NO<sub>2</sub> in benzene containing Na-carbonate


2-Nitro-1-hexene (called 2-Nitro-hexen(1) in Ger), H<sub>2</sub>C:CH(NO<sub>2</sub>)<sub>2</sub>H; liq, bp 81–2° at 50mm (Ref 1), RI at 25° 1.4462. Prepd by treating Et<sub>2</sub>NHaq with aq HCHO (37%) & then with C<sub>8</sub>H<sub>4</sub>N<sub>3</sub>O<sub>2</sub> to give 2-nitrohexylidethyamine whose HCl salt is pyrolyzed to yield the desired nitroalkene (Ref 2); also by distilling under reduced pressure 2-nitro-1-hexanol & phthalic anhydride (Ref 3)


2-Hexene, α-Hexylene or 2-Methyl-2-propyl ethylene. CH<sub>3</sub>CH:CH(CH<sub>2</sub>)<sub>2</sub>H<sub>2</sub>; mw 84.16; colorless liq, fr p -136°, bp 68°, fl p -5°; d 0.686 at 20°, RI 1.3948, insol in w; sol in alc or dil H<sub>2</sub>S<sub>4</sub>; low toxicity. Prepd by reacting 3-iodohexane with sodamide; by hydrogenation of α-ethylalcohol over finely divided nickel. Used as chemical intermediate

Refs: 1) Beil 1, (89), [192], (804) & <833> 2) CondChemDict (1971), p 444

2-Nitro-2-hexene (Called 2-nitro-hexen(2) in Ger). H<sub>2</sub>CC(CH<sub>2</sub>NO<sub>2</sub>)<sub>2</sub>H; mw 129.18, N 10.84%; yellow liq, bp 82.3° at 10mm, d 0.9824, nD<sub>20</sub> at 25° 1.4572 (Ref 2); sol in benz (Ref 4). It is an eye irritant and is moderately toxic (Ref 5). Prepd by refluxing MeCH(NO<sub>2</sub>)<sub>2</sub>CH(OAc) Pr in dry benzene containing Na-carbonate (Refs 2 & 4); by heating 2-nitrohexanol with phthallic anhydride (Ref 3)


3-Nitro-2-hexene. H<sub>2</sub>CH:C(NO<sub>2</sub>)<sub>2</sub>C<sub>2</sub>H<sub>4</sub>; liq, bp 70.9° at 10mm, d 0.9797, RI at 25° 1.4545 (Ref 3), sol in benz. Prepd by refluxing C<sub>8</sub>H<sub>7</sub>CH(NO<sub>2</sub>)<sub>2</sub>CH(OAc)CH<sub>2</sub>H in dry benzene containing Na-carbonate (Ref 2). Toxicity about same as 2-nitro-hexene

Refs: 1) Beil 1, <836> 2) Same as 2 above 3) K.F. Lampe et al, JChemEngData 7 (1) 85 (1962) & CA 57, 8414 (1962)

4-Nitro-2-hexene. H<sub>2</sub>CH:CHCH:C(NO<sub>2</sub>)<sub>2</sub>C<sub>2</sub>H<sub>4</sub>; mw 129.1, N 10.8%; yellow-green liq, bp 72° at 10mm, d 0.9833, RI at 25° 1.4572, sol in benz. Prepd by refluxing CH<sub>2</sub>CH(OAc)CH<sub>2</sub>CH(NO<sub>2</sub>)<sub>2</sub>C<sub>2</sub>H<sub>4</sub> in dry benzene containing Na-carbonate

Refs: 1) Beil, not found 2) same as 2 above 5,5 Dinitro-2-hexene. H<sub>2</sub>CH:CHCH:C(NO<sub>2</sub>)<sub>2</sub>C<sub>2</sub>H<sub>4</sub>; mw 174.18, N 16.08%; liq, bp 60–70° at 1.2mm, RI at 25° 1.4555, sol in hexane. Prepd by heating 5,5-dinitro-2-hexanol with a few drops of concd sulfuric acid

Refs: 1) Beil, not found 2) G.B. Linden, Brit P 936087 (1963) & CA 61, 6920 (1964)

3-Hexene, α-Diethylthylene or γ-Hexylene. C<sub>8</sub>H<sub>7</sub>CH:CH<sub>2</sub>H<sub>2</sub>; mw 84.16, liq, bp 70°, d 0.693 at 20°. Prepd by reacting 3,4-dibromohexane with Zn dust; or by partial hydrogenation of 3-hexyne over Raney nickel

Refs: 1) Beil 1, [192], (806) & <837> 2) CondChemDict, 1971, p 444

3-Nitro-3-hexene (called 3-Nitro-hexen(3) in Ger). C<sub>8</sub>H<sub>7</sub>C(NO<sub>2</sub>)<sub>2</sub>:CH<sub>2</sub>H<sub>2</sub>; mw 129.18, N 10.84%; yellow liq, bp 70.6° at 10mm, d 0.9736, RI 1.4510 at 25° (Ref 2); sol in benzene. Prepd by refluxing C<sub>8</sub>H<sub>7</sub>CH(NO<sub>2</sub>)<sub>2</sub>CH(OAc)CH<sub>2</sub>H<sub>2</sub> in dry benzene containing Na-carbonate (Ref 1)


3,4 Dinitro-3 hexene (called 3,4 Dinitro-hexen(3 in Ger). C<sub>8</sub>H<sub>7</sub>C(NO<sub>2</sub>)<sub>2</sub>:C(NO<sub>2</sub>)<sub>2</sub>C<sub>2</sub>H<sub>4</sub>; mw 174.18,
N16.09%; light yellow needles, mp 29-32°, bp 53-5° at 1 mm, RI at 25° 1.4640 (Ref 3).
IR spectra in Ref 2. Prepared by reacting EtCl:CeCl with N₂O₄ in ether (Ref 3). Reacting 1-chloro-2-nitropropane with 9q KOH and then acidifying to a pH of 7.9. The residue after separation and washing is distilled, under vac & nitrogen atmosphere to prevent explosion, to get additional product (Ref 4)


Trinitrotoluene, C₆H₅NO₃O₆, Tetranitrotoluene, C₆H₄N₃O₈, and 

Pentanitropy, C₅H₃N₂O₁₀, the derivatives of Hexene were not found in Beil

1,1,1,6,6,6-Hexanitro-3-hexene. (NO₂)₃CCH₂CH₂CH₂C(NO₂)₃; mw 354,18, N 23.73%, OB to CO₂ -13.5%; crystals, mp 128°, d 1.77.
Prepd from 1,4 Dibromo-2-buten and Ag-nitroform. It is an explosive of about the same sensitivity as PETN

Refs: 1) Beil, not found 2) D.V. Sickman & W.F. Sager, NAVORD Rept 486, (1954) 3) Not found in CA thru 1971

Hexite. Same as Hexanitrodiphenylamine, Vol 5, p D1434-R

Hexa. Same as Hexanitrodiphenylamine, Vol 5, p D1434-R

Hexo. One of the Ger abbreviation for Hexogen (RDX). See Vol 5, p 611-L

Hexo (S-15 and S-22). Ger “Substitute Explosives” (Ersatz-sprengstoffe), constg Hexogen. See Vol 5, p El22, Table E15

Hexocine (Fr.). Mixture of RDX and Wax

Hexogen (H). Ger for Cyclonite or RDX. Also known as W-Salz, E-Salz, K-Salz, SH-Salz and KA-Salz, depending on the method of prep

Ger WWII methods of prep are described in Vol 3 of Encycl, pp C613-R to C614-R, under CYCLOTRIMETHYLENE TRINITRAMINE

Hexolit. Same as Naplot (See below), but with Hexogen (RDX) in lieu of Pentrit (PETN)

Note: Naplot developed during WWII by the

Deutsche Spengchemie GmbH was a type of NC-DEGDN-PETN propellant or expl. Two comps are listed in PATR 2510, (1958), p Ger 117-R:
a) Naplot in tubes: NC (12.6%N) 34.1, DEGDN 35.0, PETN (unwaxed) 35.0, stabilizer 0.75, MgO 0.05, graphite 0.1%, calorific value 1300 cal/g
b) Naplot in sticks: NC 29.1, DEGDN 20.0, PETN 50.0, stabilizer 0.75, MgO 0.05 and graphite 0.1% (Ref 1)

Berkovic (Ref 2) states that Hexolit is composed of RDX 90 & TNT 10 and claims that it has low impact sens

Refs: 1) Dr. H. Walter, formerly of PicArsn; private communication (1960) 2) M. Berkovic, Explosivst 12, 207 (1964)

Hexolites (Fr.). Mixtures of RDX (hexogene in Fr) and TNT (tolite in Fr). Same as Cyclotolites

Hexone T4. Same as Cyclonite or RDX, Vol 3, C611-R

Hexonite (Swiss). Hexonites are plastic explosives, proposed by Stettbacher, consisting of RDX and NG, with or without Collodion Cotton. For instance:
a) RDX 20-50 & NG 80-50%  b) RDX 50, NG 46 & Collodion Cotton 4%

Comparative brisance tests with 50/50 HXogen and 50/50 Pentrit by the iron plate test showed that Hexonite is not as effective as Pentrit (Ref 1). It is also claimed that RDX in Hexonit does not form such a homogeneous mass with NG and NC as does PETN in Pentrit (Ref 2), and it tends to exode. NG was used by the Germans for loading torpedo heads

Refs: 1) A. Stettbacher, Nitrocellulose 4, 224 (1933) 2) A. Stettbacher, Pólvoras y Explosivos, G. Gili, Buenos Aires (1952) p 111

Hexonyl. Same as Hexanitrodiphenylamine, Vol 5, p D1434-R

Hexoplast 75. A plastic explosive, developed during WWII at the Krümmel Factory of Dynamit AG. It contained RDX 75, NC 1.2 to 1.4, liquid DNT 20.0 and TNT 3.8 to 3.6%. This mixture was prepd by heating the required amount of RDX to 90° in a Werner-Pfeiferer mixer, and blending it with a small amount of
NC. This was followed by the addition of a DNT-TNT mixture and further blending. By using this order of addition, lumping was avoided.

The mixture was put out in cylinders about 220 mm long by 28 mm in diameter. Due to difficulty with direct cap initiation, a booster was provided. It consisted of compressed, phlegmatized PETN pellets about 40 mm long by 21 mm diam and equipped with a detonator well 20 mm deep.

**Note:** This explosive was developed as a substitute for the plastic explosive, which used RDX plus American vaseline, because the latter component was no longer available in Germany. This vaseline, called "long fibrous" by Meyer, had much greater adherence than vaselines manufactured in other countries.


**Hexose, Pentanitrate.** See Glucose Pentanitrate in Vol 6.

**Hexotol,** which corresponds to the US & Brit Cycloptol, is a Swedish expl mixture usually containing 60% Hexogen (RDX) & 40% Tol (TNT).

It has been manufd by AB Bofors-Nobelkrut in the form of granules. It corresponds to US Composition B-2 which is described in Vol 3, Table on p C479. When 1% of synthetic wax is added to 40/60 mixture, the expl becomes Composition B. Hexotol has been used in Sweden for cast-loading shells at a density of 1.65-1.70. It is almost as insensitive to impact as TNT, but much more sensitive to initiation. Its deton velocity is 7800 m/s, its fragmentation power is higher than that of TNT and it is as stable as RDX in the vacuum stability test.


**Hexotonal.** A Swedish expl mixture of Hexogen (RDX), TNT, and finely divided Al. Small amounts of desensitizer (wax), and in some formulations carbon black, are added. Examples of comps: a) RDX/TNT/Al/Wax—40/44/15/1 or 40/40/15/5. b) RDX/TNT/Al—30/50/20, plus 1% wax & 1.5% carbon black. Hexotonal is usually prepared starting from Hexotol 60/40 or 50/50.

Hexotonal has been manufd by AB Bofors for use as a cast filling at d 1.68-1.74. Its sensitivity to impact and friction is moderated by an addn of wax, its sensitivity to initiation is comparable to that of Hexotol, but bursting effect and fragmentation power are higher. (Ref 3, p 203)

The following tests are described in Ref 2, pp 203-04: a) Content of RDX, plus Al b) Content of Al c) Content of desensitizer plus Al d) Content of desensitizer e) Content of TNT f) Acidity g) Explosion temperature (as described on pp 60-62 of Ref 2), and h) Moisture content.

Accdg to Ref 1, Hexotol consisting of RDX/TNT/Al—40/45/15 was used in 40mm shells.

*Ref: 1) Anon, "Ammunition for 40mm Automatic Gun L/70 MV 1000 m/sec," Sweden, (Nov 1953) 2) AB Bofors, same as Ref under Hexotol, pp 203-204.*

**HEXYL.** See Hexanitrodiphenylamine under Diphenylamine in Vol 5, p D1404-R.

**Hexylamines.** See Aminohexanes in Vol 1, p A 215-R.

**Hexylanilines 2,4,6-Trinitro.** (NO2)3C6H4NHCH2C5H11; mw 312.32, N 17.94%, OB to CO2 —133.2%; yellow plates (from CS2) mp 70-70.5°; sl sol in eth or ligroin; sol in alc or benz; v sol in chif. Prepd from Picrylchloride & hexylamine in alc soln.

*Refs: 1) Beil 12, 764 2) E.V. Behrens, Rec 14, 38 (1895).*

**Hexylene Oxonide.** C6H12O3; mw 132.18; mobile oil, bp 60° at 12mm, d 0.994 at 23°, nD 1.4059 at 25°. Sl sol in water with slow decomposition; sl sol in pet eth. Exploses if heated strongly above 60°. Prepd by reacting hexylene (in hexane or EtCl2) with 15% ozone mixture.

*Refs: 1) Beil 1, (90) 2) C. Harries & K. Haefner, Annalen 374, 331 (1910).*
Hexyl nitrate. C₅H₁₀NO₂; mw 145.18, N 9.65%; liq, bp 46° at 1 mm, d 0.9745. RI 1.481 at 25°, sol in eth & EtCl₂ (Ref 2). Prep. by reacting hexyl chloride with silver nitrate in dry MeCN (Ref 3), or directly with 98% nitric acid (Ref 5); addition of N₂O₅ in EtCl₂ to hexanol (Ref 4).


Hexyl nitrate. C₅H₁₀NO₂; mw 129.18, N 10.84%; liq, bp 50° at 48 mm (Ref 2), 64° at 76 mm; RI 1.399 at 20° (Ref 3). Prep. by heating C₅H₁₀NO₂ with HCONMe₂ & NaN₂O (Ref 4); C₅H₁₀Br + HCONMe₂ + NaN₂O (Ref 5).

4) N. Kornblum et al, JACS 78, 1501 (1956)
5) ibid, 1497 (1956)

HEXYLS. Designation for 2,4,6-Trinitro-tris (alkylnitramino) benzenes, to be described under Nitramines.

3-Hexyne and Derivatives

3-Hexyne; 3-Hexyne or Diethylacetylene. C₂H₄; mw 82.14; colorless liq, d 0.726 at 25°, refr index 1.4112; ins in water; v sol in alc or eth. Prep. by reacting 3,4-dibromo-3-hexene with Zn dust in boiling alc.

Ref: Beil 1, 229 & 980

No azido or nitro derivatives of 3-hexyne were found in Beil or CA. However, US Rubber Co & Navord reports describe:

1,1,1,6,6-Hexanitro-3-Hexyne. (NO₂)₅C₂H₄; mw 352.16, N 23.87%. OB to CO₂ 91.1%; crystals, mp 129-130°, d 1.73. Prep. from 1,4-dibromo-2-butylene and Ag-nitroform. It has about the same explosive sensitivity and power as PETN. Its hot bar ignition temp is 205°.

3) J.C. Conly et al, US Rubber Co Quart Prog Repts on Contr NOrd 10129 (Nov 1952–Feb 1954)

Heyrovsky, Jaroslav (1890-1967) received the 1959 Nobel Prize in Chemistry for his discovery of polarography and for his role in the development of this technique which has become such an important tool in electrochemistry, analytical chemistry and many other areas of science and technology. He was for many years a professor of physical chemistry at the Charles University of Prague. When the Prague Polarographic Institute was founded in 1950, Heyrovský became its first director. Obituary notices for Prof Heyrovský appeared in many scientific journals testifying to his widespread recognition and esteem.


Hidrolitas. (Span) Explosives contg either PETN or RDX, paraffin and moist NH₄NO₃. They are less sensitive to impact than either PETN or RDX, but are less powerful.

Ref: Vivas, Feigenspan & Ladreda 2, (1946), p 288

High-Blast Explosive Filler. A MAX-2 composition consisting of aluminum 62, Composition A-4 38 & graphite (added) 1.5% gave superior positive impulse output in 37mm shell. Positive impulse was determined by using piezoelectric gages. Max damage occurs if the expl detonates immediately after the shell fully penetrates the skin of an aircraft.

Ref: S. D. Stein, PATR 2523 (May 1958)

High-Blast Metal Oxygen Expls. Patented expls especially designed for aircraft destruction. They achieve enhanced blast effects through metal-O reactions and are claimed to be less affected by altitude than other HEs. The composition contains: 54(50/50 AI/Mg), 36 AP, 4 TNT & 6% RDX or 54 (50/50 AI/Mg), 28 AN, 10 Tetryl & 8% TNT
High Capacity (HC) Bombs. These British bombs used for general bombardment purposes are described in Ref.

High Density Explosives. Explosives of high density are prep'd by heating AN with freezing point depressants such as urea to effect only partial liquefaction of AN and adding to the mixture Cu-Si, Fe-Si, Al, or aromatic nitrocompounds
Ref: C. A. Woodbury & P. G. Wrightsman, USP 2,063,572 (1936), CA 31, 865 (1937)


High Explosives (HE). A high explosive is a substance that undergoes extremely rapid chemical reaction, when properly initiated, to produce gaseous products at high temperature and pressure. These products are then capable of doing useful work as they expand. It has been customary to divide HE into two categories: *primary* explosives and *secondary* explosives.
This distinction is more in kind than in degree. Small quantities of primary or initiating explosives usually detonate when exposed to flames or high temperatures while secondary explosives usually burn or deflagrate under these conditions. However under slightly altered conditions primary explosives can be made to deflagrate and secondary explosives can be made to detonate. Examples of primary explosives are: Lead Azide, Mercury Fulminate, DDNP, etc
Examples of secondary explosives are: PETN, RDX, HMX, Tetryl, TNT, as single HE compus and Comp B, Comp C, PBX 9404, Dynamite ANFO (Ammonium Nitrate/Fuel Oil) as HE mixtures

It has also been traditional to distinguish HE from "low" or propellant explosives. Here again the distinction is not clear-cut. Although a "low" explosive such as *Black Powder* probably cannot detonate, most smokeless powders can. Conversely, most secondary, explosives can be made to deflagrate. The main difference between detonation and deflagration lies in the mode of propagation. Deflagration propagates via heat transfer such as conduction or radiation, while detonation propagates via a shock wave mechanism. As already stated both modes of action—detonation or deflagration are possible in most explosives.

The above classifications are not strictly applicable to the *sensitivity* of an explosive. Although it is true that most primary explosives are more "sensitive" than most secondary explosives, certain low explosives, e.g. Black Powder, are more "sensitive" than many secondary or even primary explosives.
See also DETONATION, EXPLOSION AND EXPLOSIVES Introduction and Definitions in Vol 4, p D217-L and also History of Explosives and Related Items in this Vol

Written by J. ROTH

High Explosive Projectile (Shell): The HE projectile is one in which the percentage of filler to total weight is as high as possible, consistent with other requirements, such as good fragmentation or demolition. The projectile consists of the shell proper and the fuse. The body of the shell is generally in the shape of a steel cylinder, having a tapered portion near the nose ("ogive"). Behind the ogive there is a smoothly machined section called "bourelet", the diameter of which is a few thousandths of an inch less than the diameter across the space between grooves of the bore (lands) of the gun barrel. The bourelet rides on the lands and supports the forward portion of the projectile as it travels through the bore. In the rear of the body of the shell there is a "rotating band" usually made of gilding metal. This band acts as an after support of the shell and at the same time imparts rotation to the projectile because its diameter is somewhat larger than that across the lands and therefore becomes engaged as the projectile travels through the bore. Due to the tight fit in the bore, the band acts as a gas seal preventing the hot propellant gas from passing between the walls of the shell and the gun, so that the full expansion of gas is utilized to do the useful work of imparting projectile motion. A tapered section to the rear of the rotating band is called the "boat-tail", the purpose of which is to increase the stability of the shell during flight.
In order to prevent any hot gas from contacting the explosive charge (in the event that any porosity exists in the shell base) a thin steel disc is welded or brazed to the shell base ("base cover"). The shell is filled with high explosive (TNT, Amatol, Pentolite etc), leaving a cylindrical cavity at the upper part for a booster charge. The neck of the shell is threaded inside so that the fuze (including the booster) can be screwed on. The outer edge of the fuze is pointed and both the fuze and ogive meet to form a smooth contour.

The HE shell may be designed primarily for fragmentation effect, obtained through dispersion of the fragments at high velocity, or for demolition effect produced by the blast of the HE, or a combination of these effects.

The fragmentation shell has thicker walls than the demolition shell. The HE bursting charge is larger for the demolition than for the fragmentation shell.


High Explosives, Detection as II Complexes. We quote from a very interesting paper by Parihar, Sharma & Verma (Ref 8) about a novel method of detecting explosives.

"A rapid and convenient procedure for the detection of explosives as II complexes with aromatic amines is described. The method could be of immense value in places where normal laboratory facilities are not available.

Recently nitroaromatic compounds have been employed as II acceptors in the study of the charge-transfer complexes of a variety of organic compounds. During such investigations several chromatographic techniques were used. For example charge-transfer was found to be effective during the investigations of substituted anilines 2,4,7-Trinitrofluorenone systems in Gas Chromatography (GC) [1]. GC was helpful in the separation of aromatic compounds on columns containing 2,4-Dinitrochlorobenzene (DNCB) [2], and for olefins on 1,3,5-Trinitrobenzene (s-TNB) [3] impregnated columns. Picric acid [4] and other nitro aromatic compounds [5] have been used as complexing agents for the separation of aromatic compounds by employing Thin Layer Chromatographic (TLC) technique.

Nitro compounds form charge-transfer complexes with aromatic donors due to polarization of the nitro group as in I and II. According to Mulliken [6, 7] these II complexes involve hybrid structures with only a dative and no bond. It has been more recently suggested [8] that in the formation of 1:1 II-complexes of m-Dinitrobenzene (m-DNB) with different aminobenzoic acids only one nitrogoup takes part.

Explosives like s-Trinitrotoluene(s-TNT), s-TNB, Tetryl etc being II acceptors make charge-transfer complexes with aromatic amines. Only in a few cases the II complexes have been obtained in the crystalline state. In large number of cases these complexes are very unstable and cannot be isolated for the identification of the parent compound. The very labile physical linkages between the II acceptor and the II donor molecules are responsible for their break down. Solvent forces, steric hindrance due to groups, temperature etc weaken the charge-transfer linkages. Particularly during the study of such complexes by chromatographic techniques the absorption forces have a powerful dominating effect.

When explosives are found singly or as mixtures there is a problem of their rapid identification. It is particularly significant in field areas where normal laboratory facilities are not available and unknown explosives have to be quickly characterized. Though the II complexes of nitroaromatic compounds with hydrocarbons have been studied by TLC [5], yet suffer from the disadvantage of their colorless nature and difficulty of location on the chromatoplates.

The present paper describes a simple and convenient procedure for the identification of explosives viz: Picramide, Hexyl, 2,4-Dinitrophenol (DNP); Tetryl, s-TNT; s-TNB; 2,4,6-Trinitroanisole (TNA); 2,4-Dinitroanisole (DNA); 2,4,6-Trinitrophenol (TNP); DNCB; m-DNB and Picryl Chloride (PC) in ordinance stores as charge-transfer complexes with amines utilizing TLC.
technique. The resolved Π complexes being highly colored could be easily located from their Rf values. Thus by running the unknown samples along with the known compounds or alternatively from the known data of their resolutions as complexes with different aromatic amines, they could be readily detected up to 2-4y.

Note: The term Rf values refers to a relative rate of movement of the test sample through the chromatographic column.

The authors summarize their findings as follows:

"The resolutions of the Π complexes was found to be governed by three factors (i) adsorbent, (ii) nature of the Π donor (here aromatic amine), and (iii) irrigating solvent. In general the migrations of the explosives were in relation to the effect of electron attracting or repelling groups present in them.

On diphenylamine treated silica gel plates DNA moved less than DNP, similarly TNA had lower Rf than TNP. This is due to more electron repelling nature of the ethoxyl group compared to methoxyl. DNCB migrated higher than s-TNB, due to less electron affinity of chloro than nitro group. Similarly s-TNT had higher migration than s-TNB owing to electron repelling nature of CH3. s-TNB had lower Rf than m-DNB due to the presence of a third nitro group. Hexyl was most strongly bound compared to other polynitroamines ie Tetryl and Picramide.

The relative role of the adsorbents was found to be interesting. For example on employing the same amine diethylaniline (DEA) as Π complexing agent and same solvent system monochlorobenzene (MCB) -ethylene dichloride (EDC) 9:1, DNA gave higher Rf than TNA on alumina in comparison to magnesium silicate plates. On alumina Π complexes of TNA, TNP and PC produced tailings. Magnesium silicate gave excellent resolutions but required more irrigation time than other adsorbents.

The nature of the aromatic amine to form Π complexes was important. On silica gel G plates and MCB-EDC as solvent PC-anisidine complex had lower migrations than PC-DPA or PC-DEA complexes.

The TNA-anisidine, TNP-anisidine complexes had lower migrations than DNA-anisidine and DNP-anisidine complexes. This observation was opposite to the resolutions of the complexes of these explosives with DPA and DEA.
It was seen that irrigating solvent had pronounced effect on the relative mobilities of the II complexes eg the $R_f$ values of Tetryl and Picramide were reversed with petroleum ether-ethyl acetate (9:1) compared to MCB-EDC system.

Their data in tabular form follows:

<table>
<thead>
<tr>
<th>Names of the Explosives</th>
<th>Adsorbents Solvents</th>
<th>Silica Gel G</th>
<th>Magnesium Silicate</th>
<th>Alumina</th>
</tr>
</thead>
<tbody>
<tr>
<td>Picramide</td>
<td>.31</td>
<td>.09</td>
<td>.22</td>
<td>.04</td>
</tr>
<tr>
<td>Tetryl</td>
<td>.32</td>
<td>.05</td>
<td>.30</td>
<td>.06</td>
</tr>
<tr>
<td>Hexyl</td>
<td>.00</td>
<td>.00</td>
<td>.06</td>
<td>.00</td>
</tr>
<tr>
<td>$s$-Trinitrotoluene</td>
<td>.55</td>
<td>.34</td>
<td>.63</td>
<td>.29</td>
</tr>
<tr>
<td>$s$-Trinitrobenzene</td>
<td>.49</td>
<td>.24</td>
<td>.42</td>
<td>.11</td>
</tr>
<tr>
<td>1,3-Dinitrobenzene</td>
<td>.54</td>
<td>.28</td>
<td>.59</td>
<td>.41</td>
</tr>
<tr>
<td>Picryl chloride</td>
<td>.62</td>
<td>.30</td>
<td>.57</td>
<td>.38</td>
</tr>
<tr>
<td>2,4-Dinitrochlorobenzene</td>
<td>.66</td>
<td>.40</td>
<td>.70</td>
<td>.61</td>
</tr>
<tr>
<td>2,4,6-Trinitrophenetole</td>
<td>.52</td>
<td>.29</td>
<td>.60</td>
<td>.27</td>
</tr>
<tr>
<td>2,4-Dinitrophenetole</td>
<td>.39</td>
<td>.12</td>
<td>.39</td>
<td>.17</td>
</tr>
<tr>
<td>2,4,6-Trinitroanisole</td>
<td>.48</td>
<td>.23</td>
<td>.54</td>
<td>.34</td>
</tr>
<tr>
<td>2,4-Dinitroanisole</td>
<td>.35</td>
<td>.09</td>
<td>.31</td>
<td>.21</td>
</tr>
</tbody>
</table>

and for two other solvents the $R_f$ values for the same explosives arranged in the same order as above are:

<table>
<thead>
<tr>
<th>Silica Gel G</th>
<th>Magnesium Silicate</th>
<th>Alumina</th>
<th>Silica Gel G</th>
<th>Magnesium Silicate</th>
<th>Alumina</th>
</tr>
</thead>
<tbody>
<tr>
<td>.33</td>
<td>.19</td>
<td>.36</td>
<td>.11</td>
<td>.41</td>
<td>.14</td>
</tr>
<tr>
<td>.37</td>
<td>.13</td>
<td>.40</td>
<td>.06</td>
<td>.45</td>
<td>.08</td>
</tr>
<tr>
<td>.00</td>
<td>.00</td>
<td>.00</td>
<td>.00</td>
<td>.00</td>
<td>.03</td>
</tr>
<tr>
<td>.65</td>
<td>.46</td>
<td>.76</td>
<td>.49</td>
<td>.72</td>
<td>.53</td>
</tr>
<tr>
<td>.55</td>
<td>.39</td>
<td>.53</td>
<td>.20</td>
<td>.58</td>
<td>.43</td>
</tr>
<tr>
<td>.58</td>
<td>.34</td>
<td>.70</td>
<td>.47</td>
<td>.70</td>
<td>.47</td>
</tr>
<tr>
<td>.68</td>
<td>.45</td>
<td>.71</td>
<td>.51</td>
<td>.52</td>
<td>.22</td>
</tr>
<tr>
<td>.70</td>
<td>.43</td>
<td>.75</td>
<td>.67</td>
<td>.73</td>
<td>.55</td>
</tr>
<tr>
<td>.61</td>
<td>.42</td>
<td>.68</td>
<td>.44</td>
<td>.18</td>
<td>.13</td>
</tr>
<tr>
<td>.48</td>
<td>.18</td>
<td>.50</td>
<td>.17</td>
<td>.56</td>
<td>.34</td>
</tr>
<tr>
<td>.54</td>
<td>.38</td>
<td>.59</td>
<td>.39</td>
<td>.25</td>
<td>.19</td>
</tr>
<tr>
<td>.40</td>
<td>.14</td>
<td>.42</td>
<td>.22</td>
<td>.60</td>
<td>.21</td>
</tr>
</tbody>
</table>
The word "detonics" in the title is chosen in preference to the word detonation to indicate the physics of detonating high explosives and their mechanical effects. The major emphasis is on commercial high explosives for rock blasting, with the exclusion of "Permitted Explosives" and the many complicated problems connected with these.

The first three chapters deal with the mechanism of detonation and the initiation of detonation by means of strong shock waves or mechanical impact at low velocity. Burning and the effects of heating are treated in Chapter 4, and in Chapter 5 is given a systematic discussion of the various phenomena of light emission during detonation. Chapter 6 is devoted to a fairly detailed survey of the mechanical effects in surrounding media and a discussion of the work performed by the reaction products of a contained charge. Particular attention is paid to the effects of a charge in rock. The final chapter is an introduction to such methods and results of the physics of high dynamic pressures in condensed media as have appeared essential to the understanding of the mechanical effects of detonation.

The authors have felt in their own activity a need for a systematic survey of current experimental facts and data of real explosives as a complement to existing theoretical treatments of idealized explosives and detonation phenomena. The theoretical treatment has therefore been deliberately limited and is included mainly with the purpose of simplifying the account of experimental results.

Of necessity, the choice of results and data has been determined by the experimental work and applications in which the authors have been engaged. Wherever possible, results published by workers outside Sweden have been included, but the authors are fully aware of the difficulty in doing justice to the great number of scientists in many parts of the world who have made important and beautiful contributions to this field of research.

The major part of the experimental work described has been done since 1946 in the Physics Research Laboratory (Detoniklaboratori) of Nitro Nobel AB and from 1953 at the Swedish Detonics Research Foundation. Some of the work has been done since 1960 at the Swedish Research Institute of National Defense"
High Explosives, History of. See under HISTORY OF EXPLOSIVES AND RELATED ITEMS

High-Explosive Train. See Vol 4, p D838 & D839. Figs 1-21a & 1-21b

High Grade Nitrocellulose. Same as Guncotton described under Cellulose and Derivatives in Vol 2, p C106-R

High Nitrogen Nitrocellulose. See under Cellulose and Derivatives in Vol 2, p 108-R

High-Low Pressure Gun (H/L Gun). See Hoch- und-Niederdruckkanone in PATR 2510 (1958), p Ger 90-R & in this Vol

High and Low Temperature Tests for Small Arms Ammunition. The purpose of these tests is to determine the effect of high (as high as +165°F) or low temperature (as low as -70°F) storage upon the ballistic performance of ammunition, or components thereof.

The detailed description of the test is given in Ref

Ref: Anon, Ordnance Proof Manual 7-24 (1945), p 7

High Mechanical Strength Explosives. Hopper (Ref 1) evaluated HMX bonded with "Exon 461," a Firestone Plastics Co, Pottstown, Pa, fluorocarbon polymer, for an application requiring a high-energy, high mechanical strength expl. Comps in the range 85-95% HMX & 5-15% Exon 461 were prep'd as granular molding powders and compressed into large cylinders in a heated mold. As the percentage of HMX was increased, the d increased from 1.84 to 1.86 g/cc, and the deton vel increased from 8388 to 8838 m/sec. The compn 85/15 HMX/Exon gave the greatest mechanical strength, at RT its compressive strength was 11000-12000 psi and tensile strength of 1100-1500 psi. Storage at 160°F for 100 days had no appreciable effect on mechanical props measured at RT. Especially noteworthy is the excellent thermal stability of HMX/Exon in the 120°C vacuum stability test.

A number of plastic-bonded expls based on HMX & RDX plus polymeric binders has been developed at the Lawrence Livermore Lab, University of California, Livermore. Their props have been compiled by Dobratz (Ref 2)

Refs: 1) J. D. Hopper, PATR 2592 (Feb 1959)
2) B. M. Dobratz, "Properties of Chemical Explosives and Explosive Simulants," UCRL-51319 (Dec 1972)

High Nitrogen Content Compounds as Flash Reducing Agents. Some aliphatic nitramines of low carbon content were prep'd & studied for use as ingredients of flashless propellants. Nitro-urethane, Dimethylitramine (See Vol 5, p D1306-L) & Cyclotrimethylene trinitromamine (See Vol 3, p C630-R) were found to be unsuitable for use in proplnt comps. Dinitrodicyclemethylene-diamine had properties which indicated it offered promise and tests were recommended to determine the practicability of using it in proplnts (Ref 1)

Benson (Ref 2) also reported that some success had been achieved in the reduction of muzzle flash by incorporating high nitrogen comps such as Dicyandiamide (See Vol 3, p C587-L) & Nitroguanidine. In order to extend this approach and to investigate other comps; selected hydra- zine derivs were prep'd & studied. Cyanuric Hydrazide & 5-Amino-tetrazole (See Vol 1, p A258-L) were of greatest interest because of their high nitrogen content and other props

Since 5-Amino-tetrazole (ATZ) was not available commercially at that time, Benson (Ref 3) also reported a method for the synthesis of ATZ of 99% purity in 70-85% yield by reaction of Na nitrate & amino-guanidine bicarbonate dissolved in nitric acid, followed by treatment with Na acetate. Davis (Ref 4) prep'd Cal .50 gun proplnts and determined the optimum percentage of 5 to 10% ATZ without DNEB (Dinitroethylbenzene), but coated with 5 to 0.8% DNT, gave acceptable service ballistics. The proplnts gave completely flashless rounds in 45-inch barrel rifle and machine gun with an APG-flash hider

Nine new materials of high nitrogen content were studied by Sheffield (Ref 5) and attempts were made to reduce the hygroscopicity of ATZ. Of the new comps investigated, Melamine was found to be the most promising from the standpoint of its props & availability, but Cyanuric Hydrazide was only slightly inferior in these respects. Physical treatment of ATZ had no significant effect on its hygroscopicity, and chemical treatment did not yield material of promise.
H 102

Helf (Ref 6) investigated ten new materials as flash & smoke suppressants in propellants. Guanylurea Nitrate was the most promising compd with regard to physical & chemical props, stability, and O balance. GuanylNitro Urea was unsuitable for propells contg NG. Carbohydrazole-N-carboxyamide & Carbohydrazole-N,N'-dicarboxyamide showed desirable promise but depended on a special starting material. Ammonium Nitrourea, Diammonium -methylene-bis-(nitrosohydroxylamine) & Guanidine Oxalate were investigated and found unsatisfactory because of their chemical & stability props.

Continuing this investigation Helf (Ref 7) found Carbohydrazole Oxalate, Dinitrobiuret, Diaminobiuret, Hydrazine Oxalate, & Hydrazine Nitrate were either unsuitable or not compatible with Nitrocellulose. Urazine, N-Guanioxamic Acid and Cyamelide were stable compds of good physical props but their methods of prepn limited large scale production. Nitrobiuret, although decomposed at 100°C, was stable & compatible with NC, and was of particular interest as a non-smoky flash reducing agent. Guanylamino-tetrazole & Guanylamino-tetrazole Nitrate possess good physical props & are compatible with NC. Both should yield cool propell compns with good ballistic potential. Nitroguanilamino-tetrazole, not previously described in the literature, appeared to have excellent potential as an ingredient of low-temp, high-force propellants.

Audrieth et al (Reps 8, 9 & 10) also prep & investigated compds of high nitrogen content with a view toward their use in propell & HE compns for PA.

Muzzle flash, which is undesirable in gunnery because it discloses position, interferes with vision of gunners, and interferes with guidance and telemetry, is due to the combustion of product gases CO & H₂. Current US composite nonflashing propells are based on British Cordite N, containing Nitroguanidine, while German Nitroguanidine Propellant is considerably cooler because it employs Diethylene glycol Dinitrate instead of NG as the plasticizer for NC.

See Flashless Cordites in Vol 3, p CS22-R.

Reps: 1) F. R. Benson, PATR 1174 (June 1942) (Aliphatic nitramines of low carbon content)

2) F. R. Benson, PATR 1409 (April 1944) (Hydrazine derivs)

3) F. R. Benson, PATR 1548 (July 1945) (Synthesis of 5-Amino-tetrazole)

4) C. S. Davis, PATR 1615 (July 1946) (Propellents contg ATZ)

5) O. E. Sheffield, PATR 1694 (May 1948) (High nitrogen compds)

6) S. Helf, PATR 1752 (Nov 1949) (High nitrogen compds)

7) S. Helf, PATR 1841 (Oct 1951) (High nitrogen compds)


High Pressure Effect on Explosives. Bridgeman (Ref 1) determined that purely hydrostatic pressures as high as 10⁵ atms will not initiate explosives even as sensitive as Lead Azide. In fact there is evidence that high pressures slow down thermal decomposition of explosives. Bowden et al found that for Cyancric Triazine, Lead Azide & PETN the time to explosion at a fixed temp, or conversely the temp at a fixed explosion time, increased as the pressure on the system increased from ambient to 22,500 atms (Ref 2). For Cyancric Triazine this effect was appreciable but for PETN it was very slight. Bowden suggests that these observations are understandable in terms of LeChatelier's Principle since all of these explosives produce large vols of gas on explosion so that application of external pressure is expected to repress the explosion reactions. The larger degree of repression for Cyancric Triazine than for PETN may be rationalized by the lower explosion pressure of the former than the latter. Thus 22,500 atms may be a significant fraction of the explosion pressure of Cyancric Triazine while it is only a relatively small fraction of the explosion pressure of PETN.
Recently Lee et al (Ref 3) re-examined the behavior of PETN under 10 to 50 kilobars of external pressure. They also find a reduction in decomposition rate with increasing applied pressure. HMX behaves similarly to PETN. TNT whose explosion products contain a high proportion of solid carbon, as expected from LeChatelier’s Principle, shows little pressure effect on its thermal decomposition. Nitromethane, however, appears to decompose more rapidly under an external pressure of 50 kilobars than 10 kilobars. This effect is not completely understood but Lee et al suggest that high pressure may favor the formation of the thermally less stable aci form of Nitromethane:

\[ \text{OH} \quad \text{H}_3\text{C}^+\text{NO}_2 \quad \text{pressure} \quad \text{H}_3\text{C}^+\text{N}^+\text{O} \]

Written by J. ROTH


High Pressure Pump. See Hochdruckpumpe (HDP) oder V-3 in this Vol

High-Pressure Technique (Hochdrucktechnik—in German). The term “high pressure”, according to Dodge (Ref 19), refers to any pressure above a lower limit of about 50 atmos (750 psi)

Applications of high pressures in chemical industries dates from the beginning of this century, when the industrial synthesis of ammonia from the elements was first achieved by Haber, LeRossignol and others. The Burton process for oil cracking dates from about 1913

At present, pressures of the order of 1500 atmos are widely used, but the technique for carrying out reactions under pressures up to about 10,000 atmos is fairly well developed. In research laboratories, pressures as high as 150,000 atmos (3,000,000 psi) have been used to a limited degree

One of the principal applications of high pressure technique in industry is to increase the chemical reaction rate

High-pressure operations are conducted in special reaction vessels which may be divided into batch reactors, such as autoclaves (qv) and continuous reactors. Continuous reactors consist of long cylindrical tubes, constructed of steel in a manner to resist pressures as high as 1500 atm. Inside the thick-walled reactor, a thin-walled vessel containing a catalyst is placed. The incoming gaseous reaction components, previously preheated (by allowing them to flow along the outside of the reactor), enter the reaction chamber, pass through the catalyst where they react, forming the desired product, and exit through the other end of the reactor into a cooler where they are collected as a gas, liquid or solid


High-Pressure Test (HPT) Cartridge. See under Cartridge, Ammunition, Vol 2, p C74

High Rate Detonator Production Study. As part of the overall program to modernize the US Govt owned, company operated, Army Ammunition Loading and Assembly Plants, it is planned to develop fully automatic equipment to manuf non-electric detonators at the rate of 1200 per minute. The survey of literature sources and industry was undertaken to discover techniques and equipment that may be applicable, as described in Ref 1


HIGH SPEED PHOTOGRAPHY

Introduction

High-speed photography provides the main observational facility in the study and measurement of explosive phenomena. In Vol 2, pp C13-19, many high-speed cameras are described & some of the techniques for using them are discussed. The present article deals with broader aspects of high-speed photography. Not only is the information on high-speed cameras brought up to date and extended, but other aspects of high-speed photography, such as illumination, films & film processing are discussed

Advances in Shuttering Systems

High-speed photography serves to extend and help calibrate other even more rapid systems of data recording. The use of cameras and other optical and electronic devices has kept pace with advances in high energy physics. The extremely brief exposures required for photographing the effects of explosion phenomena are achieved by regulating the optical, electrical and mechanical design of the camera's shuttering system

The picture taking rate for photographic recording of rapid moving or brief duration phenomena has steadily increased from "slow-motion" techniques at 64 pictures-per-second (pps) to more than millions of pps. Time resolution has now been extended down to the order of 10 picoseconds or less. Three classes of high-speed photography are recognized. According to their time and rate of picture-taking, they are called: high-speed, very high-speed and ultra-high-speed photography

High-Speed & Very High-Speed Cameras

The first high-speed cameras accomplished action stopping exposures at more than 1000 pps. These rotating prism cameras (Ref 2) eliminated the start-and-stop clawing action of conventional motion picture cameras. The 8mm size film moved continuously and had relatively poor resolution. Later models, using 16mm films, reached speeds of more than 40,000pps. By increasing the number of faces in the prism shuttering device each standard 16mm frame is split into 2 or 4 frames, thereby increasing the pps. Greatly increased detail resolution was achieved at first by going to larger film formats such as 35mm and finally 70mm. In recent years newly developed anti-reflection lens and prism coatings produced substantially better contrast, higher resolution and more effective light transmission by reducing glass interface glare and reflection

Prior to World War II, Dr. C. N. Hickman of Bell Labs invented the "ribbon-frame" camera using the inexpensive 122 size (post-
Film advance sprocket and image motion compensation prism are mounted on a single shaft to improve precision of registration.

These cameras, generally using standard 16mm motion picture film (although Fastax also made 35mm film models as well as the original 8mm size) are efficient at picture taking rates from 100pps to 10,000pps. The Photec IV, through an improved yet simplified optical design, permits an effective aperture of f:2.8 to be attained. This 2-stop gain over the "competitive" models allows four times as much light to reach the film. The optical/mechanical design advance allows the picture-taking rate to be increased, the shuttering time to be reduced substantially (to one/fiftieth of the picture-taking rate). An additional benefit is the reduction in heat reaching the subject because weaker light, and consequently "cooler," sources can be used. Thus the Photec IV brings motion picture cameras into the very high-speed photography group. Further references to the subject of very high-speed cameras are covered in the Proceedings of the International Congress on High-Speed Photography (HSP), 1 through 10. These excellent surveys have been published nearly every two years since the First International Symposium on High-Speed Photography in Oct 1952.

**High-Speed Video Systems**

The next few years should produce effective high-speed television image sequences. At present, video systems are limited to a picture-taking rate of less than 200pps and are consequently classified as slow motion photography, although the data recovery rate, or read-out, is instantaneous. It is expected that an increase in the framing rate by an order of magnitude will be forthcoming soon. This will put video systems in high-speed category.

**Ultra High-Speed Cameras**

It is in the area of ultra high-speed photography that the most significant advances are now taking place. Exposure time has steadily been reduced from microseconds, through nanoseconds to 10 or less picoseconds ($10^{-12}$). A pulse of light has now been photographed with a movement blurring of the image of less than two millimeters (Ref 26). This achievement in ultra high-speed photography is due to a unique
electronic shuttering system, namely a Kerr cell that opens and closes (optically) in 10 picoseconds. No electrodes are required and carbon disulfide has replaced the heretofore traditional nitrobenzene. A first-harmonic pulse of light from a Nd:glass laser is used to drive the shutter.

**Image Dissection Cameras**

This type of ultra high-speed camera (Ref 3) also called lenticular camera, high-speed raster camera (in the USSR), continuous-access camera and, more informally, “scrambled” image camera, seems to have the greatest potential for high-speed photography in the future. Resembling holographic recording (Ref 1) in one sense, inasmuch as it permits the concentration of many sequential images in a limited space, it offers the prospect of ultra high picture-taking rates (over 10^10 pps). After an ordinary development process the plate or film can be shown immediately as a motion picture or video image. Eventually, when the resolution of image converter tubes improves, the principle of image dissection should provide the maximum rates of high-speed photography (Ref 23).

This is emphasized in the following table, taken from an excellent summary of state of the art of high-speed photography by Courtney-Pratt. The data in this table also shows that (Ref 26, p 172) there were no spectacular advances in image dissection camera performances in the period 1957 to 1972.

**Table: Image Dissection Cameras**

<table>
<thead>
<tr>
<th>Type</th>
<th>Reposition Rate, P.p.s.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clear lines or holes in opaque plate</td>
<td>&lt;10^10</td>
</tr>
<tr>
<td>Air plate and theseing mirror</td>
<td>&lt;10^10</td>
</tr>
<tr>
<td>Lenticular plate, aperture scanning</td>
<td>&lt;10^9</td>
</tr>
<tr>
<td>Lenticular plate, mechanical traverse (including cinemicroscopy up to 2000 X)</td>
<td>10^7</td>
</tr>
<tr>
<td>Fibre optics, long series, low resolution</td>
<td>10^6</td>
</tr>
<tr>
<td>Fibre optics, short series, evaluation of 200 samples per frame</td>
<td>10^4</td>
</tr>
<tr>
<td>Lenticular plate, rotating mirror, in USSR</td>
<td>10^4</td>
</tr>
<tr>
<td>Lenticular plate or air plate and image converter tube</td>
<td>&gt;10^5</td>
</tr>
</tbody>
</table>

Other ultra high-speed cameras using rotating mirrors, rotating slotted discs and optical image dissection systems ranging from pinholes to lenticular plate arrangements remain in constant use due to the greater number of frames or pictures produced in sequence. If somewhat less resolution can be tolerated, fiber optic light guides can be used to provide still longer series of pictures.

**Flash X-rays**

Extremely brief flashes of X-rays are useful in the study of the movement of dense materials, such as bullets, shell and bomb fragments, metal liners and containers, through smoke, turbulence and explosive debris (Ref 9). Unaffected by glare, sunlight and other atmospheric conditions, X-rays (radiographs) penetrate, as they do in industrial and medical applications, to detect and reveal the behavior of explosively driven material (Ref 6). Multiple flashes permit stereo or sequential pictures to be made depending on the synchronization, arrangement and number of X-ray sources used. Cineradiography is particularly useful in investigations of such phenomena as explosive welding, where the high-speed action takes place within an optically opaque substance (Ref 10). In addition, obscuration by the very bright luminescence of the detonating explosive charge is eliminated.

Along with advances in X-ray technology, progress in electronic image enhancement (see next section on Image Enhancement) permits gains up to fifty times. This permits the extension of flash X-rays into diffraction recording as well as higher rate cinematography (Ref 11).

**Electronic Amplification of Light Images & Techniques of Image Enhancement**

Perhaps the most important trend at this time in ultra high-speed photography is in the increased use of image amplification (Ref 25). Image converter cameras were developed in recent years not only in the United States but also in England, France, Germany, Holland & the USSR (Ref 5).

The benefits of electronic image enhancement are as follows:

1. Increased sensitivity—2000 to 10,000-fold over the primary image.
2. Faster shuttering—sub-nanosecond or less.
3. Memory tube delay permits millisecond camera synchronization.
4. Reduced distortion—special flatfield, distortion-corrected lenses are used.
5. Variable contrast control produces higher signal/noise ratio.
6. Eventual application to video systems of high-speed photography.
At times the image produced in high-speed photographic studies dealing with explosive phenomena is less than satisfactory. The nature of the subject, the available instrumentation, the cost, unforeseen accidents, such as the failure of a light source to synchronize; all can contribute to substantial underexposure. This results in a low signal to noise ratio making precise film or picture analysis difficult or impossible.

It is customary, when weak images are expected, to employ the most sensitive films available, the so-called “superspeed” films, and if necessary to use chemically vigorous or “forced” development (Ref 6a). The gain in sensitivity generally is in the range of two to ten times the original (manufacturer’s) rating of the film’s speed. Even color film, such as High-Speed Ektachrome can be exposed at an exposure speed index (EI or ASA speed) of 5000 or more when processed as a negative rather than the customary reversal into a positive film. Even with the resultant increase of coarse grain, the color differences in the subject help reveal useful data.

A technique of “post-fogging” called intensification (Ref 2a) can frequently increase the maximum recording rate up to four times. This is impractical to attempt with long lengths of motion picture film but can be successfully employed with a high-speed still, or short strips of film. Although the procedure raises the threshold speed of films by this pre-development, extra exposure it is not the same as “pre-fogging” which merely lowers the inherent contrast range of films. The difference is in the employment of 5 to 30 minutes pre-development exposure to weak green light, rather than an instantaneous flash exposure to low level white light. It is particularly useful with fast Polaroid photographic material.

In many instances image rescue can be accomplished after the act, e.g in case of extreme overexposure. This can be corrected easily by reprinting the film original on to highspeed reversal or negative film stock using an intense print light source. Little or no detail loss will result. An alternate procedure, having some minimal risk to the original film, would be to employ photographic chemical reduction. Standard photographic darkroom technique (Ref 1a) handbooks describe this technique and include recommendations for proportional reducers.

These same references also discuss photo chemical image intensification. The process of intensifying a film image is most satisfactory when the image is weak due to underdevelopment in processing. However, even when the faint image is due to underexposure, frequently a much stronger and more detailed image can be produced by use of chromium and mercury type chemical intensifiers.

Occasionally the original, showing an extremely weak image can best be made useful by re-printing or rephotographing the film on to extremely high contrast film such as used in graphic arts. These “process” or “lithographic” films are available in all sizes and are convenient to use as a contrast and detail enhancing agent.

Several photographic feedback techniques were used with these procedures. One is to re-photograph the original film by reflected light as it rests against a very black, low reflective background. The positive image seen usually shows much more detail than direct observation of the original film can reveal. Secondly, there is a device on the photo market that presses the weak original film against a front surface mirror in order to gain a doubling of the “signal” strength. The third and most effective and unique of these techniques is Michel Cloupeau’s “optical contraster.” (Ref 7a) This system places the badly underexposed original film between semi-reflecting plates, and, by means of collimated light, enables one to choose an image which is the sum of multiple passes through the original film, thereby raising the information signal level substantially. This permits either direct observation or photo reproduction.

Image intensifier systems have developed from the first electronic image converters that were used as fast-acting camera shutters (Ref 3a). Multi-stage image-intensifying circuitry now permits extremely high gain (Ref 5a). Signals due to individual photoelectrons leaving the photocathode of the intensifier have been photographed. Nearly all the information available in low-density electron images can be recorded. In recent years multistage electronic image converters have been developed that produce a light gain from 8000 to 50,000X (Ref 4a). Film exposure times can be as large as the interframe times, or as short as $5 \times 10^{-9}$ sec. Picture taking rates from 1 to 100 million pps are easily achieved.
Streak Cameras

Also known as "smear" cameras and continuously-writing cameras, they generally serve to record movement velocities (Ref 26). Two basic principles are applied to the camera design in order to record a continuous section of a moving image. Either the image is optically "wiped" along a strip of photographic film lining a drum by a rotating mirror or prism system or the film itself is moved at a known rate past the image. Since only a point of light is necessary to be observed as it is displaced or moved normal to the film's directional axis, the real image of the subject is focused on to a slit nearly in contact with the film. As the image of the action being observed travels along the slit, this multiplicity of points photographically are "streaked" on to the film at a known rate. Frequently two or more slits are used when there is more than one action to be observed.

Spatial resolution is determined by the narrowness of the slit, the velocity of the image/film travel rate, and the optical resolution of the lens system.

As shown in the table below (from Ref 26), there has been a considerable improvement in the time-resolution capability of streak cameras in the last two decades.

<table>
<thead>
<tr>
<th>Characteristics of Streak Cameras.</th>
<th>Time resolution, sec</th>
</tr>
</thead>
<tbody>
<tr>
<td>Streak cameras</td>
<td>( \leq 1972 )</td>
</tr>
<tr>
<td>Streak record, drum camera</td>
<td>( 5 \times 10^{-8} )</td>
</tr>
<tr>
<td>Streak record, rotating mirror, single rotor</td>
<td>( 2 \times 10^{-9} )</td>
</tr>
<tr>
<td>Streak record, rotating mirror, multiple reflection</td>
<td>( 10^{-10} )</td>
</tr>
<tr>
<td>&quot;Schardin Limit&quot;</td>
<td>( 0.25 \times 10^{-9} )</td>
</tr>
<tr>
<td>Deflecting image converter tubes</td>
<td>( (2 \text{ or } 3) \times 10^{-12} )</td>
</tr>
</tbody>
</table>

The format or film size, shape and dimension, runs the full range of photographic recording. Early streak cameras consisted of a large strip of film wrapped around a drum electrically driven to rotate at a high speed. (See diagram taken from Ref 26)

Diagram of a simple drum camera. From measurements of the streak record, velocities of movement of the object can be determined relative to the speed of movement of the film. The image-illuminated slit recorded when a steady rate of film movement was achieved. Motion picture films were used in the 35mm and 70mm sizes in somewhat similar arrangements. Most of the more recent designs of streak cameras substitute rotating mirrors for the moving film and can give a time resolution at least an order of magnitude greater than the moving film type (Ref 12).

A somewhat unusual approach is encountered when high-speed motion picture cameras of the Hycam type are used as a streak camera. This is accomplished by filming through the camera's viewfinder which bypasses the prism system that normally cuts up the scene into frames or pictures. After nearly one-fourth of the film has run through the camera the voltage-controlled rate of film travel is reasonably steady and can be used for velocity measurements.

Variations in film travel speed are easily ascertained by the millisecond markings imprinted along the edge of the film as the camera runs. Two advantages are gained by this technique: one, the camera can be set to initiate the subject action at any pre-determined delay; and two, since the film never is exposed again to
the subject as with a rotating mirror system, danger of re-writing is eliminated. Film lengths up to 1000ft can be used in this manner. However, most of the time, continuous-running cameras, using short lengths of film, are used and double exposure caused by the over-run or rewriting is avoided by high-speed shuttering.

External Shuttering

Frequently it is as necessary to cut off the illumination as quickly and precisely as it is to turn it on. Internal electro-shuttering, such as accomplished with the Faraday and Kerr cells, can be synchronized with the camera and event exactly as needed (Ref 17). However, as mentioned above, cameras such as the Beckman/Whitley framing camera as well as most continuously writing “streak” cameras, run the risk of incurring double exposures (Ref 13). This is due to the image re-writing propensity of the rotating mirror and many drum-camera systems. Small auxiliary detonating charges can be used in a number of ways to block off light and image exactly when required. Some of the methods used are listed below:

1. Explode a small charge in contact with a block of glass in the image path. The internal multiple cracking caused by the charge makes the glass effectively opaque.
2. Blast a mirror out of the optical path to terminate recording of the event.
3. Implose a ring-shaped explosive charge of Primacord wrapped around an aluminum cylinder through which the light travels to the film.
4. Wrap a coil of Primacord explosive, sometimes covered with black grease, around the camera observation port window safety glass outside the instrument shelter. Detonation of the explosive coil creates a dense black smoke cloud. This method is usually called a smoke shutter.

Illumination for High-Speed Photography

Despite the presence of brilliant light produced by an explosion, additional high-intensity illumination is often necessary to record high-speed phenomena. Also, it is frequently required to regulate the length of the picture taking sequence by controlling the duration of a supplementary light source.

Spark Light Sources

The original “instantaneous point” light sources was an electric spark discharged from primitive capacitors of the Leyden jar type. Thanks to General Liebessart and other early experimenters (Ref 4), the spark evolved to become a controlled, channeled efficient light source. However, at present it is more useful in ballistic and wind tunnel studies of aerodynamic phenomena than for studying explosives. Confined sparks are still used as a primary point source in Schlieren high-speed photography. This is particularly useful in shockwave studies. Frequently, inexpensive expendable fresnel lenses (plastic) are used in conjunction with spark light sources to redirect much of the spark light into a small lens aperture on the camera. This application of high-speed photography, usually referred to as shadowgraphy, is useful in studying strong disturbances in a transparent medium such as air, water and glass when a powerful shockwave or high temperature boundary is present.

Flash Lamp Light Sources

Electronic flash lamps are used when the light duration has to extend into milliseconds rather than the microsecond range (Ref 7). Ultra high-speed cameras such as the Beckman/Whitley 189 framing camera can produce 25 or more sequential photographs during one flash of a small electronic flash; commonly, but incorrectly, called a “strobe” light. Repeated electronic flash units have been made in Germany that permit a picture taking rate up to 10,000 pps at quite high intensities. However, in the United States when a nondestructive extended light source is required, conventional photo-flash lamps are generally used to provide intense illumination. Arranged in overlapping relay sets they supply light enough to cover a full length motion picture roll in a rotating prism type of camera such as the regular high-speed cameras mentioned at the beginning of this article.

Explosive Light Sources

Still higher energy light sources can be fabricated. These use blocks of high-explosive such as Comp B detonated at one end of a tube or funnel-shaped cone filled with argon, krypton or xenon (Ref 8). These ionizable gases serve as the light emitting source. Temperatures in the vicinity of the shock wave in argon can exceed 28,000°K (Ref 15), but strangely enough
it affects color film much as a pulsed daylight quality source. The duration and luminosity of this type of explosive “candle,” using free-flowing argon, can range up to more than 500 microseconds at intensities much exceeding sunlight. Lining the light source tube with aluminum foil covered sheet explosive may help in achieving the longer duration (up to 500μsec) luminosities. Reflecting surfaces on the inside of the argon container have the effect of making the light source have an apparent radiating area greater than its actual cross-section. Thus, an argon light source with a reflecting internal surface illuminates a target more effectively than an identical light source with matte surfaces. The length of the light path is roughly proportionate to the duration of the luminosity, if rarefaction effects can be minimized.

**Laser Light**

New techniques in high-speed photography have been aided by recent developments in laser technology (Ref 19). In some instances the monochromatic nature of laser illumination has been useful in replacing fast X-rays in order to perceive important details in self-luminous events (Ref 18). The steady increase in use of lasers permits simpler interferogram systems to be arranged. In addition, hologram techniques are now practical with sub-nanosecond exposures. This extends interferometric procedures into a true 3rd-dimensional recording system.

Non-expendable light sources, such as a Q-switched pulsed laser can be protected from destructive forces encountered in the photography of explosive material by piping the light through fiber optics to the experimental zone. Occasionally lens systems are used to relay the light from mirrors located near a protective barrier shielding the laser (Ref 16).

**Photographic Films and Processing Procedures**

High-speed photography benefits by certain changes in modern films. Many of the newer film emulsions are now coated on polyester plastic bases. In addition to attaining dimensional stability approaching that of glass, these films are thinner and stronger than the conventional acetate film supports hitherto used. This permits more film, thereby a longer run, on a given motion picture size spool. The increased strength and resistance to tear allows more rapid acceleration with no torn sprocket holes in all moving film type high-speed cameras.

One detriment with the new mylar type films is that conventional splicing can not be used—pressure sensitive clear tapes seem to be the most practical approach to the joining of film lengths.

A still more useful improvement is in increased sensitivity. Exposure indices (EI) or film “speed” now easily attain the range of 2000 to 5000, an increase of 10 fold over former high-speed films. With vigorous film processing at high temperatures, color and black & white films have been successfully exposed at an EI of 6400. The use of Polaroid film continues to expand with a soon-to-be available fast film effectively equal in “speed” to 50,000.

High temperature film processing not only now permits film to be processed in the tropics and in high temperature locations but allows the specially hardened emulsions to develop very rapidly with no detrimental changes in their characteristics.

**Color Photography**

The use of color film in many instances extends the usefulness of high-speed photography. When more than one explosive subject is being studied simultaneously, it is convenient to observe each through complementary or contrasting color filters. With streak cameras using more than one slit on the same experimental shot, the color contrasting hues can overlap and intermingle and still be free of double exposure confusion. This is done simply by placing a clear bright-colored gelatin filter over each camera slit so that the resulting streaks are rendered in the hue pertaining to a particular slit.

Specially made color filters can be inserted in place of the conventional knife-edges in Schlieren systems (Ref 20). Frequenty this produces higher resolution because of the avoidance of knife-edge diffraction. Restricting parts of the subject image to monochromatic recording also presents additional gains in sharpness. Composite color filters placed at field stops in the optical system can add considerable information to the photo image when captured on color film.

**Special Color Films**

A Kodak color film with the blue layer
quite sensitive to infra-red radiation is now commonly available. This enables dynamic thermography to be included in high-speed photographic recording.

The use of color film for high-speed photographic studies of explosions underwater has been made more feasible by GAF’s new Blue-insensitive Anscochrome Aerial Film which does not have a blue-sensitive top layer (Ref 21). This high-speed (EI nearly 1000) color film has high resolution and eliminates the need for a minus-blue filter on the camera. Schlieren and photoelastic stress analysis can conceivably gain in contrast with this new film. In addition, it should be less affected by smoke and other explosive debris which scatters the shorter wavelength light (blue) more than “red” light.

Lastly, a somewhat unorthodox color film, referred to as Extended Range (EG & G) film permits photographing an explosive subject over an extremely wide exposure range. This multiple layer color negative film permits photography requiring effective exposure ratings from approximately 0.03 or smaller to a maximum equal to EI 400. Originally developed to photograph the sun’s corona, it has found use with high-speed photography when the correct exposure is difficult to determine. This film, however, produces colored images in blue, red, and yellow that bear no relation to the subject’s original colors.

An experimental new color film called XRC (for extended response color) has been introduced by Applied Photo Sciences Inc (Ref 22) that is a “true” color film. It too has a remarkably extended range of responses to scene brightness, said to “nearly match the human eye”.

An excellent review of advances in high-speed photography is presented by Courtney-Pratt in Ref 22, listing 115 references. Additional source material, taken from Courtney-Pratt’s article, is listed at the end of the references to this article.

Written by ZEV PRESSMAN


* J Soc Motion Picture & Television Engr

** Proc Intl Congress of High-Speed Photography


Attention is drawn also to the following books, etc which provide valuable source material:

P. Payolle & P. Naslin, Photographie Instan
tée et Cinématographie Ultrarapide, Editions de la Revue d’Optique, Paris, 1950

A.A. Sakharov, High-Speed Photography in Science & Technology, Publishing House of Foreign Literature, Moscow, 1955
J.H. Waddell, Photographic Motion Analysis, Industrial Laboratories Publishing Co, Chicago, 1955
C.H. Greenewalt, Hummingbirds, Doubleday, NY, 1960
The bibliography on image dissection in high-speed photography by J.S. Courtney-Pratt
The bibliography on Holography compiled by R.P. Chambers & J.S. Courtney-Pratt
K. Vollrath & G. Thoman, High-Speed Physics, Springer, Vienna, 1967
The bibliography on high-speed photography (to 1960) compiled by Eise Gavrin, Eastman Kodak, Rochester, NY (Distributed at HSP 5)
Bibliography on High-Speed Photography 1960-1964 (Kodak Pamphlet P-27) compiled by Elizabeth W. Tapia, Eastman Kodak Co, Rochester, NY, 1965 (Distributed at HSP 7)
The bibliography on high-speed photography 1964-1970 compiled by Elizabeth W. Kraus, ed W.F. Walker, Eastman Kodak, Rochester, NY (Reprinted HSP 9, 548-595)
The volumes comprising the Proceedings of the International Congresses on High-Speed Photography, as described above

High Temperature Behavior of Explosives. See Wenograd Test

High Velocity Test Gun. A high-velocity gun facility, 87 feet long and with a 3.5-inch bore, is being used to study material behavior under extreme impact conditions. Impact velocity will reach a maximum of 8000 to 10,000 fps and will exert pressures of several million pounds per square inch upon experimental materials.

Studies of the reaction of materials to extreme mechanical shocks are important in the design of components and systems which may experience severe impact. Experiments with different metals, alloys, ceramics, plastics and composites help engineers choose the ideal material for different applications under shock loading. They also provide the scientist with a more fundamental understanding of the materials themselves.

The gun will be fired by igniting a charge of conventional artillery propellant weighing up to 20 pounds. The explosion propels a two-pound projectile (Sabor) and impactor through the barrel to the target. Since the experiments require precisely controlled impact conditions, the barrel must be honed to a tolerance of 0.0001 inch and aligned to 0.020 inch along its entire length before each test. As in similar guns, the bore is evacuated to about one micron
of mercury to prevent undesirable effects during impact.

The test material may be mounted either on the sabot or placed as a target at the muzzle. Instrumentation which will record data on material deformation will provide a resolution of a few billionths of a second in time and a few millionths of an inch in displacement.

Refs: 1) Sandia Science News, Vol 2, No 4 (March 1969), Sandia Corp, PO Box 5800, Albuquerque, New Mexico 87115 2) G. Cohn, Ed, Explos & Pyrots 2 (11), 1969

Highways and Byways in Combustion. Title of a paper by D.T.A. Townsend, JInstFuel 27, 534-44 (1954) & CA 49, 2057 (1955). In this paper investigations of high-pressure explosions, flame propagation and combustion of higher-mol-wt hydrocarbons are reviewed.

Hill Powder. Has been used for loading shells: NH₄ Picrate 43.0, K Picrate 53.5 & charcoal 3.5
Ref: Daniel (1902) 375

Himalaya’s Explosives (Portugal). KClO₃ 60-80; starch, charcoal etc 10-25; oil 4-10, metallic dust or powder 2-10, metallic oxides, or dehydrated CuSO₄ 2-5%
Ref: M.A.G. Himalaya, BritP 22030 (1910) & CA 9, 3158 (1911)

Himley Explosive. KClO₃ 45, KN₀₃ 35 & coal tar 20
Note: This explosive was prepared by impregnating the first two ingredients with tar dissolved in petroleum ether and allowing the ether to evaporate.
Refs: 1) Daniel (1902) 375 2) Pérez Arz (1945) 216 3) Giua, Trattato 6, 400 (1959)

Hinde Explosive. NG 64.0, pulverized coal 23.0, Amm citrate 12.0, ethyl palmitate 0.25, Ca carbonate 0.25 & Na carbonate 0.50%
Refs: 1) Daniel (1902) 375 2) Giua, Trattato VI (1) (1959), 344

Hinshelwood, Sir Cyril, 70, retired since 1965 as professor of chemistry at Oxford University, died Oct 9, 1967 in London. Sir Cyril, who was a fellow of the Royal Society and its president (1955-60), in 1926 published “Kinetics of Chemical Change,” a subject in which he was already world-renowned. Continuation of his research into chemical kinetics later earned him the Nobel Prize in Chemistry (1956), along with N. N. Semenov of Russia. Sir Cyril was also noted for his breadth in fields other than science. He spoke six languages, including Chinese, and was a serious student of the literature of many countries, as well as being a painter. He served on many government advisory bodies and boards and was, for these contributions, knighted by King George VI in 1948. He was president of the Chemical Society (London) in 1946.
Ref: C&EN 45, No 46, 63 (1967)

Hippuric Acid and Derivatives
Hippuric Acid, also called Benzyalminocacetic acid, Benzoylglycine, Benzaminocacetic acid, C₆H₅CONHCH₂COOH, mw 179.19; colorless crystals, mp 188° (decomposes on further heating), d 1.371, sol in hot water, alc & eth. Used in org synthesis & medicine.

Hippuric Acid Azide or Hippuroyl Azide, C₆H₅.CO.NH.CH₂.CO.N₃; mw 204.19, N 27.44%; nds (from benz or eth), mp 98°; sol in alc, chl & glacial acetic acid; sl sol in eth & cold benz; insol in w; was prep'd by the action of Na nitrite on hippuryl hydrazine in warm water.
Refs: 1) Beil 9, 247 2) T. Curtius, Ber 23, 3031 (1890)

Mononitrohippuric Acid, O₂N.C₆H₅.CO.NH.CH₂.CO.OH; mw 224.17, N 12.50%. Three isomers are known:
1) Nitrohippuric Acid, crysts (from w), mp 190° (Ref 1)
2) 3-Nitrohippuric Acid, nds (from w), mp 166° (Ref 2)
3) 4-Nitrohippuric Acid, colorless prisms (from w), mp 129-30° (Ref 3)

Other props & methods of prepn are given in the refs.


Mononitrohippuric Azide, O₂N.C₆H₅.CO.NH.CH₂.CO.N₃; mw 249.21, N 28.11%. Two isomers are known:
1) Nitrohippuric Azide, yel powd, very un-
stable especially in daylight, decomposes explosively above 74° if heated rapidly. Moderately sol in eth; sl sol in alc, chif & benz; insol in w; prep from 3-nitrophippuric hydrazide & Na nitrite in glacial acetic acid (Refs 1 & 3) 4-Nitrophippuric Acid Azide, yel powd, mp 70-72° (dec), degr on heating & explodes on heating in light; sol in eth; sl sol in alc & benz; insol in w; can be prep from 4-nitrophippuric hydrazide & Na nitrite in HCl (Refs 2 & 4) 


Refs: 1) Beil 9, <1937> 2) B.C. Saunders et al, BiochemJ 36, 368, 371 (1942) 4-Nitrophippuryl-aminoacetyl Azide, O₃N.C₆H₄.CO.NH.CH₂.CO.NH.CH₂.CO.N₃; mw 306.24, N 27.45%; yel powd, mp 91-92° (dec), explodes on heating in the light; sol in NaOH; sl sol in eth; was prep by treating 4-nitrophippuryl-aminoacetyl-hydrazide, Na nitrite & HCl 


Hispano-Suiza Gun. A 20mm aircraft gun used by the British during WWII and which proved to be a successful weapon. It was mounted either for firing thru the propeller hubs or from fixed mounts in the wings of an airplane. It was capable of delivering fire at the rate of 600-700 rounds per minute 

Refs: 1) War Dept Tech Manual, TM 9-1901, p 323 2) J.A. Solomon, PATR 949 (1939) (20mm CRA for Hispano-Suiza gun) 3) M.L. Mathesen, PATR 994 (1939) (20mm CRA for Hispano-Suiza gun) 4) M.L. Mathesen, PATR 1025 (1940) (20mm CRA for Hispano-Suiza (Birkigt) gun) 5) A.B. Schilling, PATR 1627 (1946) (20mm HE CRA for Hispano-Suiza gun) 6) A.B. Schilling, PATR 1628 (1946) (20mm AP CRA for Hispano-Suiza gun) 

HISTORY OF EXPLOSIVES, AMMUNITION AND WEAPONS

Introduction

Prior to describing history of explosives, it is desirable to say a few words about cold weapons which were used in wars many centuries before explosives or incendiaries were invented

Greener (Ref 7, pp 1-12) described such weapons and gave many illustrations as did Dupuy & Dupuy (Ref 69, pp 2ff) 

Primitive cold weapons can be subdivided into two major categories: shock and missile, of which missiles could be operated by hand or mechanically. The earliest shock weapon was a heavy club, while the first missile was a stone or other heavy object hurled by hand. The next development in missiles was a sting (leather or string) which could be used for hurling smooth stones at longer distances than by hand. The stones were later replaced by lead balls. This weapon was used as late as AD 1572 and is still used by some savage tribes (Ref 69, p 2). In some weapons a stick was hurled and this, in turn, evolved into darts and javelins. In Australia appeared the boomerang which is still used for hunting. The club was modified by a number of ways, one of them was to attach a heavy stone or metal piece at its end. One such weapon was the Russian politsa. The club with a sharp ax-like end was the tomahawk of the American Indians. The shock action counterpart of the javelin was the heavy pike or thrust spear. The spear with an ax-like end was halberd invented in 15th century and used since then by Vatican Guards

The bow and arrow, invented late in the Stone Age (ca 3500 BC), became an invaluable weapon for many centuries (See Vol 1 of Encycl, p A488-R) (Illustrated in Ref 7, p 3)

At the Bronze Age (sometime before 2000 BC), which followed the Stone Age and then later at the Iron Age (ca 1000 BC), the most important contribution was the adoption of metal for points of arrows & spears and also for edges and smashing surfaces of other weapons. The new weapons of metallic era were the dagger and the sword. They were probably first made by Assyrians (Ref 69, pp 2 & 8). Their chariots also were made of iron

The most important protective armor developed in ancient times was the shield, which was
usually held in the left hand or on the left arm, leaving the right arm free to wield a weapon.

Emergence of historical China, as a small country around Yellow River Valley was around 1600 BC and the first known ruler was the Shung Dynasty (1523 to 1024 BC), which was followed by the Chou Dynasty (1027 to 500 BC). Their weapons were metallic but not as good as those of the Middle East. The same may be said about early Hindu weapons (Ref 69, pp 14–15). More about China is below.

Substantial advances in tactics and doctrine of land warfare were made in the middle of the 6th Century BC by Cyrus the Great of Persia, but he used the same cold weapons as listed above with only slight modifications (Ref 69, p 16).

The Greeks at that time developed an Army outfit consisting of well trained soldiers known as hoplites. They carried a pike, a short sword and a shield (See Fig on p 17 of Ref 69).

A very important improvement in hurling missiles took place ca 400 BC when ballista (See Vol 2 of Encycl, p B5-L) and catapult (Vol 2, p C91-L) were invented. They are illustrated in Ref 7, p 6 and Ref 69, p 38.

They hurled missiles up to 500 yards.

In siege warfare invented after 350 BC by engineers of Alexander the Great of Macedonia were the battering ram and the famous movable tower (turns on wheel) (See Fig on p 39 of Ref 69). Other inventions were mantelet, telenon and mural hook (See Figs on p 40 of Ref 69).

The above described weapons were also used by Macedonians under Alexander the Great (356, 323 BC) during their conquest of Europe and half of Asia. No new weapons were used, but they encountered in India a new weapon which will be described below.

Next to Alexander, Julius Caesar (100–40 BC) was the outstanding director of siege operations, bringing systematic procedures to operations. Fig on p 83 of Ref 69 shows fort, tower, mound and mantelets designed by him, while on p 84 is shown testudo, which consisted of long shield raised and interlocked over the soldiers’ heads and backs.

One important cold weapon, the crossbow or arbalest, appeared in the 11th century (See Vol 1 of Encycl, p A477–L, where it is erroneously shown that it was invented in the 4th century).

It is illustrated on p 7 of Ref 7 and p 279 of Ref 69. It was used successfully by the Normans in invasion of England and one of the crossbow’s arrows killed King Harold.

In siege operations, one new weapon was introduced. This was the trebuchet or mangonel, a missile-hurling machine for battering fortifications or for throwing rocks or other objects over walls. Unlike the ballista and catapult, which obtd their power from tension or torsion, the propelling force of the trebuchet was provided by a counterweight (up to 10 tons) (Ref 69, p 281 and Fig on p 280).

One very important cold weapon appeared shortly before the introduction of Gunpowder (ca 1200 AD): the so-called English longbow, which was originally a Welsh weapon. It was claimed to have a range of up to 400 yards, maximum, and effective range ca 250 yards.

It is described and illustrated on pp 5 & 12 of Ref 7 and on p 332 of Ref 69.

Crossbow and longbow continued to be in service up to ca 15th century, competing with early firearms such as handguns (Ref 7, p 4).

History of Incendiaries, Fireworks and Black Powder up to the Middle of 19th Century.

In addition to a brief history of Black Powder (also known as Gunpowder) given in Vol 2 of Encycl, pp B165–R to B168–L, we are giving here chronological listing in the manner done in a Japanese article by Dr Heizo Nambo (Ref 66). Its English translation was obtd thru the courtesy of Mr Gunther Cohn of the Franklin Institute, Philadelphia, Pa, 19103.

We also used the following books, full titles of which are given in the list of Refs, which follows the description: Daniel (1902) (Ref 3, pp I to X, written by M Berthelot); Greener (1910) (Ref 7, pp 13–17); Marshall 1 (Ref 11, pp 11–50); Colver (1918) (Ref 12, pp 1–29, 496–505 & 563–64); Marshall 3 (1932) (Ref 20, pp 1–2); Stettbacher (1933) (Ref 21, pp 3–18); Newman (1943) (Ref 33); Dutton (1960) (Ref 58, pp 5 to 174); Dupuy & Dupuy (1970) (Ref 69, pp 322–33 and other pp listed in General Index on p 1322–R, under Gunpowder); Gorst (1972) (Ref 71, pp 5–12).

We are referring also to historical information given under individual items in Vols 1, 2, 3, 4, 5 and 6.

Accdg to Greener (Ref 7, pp 13 & 14):
There seems little doubt that the composition of gunpowder has been known in East from times of dimmest antiquity. The Chinese and Hindus contemporary with Moses are thought to have known of even the more recondite properties of the compound. It is very possible that Alexander the Great did absolutely meet with fire-weapons in India, which a passage in Quintus Curtius seems to indicate".

The introduction of explosives into Europe followed the Mohammedan invasion. Greek fire, into the composition of which nitre and sulfur entered, was used prior to the fall of the Western Roman Empire (prior to AD 284).

It is known that prior to invention of explosives, incendiaries and fireworks were known, especially to Chinese and Hindus and later to Arabs and Greeks. That is why the chronological list given below starts with them:

1190 BC. This is the earliest date for incendiaries used in warfare listed in Ref 66, p 10. It concerns Trojan troops attacking Greek Navy with incendiaries

Note: The siege of Troy is listed in Ref 69, p 11 R as taking place ca 1184 BC

850 BC. In this year incendiaries were used in the battle of Mesopotamia, Irak (Ref 66, p 11)

Note: This battle is not listed in the book of Dupuy & Dupuy (Ref 69)

500 BC. Accdg to Dupuy & Dupuy (Ref 69, p 36-L), Sun Tzu, a native of the state of Ch' i (Modern Shantung) wrote the "Art of War", the first known military treatise, which provided valuable lessons for military men even today

500-470 BC. In the military tactics of Fan Li of Yuen (Che Chiang province), ballista of stone (Pao of stone) was described (Ref 66, p 11). There were several ballistas of stone listed on p 25 of Ref 66. All of their names contained the word Pao, such as "Pao Che " , "Tan Shao Pao", "Suang Fen Pao" and "Hu Tung Pao" (See Fig on the left side of p 38, Ref 69)

500-450 BC. In the military tactics of Sung Si in Wu of Kiang Su, soldiers used "pile of fire" and fireworks in the attack (Ref 66, p 11)

431-404 BC. Chemical Warfare. The earliest recorded use of chemical gas during the Peloponnesian War (431-404 BC) when Spartans directed the fumes produced on burning of green wood mixed with tar and arsenic towards the besieged Athenians in towns of Plataea and Delium. Another use of poisonous gases was recorded in 187 BC when the Romans were driven from the town they besieged by means of fumes produced by burning feathers and coal. Poisonous fumes produced by burning green wood were used during Franco-Algerian War. More effective chemical agents were proposed during Crimean War (1853-56) and American Civil War (1861-65), but the real chemical warfare began in 1915 (qv) (See Vol 2 of Encycl, p C166 and also next item)

400 BC. Accdg to Dutton (Ref 58, pp 13-14), Spartans soaked blocks of wood in hot pitch & sulfur, piled them against the walls of enemy's fortresses, and applied the flame of a torch. As the walls were then built of wood the device was very effective. This served as a prototype of wild fire used much later. It seems that Spartans also invented a prototype of flame throwers. The original device was in the form of a huge iron caldron filled with burning pitch, sulfur & charcoal. This was mounted on top a movable platform which the attackers moved up to the defenders' walls. The hollow trunk of a tree was then tilted to the edge of caldron, and air was pumped thru the trunk with a big bellows. Thus tarry-sulfurous smoke and tongues of fire were air-blasted upon the enemy to suffocate them. It is probable that fire pots were invented as early as that. They were earthenware jugs of various sizes, some thrown by hand, others for hurling by ballista or catapult. Filled with burning brews and acids, the jugs broke upon striking and showered their contents on the enemy. The most fiendish and destructive of the fire weapons was the Greek Fire, invented in the 7th Century AD

410-304 BC. Allied Navy of Sparta threw shells of sulfur, pitch and pine resin by means of ballista to fire the Athenian Navy (Ref 66, p 11)

325 BC. Alexander the Great's troops which invaded India were defeated by means of "thunder bolts" thrown from the castle wall in Lahore by its inhabitants called Oxydarae (Ref 66, pp 4 & 11)

Note: Accdg to Greener (Ref 7, p 13), Alexander met with real "fire-arms" in India

249 BC. Spartans used incendiaries, consisting of small wood pieces impregnated with sulfur and pitch, in the battle of Plataea (Ref 66, p 11)

160-122 BC. Three components of Black Powder nitre (saltpeter), charcoal and sulfur were de-
scribed by Chinese alchemists in the Chuan Nan ’Tso of Liu An (Ref 66, p 3)

220 BC & 160–122 BC. When several Chinese alchemists were separating gold from silver at low-temp reaction adding saltpeter and sulfur to the gold ore in the alchemist’s furnace, they forgot to add charcoal in the first step. Trying to rectify their error they added charcoal in the last step. This resulted in tremendous explosion. This accident indicates that the mixture of saltpeter, sulfur & charcoal, exploded at that time, was similar to present Black Powder. No advantage of this accident was taken at that time (Ref 66, pp 3 & 29–31)

141–87 BC. In Robert Norton’s “Gunner” published in 1628, it was reported that BkPdr was invented at the time of King Vitye (Wu Ti) of China and used in battles (Ref 66, p 11)

Note: More detailed description of this subject is given in Greener (Ref 7, p 17)

200 AD. Tseng Tsao of Wei at the San Quo Dynasty threw stones in the battle of Kuan Tu by means of the “thunder vehicle” (Ref 66, p 12), which was actually stone ballista (Ref 66, p 24)

222–235 AD. Alexander VI of Roman Empire called the “automatic fire” a ball (consisting of quicklime and asphalt) which spontaneously ignited on coming in contact with water (Ref 66, pp 6 & 12)

275 AD. Accdg to Greener (Ref 7, p 14), Julius Africanus mentioned “shooting Powder”, but its compn was not given

350 AD. Aeneus of Rome put sulfur, pitch, incense, pine-resin, and tow (crude flax fibers) into an oval wooden container, ignited the mixture and threw it on the decks of enemy ships (Ref 66, p 12)

668 AD. On the basis of advice given by Kallinikos of Heliopolis in Syria, an incendiary of secret formula known now as Greek Fire or Sea Fire. Fire was used by the Byzantine fleet to annihilate the Caliphate Navy (Ref 11, p 12)

Note: Accdg to Marshall 1 (Ref 11, p 12), Col Hime in his book listed here as Ref 4, came to the conclusion that, besides naphtha, sulfur and pitch, there must have been quicklime in mixture, which on coming in contact with sea water turned into slaked lime whilst raising the temperature to the point of ignition of sulfur. Greek Fire mixture was discharged from tubes or spigots located in the bows of the ships, against the enemy. Marshall prep'd a similar mixture but was not able to ignite sulfur. He thought that the naphtha was ignited by a flame located at the orifice of the discharge tube (Ref 11, p 13)

Later the name “Greek Fire” was given also to combustible materials which were ignited and then thrown by ballista or other machines. These comps were solid masses of sulfur, pitch, naphtha, and other combustibles, and when saltpeter was discovered it also was included. Such mixtures may more correctly be called Wild Fire. Such mixts were used by Moslems in the Crusades (See under year 1250 AD) (Ref 11, p 13)

673–678. Moslem Arabs (of Caliphate) were defeated at the walls of Constantinople by defenders using Greek Fire (Ref 58, p 14)

683. In the battle of Mecca (Saudi Arabia), Kaaba was fired by the incendiary missiles of Syrian Army (Ref 66, p 13)

690. Accdg to Greener (Ref 7, p 14), the Arabs or Saracens are reputed to have used Gunpowder at the siege of Mecca; some writers even affirm that it was known to Mahomet. Marcus Graeco described in “Liber Ignium” an explosive composed of 6 parts saltpeter and 2 parts each of charcoal and sulfur

716–718. The Arabs of Caliphate again appeared before Constantinople with 1800 ships, but again were defeated by Greek Fire (Ref 11, p 12)

904. Accdg to Greener (Ref 7, p 14), it was recorded that Gunpowder was used by the Saracens at Thessalonica

904. The Caliphate troops (Baghdad, Iraq), when attacking Salonica (Greece), threw ceramic pots charged with a mixture of pitch, pine resin, and quicklime over the heads of enemy troops to suffocate them (Ref 66, p 13)

904. Before the end of Tang’s rule, Cheng Fan attacked Yu Chang by means of “flying fire” hurled by ballista (Ref 66, p 13)

940. The Chinese device, called “fire pao”, consisted of ballista throwing a “fire ball”, made of explosive similar to BkPdr, tied to the top of an arrow (Ref 66, p 13)

941. The East Roman (Byzantine) Navy poured from the tube or syphon located at the boat’s stern a liquid (“sea fire”) made from naphtha, quicklime and sulfur, over the Russian ships attacking Constantinople (Ref 66, p 14)

1000. Tung Fu of Sung Dynasty invented “fire arrow”, “fire ball” and “fire dart rocket”
(Ref 66, p 14)

Note: In the book of Dutton (Ref 58, p 5), it is stated that during Sung Dynasty (960–1280) some unknown man described the invention of To-lo-iskang which consisted of a long bamboo tube in which a handful of explosive similar to BkPdr was placed. When set on fire a strong flame came out accompanied by ejection of some grains of pdr (to a distance up to 150 paces) and creating thunderous noise similar to that of “paos.”

1002. In the reign of Chen Tsung of Sung, Lo Yung Si invented a “portable ballista” (Ref 66, p 14)

1040. At the time of Chen Tsung of Sung, Pe Sung built a Gunpowder Plant in Pien King ( Hai-feng, Honan) (Ref 66, p 14)

1043. Russian fleet was defeated at the siege of Constantinople by means of Greek Fire (Ref 11, p 12)

1045. Pe Sung Govt issued permit to Wu Ching Tsung Yao for manuf of “firing arrow”, “thorn fire ball”, and “poisonous smoke ball” (Ref 66, p 14)

Note: Brief description of these and other Chinese devices is given in Ref 66, pp 32–35 & 41)

1045. Accdg to Nambo (Ref 66, p 38), the compn of powder for “fireballs”, as described by Wu Ching of Sung, was: saltpeter 48.5, sulfur 25.5 & other ingredients 26.0%.

1045. Accdg to Nambo (Ref 66, p 38), the compn of powder for “thorny fire balls”, as described by Tsung Yao of Sung, was saltpeter 50.0, sulfur 25.0, charcoal 6.25 & other ingredients 18.75%, while the compn of powder for “poisonous fire balls” was saltpeter 38.5, sulfur 19.25, charcoal 6.4 & other ingredients 35.85%.

1073. Accdg to Greener (Ref 7, p 14), Gunpowder was used by King Salomon of Hungary at the siege of Belgrade.

1098. Accdg to Greener (Ref 7, p 14), in a sea conflict betw the Greeks and Pisanians, the former had fire-tubes fixed at the prows of their ships.

1096–1099. In the battle of Nice of the 1st Crusade, the Caliphate troops threw burning pitch and fatty balls. They also shot fire arrows with pitch, sulfur and tow from the walls of Jerusalem (Ref 66, p 14)

1126. In a triple war among the Sung, Kin and Mongols, explosive arms were used, first by Sung and then by others. The devices included: fire arrow, radiation firearm, fire burster, fire lance, fire ball, thunder ball, etc. They are described in the paper of Nambo (Ref 66, p 14)

1129. It was suggested equipping Chinese fleet with weapons such as ballista, fire arrow, etc (Ref 66, p 15)

1132. Li Heng besieged Te An using long ladders. The General of Defense Army used 20 fire lances, long bamboo tubes charged with Gunpowder (Ref 66, p 15)

1135. During Chinese Civil War, Kin’s fleet used thin, brittle pottery bottles charged with quickline, poison and iron pickles (?). The resulting smoke injured the eyes of enemy (Ref 66, p 15)

1147. Accord to Greener (Ref 7, p 14), Arabs used firearms against the Iberians (Spanish) 1169–1189. In the reign of Haiao Tsung of Nan Sung, true fireworks, Yan Huo, made their debut. They were similar to those used today (Ref 66, p 16)

1214. When the Mongols captured Chung Ching, Kin moved his capital to Pien Ching and the Mongols came into possession of techniques for manuf of Gunpowder and firearms (Ref 66, p 16)

1218. Accord to Greener (Ref 7, p 14), there was artillery at that time in Toulouse.

1218–1258. In the reign of Nan Sung, Gunpowder and fireworks were manufd in quantity sufficient not only for local consumption but also for export, mostly to Caliphate country (Irak) (Ref 66, p 19)

1220. Accord to Sancho (Ref 30, p 254), Moors used machines for hurling stones.

1221. During the siege of Chin Chou, Kin arranged 13 sets of ballistae to shoot “iron bombs” into the castle. The iron bomb (Tie huo pao), was cast, hollow ball, 6cm in diam, filled with Gunpowder and equipped with a fuse cut to the desired length and ignited before launching the bomb (Ref 66, p 16)

1221. When Chings (Chengchiz) Khan and his Mongolian troops invaded Europe and Russia, they threw stones by crossbow ballistae to break fortification walls; also used poisonous powder flasks (Du huo fou), fire arrows (Huo jian) and fire stone balls (Hua pao). Such weapons were used at the battle at Amu Dariya river against Khorasan Empire (present Russian
Turkestan) and in other battles (Ref 66, p 18)

1225. Saltpeter became known in Europe
(Ref 11, p 21)

1231. When Mongolian general Toulou captured
Hochung Fu, the Chinese garrison escaped by
crossing the channel on old ships; then they
destroyed the ships by iron bombs (Chen tien
lei) (Ref 66, p 16)

1232. During the 2nd attack by Mongols on
Nan Ching, Kin used iron bombs (Chen tien lei)
charged with powder and provided with a fuse.
They flew off a long way to burst. He also
used flying fire lances (Fei huo quiang) (Ref
66, p 16)

1232. Acceg to Marshall 1 (Ref 11, pp 14–15),
in the Chinese chronicle “Tun Klang Kang-mu”,
translated into French in the Journal Asiatique
for Oct 1849, there was a description of a
weapon used at the siege of Pien King (now
Kai-fung-fu). It was “Ho-pao” or “Fire-pao”,
called “Tchin-tien-loulu”, translated as “thunder
that shakes the sky”. The device consisted of
an iron pot filled with “yo” (medicine). As
soon as light was applied the “pao” rose and
fire spread in every direction. Its noise re-
sembled that of thunder and could be heard
more than 100 lis (33 Engl miles). It could
spread fire over more than a third of an acre.
This fire even penetrated the breast plates on
which it fell. These “bombs” were probably
thrown by ballista

Note: This item described in Marshall (Ref 11)
is identical with that described by Nambo (Ref
66, p 16)

1232. Mongolian soldiers constructed with ox-
ides a passage to the ramparts of Nan Ching
castle. They commenced to sap the walls and
made holes in them in which they could remain
sheltered from the besieged men above. Then
the besieged Chinese hung “fire pao” on the
chains down the face of the wall. When the
“pao” reached the levels of hidden Mongols,
they were ignited to explode, thus killing the
invaders. In addn the besieged used “Fei ho
tiang” (arrows of flying fire), which consisted of
arrows with attached substance which was
susceptible to taking fire in flight. While flying,
such arrows spread flames over a width of 10
paces. These Chinese devices were much feared
by Mongols (Ref 11, p 15 & Ref 66, p 17)

Note: Marshall thought that the effects of the
above Chinese devices could hardly have been
produced without the use of saltpeter, nor the
great noise without an explosive (Ref 11, p 15)

1233. In the battle of Nan Ching, Kin defeated
the Mongols by means of “flying fire lances”
(Huo quiang), which were tubes, 0.8m long
made of 16 piled sheets of strong paper charged
with an incendiary mixture and porcelain frag-
ments. On ignition the flame shot no less than
3 meters (Ref 66, p 17)

1234. Allied Army of Mongols and Sung
against Kin used stone fire balls, called “Huo
pao”. Kin was defeated and committed suicide
(Ref 66, p 17)

1241. When the Mongolian Khan Bhatu (Batyi)
invaded Russia and Western Europe, and went
as far as Nahlstadt near Lignitz, his troops used
poisonous smoke bombs (Ref 66, p 18)

1241. When Bhatu troops besieged castle
Olmütz in Moravia (present Slovakia), they used
fire arrows to burn the temple (Ref 66, p 19)

1247. Acceg to Greener (Ref 7, p 18), Seville
had been defended by a “cannon throwing stones”

1249. English Monk Roger Bacon (1214–1292
or 1294) gave in his book “Opera Artis et
Magie” the compn of Gunpowder as saltpeter
7, sulfur 5 and coal 5 parts (Ref 66, p 17)

Note 1: Acceg to Greener (Ref 7, p 14), it
is presumed that Bacon obtd his knowledge
about BrPdr from the Treatise found in the
Escorial (Spain) Library collection

Note 2: Acceg to Marshall (Ref 11, p 17). Bacon
published ca 1250 his “Opus Tertium” and a
part of its translation (from Latin) was published
by Col Hime in the Journal of the Royal Arti-
lery, July 1911

1250. Compn of Roger Bacon’s Gunpowder is
given in Ref 11, p 26 as: saltpeter 41.2, char-
coal 29.4 & sulfur 29.4%

1250. It was mentioned under the year 668 AD
as the incendiary which Marshall proposed to
name Wild Fire. Such incendiaries were used
by Arabs against the Crusaders. Thus Joinville,
participating under St Louis in the disastrous
Sixth Crusade wrote that this incendiary device
came flying thru the air like a winged long-tailed
dragon, about the thickness of a hoghead, with
the report of thunder and velocity of lightning;
and the darkness of the night was dispelled by
this deadly illumination. These devices terrified
the crusaders, because they believed them to be
the products of Devil. Fortunately the damage was only slight (Ref 11, p 13)
1257. In the reign of Li Tsung of Sung, the invading Mongolian troops were held back by means of big supply (found in Chinese arsenals) of iron bombs (Tie huo pao) and flying fire lances (Huo quiang) (Ref 66, p 17)
1259. In the battle of Niebla, Spain the Moors (Arabs) threw (by means of ballista) stones and filth and also “missiles with thunder” (Ref 66, p 19)
1259. In the battle of Melilla, Morocco, the Moors used cannons and firearms (Ref 66, p 19) Note: It is more probable that they threw “Wild Fire” missiles by means of ballistae Note: In the opinion of Marshall 1 (Ref 11, p 13), it is probable that the “missiles with thunder” used by Moors in Spain and Morocco were the same as used in 1250 AD by Arabs against the Crusaders of Saint Louis (See above under the year 1250 AD)
1259. Acqdg to Marshall 1 (Ref 11, p 15) and Dutton (Ref 58, p 3), it was stated in Chinese annals that in the 1st year of Kai-King was made an appliance called “Tho-ho-tsiang”, meaning “lance with violent fire”. A nest of grains of powder was introduced into a long bamboo tube, and when the pdr was lit the flame shot forth from the tube with a noise like that of a “pao”, which could be heard at a distance of about 300 paces (Dutton gives 150 paces). It was a device resembling current Roman candle
1259. In the reign of Li Tsung of Sung Dynasty was invented “fire lance with projectile” (Shi huo quiang). It consisted of a bamboo tube charged with powder and contg on top of it a ball serving as a projectile. When the powder was ignited the flame came out and the ball flew away. It was accompanied by a loud noise which was heard at a distance of 270 meters (Ref 66, p 17) Note: It is not stated whether the lower end of bamboo tube was open or closed. In case of closed tube it was necessary to attach it to some firm object because the recoil will kick it in reverse direction, and in this case it could be considered as a prototype of modern rocket. Similar bamboo weapons were used by Arabs (Ref 58, p 9)
1268. In order to relieve the 5-year siege of Sing Yang by Mongolian troops, 3000 Chinese were sent on 100 boats and they routed the Mongols by means of fire lances and other fire arms (Ref 66, p 18)
1273. Acqdg to Greener (Ref 7, p 18), Abou Yuesof used cannon firing stone shots at the siege of Sidgil-messa
1274. When Chinese troops under Yang landed at the Hakata Bay (Japan) they shot iron bombs (Tie pao), which surprised and defeated Japanese troops. They called the bombs “teppo” (Ref 66, p 20)
1277. When Ching Chiang fell, the defending general of Sung fired a large cannon (Huo pao). This destroyed the castle walls and burned to death a 200-man garrison (Ref 66, p 18)
1279. In the Naval battle near Asmen (Huan Tang), betw Chang Shih Chieh of Sung and Chang Hung Fan of Yuan large cannons (Huo pao) were used by both sides. The Navy of Sung was defeated (Ref 66, p 18) Note 1: Chronological list given in the article of Nambo (Ref 68) does not go later than 1279 AD
Note 2: Acqdg to Marshall 1 (Ref 11, p 18), the Chinese do not appear to have developed expls beyond this point, or to have made the next step, namely of causing the pdr to throw a heavy metal projectile instead of a ball of fire. This step could only be taken by a nation which was at once progressive and well acquainted with the working of metals Note 3: Marshall forgot to mention that progress in China was impeded by Chinese Civil Wars and Mongolian invasion Prior to 1300. The Arabs introduced saltpeter in their Greek Fire and other incendiaries. In Europe saltpeter was more scarce than in Africa and Asia (Ref 11, p 17)
1300. In the “Liber Ignium” of Marcus Graecus, which was probably translated into Latin from an Arabic source ca 1300, were several references to incendiaries used by Arabs. One of them “ignis volatilis” (flying fire) consisted of resin 1, sulfur 1 and saltpeter 2 parts, dissolved into a hollow reed or wood. This was in the opinion of Marshall 1 (Ref 11, p 17) an incendiary (“wild fire”)

The compn listed on p 18 of Ref 11 consisted of sulfur 1, vine or willow charcoal 2 & saltpeter 6 parts. The ingredients were rubbed down together on a marble slab and put into a case, which was short & strong and filled only half-full
when it was desired to make a loud noise. On the other hand, a thin, long container, known as "tunica" was completely filled with the above powder and flew on being lit. This "flying tunic" was an imperfect rocket (Ref 11, p 18)

14th Century. Early. Acdept Greener (Ref 7, Chap IV, "Early Hand Fire Arms", pp 44-51): "No distinction can be drawn between the small cannon or "crush-gun" of the fourteenth century and the earlier fire-arms. A pyrotechnical piece developed into a variety hand weapon and used for military purposes, especially for causing disturbances among troops, frightening horses and stampeding cattle, was employed by Eastern nations and by Arabs in Northern Africa". Such a device which was also used by incendiaries, pillagers, and outlaws is shown in Fig on p 44 of Ref. The gun consisted of an iron tube ca 6 ft long, covered with two hollowed pieces of wood, and bound round with hemp, hair or leather. The tube was fastened to a stick, which could be held under arm or placed on a shoulder. The compn of the charge resembled a Roman candle

14th Century. Acdept to Marshall 1 (Ref 11, p 25), some early powder makers added camphor, sal ammoniac and gum to their Gunpowder to prevent its crumbling. In the "Codex Germanicus" of the 14th Century, the following recipe is given: If you want to make a good, strong powder, take 4 lbs of saltpeter, 1 lb of sulfur, 1 lb of charcoal, 1 lb of "salpratica" (mixture of saltpeter, camphor & ammonia dissolved in alcohol), 1 oz of sal ammoniac and one twelfth part of camphor. Pound it all together with some aze until well mixed and then dry in the sun. When "corning" of powder was invented in 1425, there was no necessity to incorporate camphor

1304. Acdept to General Fuller (Ref 50), this is the earliest year when a weapon resembling a cannon was documented. It was in an Arabic manuscript. Acdept to Dutton (Ref 58, p 6), it was preserved in the Asiatic Museum at StPetersbourg, Russia, a fragile and faded Arabic parchment, which presents a sketch drawn by Shems Eddin Mohammed in 1304 of a new weapon. It is described as a "spear that hurls arrows at the enemy. The spear was a bamboo tube which had been reinforced by metal bands. Among flame and smoke, an arrow is shown on the sketch flying from the muzzle. Marshall 1 (Ref 11, p 19) said: "There is a manuscript in Asiatic Museum at Petrograd probably compiled by Shems ed Din Mohammed about 1320, which shows tubes for firing off both arrows and balls by means of powder"

Note: It seems that the manuscript listed by Marshall as of 1320 is the same as described by Fuller and Dutton as of 1304. It is also probably the same manuscript as listed by Sancho (Ref 30, pp 253-54), saying that the manuscript written in 1304 in Arabic, discovered by Reinaud & Fave in a library at StPetersburg, contd the following compn: saltpeter 10 drachmas, charcoal 2 & sulfur 1.5, called Medfraw

Note: Historical description of development of cannons is given in Vol 2 of Encycl, pp C26 to C29

1308. Acdept to Greener (Ref 7, p 18), Ferdinand IV of Castile employed guns (marquenas de trueñas) in the siege of Gibraltar

1308. A cannon was used for defense of the castle of Heyer on the Rhein. The castle was destroyed and the cannon remained in its ruins until it was found in 1560 (Ref 7, p 18)

1310. Acdept to Greener (Ref 7, p 18), firearms were used by the English at the siege of D'Eu

1311. Acdept to Greener (Ref 7, p 18), Ismail attacked Bazas in Granada, Spain, with "machines throwing balls of fire, with noise like thunder"

1313. Acdept to Col Hime (Ref 10, p 2), a weapon resembling cannon was invented by an unknown German monk. The weapon was used in 1314 at the battle of Bannockburn (Quoted from Marshall 3 (Ref 20, p 1)) (See also Vol 2 of Encycl, p C26)

1313. Acdept to Greener (Ref 7, p 18), the city archives of Ghent (Belgium) stated that the city was in possession of a small cannon

1313 (About). Acdept to Newman (Ref 33, pp 71 & 75), the gunmakers of Flanders constructed (no year is given) the famous Giant Bombard of Ghent, also known as "Dulle Griete". Its caliber was 25 inches, while its length was 40 feet

Note: Acdept to Greener (Ref 7, p 27), Sir John Froissart stated that this gun was used by D'Aravelle at the siege of Oudenarde (no date is given)

1320. Acdept to Greener (Ref 7, p 17), Berthold Schwartz, a monk of Friburg in Germany, studied the writings of Bacon regarding explosives, and manufactured gunpowder while experimenting. He has commonly been credited as an inventor, and at any rate the honor is due to him for
making known some properties of gunpowder; its adoption in Central Europe quickly followed his announcement, which is supposed to have taken place about 1320. It is probable that gunpowder was well known in Spain and Greece many years prior to its being used in Central Europe.

1323. Accdg to Sancho (Ref 30, p 254), “balls of fire with noise” were used at the siege of Baeza (or Baza), Spain.

1324. Accdg to Marshall 3 (Ref 20, p 1), the town of Metz, France appears to have cannons at that time.

1325. Accdg to Greener (Ref 7, p 18), in the records of Florentine Republic was stated that two officers were ordered to manuf cannons and iron balls for the defense of castles and villages.

1326. Accdg to Sancho (Ref 30, pp 254–55) in Florence there were used cannons of bronze to shoot iron balls and the existence of plant manuf cannons is mentioned in chronicles.

1327. Accdg to Green (Ref 11, p 19), in an illuminated manuscript entitled De Officiis Regum, written in Latin by Walter de Millemette and preserved in Christ Church Library, Oxford, England, there is a drawing reproduced here of a rudimentary gun shaped like a bottle, showing a dart used as a missile and a man applying a fire to a touch hole. The same illustration is reproduced in the book of Dupuy & Dupuy (Ref 69, p 333). The beautiful manuscript (in color) was seen by the senior author of this Encycl during his trip to England.

Note: Accdg to Dutton (Ref 58, p 32) this type of weapon was made of brass.

1326. Accdg to Sancho, the French attacked Gravelines using cannons & bombards (Ref 30, p 255).

1327. Accdg to Marshall 1 (Ref 11, p 19), the Republic of Venice ordered the provision of metal cannons and iron bullets for the protection of castles and villages. Accdg to Sancho (Ref 30, p 255), the Venetians took Guerra employing cannons.

Note: Definition of term “cannon” is given in Vol 6 of Encycl., under GUN. It is preferred to that given in Vol 2 under CANNON. We are using the word “cannons” for plural in order to avoid confusion.

1327. Accdg to Greener (Ref 7, p 19), John Barbour wrote in 1372 that in 1327 at the battle of Warwater, the Scotch first saw the firearms.

1330. Accdg to Sancho (Ref 30, p 254), the first firearms appeared in France during the reign of Philippe de Valois (1293–1350).

1331. Cannons were used by Moors at the siege of Alicante (Spain) to open a breach in the wall (Ref 11, p 19 & Ref 30, p 254).

1338. Accdg to Green (Ref 7, p 18), the English contemporary record mentions that King’s ships “Bernard de la Tour” and “X’ofre de la Tour” were equipped with cannons and handguns.

1338. Accdg to Nambo (Ref 66, p 39), the compn of French powder of unknown use was: saltpeter 50, sulfur 25 & charcoal 25%.

1338. Accdg to Green (Ref 7, p 19), powder and cannons were provided for the protection of the ports Harfleur and l’Heure (France) against Edward III.

1339. Accdg to Marshall 3 (Ref 20, p 1), there was mention of six brass “gennes” in the London Guildhall, together with “petelet” weighing 4.5 lbs and 32 lbs of powder, showing that the darts used in the very earliest guns were replaced by small balls.

1340. Sir John Froissart in “Chronicles of England, France and Spain,” Bk 1, Chap 72, stated that Scots used cannons in the siege and capture of Stirling and in 1340 or 1341 Gedymyn, Grand Duke of Lithuania was killed by a cannon ball at the fortress of Walona, which was being besieged by the Germans. It is evident that at that time the guns were in use for siege warfare but they not sufficiently mobile for the field (Ref 20, pp 1–2).
14th Century. Accdg to Nambo (Ref 66, p 39), the compn of German powder of unknown use was: saltpeter 66.7%, sulfur 22.2% & charcoal 11.1%. A slightly different compn is given in Marshall 1 (Ref 11, p 31). It was: saltpeter 66.6%, charcoal 16.6% & sulfur 16.6%

1340. Accdg to Colver (Ref 12, p 496), solid spherical projectiles for cannons started to replace the arrow-shaped projs. The material was stone, bronze or iron

1340. In two frescoes in the church of former monastery of San Leonardo in Lecce, near Siena, painted by Paolo del Maestro Neri, are shown: a large cylindrical cannon discharging a spherical ball, and several hand guns (Ref 11, p 19)

1341. Accdg to Sancho, the Scots took from the English the town of Stirling employing artillery (Ref 30, p 254)

1342. Cannons were used by Moors in the defense of Algéciras (Spain) against Don Alfonso XI de Castilla (Ref 11, p 19 & Ref 30, p 254)

1342. Accdg to Marshall 1 (Ref 11, p 19), Counts of Derby and Salisbury, who were present at the above battle of Algéciras, supposedly introduced guns and gunpowder into England. In the following years there are several refs in the accounts of the Wardrobe of Edward III of payments for saltpeter for manuf of Gunpowder

Note: Accdg to Greener (Ref 7, p 15), ingredients for Gunpowder manuf'd in England were usually separately purchased abroad and mixed when required. Mr Oliver of Boklesberry appears to have been one of the first dealers in Gunpowder (Ref 7, p17)

1344. Accdg to Dutton (Ref 58, p 32), Edward III of England was the first to add cannons to his Army. There were at least two clumsy bronze cannons with him when he crossed the channel to France in the War against Philippe de Valois

1346. Accdg to Dutton (Ref 58, p 34), the battle of Crécy, fought on August 26, 1346 was won by English longbows, but it is not known if the cannons took action, because some accounts do not mention them at all

Note 1: Accdg to Greener (Ref 7, p 18), John Cook, a clerk of the King's wardrobe, to which department the arms and munitions of war belonged, stated that 912 lbs of saltpetre and 846 lbs of sulfur were provided for the use of the army in France; later in the year, before Calais, he obtained a further supply. That firearms were used by the English at Crécy in 1346 is a well ascertained fact

Note 2: Accdg to Marshall 3 (Ref 20, p 2), it was improbable that cannons were used at the battle of Crécy, because at that time no light weapons existed but there were only heavy siege cannons

Note 3: Gen Fuller stated in Vol 1 of his book (Ref 50) that accdg to Sir Charles Oman the weapons used at Crécy were ribaudesquins, which were invented in 1339 and may be considered as ancient machine guns. They consisted of several iron tubes clamped together to form a device resembling a multiperforated cylinder. The tubes could be fired separately or simultaneously

1347. Accdg to Dutton (Ref 58, p 39), cannons were definitely used by the British at the siege of Calais, France. They also had ribaudesquins, mentioned above under Note 3

14th Century, 2nd Half. A brief description and illustrations of 14th century cannons are given in Greener (Ref 7, pp 20–5). The illustrations include: p 21 – Early breech-loading cannons; p 22 – Iron breech-loading cannon, Cast breech-loading cannon, Italian bombard, and Early English breechloader; p 23 – Italian muzzle-loading cannon called “Cerbottain”; p 24 – German Artillery; p 25 – Elbow-joint bombard

1350. Bullets. The primitive bullets, which were either stones or metallic slugs of irregular shape were gradually replaced by cast spherical lead slugs (balls) which were smaller in diam than the bores of the guns which at that time had irregular calibers. When the calibers were standardized the bullets were made of the same caliber as the barrel. All early bullets were loaded thru the muzzle on top of a wad previously placed on top of a charge of BKpdr. There were no cartridges until the time of Gustavus Adolphus, born in 1594, King of Sweden, 1611 to 1632, when a successful paper cartridge was invented. The first elongated bullet was invented in 1823 by the British Officer Norton. The other elongated bullet was invented in 1836 by W. Greener, the famous Brit gunmaker. More detailed description of historical development of Bullets is given in Vol 2 of Encycl, pp B324 to B331
charcoal 22.2% (See also Ref 11, p 26)

1360. Accdg to Marshall 1 (Ref 11, p 23), as quoted from paper of F.M. Feldhaus in SS 4, p 275 (1909), the Rathaus at Lübeck was destroyed by fire thru the carelessness of Gunpowder makers

1372. Accdg to Feldhaus (see above) there was no mention of Gunpowder or firearms in the archives of Augsburg before this year, although Berthelot claimed that a powder mill existed there since 1340 and that a mill was at Spandau in 1344 and at Liegnitz in 1348 (Ref 11, p 23)

1378. Accdg to Greener (Ref 7, p 22), the earliest cast cannons made of copper and tin were produced by a founder named Aran at Augsburg, Germany. They were primitive breech-loaders, built up of iron strips surrounded by iron rings — a method which continued for several centuries (See illustrations on p 22 of Ref 7)

1379. Cannons and bombards were employed at the siege of Puil de Guillen in Perigord (Ref 30, p 254)

1379. The French possessed at Cambrai 5 iron cannons & 5 of other metal (Ref 30, p 254)

1381. Accdg to Greener (Ref 7, p 45), the first account of hand-cannon being used in Germany was in 1381, when the town of Augsburg supplied thirty men armed with them to the contingent of the Swabian towns in their war against the South German nobles

Note: In the same book it is stated that hand guns were used in the 14th century by French, Italians and Netherlanders (See also Vol 2 of Encycl, pp B324—B325)

1382. Accdg to Gorst (Ref 71, p 6), the first recorded use of Gunpowder and artillery in Russia was in 1382 at the defense of Moscow against the invading Mongols (Tartars) (called by Russians Tatary), under the Khan Takhtamys

1386. Accdg to Greener (Ref 7, p 49), handguns were used in Padua, Italy

1388. The fire-arm of Rouen was, accdg to Greener (Ref 7, p 20), a small iron-forged weapon. It was shooting feathered iron arrows, loaded from muzzle, propelled by a charge of Gunpowder, which was put in a separate movable breech-block or chamber (See illustrations on pp 21 & 22 of Ref 7)

1389. Accdg to Sancho (Ref 30, p 255), the manuf of firearms and Gunpowder started in Russia

1391. Accdg to Fuller, Vol 1 (Ref 50), solid iron balls partly replaced stones as cannon missiles, but hollow iron projs filled with Gunpowder (to act as an explosive) were not invented (in Holland) until the 16th century (Greener, 1st edn, Ref 2, p 27)

14th Century, Late. The smallest among the early fire-arms were the Italian bombards, one of which is shown in Fig on p 23 of Ref 7. The bombards were muzzle-loading, and had the powder chamber of much smaller caliber than the forward portion. This fore part was usually more or less tapered so that shot of different diameter might be fired

Another early weapon was the “bombardo cubito” or “elbow-joint gun” which is described and illustrated on p 25 of Ref 7

Large cannons were made at a very early date, even if they were never used. The fact that such a weapon was possessed by a town, possibly terrorized opponents. One of such huge weapons was the “Faust Bucaleus” of Frankfurt, made in 1399 and used at the siege of Tannenburg Castle (Ref 7, p 27). Method of manuf of Gunpowder at the end of 14th century is shown in illustrations on p 16 of Ref 7. It is briefly described on p 26 of Marshall 1 (Ref 11)

1399. Accdg to Sancho (Ref 30, p 255), the English used 10 cannons during the siege of St Malo (France)

1400. Cristina de Pizana employed “balls of fire” and other incendiary projectiles (Ref 30, p 255)

15th Century, Early. Accdg to Dutton (Ref 58, pp 39–40), by 1400 iron cannons, bound by iron hoops to keep them from bursting, and iron cannon balls were coming into use. At the end of the 14th century a handgun was invented which weighed 10 lbs and fired lead shots. One man carried it mounted on the stand and aimed at the enemy. A similar gun but of 16th century is shown in Fig on p 54 of Greener (Ref 7). A cavalry handgun, called petronel is shown on p 46 of Greener and a larger semi-portable gun on p 42. Some lighter hand guns did not need any stand — they could be fired from the shoulder, as shown in Figs on pp 45 & 51 of Greener. A simpler hand gun is shown and briefly described in Vol 4 of Encycl, p D755-R with Fig on p D755. Its stock was a long piece of wood held under the arm of the soldier. The first improvement of hand gun was the introduction of bent stock and
the resulting gun was known as arquebus. It is briefly described in Vol 1 of Encycl, p A488-L and illustrated on p 54 of Greener (Ref 7). This weapon was known accdg to Newman (Ref 33, p 33) before 1503, although Dutton (Ref 58, p 60) claimed that it was invented in 1521.

Each hand gun or arquebus was provided at the rear, closed end (breach) with a touch hole serving for igniting by means of a match of BkPrd propIpt located in the barrel (See Vol 2 of Encycl, p C73, and Vol 4, p D753-R). When the match was mounted later (after 1460) on a S-shaped lever, called “cock”, the device became known as matchlock (See also under years 1460–1480).

1410. Accdg to Gen Fuller 2 (Ref 50), canister or case-shot was invented in 1410 and although it was partly replaced by shrapnel (See Ref 11, p 30), it is still used (See Vol 2 of Encycl, pp C24-R & C25-L).

1413. Accdg to Greener (Ref 7, p 27), Mahomet II of Ottoman Empire used at the siege of Constantinople a giant cannon which was 48 inches in diameter and fired stones weighing 600 lbs.

1415. Accdg to Marshall 1 (Ref 11, p 33), the English used at the siege of Honfleur, France, mines charged with Gunpowder for blowing up the walls of fortifications.

1418. Accdg to Greener (Ref 7, p 33), this seems to be the first year when mortars were used. It was in defense of Cherbourg, that these weapons hurled red-hot shot. Three illustrations of mortars are on pp 35, 36 & 37 of Greener. Current mortars are briefly described in Vol 2 of Encycl, p C27-R, under CANNON. Mortars were also used at the siege of LaFère in 1580 and at Gibraltar in 1780.

1419. Accdg to Dutton (Ref 58, pp 41 & 46), the earliest cannons were not mounted on wheels but were just dragged by beasts or men over the roads or fields. For the 1st time in history cannons were not put on wheeled gun carriages, but on stout four-wheeled farm carts. This was done by Jan Žiska (pronounced Zhiska) in 1419 during the insurrection of Czechs against Sigismund of Germany who was the ruler of country known as Bohemia at that time. Žiska’s supreme stroke was a big “wagon-fort”, called Wagenburg. Zhiska armored his forts, pierced iron sides with loop-holes and put at each hole a soldier with a handgun. This may be considered as a predecessor of present “armored vehicles” and of tanks. With the help of wagon-forts, Zhiska won many battles against the Germans until his death which took place in 1324. This insurrection is known as Hussite War and it lasted fourteen years.

Accdg to Greener (Ref 7, p 49), handguns figured conspicuously during Hussite War.

Note: Previous to this war the “wheeled forts” were used by the Tartars against the Russians, but they were manned by bowmen instead of gunmen. The idea was resurrected in 1919 during Russian Civil War (1919–1920) when the two-wheeled carts (known under the name of tachanka) were equipped with light guns and machine guns.

1425. In this year is the earliest known reference to corned powder. It was in the “Firebook” of Conrad von Schongau and listed in Marshall 1 (Ref 11, p 25). “Corned” means that the BkPrd cake was broken into small grains instead of preparing it in fine powder, known as “serpentine”. The process was essentially the same as described in Vol 2 of Encycl, p B169-L.

Note: Apparently, corned powder came gradually into use for small arms and hand grenades during the 15th century and for big guns in the 16th; the construction of these being sufficiently improved by that time (Ref 11, p 26).

1425. French Gunpowder mills began to grain and classify their pdr by passing thru-sieves (Ref 11, p 26).

1453. Accdg to Dutton (Ref 58, pp 49–55), the next important step in the use of firearms came during the siege of Constantinople by Ottoman Turks. In 1453 the big Army led by Sultan Mahomet II, lined up against the walls of Constantinople, fourteen batteries, totaling 69 cannons. Thirteen of the cannons were of gigantic size and could project stone balls weighing more than ½ ton each. Two of such stones measuring 46 inches in diameter may be seen now at the Turkish Museum at Istanbul. The biggest of these cannons was the Basilica. It required 60 oxen for dragging it on a sled. After two months of constant bombardment of all 69 cannons aimed at one section of the wall, a big gap was made and this allowed the Turkish Army to pour thru.

Note: Mahomet’s great cannons were cast
by a Hungarian named Urban, because such industry did not exist outside of Europe

15th Century, Middle. Accdg to Greener (Ref 7, p 27), the production of large muzzle-loading cannons became quite common in Germany and several of these huge weapons were often referred to by name, such as "Foulenette" and "Helfant". Of these, the most known was Mons Meg constructed in city of Mons, Belgium. It weighed nearly 4 tons and threw stones weighing over 350 lbs. It was installed at Edinburgh Castle and was supposed to have been of the same general construction as the cannon which in 1460 killed King James II. Its illustrations are on p 26 of Ref 7, together with illustration of large German cast gun.

Another huge cannon was "Endorferien" made in 1487 for Sigismund of Tyrol. It was a pair with "Bassina" at the Paris Museum (Ref 7, p 27). Illustration of French "Orgue des Bombardes" is on p 27 of Ref 7.

During the reign of Henry VI, who lived between 1421 & 1471, English artillery was inferior to that of France, and many French towns previously occupied by English fell back to French. This situation was remedied later by Henry VIII (1491-1547) (Ref 7, p 30).

Accdg to Fuller 2 (Ref 50, p 12), a light, long muzzle-loaded artillery weapon, called Culverin was developed in the 15th century (See Vol 3 of Encycl, p C573-L). Culverin fired iron balls weighing 17 lbs each, while its lighter model, demiculverin, was 9-10 pounder. Still lighter were saker, minion, falcon and falconet (See Vol 6, p F5-L).

In the book of Greener (Ref 7, pp 48-9) are given illustration of: "Early Culverin", "Hand Culverin with Bent Stock" and two "Culverins with Side Flash Pans".

1460–1480. Matchlock was developed sometime in the 15th century and Greener (Ref 7, pp 51ff) gives a brief description and illustrations taken from German MMS of 1460-80 which were located in the University Library at Erlangen. On p 52 are four figs showing the development of matchlock; on p 54 the mechanism of matchlock and on p 68, German Matchlock Gun. The guns equipped with matchlocks were known as muskets and the soldiers using them as "musketeers" (See Vol 4 of Encycl, p D753)

Note: Accdg to Newman (Ref 30, pp 36-7), arquebus was replaced by musket about 1600 and at about the same time crossbowmen and longbowmen were finished, while the number of pikemen was reduced to six for every four arquebusiers or musketeers. Some pikemen were necessary to protect musketeers from heavy-armed cavalry. The pistol was developed so that armored cavalry could ride into the thick of the pikemen and shoot them down. The cavalry pistol was at first half size arquebus or demi-bague and because they were held against the chest, they were called poitrinales.

In the book of Greener (Ref 7, pp 48-9) are given illustrations of the following early pistols: "Iron Club Pistol of 15th Century" and "Pistol Battle-Axe".

1480. Accdg to Nambo (Ref 66, p 39), compn of Swedish Gunpowder for common use was: saltpeter 57.0, sulfur 21.5 & charcoal 21.5%.

1492. Accdg to Dutton (Ref 58, p 57), the success of firearms at Constantinople created big demand for them and European producers of Gunpowder and cannons became very busy. This required money to pay for them, and fortunately for Europe newly discovered America started to supply gold.

1497. The first successful attempt to use anything but simple, solid, spherical projectiles (See year 1340) seems to have been made at the siege of Weissenburg. There, use was made of stone balls of slightly smaller diam than the bore of the cannon, but coated with an incendiary material and wrapped in a cloth soaked in the same material. The proj was placed thru the muzzle on top of Gunpowder and on firing the wrapping ignited and the flame did not extinguish until the target was reached. This was the first successful incendiary projectile (Ref 12, p 497).

15th to 20th Centuries. Hand Grenades probably existed as early as 15th century (See Vol 6 of Encyc, p G134-R, under GRENADERS). Accdg to Marshall 1 (Ref 11, p 32), the early grenades, probably made of earthenware, were used at the siege of Arles in 1536. Whitehorn, writing in 1560, recommended replacing earthenware grenades by hollow balls, cast from molten mixture of brass 3 parts and tin 1 part and loading them with serpentine powder 3 parts, fine corned pdr 3 parts and rosin 1 part. A small quantity of fine corned pdr was used for priming.
The term “grenades” was coined on account of their resemblance to pomegranate. By the middle of the 17th century, special military units were organized consisting of tall, strong soldiers called “Grenadiers”. Grenades thrown by them consisted of hollow, cast iron balls (spheres) ca 2½ inches in diam, contg chge of BkPdr and provided with a fuse. Before throwing the grenade, the fuse was ignited. The hand grenades practically disappeared beginning with the middle of the 18th century. However, their modified versions, revived during Russo-Japanese War (1904–1905) and during WWI. As the range of grenades thrown by hand was rather short (max ca 35 yards), attempts were made to increase it by using a sling or catapult, but by throwing hand grenades by means of rifles (or using specially designed rifle grenades), ranges up to several hundred yards were achieved. More detailed information on this subject is given in Vol 6 of Encycl, pp G134-R to G139-R. Description of “Grenade Launcher” is on pp G139-R & G140-L of Vol 6

15th Century, Late. Acdg to Dutton (Ref 58, p 8), Leonardo da Vinci (1452–1519) invented a steam cannon. Such a cannon was actually built and tested during American Civil War (1861–1865) but rejected as being too cumbersome. He also invented multibarreled machine-guns, three types of them are shown on plate VI, opposite p 51 of Newman (Ref 33)

16th Century, Early. Henry VIII improved the armament situation in England, and by 1515 there were assembled in Tower of London 400 cannons, most of them mounted on wheels. Further enlargement of supply came when Henry VIII seized in 1522 some firearms from Venetian galleys trading them to Flanders (Ref 7, p 30)

Acdg to Greener (Ref 7, pp 39–40), the war vessels of the early 16th century were furnished with small cannons which were fired from taffrail, and others which were fixed to the decks and fired thru portholes, as may be seen from illustration on p 39, showing the deck of Mary Rose (sunk by French in 1545) with big breech-loading cannons

1500 is the date of birth of Benvenuto Cellini (who died in 1571). It is given to indicate that at that time sporting firearms were already known. Cellini, when a very young man, was very fond of shooting. The firearms were of smooth-bore (Ref 11, p 28)

1515–1547 are the years of rule of French King François (Francis) I, born in 1494. He created permanent Army and equipped it with various cannons and small arms. Illustrations on p 32 of Greener (Ref 7) include: Cannon, Great Culverin, Bastard Culverin, Culverin and two Falcons. On p 33 is shown a cannon mounted on wheels

1517. Acdg to Marshall 1 (Ref 11, p 30), the doubtful honor of having invented infernal machine was ascribed to a Nuremberg citizen in 1517, but there is a drawing of one by Leonardo da Vinci who lived from 1452 to 1519. In 1645 attempts were made to blow up Swedish ships in Wismar harbor by means of clock-work bombs. In them the clock-work actuated a flint lock with a revolving steel wheel. Clock-work infernal machines contg a NG explosive were used also by the Irish-American Fenians in 1882 and 1884

Note: Daniel (Ref 3, pp 268–83), under the title “Engins Criminel”, described them as “devices intended for committing criminal attempts against persons or property”. The following attempts against important persons are described by Daniel: 1) Dec 24, 1800 an attempt in Paris against Bonaparte, while he was First Consul resulted in death of many persons, but Bonaparte was just shaken 2) July 28, 1835 an attempt in Paris against King Louis Philippe resulted in death of many persons but left the King unhurt 3) Jan 18, 1858 an attempt by Ital patriot Count Orsini against Napoleon III left Emperor unhurt but killed and wounded many of the escort 4) March 13, 1881 an attempt against Czar Alexander II of Russia resulted in his death. Two bombs were thrown, each of thick glass loaded with Dynamite and initiated by Ag azide. The 1st bomb wounded some of the soldiers escorting the Czar. Then he ordered the carriage to stop and went to see the wounded in order to comfort them. Just before he intended to remount the carriage, the 2nd bomb hit at his feet causing most serious and painful wounds which resulted in great suffering and death. He was the best Ruler in the more than 1000-year history of Russia

Stettbacher (Ref 21, pp 417–19) described under the title “Anarchistenbomben und Höl-
lenmaschinen”, construction of anarchist’s bombs and infernal machines and mentioned attempts against Napoleon III and Alexander II.

In his later work, Stettbacher (Ref 41, pp 130–32) described under the title “Sabotage-ziender”; chemical igniters used in anarchists’ bombs and other devices.

The same information is given in Spanish version of his book published in Argentina (Ref 46, pp 164–66), under the title “Bombas de Sabotaje”.

1820. Rifled Small Arms are said to have been invented by Augustus Kotter of Nuremberg, but for a long time they were used for sporting purposes, because the necessity of ramming the bullet down the barrel with the spiral grooving made the loading very slow. Moreover, the powder left much fouling in the grooves, and consequently it was necessary to clean the gun after a few rounds. With the old musket which was smooth-bore, the bullet was smaller than the bore and there was no difficulty in loading, but it was not as accurate as rifles. As for sporting purposes, accuracy was more important than rapidity of fire, therefore the rifle was able to hold its own. Sporting rifles were successfully used by American colonists during the War of Independence. Then the English created the “Rifle Brigade” to fight the American marksmen (Ref 11, pp 28–9).

Note: In the Crimean War (1854–1856), rifles used by the English and French (called Stutzers by Russians) against Russian smooth-bore arms, contributed greatly to defeat of Russians.

1525. Beginning with this year, bronze cannons were cast in England by Peter Bawde, French artificier. Later, in 1535, John O’Ewen was engaged in the work, and by 1543 the cannon industry was established in Uckfield, Sussex, then the center of iron industry in Britain (Ref 7, p 30).

1525. Important improvement in BkPdr was achieved in France by control of the grain size (Ref 64, p 2–2).

1533. In order to create a big Navy, Henry VIII of England needed money which he got by confiscating the properties of Roman Catholic Churches and Monasteries. This money permitted him to create a Navy, which at the time of Henry’s death (1547) amounted to 53 ships, all equipped with bronze cannons. With the aid of these ships, Spanish galleons were attacked and their gold seized. This created bad relations with Spain (Ref 58, p 61).

1543. Japanese Gunpowder “Tanegashima”, early period was, accdg to Nambo (Ref 66, p 38): saltpeter 75.0, sulfur 8.3 & charcoal 16.7%.

1548. Chinese Gunpowder of Ming Dynasty was accdg to Nambo (Ref 66, p 38): saltpeter 75.75, sulfur 10.6 & charcoal 13.65%.

16th Century, Middle. In addition to early breech-loading cannons in which the charge of powder was loaded into a separate breech (See illustrations on pp 21–22 of Ref 7) and wedged into the cannon, there were numerous methods employed for closing the breech after inserting the charge. One of these is shown in illustration on p 37 of Greener which is not reproduced here.

16th Century, Middle. Up to that time, no cartridges of any kind existed for propelling a charge of BkPdr and it was loaded loose thru the muzzle. Then the paper cartridges started to be invented and by about 1590, three types were known. They are shown in Fig 17 of Vol 4 of Encycl, p D754-R. The really successful paper cartridge was adopted by King Gustavus-Adolphus of Sweden (1594–1632) for his Army (See Fig 18 on p D755). None of the paper cartridges contained primers as we know them now (See under year 1805).

Paper cartridges were not replaced by metallic cartridges provided with caps until the beginning of the middle of the 19th century (See under year 1805).

The historical development of cartridges from paper to metallic is also described in Vol 2 of Encycl, p C73-R.

1554. Accdg to Marshall 1 (Ref 11, p 23), a Gunpowder mill was erected at Rotherlith, England.

1558–1603. During this period Elizabeth I (born in 1533) ruled in England. She succeeded Mary I, during whose rule the Henry VIII Navy was neglected and many of the ships deteriorated.

At that time Philip II of Spain (born in 1527 and King 1556–1598) was planning to attack England by means of newly constructed ships which formed the “Invincible Armada”. Elizabeth then ordered John Hawkins to restore the Navy and Francis Drake to construct a large Gunpowder Plant at Waltham Abbey (Ref S8,
Several other mills were installed at that period and the monopoly was conferred upon the Evelyn family (Ref 7, p 17). This took place, accdg to Marshall 1 (Ref 11, p 23), a few years later than 1561.

1560. Comps of two Swedish Gunpowders as listed by Nambo (Ref 66, p 39) were:

- a) saltpeter 50.0, sulfur 16.7 & charcoal 33.3%
- b) saltpeter 66.7, sulfur 16.7 & charcoal 16.7%

1560. Comps of English Gunpowder “Whitehorse” were, accdg to Marshall 1 (Ref 11, p 20): saltpeter 50.0, sulfur 16.6 & charcoal 33.3%

1569. Comps of two Japanese Gunpowders are given by Nambo (Ref 66, p 38): a) saltpeter 77.6, sulfur 10.7 & charcoal 11.7%
- b) saltpeter 76, sulfur 12 & charcoal 12%

1578. Accdg to Marshall 1 (Ref 11, p 27), the first instrument for testing Gunpowder was devised by Bourne. Up to that time, the only test was to burn a small quantity to see how much residue would be left. The instrument of Bourne consisted of a small metal cylinder with a heavy metal lid on a hinge. The lid was prevented from falling by a ratchet, and the angle to which it rose when powder was fired inside the cylinder measured the strength. A much better instrument was invented in 1627 (qv)

1587. English Navy, under Admiral Drake, attacked and destroyed the Spanish galleons assembled in Cadiz in preparation for the attack against England (Ref 58, pp 66–7)

1588. After repairing ships damaged at Cadiz and rounding up more of them, Philip II sent, early in 1588, 132 ships forming the “Invincible Armada” against English 40 warships and 150 merchant ships equipped with guns, under Lord High Admiral Lord Howard of Effingham. The Armada came late in July to Plymouth, where it was attacked by the English, who destroyed many ships by burning them. The remaining ships escaped across the channel to Calais, but they were pursued and attacked again. Finally, the Spanish Admiral decided to escape to Spain thru North Sea. This happened to be very rough and it contributed to destruction of many ships. By the time Armada arrived at Spain, there were less than 50% of ships, while English lost no ships (Ref 58, pp 67–8)

1588. Accdg to Colver (Ref 12, p 497), early hollow shells were used by Chinese. They were made of earthenware. They were filled with incendiary mixts and provided with a fuse which was supposed to ignite on firing of proplnts. These shells were dangerous to use because they often broke inside the bore. The first successful hollow shells and shrapnel are briefly described in next item

1588. Accdg to Marshall 1 (Ref 11, p 31), the first really successful explosive shell was employed at the sieges of Wachtendonck and Bergen-op-Zoom, Holland. This was due to the employment of newly invented igniter (fuse) by an Italian master gunner refugee from Parma. The fuse apparently consisted of a tube (or pipe), filled with slow-burning powder, which was driven into fuse-hole of the shell. This type of fuse was made to burn 14 to 20 seconds, corresponding to ranges 1000 to 2000 yards in the mortars, which were always used instead of cannons for throwing shells. Such shells were used for the destruction of stone fortifications and ships; against men they were not very effective, as there was usually time to get away from them or to extinguish the flame of fuse by water or sand. Until after the introduction of watches, which were invented in 1674 by Christian Huygens of Holland (1629–1695), no convenient means existed for testing the time of burning of a fuse. In the middle of the 18th century fuses were made of beechwood with a hole down the middle filled with fuse powder. They were of different lengths. Great accuracy was not required of them until Capt Mercier, during the siege of Gibraltar in 1779, proposed to fire shells from cannons instead of mortars or howitzers. Accurate fuses were also required for the Shrapnel, invented by Lt Strapnel ca 1784 and officially adopted by British Govt in 1803. These fuses were replaced after 1831 by Bickford fuse, described in Vol 2 of Encyl, p B112-L (Ref 11, pp 32 & 38)

1595. Accdg to Marshall 1 (Ref 11, p 31) and Nambo (Ref 66, p 39), compn of German Gunpowder was: saltpeter 52.2, sulfur 21.7 & charcoal 26.1%

1597. Chinese Gunpowder of Ming Dynasty contd: saltpeter 80.66, sulfur 5.64 & charcoal 13.70% (Ref 66, p 38)

16th Century, End. Accdg to Chalon (Ref 8, p 228), quoted by Marshall 1 (Ref 11, p 27), the French compn of “Poudre noire” of proportion 6:1:1 (meaning saltpeter 75.0, sulfur
12.5 & charcoal 12.5%) has been adhered to there more or less ever since. It is also listed by Nambo (Ref 66, p 39) as of year 1598

Note: It is mentioned by Marshall 1 (Ref 11, p 27), that German powder contg saltseeter 66.5, sulfur 16.5 & charcoal 16.5% was more satisfactory than French powder and that the German Cocoa Powder was ballistically the best BkPdr ever made

1606. Accdg to Nambo (Ref 66, p 23), the Chinese Gunpowder for general purpose, described by Wu Pei Chih of Ming Dynasty contd: saltseeter 75.0, sulfur 12.5 & charcoal 12.5%

1608. Accdg to Marshall 1 (Ref 11, p 27), Danish BkPdr contd: saltseeter 68.3, sulfur 23.2 & charcoal 8.5%

1611–1632. During the rule of Swedish King Gustavus Adolphus (born in 1594) many improvements in military weaponry were introduced. He is considered by Dupuy & Dupuy (Ref 69, pp 522–24) as the father of modern field artillery and of the concept of massed, mobile artillery fire. His 3-pounder regimental cannon, illustrated on p 524, looks like a modern weapon. He was also the inventor of an improved paper cartridge which is illustrated in Vol 4 of Encycl, p D75S. This invention permitted an increase in the rate of fire. For further development of cartridges, see year 1826

Gustavus Adolphus was the initiator and successful participant of "The Thirty Years' War" (1618–1648), which started as a religious war between Protestants and Catholics. The War, which involved several nations, is described in Ref 69, pp 533–46. The King personally was engaged in many battles and was killed in 1632 at the battle of Lützen. Then Bernard took command and the war continued until 1648. As result of peace treaty of Osnabrück, Sweden received an indemnity, and substantial Baltic coastal territories belonging to Russia, and to Catholic part of Germany

1615. Accdg to Dupuy & Dupuy (Ref 69, p 523), the true fusil or flintlock musket was invented by French gunsmith LeBourgeois. It was perfected as a sporting weapon about 1630, but was not adopted for military purposes until 1670 in France and then later in other European Armies. By ca 1680, someone – possibly S. Vauban (1633–1707) – invented the ring bayonet, which left the bore clear for firing

1621. Accdg to Nambo (Ref 66, p 38), the Chinese Gunpowder for propulsion described by Wu Pei Chih of Ming Dynasty, contained: saltseeter 75.0, sulfur 9.5 & charcoal 13.5%

1627. According to Marshall 1 (Ref 11, p 27), a much better instrument for testing BkPdr than that of 1578 (qv) was devised by Curtenbach. This consisted of a heavy conical shot which rested on the mouth of small mortar and could travel vertically upwards along a stretched wire provided with series of catches to stop the shot from falling down

1627 and Later. Black Powder in Mining. Accdg to Dutton (Ref 58, p 90), a Hungarian engineer, Kaspar Weindi, experimented in blasting ore near the town of Selmezbenya by means of BkPdr. From that time, use of BkPdr for breaking ores spread to Germany, Sweden and other countries. Accdg to Marshall 1 (Ref 11, p 33), blasting was probably introduced to England in 1670, and in 1689 Th. Epsly Sr started to use BkPdr in Cornish mines

Accdg to O. Gutmann, as quoted from Marshall 1 (Ref 11, pp 33–4): “When bore-holes first came into use they were made with iron-mouthed borers, fairly large – nearly 3 inches in diameter, and then closed with a wooden plug termed the shooting plug. Henning Hutmann in 1683 employed a kind of drilling-machine. In 1685 clay tamping and in 1686 firing-tubes began to be used. In 1689 paper cartridge tubes were used to replace the older form of leather, and in 1717 bore-holes of smaller diameter came into vogue. The use of the chiseiborer dates from 1740, blasting the untouched breast from 1767 (first at Zimmwald)”

1635. Accdg to Marshall 1 (Ref 11, p 26) and Nambo (Ref 66, p 39), English Gunpowder of Government Contract contd: saltseeter 75.0, sulfur 12.5 & charcoal 12.5%

1637. Accdg to Nambo (Ref 66, p 38), the Chinese BkPdr for bursting charges, described by Tien Hung of Ming Dynasty contd: saltseeter 82, sulfur 9 & charcoal 9%

1643–1715. During the rule of Louis XIV, the Great of France, an Army of nearly 400 thousand men, serving as volunteers, was organized and its frontiers were fortified by famous military engineer Vauban (1633–1707). Louis XIV equipped his infantry with flint
muskets, which replaced arquebus and matchlock. His muskets were equipped with bayonets and paper cartridges, which by 1700 allowed the firing of two rounds per minute instead of previously one round. Some infantrymen were put on horses and the resulting cavalry became known as dragoons. At Essonne, near Paris, a Gunpowder plant was constructed which manufd an improved product in seven sizes from finely ground grains for handguns to walnut size pellets for the large cannons. The use of large-grain pdr increased the range 1 or 2 miles. Towards the end of his rule, corruption, abuse & inaccurate payment developed and the number of available volunteers stopped. These corruptions and huge debts were passed in 1715 to Louis XV (1710–1774) and finally to Louis XVI (born 1754, deposed in 1792 due to revolution and guillotined in 1793). The first universal conscription was introduced in August 1793 (Ref 58, pp 72–88)

1646. Bofors Industries (Aktiebolaget Bofors) of Sweden was established to become one of the world’s largest companies manuf exps, proplnts, ammo and weapons. A brief description of its activities is given in Vol 2 of Encycl, pp B218 to B220

1647. Accdg to Nambo (Ref 66, p 39), the compn of Swedish Gunpowder of Nye was: saltpeter 66.6, sulfur 16.7 & charcoal 16.7%

1647. Master-Gunner Nye, in his “Art of Gunnery”, described the instrument of 1627 (qv), and also proposed that the strength of powder be measured by firing bullets from a pistol into clay, or by firing a ball from a mortar and finding out how far it travelled. This mortar test was adopted by French as “mortier éprouvette” and was also used by other countries (Ref 11, p 28). Further improvement in testing was done in 1742 (qv) 17th Century, Middle. Accdg to Marshall 1 (Ref 11, p 34), it seems that K chlorate was known to Glauber (1603–1668), but it was not investigated until about 1786 (qv), when Count C.L. Berthollet lived (See Vol 2 of Encycl, p B106-R)

1654. Ammonium Nitrate (AN) was first prepd by J.R. Glauber (1603–1668) who also prepd Na sulfate. It was not until beginning of the 19th century that it started to be used as a replacement for K nitrate in BkPdr. In 1840 Reise & Millon reported that a mixt of powdered AN and charcoal exploded on being heated to 170°. In 1867 Ohlsson & Norrbom patented its use in mining expls, and in 1870 A. Nobel purchased their patent (Vol 1 of Encycl, p A342-L). Nobel then introduced a series of AN expls, called Extra Dynamites and Ammonium Nitrate Galatins. During WWI, AN found extensive application in its mixt with TNT (such as Amatol and Ammonal) or with other HE’s such as Di- and Tri-nitronaphthalene (Vol 1 of Encycl, p A341 and Tables on pp A355 & A356 (See also Vol 5, under DY-NAMITE, p D1583 and as “Do-it-Yourself” Ammonium Nitrate–Fuel Oil Explosives in Vol 5, pp D1528 & D1529

The latest development in AN-contg expls is its mixtures with fuel oils. Such expls are known as AN-FO and their brief description is given in Vol 6 of Encycl, pp F211-R to F213, under Fuel Oils

1666. Chain Shot was invented by DeWitt of Holland, for use against the rigging of ships. It consisted of two balls or half balls united by a chain (Ref 11, p 31) (See under Grape Shot in Vol 6 of Encycl, p G126-L & R)

1669. Accdg to Dutton (Ref 58, p 114), phosphorus was discovered by a German chemist named Brand, who prepd it from urine

1689. Accdg to Feldhaus, SS 2, 218 (1908), as quoted in Marshall 1 (Ref 11, p 22), Thomas Epsly Sr started the use of Gunpowder in Cornish mines, England

1689–1725. Accdg to Dupuy & Dupuy (Ref 69, p 645), the Czar Peter I (The Great, born in 1672) completely reorganized Russian Army and Navy, equipping them with latest arms. This permitted him to win the “Great Northern War” (1700–1721) against Sweden, lead by the greatest warrior of that time, King Charles XII, born in 1682, ruling during 1697–1718 (Ref 69, pp 614–617). By the treaty of Nystadt (1721), Russia recaptured the Baltic provinces ceded to Sweden during “The Thirty Years’ War” (1618–1648), lead until his death in 1632 by the great Swedish King Gustavus Adolphus (born 1594, ruling 1611–1632) (Ref 69, pp 533–46)

1696. The Swiss used drills of tempered iron for drilling boreholes in rocks during blasting with BkPdr thru Alps in constructing the Albula Road, the main thoroughfare for wagon travel (Ref 58, p 93)
18th Century, Beginning. Accdg to Marshall 1 (Ref 11, p 36), flintlock was invented, but accdg to other sources it was in the middle of 17th century. Its other name was firelock and it replaced the wheellock. A brief description is given in Vol 4 of Encycl, pp D754–55.

The improved types of flintlocks were used by the British as late as 1850 and by the Americans as late as the end of Civil War (1865)

18th Century. Cannons, which since the 16th century were cast in bronze, started to be cast in iron. Until the 2nd half of the 19th century, a cannon consisted simply of a block of cast metal with a smooth-bore machined out and a vent drilled near the breech. It is true that the breech-loading guns were made at a much earlier date (See illustrations on pp 21, 22 & 37 of Greener's book of 1910), but they were not adopted because the breech leaked (Ref 11, p 29)

1707. Chronographs. The earliest device was conceived by Cassini, but the first practical instrument based on his idea was constructed in England in 1740 by Robins. This instrument underwent in later years many improvements, such as by Hutton (1775), Didion-Morin-Plombert (1836), Piombert (1860) and Minarelli-Fitzgerald (1900–01). Other older devices are listed in Vol 3 of Encycl, p C305, while more recent devices, such as of LeBoulangé, Aberdeen, Mettegang, Pin Chronograph, etc are described on pp C306 to C318

1738. Closed Bomb or Vessel. The first recorded attempt to measure the pressure generated when gunpowder was fired in a closed space (such as bomb or a gun barrel) was made in 1738 by d'Antony in Italy. This was followed in 1742 by Robins in England, d'Arcy in France (1760) and by Count Rumford in Germany (1792). More accurate measurements could be obtd in the 19th century, such as using instruments invented ca 1860 independently by Maj T.J. Rodman in US and by Gen N. Mayevskii in Russia. Other devices were of Sir A. Noble & Sir F. Abel of England and of M. Deprez, E. Sarrau & P. Vieille of France. Bichel Bomb was invented in 1898 in Germany. In Russia the Dolgov Bomb has been used. It consists of a vertical cylinder. All of these bombs are described in Vol 3 of Encycl, pp C330 to C345

1742. Accdg to Marshall 1 (Ref 11, p 28), Robins placed the mortar described by Nye in 1647 (qv) on a more scientific basis by the invention of the ballistic pendulum by means of which the actual velocity of a projectile could be measured

1745. Electric Cap was invented by Dr Watson of England and improved in 1749 by B. Franklin of US, but the device was forgotten until 1830 when Moses Shaw was granted a patent for a similar device. It was a “high tension” cap. The first “low tension” cap was developed by M. Shaw with the assistance of Dr R. Hare (Vol 1 of Encycl, p B186-R)

The biggest contribution to the construction of modern cap was done by L.A. Burrows of duPont Co beginning about 1937 (Vol 1 of Encycl, p B187 & p B363-L, giving a list of his inventions)

1765. Accdg to Dupuy & Dupuy (Ref 69, p 665), J.B. deGriveauval (1715–1789) began to revolutionize French Artillery, and his 4-, 8- & 12-pound guns and 6-inch howitzers were later used by Napoleon

1766. Henry Cavendish (1731–1810), the British physicist and chemist, was the first to identify hydrogen (Ref 58, p 113)

1771. Picric Acid (PA) was first prep'd by Woolfe on treating silk with nitric acid. It was used as a yellow dye, and its expl props were not discovered until 1871 (See also 1843) (Ref 11, p 49 & Ref 12, p 6)

1772. Daniel Rutherford (1749–1819), Scotch botanist, first identified nitrogen (Ref 58, p 113). Do not confuse with Lord Ernest Rutherford (1871–1937), Brit physicist noted for his research on radioactive transformation and disintegration of nitrogen

1774. Swedish alchemist K.W. Scheele (1742–1786) separated from saltpeter its “fire-air”, now known as oxygen. He also obtd glycerin and identified chlorine and fluorine (Ref 58, p 112)

1780. Potassium Chlorate was discovered by C.L. Bertollet, and he prep'd some explmixts contg it (See Vol 2 of Encycl, p B107-L). Two disastrous expls in 1788 & 1792 during pulverizing and mixing the expsmixts contg it stopped its use for a while (See Vol 2 of Encycl, pp C190 to C197 & C202, and also under 1850. Chlorate Explosives
1782. Ballons & Airships and Their Application in War. Under this title there is described in Vol 2 of Encycl, pp B10–B11 their historical development. The first successful balloon was invented in 1782 by brothers Mongolfier of France, and ever since then attempts have been made to use them for military purposes, such as for observation and later for dropping bombs (See under year 1849, Bombs, Aerial).

The most ingenious application of balloons guided only by prevailing winds was made by the Japanese against US territory in 1944. It is briefly described in Vol 2, p B11-L and more fully by R.W. McKay in Engineering Journal 28, 263–67 (1945), under the title “Japanese Paper Balloons”.

1784. Accdg to Guittmann, as quoted from Marshall 1 (Ref 11, p 26), presses were introduced to compress the mill-cake BkPdr before coming. It was in order to obtain really compact grains of BkPdr.

1784–1785. Shrapnel shell was invented by Brit Lt Shrapnel to use against troops in the open (Ref 11, p 31). It partly replaced case-shot, described in Vol 2 of Encycl, pp C24-R & C25-L. A brief description of Shrapnel shell is given in Colver (Ref 12, p 497). It was adopted in 1803 by British Govt for military purposes.

1786. Potassium Chlorate, supposed to be discovered by Glauber in the 17th century (middle) was prepd by C.L. Bertollet in pure state and its props detd. He proposed using it in lieu of K nitrate in BkPdr, but abandoned the idea after disastrous expln during its manuf in 1788, which killed several persons. The first successful mixts using K chlorate were Cheddites (See Vol 2 of Encycl, p C155ff and Marshall 1 (Ref 11, pp 35–36).

1789. Accdg to Marshall 1 (Ref 11, p 26), at Faversham, England the powder was compressed by means of a screw-press, while the hydraulic presses were introduced in the 19th century.

1789. Uranium was obtd by German chemist M.H. Klaproth (1743–1817), who also obtd titanium and zirconium. Uranium was obtd by roasting radioactive pitchblend (Ref 58, p 114).

1792. Accdg to Marshall 1 (Ref 11, p 33), the Indians fired rockets in the defense of Seringapatam, but they did not appear to have done much damage, while at the siege of the same town in 1799, explosive rockets were used with some success.

18th Century, End. Accdg to Marshall 1 (Ref 11, p 28), practically every country had come to use Gunpowder of compn: saltpeter 75.0, sulfur 10.0 & charcoal 15%.

19th Century, Beginning. Accdg to Colver (Ref 12, p 498), considerable use was made of red-hot iron shot particularly in combat with wooden ships.

19th Century, Beginning. Ammines or Ammoniats were invented, but they were not systemized and explained until 1893 when Alfred Werner introduced his theory of coordinated compounds. Many of them are explosive, as can be seen from Tables in Vol 1 of Encycl, pp A277 to A282.

1800. Accdg to Marshall 1 (Ref 11, p 37), Mercury Fulminate was invented by Edward Howard, who described it in a paper before the Royal Society.

1802. Eleuthère Irénée du Pont de Nemours, French immigrant, began the manuf of BkPdr at a mill situated on the banks of Brandywine Creek, near Wilmington, Delaware. The compn was approx: saltpeter 75, sulfur 10 & charcoal 15%. The manuf was continued until 1921 (Ref 52, p 3).

1805. Accdg to Marshall 1 (Ref 11, pp 36–37), the Rev A.J. Forsyth, a Scotch clergyman, made a sporting gun with detonator lock. He patented the device (which is described in Vol 4 of Encycl, p D755). As the device was initiated by the blow of a hammer, it was the original percussion lock. The device was not adopted by the British Ordnance because it was not considered suitable for military purposes. Before many years elapsed, Forsyth's device was displaced by a copper tube or cap contg Mercuric Fulminate. It is not certain who invented the MF cap. Accdg to H. Wilkinson, J. Shaw of Philadelphia invented a steel cap in 1814, a pewter (alloy of tin with some lead) cap in 1815, and copper cap in 1816. J. Egg of London seems to have adopted the idea from Shaw. Prefet and Debouhert of Paris patented in 1820 caps filled with MF and AgF, respectively. E.G. Wright of Hereford published in 1823 a paper on the MF cap, contg much information. Frederick Joyce was the first to make a real success of percussion cap ca 1824.
The next important step was to combine shot, powder and cap in one cartridge, which could be inserted in the breech of the arm. In 1836 Lefaucheux introduced his pin-fire breech-loading shotgun, the barrels of which were made to drop as in the modern weapon to allow the cartridges to be introduced. About 1853 the English and French gun-makers introduced the center-fire hammer gun, which fired cartridges having a cap in the middle of the base of the cartridges, but the first really successful center-fire gun was that made by Daw in 1861 (Ref 11, p 37).

More detailed description of capped cartridges is given in Vol 4 of Encyc, p D756-L. 1807 to 1815. Accdg to Marshall 1 (Ref 11, p 33), Col Congreve of the Hanoverian Army (later Sir W. Congreve), independently from Indians (See 1792AD) developed, sometime before 1807, a rocket, the most powerful device of its kind that had been used in warfare. It proved very effective at Copenhagen and Walcheren in 1807 and at the passage of the Adour in 1813, but it was at the battle of Leipzig in 1813 that it achieved its greatest renown. It also rendered good service at Waterloo in 1815 and during the War of 1812 between the US and Gt Britain.

Since the Napoleonic Wars, the improvements in ordnance have been so great that the rocket ceased to be used as a weapon. However, it continued to be used for signalling and illuminating purposes, until it was displaced to a great extent by star shell.

1819. Accdg to Marshall 1 (Ref 11, p 26), Col Congreve introduced his granulating machine for BkPdr which is described on p 82. It has been used in England for many years.

1823. Accdg to Culver (Ref 12, p 4), MF (Mercuric Fulminate) was proposed by Wright as an igniter of Gunpowder.

1825. Accdg to Dutton (Ref 58, pp 156–57), the Rev Dr Clayton began in England investigation of products on distillation of coal tar, which was obtd from bituminous coal. He isolated benzene, creosote, naphthalene and other chemicals. Most of his products were later nitrated, giving exps. The investigation was continued by A.W. vonHofmann (1818–1892), who came to England from Germany and by W.H. Perkin (1838–1907). Many dyes were produced, such as mauve and alizarin. Aniline was also prepd and this gave on nitrating the very powerful Tetranitroaniline which is described in Vol 1 of Encyc, pp A425ff.

Other important exps on nitration of products of distillation of coal tar were Di- and Trinitrobenzene (Vol 2 of Encyc, pp B46ff and Nitronaphthalenes, which will be described under Naphthalene and Derivatives.

Dr vonHofmann moved back to Germany in 1874 where he established a large scale coal tar industry.

1826. Cartridges for Ammunition. It was mentioned under 1350. Bullets and-under 1611–1632 that the first successful paper cartridge was invented during the rule of King Gustavus Adolphus of Sweden (1611–1632). This type of cartridge was not, however, self-sustained because it was not provided with a primer which was not known at that time. The first self-contained paper cartridge was invented in 1826 and improved on in 1831. The first metallic cartridge was invented in 1846 by Houiller of Paris. It is known now as "pin-fire cartridge". He also invented the "rim-fire cartridge". The first successful "center-fire cartridge" was invented in 1858 by Morse and improved by Col Berdan. It was adopted in 1865 by the US Army. Previously to this (1854) Smith & Wesson developed a successful metallic cartridge for revolvers. Further development of cartridges and their construction are described in Vol 2 of Encyc, pp C73 to C76.

1826. Aniline was first prepd by Unverdorben (Vol 1 of Encyc, p A406-L)

1831. Safety Fuse invented by W. Bickford (See Vol 2 of Encyc, p B112-L) and in 1840 the Bickford fuse was adopted by the English military authorities. In 1836 a factory was started in US, in 1839 in France, and in 1844 in Germany. Before 1840 gutta-percha-covered fuse had been adopted for blasting under water. Various modifications have since been invented, including fuse cased in metal "Colliery Fuse" (patented in 1886 by Sir G. Smith), which emits no sparks; and various sorts of "instantaneous fuses", which burn very rapidly and enable many shots to be fired simultaneously (Ref 11, p 38)

1832. Ammonium Perchlorate was first prepd by Mitscherlich (Ref 70, pp 26 to 28)

1832–1833. French chemist H. Braconnot of
Nancy found that on treating starch (in cold) or wood (in hot) with coned nitric acid there was obtb a viscous liquid, which on addn of water deposited a white powder which, on drying, dissolved in ether-alcohol mixture and easily burned when ignited. It was named xyloidine but is actually Nitrostarch (Ref 44, p 240 & Vol 2 of Encycl, p C101-L) (See also Ref 31a, p 245) 1833. Nitrostarch (NS) was first prepd by Braconnet, who called it xyloidine. Pelouze prepd it, in 1838, in purer condition and studied its props. It was used in the US during WWI and WWII as demolition expl and for bursting charges (Ref 70, pp 246 to 248) 1834. The earliest attempt to nitrate aromatic nitrocompounds was by E. Mitscherlich (1794–1863), in Germany, who nitrated benzene to m-DNBz and toluene to MNT’s (Ref 12, p 18) 1837. Accdg to Dutton (Ref 58, p 125), Emanuel Nobel, father of Alfred Nobel, invented, at 35 years of age, a torpedo. It exploded and wrecked his home. He was ordered to leave Stockholm. So he went to Russia to operate a torpedo factory near St Petersburg. He settled in Russia and his sons, including Alfred, were educated there. When Crimean War broke (1853) his torpedoes (or rather mines) loaded with NG-NC prevented the British fleet from approaching Cronstadt, which was the fortress guarding St Petersburg. After the war, Emanuel, with Alfred and Oscar, returned to Sweden, while Robert & Ludwig remained in Russia to work in Baku oil fields. They became millionaires 1837–1838. Prof T.J. Pelouze of Paris (1807–1867) nitrated (other than starch) cellulose substances, such as paper, and called the resulting product xyloidine, which was sol in eth-alc. He also treated cotton fibers with coned nitric acid, without allowing them to be destroyed. He plunged the nitrated fibers in large amt of cold water, followed by several washings and then drying. The product was highly combustible but insol in eth-alc. This was not xyloidine but a higher nitrated product as was proved by Pelouze in 1846. He named it pyroxylone or pyroxyle (Ref 31a, pp 245–46; Ref 44, p 240 & Vol 2 of Encycl, p C101-L) 1840. Accdg to Marshall 1 (Ref 11, pp 37–38), the Prussians adopted a breech-loading rifle, the Zündagelgewehr, invented in 1838 by Dreyse. In this ignition was effected by a needle being driven right thru the base of the cartridge into a disk of fulminating material. In spite of some defects, such as escape of gas and rusting of needle, the gain in rapidity of fire caused it to be maintained in general use during the wars of 1848, 1866 and 1870. The French adopted Chassepot in 1866, which was a considerable improvement upon the Prussian rifle. At about the same time, the English rifle of Mr J. Snider, an American, was invented. This rifle was considerably improved by Col Boxer by adopting a brass case instead of paper as in previous rifles (Ref 11, p 38) 1841. Ammonium Picrate was first prepd by Marchand and used in 1869 by Brugère as proplnt (Ref 70, p 135) (See also 1909) 1843. Picric Acid (PA), discovered in 1771 by Peter Woulfe (inventor of 2- or 3-necked bottle) but not identified, was rediscovered by A. Laurent (1807–1853). He prepd it by nitrating phenol or Dinitrophenol with nitric acid and identifying it as Trinitrophenol. He did not recognize it as an expl, but found that some of its salts are expl. When the price of phenol became reasonable, PA started to be used as a fuel in expl mixts, while its salts were used as expl additives (See also under 1871 and 1885) (Ref 11, p 49) 1844. Prof C.F. Schönbein of Univ of Basel (1799–1866), discovered ozone (Ref 11, p 39) 1845. Schönbein on treating cotton with mixed nitric-sulfuric acid during a short time (in order not to destroy fibrous structure) obtb, after washing with large amt of w and drying, a very combustible product which was named “coton-poudre”. The product was tried in mining (Ref 31a, pp 241–48; Ref 44, p 240 and Vol 2 of Encycl, p C101-L, under CELLULOSE NITRATE) Note 1: Dutton (Ref 58, pp 119–20) said that Schönbein discovered NC by accident. He wiped nitric-sulfuric acid spilled in the kitchen with his wife’s cotton apron. Then he rinsed the apron with water, hung it to dry over a hot stove and then he heard a loud expln. He repeated the nitration using another apron, which he rinsed and dried more carefully than the first one. Then he cut from the apron a strip and lighted it with a match. The strip rapidly burned without leaving any ash
**Note 2:** Acedg to Davis (Ref 31a, p 247), Schönbein also described the nitration of *cane sugar*, which was performed in December 1845, but he deliberately refrained from telling how he prep'd NC. His *Nitrosugar*, prep'd as described by Davis (Ref 31a, pp 247–48), was dried at low temp. At 100° it was semi-liquid and at higher temps gave off red fumes. When heated more strongly, it deglazed suddenly and with violence.

1846. Otto of Brunswick described in Allgemeine Zeitung of Oct 5, 1846 procedure for making NC very similar to coton-poudre of Schönbein (Ref 44, p 240)

1846. W. Knopp of Leipzig, Heerben & Karmarsch of Hanover and J. Taylor of England described prep'n of coton-poudre by nitrating cellulose with mixed nitric-sulfuric acids (Ref 31a, p 250 & Ref 44, p 240)

1846. Italian chemist, A. Sobrero, obtd on treating gelatin with mixed nitric-sulfuric a very expl liquid which he called *glycérine fulminante* or *pyroglycérine* (Ref 44, p 241)

1846. Sauerbrey of Switzerland started to manuf rifles using *fulmicotton* and exported them to various countries, chiefly to Austria, but this stopped in 1848–1850 (Ref 44, p 241)

1846. Schönbein proposed using coton-poudre as more powerful replacement of *poudre-noire* (BkPdr) (Ref 44, p 241)

1846. Dr Bley of Bernberg prep'd NC by nitrating sawdust and suggested its use in lieu of BkPdr (Ref 31a, p 251)

1846. John Hall & Sons of Faversham, England started to manuf coton-poudre by the method of Schönbein, but this was abandoned after a terrible expln on July 14, 1847 which destroyed the plant (Ref 44, p 241)

1846–1847. R. Böttger of Frankfurt on Main (1806–1887) prep'd "coton-poudre" independently of Schönbein. They joined in obtaining a patent, including one for US, under the title: "Improvement in Preparation of Cottonwool and Other Substances as Substitutes for Gunpowder" (Ref 31a, p 249; Ref 44, p 240 & Vol 2, p C101-L)

1846–1850. Acedg to Colver (Ref 12, p 498), the first real advances in fusing of shells were made at that time. Early shell fuses have been made with the intention of their being ignited by the flame of the propitg chge. Then Fre-born designed, by an adaptation of the per-

cussion cap, a time-fuse (or rather *time-fuze*) which was initiated by the shock of discharge of the projectile from the gun. On the latter date Moorsom introduced the first shell fitted with a fuze actuated by the shock of impact of the shell

1847. Mannitox Hexanitrate or Hexanitromannitox was first prep'd in Italy by A. Sobrero. N. Sokolov of Russia examined in 1878 its expl props. Used as secondary chge in detonators and blasting caps (Ref 70, pp 197 to 200)

1847. A big expln killing 21 men took place at Faversham (England) factory manuf Guncotton by the Schönbein method. After this Hall & Sons, who were the contractors, refused to proceed with the manuf. About the same time there were disastrous Guncotton explns at Vincennes and Bouchet, and these produced such an effect that no more Guncotton was manufd in England & France for the next 16 years (Ref 11, p 39)

1847. In France, the “Commission du Pyroxyle” was formed, charged with investigating the possibility of replacing BkPdr with NC (plus some dextrin) as rifle propmt, but the idea was abandoned after violent explns at Vincennes and Bouchet (Ref 44, p 241)

1847. Gladstone of England, by exercising special precautions, was able to determine nitrogen content of pyroxyline as 12.75%, which corresponds to pentanitrate, while the xyloïdine corresponded more nearly to trinitrate (Ref 44, p 252)

1847. Crum of England nitrated cotton (previously bleached by special process) until he could introduce no further nitrogen into the molecule. After washing and drying the product it was analyzed by nitrometer method which gave 13.69% N. Nitration was done by mixed nitric-sulfuric acid as described in Davis (Ref 31a, p 252)

1847. Cyanuric Triazide, first prep'd by Cahors, and in 1887 the process was improved by James. Taylor & Rinkenbach prep'd it in 1923 & 1927 in pure state and detd its expl props (Ref 70, pp 66–68). See also Vol 3 of Encycl, pp C590-R & C591-L

1849. Bomb, Aerial. After the invention of balloon (aerospace) by Brothers Mongolfier in 1782, several attempts were made to use balloons for throwing bombs from the air. One of the
first such attempts was made during the siege of Venice by Austrians in 1849. A brief description of this operation is given in Vol 1 of Encycl, p B225-R. Not much was done after this in the 19th century because in 1899 aerial bombing was forbidden by the Hague Convention. In 1903, the Wright Brothers constructed a successful aeroplane, and in 1907 the restriction on aerial bombardment was lifted. By 1910 nearly all nations started experimenting with planes dropping such bombs. The earliest recorded use of bombs dropped from a plane was by the Italians ca 1911 during the war in Tripoli. More successful was the use of improved bombs by the Spanish during the war in Morocco. During the first 3 months of WWI no bombs were dropped because the planes and dirigibles were not equipped for this purpose, but on Aug 30, 1914 a single German plane dropped bombs on Paris and one of the bombs penetrated into a crowded subway station, killing and mortally wounding 1000 persons. Then Germans also used Zeppelins in night raids on London. Considerable aerial bombing was done during the Spanish Civil War (1936–1939), but the greatest use was during WWII. For further development of aerial bombing, see Vol 2 of Encycl, pp B226–B232. German bombs of WWII are described and illustrated in PATR 2510 (1958), pp Ger 14 to Ger 20 (See also under the year 1782).

1850. Chlorate Explosives. After disastrous exps of 1783 & 1792 with mixts using K chlorate (discovered in 1780 by C.L. Berthollet), there were no applications of chlorate until about 1850. One of the earliest chlorate expts was Augendre Powder listed in Vol 1 of Encycl, p A507-L. The mixt was, however, too sensitive for use. The problem of desensitization was solved in 1871 by H. Sprengel. Still better exps were developed in 1881 & 1888 by E. Turpin and in 1897 by Street. A fairly complete list of chlorate explosives is given in Vol 2 of Encycl, pp C202 to C209.

1850. Accdg to Nambo (Ref 66, p 39), the French Gunpowder of the Napoleon III Army contd: saltpeter 75.6, sulfur 10.8 & charcoal 13.6%.

1850. Cameras, High-Speed Photographic. The first attempt to photograph a rapid event was made by H.F. Talbot of England, and his technique was applied in 1884–85 by E. Mach & P. Salcher to ballistic studies. In 1909 C. Cranz designed an apparatus called “Ballistische Kinematograph” capable of taking 50000 frames per second and an improved app capable of taking 100000 fps was designed in 1912 by C. Cranz & B. Glarzel. A description of Cameras is given in Vol 2 of Encycl, pp C13 to C19 and in this Vol under High Speed Photography, pp H104f.

1850. Electric Friction Machine suitable for field use was invented by Baron von Eben of Austria. After being improved in 1869, it was used in the blasting of Hoosac tunnel (Vol 2 of Encycl, pp B187-L & B212-L).

1852. The Committee appointed by Confederation Germanique, to which Schönheim & Böttger proposed in 1846 the cessation of their method, refused the proposition (Ref 44, p 241).

1853. Baron von Lenk, representing Austria, was happy that Prussia did not obtain the patent of Schönheim & Böttger dealt directly with them to obtain the method. He started to work on improvement of stability and possibility of adopting the improved powder as cannon propellant (Ref 44, p 242).

1853. Emmanuel Nobel (1801–1872), father of Alfred Nobel, installed sea mines of his invention in Baltic Sea to protect Russian fortress Cronstadt and City of StPetersburg against British Fleet during the expected Crimean War (See Vol 5 of Encycl, p D1585-L).

1853–1856. During Crimean War, K.I. Konstantinov constructed in Russia several pyrotechnic rockets (Ref 71, p 16).

1853–1865. Beginning with one artillery battery using NC as propellant, the number increased to thirty, although the powder caused erosion of the bore and occasional cracks in the walls of projectiles. While these problems preoccupied the artilleryists, a terrible expn took place in 1863 at Simmering, near Hitenberg (of the Gunpowder factory erected in 1853). A 2nd expn took place in 1865 at Simmering near Vienna, and this provoked the Imperial Edict of 11 Oct, 1865 prohibiting manuf of NC in Austria. This expn broke the negotiations with France and US, to whom von Lenk proposed his method (Ref 31a, p 253 & Ref 44, pp 241–42).

1854. Introduction of the earliest Dynamite.
called *Magnesiat'nyi* by Russian officer V.F. Petrushevscki. It consisted of NG absorbed by powdered magnesit or magnesia. It was used in Siberia for blasting in gold mines (Ref 71, p 10 and Vol 5 of Encycl, p D1586-L)

1854. Epichlorohydrin was prep'd by M. Berthelot (Ref 49, p 74)

1854. F. Pisani prep'd 2,4,6-Trinitroaniline, also known as Picramide (Vol 1 of Encycl, pp A409-R & A410-L)

1857. Lammot du Pont introduced a new formula, less expensive BkPdr by replacing the Indian saltpeter (K nitrate) with the much cheaper Chile saltpeter (Na nitrate) (Vol 5 of Encycl, p D1587-L)

1858. Peter Griess discovered Diazocompounds (Ref 12, p 5) and in 1860 he prep'd Diazodinitrophenol, also known as Dinol (Vol 2 of Encycl, p B59; Vol 5, p D1160-R and Ref 70, pp 99 to 102)

1858. Accdg to Marshall 1 (Ref 11, p 29), a Committee recommended the introduction of rifled ordnance into the British Navy, and from that time there has been rapid and continuous improvements in all sorts of guns. The introduction of the buffer has made the guns much steadier. Breech-loading guns were reintroduced, and the mechanism of breech greatly improved. To meet the requirements of the longer and more accurate cannons, the grains of Gunpowder were gradually increased in size so as to make them burn more slowly

1859. Ethylene glycol, Ethylene Oxide and Ethylene glycol Ether were prep'd by C.A. Wurtz (Ref 49, pp 7, 74 & 114)

1859. Diethyleneglycol prep'd simultaneously by A.V. Laurencio & Ch.A. Wurtz (Ref 49, p 158)

1859. Polyethylene glycol was prep'd by A.V. Laurencio (Ref 49, p 177)

1860. Baron von Lenk introduced bronze guns in Austria (Ref 11, p 40)

1860. Accdg to Gorst (Ref 71, p 7), rifled weapons started to be used in Russia

---

**Note:** British and French troops had such weapons already during Crimean War

**1860 and After. Cannon Propellants.** Up to about 1860's Black Powder was used exclusively as cannon and rifle propellant. Then attempts were made to replace it with smokeless propclnts made from incompletely cloilded Nitrocellulose. These propclnts were too fast burning (See descriptions under the year 1865). Some success was achieved ca 1882 by using Brown-, Chocolate-, or Cocoa Powder in larger caliber cannons. Their use was discontinued ca 1900, when fully cloilled NC smokeless propclnts replaced them (Vol 2 of Encycl, pp C29-C30) (See also year 1888)

1861. Guanidine (Gu) was first prep'd by Strecker (Ref 31a, p 374)


1862. John Tonkin of England proposed to pulp Guncotton using a beater similar to "hollandier" of paper industry. This produced, after thorough washing with water, more stable NC than the unpulped material (Ref 31a, p 253 & Ref 44, p 242)

1862. Tonkin proposed using pulped NC as an ingredient of the following compn: Na nitrate 65, charcoal 16, sulfur 16 & NC 3% (Ref 31a, p 254)

1862 & 1863. Baron von Lenk took out patents in England in the name of Révy to protect his method of purification of Guncotton. A factory was erected in 1863 at Stowmarket to make GC by his method, but it soon exploded (Ref 11, p 40)

1863. Frederick Abel, the chemist of English War Dept, started experiments and manuf on a small scale of GC in the Royal Gunpowder Factory at Waltham Abbey and succeeded in obtaining by 1865 more stable product than that of von Lenk (Ref 11, p 40) (See also below under 1865)

1863. Accdg to Colver (Ref 12, p 18), Wilbrand was the first to prepare TNT, but it was not as pure as that of Hepp, prep'd later (1880)

1863. Dioxane was prep'd by A.V. Laurencio (Ref 49, p 119)

1863. Triethylene glycol prep'd by A.V. Laurencio & Ch.A. Wurtz (Ref 49, p 170)

1863. The 1st Nitroglycerin Plant was built by A. Nobel at Helensborg, Sweden. It blew up in 1864, killing the youngest Nobel and a chemist (Ref 58, p 126 & Vol 5 of Encycl, p D1586-R)

1864. After expln at Helensborg, Nobel was forbidden by Swedish Govt to build another plant near any habitation. So he constructed a small plant on a barge moored in the center
of Millarsee (Ref 58, p 126 & Vol 5, p D1586-R)  

1864. Baron von Lenk took out a US patent for manuf of his NC (Ref 11, p 40)  

1865. Nobel obt’d permission of Ger Govt to construct a plant at Krakimel, near Hamburg (Ref 58, p 126 & Vol 5, p D1586-R)  

1865. After obt of permission from Swed Govt, Nobel constructed a new plant at Vintervicken and another plant in Norway, which at that time was united with Sweden and Denmark (Ref 58, p 126 & Vol 5, p D1586-R)  

1865. Jaworsky nitrated toluene with better results than did Mitscherlich in 1834 (Ref 12, p 18)  

1865. Sir Frederick Abel of England proved that pulping followed by thorough washing (such as done by Tonkin) improved the stability of NC because it helped to remove traces of acids (and of other impurities) found inside the fibers. He also showed that pressing of pulped material into sheets, discs, cylinders and other forms would cause NC to burn less violently in the gun. The compressed blocks of Guncotton prepd by him were an improvement over the yarn of von Lenk, but they still were too fast, not ballistically uniform and damaged the guns. The compressed blocks were, however, very suitable for blasting purposes (Ref 31a, p 254 & Ref 44, p 242)  

1865. Prussian Officer Schultz invented a sporting proplnt consisting of small pieces of nitrated wood previously treated by special process. The nitrated wood was washed with water contg small amt of Na carbonate, followed by concd soln of K & Ba carbonates and drying (Ref 31a, p 287 & Ref 44, p 242)  

Note 1: Accdg to Marshall (Ref 11, p 47), the Schultz powder was the first successful smokeless powder  

Note 2: A detailed description of “Sporting Smokeless Propellants” is given in Vol 2 of Encycl under “Bulk and Condensed Powders”, pp B322 & B323-L  

1865. Sir F. Abel patented a process for manuf of Guncotton, granulated by means of an aq soln of gum arabic. He also proposed to granulate mixts of Guncotton and Collodion Cotton using a small amt of methanol or eth-alc to gelatinize CC in the mixture (Ref 31a, p 254 & Ref 44, p 242)  

Note: Sir Abel also later patented (in collabo-ration with Dewar) a smokeless proplnt described in Vol 1 of Encycl, p A1-R. Another Abel’s smokeless proplnt is listed on p A2-L  

1866–1867. A. Nobel invented Guhrdynamit (Vol 5 of Encycl, p D1586)  

1866–1867. Abel’s “Researches on Guncotton” demonstrated that the material after proper purification is far more stable than thought to be. Moisture or exposure to sunlight did not harm it, and only exposure to elevated temp slowly decomp d it. Abel found that in NC produced at Waltham Abbey by nitration with 18 parts of mixed acid, there was ca 1.62% soluble in eth-alc. He detd its N content as 11.87%, while insol NC contd N=13.83%. He calc’d that the highest nitrated substance must contain 14.14% N, as was previously suggested by Crum (Ref 31a, pp 255–56)  

Note: Abel’s researches were described fully in JChem Soc 20, 310–57 & 505–76 (1867)  

1867. Borlinetto proposed a mixt of PA (Picric Acid) 10 parts, Na Nitrate 10 & K Chlorate 8.5 parts as substitute for Gunpowder (Ref 12, p 7)  

1867. Swed inventor J.H. Norrbin & C.J. Ohlsson proposed mixtures of pulverized AN with sawdust or charcoal as absorbent for NG (Vol 5 of Encycl, p D1587-R) (Also Ref 11, p 42)  

1867. Swed inventor Björkman patented the mixture called Seranin (Vol 5 of Encycl, p D1587-R)  

1867. Abel’s Stability Test using KI-starch paper is described in Vol 1 of Encycl, p A2-L. It was modifd in 1907 by Dupré, becoming known in England as “Heat Test” (Ref 44, p 250)  

1867. Abel of England and Trauzl of Austria proposed to replace kieselguhr with Guncotton as absorbent for NG. The resulting expl was exudable because GC was not gelatinized (Vol 5 of Encycl, p D1587-R)  

1867. Nobel obt’d BritP 1345 covering the device called fulminate blasting cap, which contd MF (Mercuric Fulminate). This cap was cramped to one end of safety fuse and then inserted in Dynamite cartridge (Vol 5 of Encycl, p D1588-R) (See also Ref 12, p 4)  

1868. A company was formed in England to manuf Schultz’s sporting smokeless powder, invented in 1865, and a factory was constructed in 1869 at Eyworth in the New Forest. The greatest success was achieved after the method
of prep was modified by Griffiths. The powder was very popular with sportsmen on account of the light recoil and absence of smoke as compared with BkPdr (Ref 11, p 48)

1868. Schultz of Prussia patented the expl Duanin, consisting of NG and partly nitrated wood (Nitrolignin), but this mixt was exudable (Vol 5 of Encycl, p D1587-R)

1868. Acq to Dutton (Ref 58, pp 135–37), the first Amer Dynamite plant was constructed in California at a spot called Rock House Canyon near San Francisco. It was manned by Chinese, but supervisors were American. The plant, consisting only of shacks, was called “Giant Powder Co”. Great opposition to Dynamite was encountered from a plant constructed at Santa Cruz, “The California Powder Works”, manufacturers of BkPdr. When, however, the Great Comstock Lode was discovered BkPdr was found to be too weak to blow the hard rock of Mount Davidson and only Dynamite could do it. The increased demand made the management of Giant Powder Co push their Chinese workers harder and harder to meet the demand, and as a result of this hurry the plant blew up in Nov 1869, killing two Americans and all Chinese. This accident did not discourage the Nobel Co, and another larger and safer plant was constructed on the sand dunes south of what is now known as Golden Gate Plant. A large Chinese crew was assembled and a German chemist was sent from Europe as Superintendent. The first 23 years of Giant Company exemplify in dramatic terms the introduction of Dynamite in the US. During that period, five successive plants, each larger than the previous ones, were wrecked by expls and in all 83, mostly Chinese men died. This did not matter because the Giant Co’s capital amounted to 2 million dollars

A detailed history of development and use of explosives in the US is given in the book of Van Gelder & Schlatter (Ref 14)

1868. E.A. Brown discovered that dry compressed Guncotton could be made to detonate very violently by initiation with a fulminate detonator such as Nobel already used for exploding NG. Shortly afterwards he made the further important discovery that wet Guncotton could be exploded by the initiation of a small quantity of dry Guncotton (the principle of the booster). This made it possible to use large blocks of wet Guncotton in Naval mines with comparative safety (Ref 31a, p 256; Ref 44, p 242 & Ref 12, p 5)

Note: The compressed Guncotton contg about 18% moisture proved to be so successful that it was adopted by several countries (especially by Russia) for loading not only of sea mines but also of Whitehead torpedoes and cannon shells. Russian Artillery used it during the Russo-Japanese War (1904–1905), but it proved to be inferior to Japanese shells loaded with Shimose (cast Picric Acid). Senior author of this Encycl, who served in 1917 in the Russian Navy, remembers that torpedoes on his destroyer of 1905 vintage were loaded with compressed, moist Guncotton

1869. Nobel was granted Engl patent for use as absorbent for NG mixts of combustibles (woodmeal, charcoal, rosin, starch) with oxidizers (K or Na nitrates) (Vol 5 of Encycl, p D1588-L). The resulting expls are now known as Dynamites With Active Base

1869. Désignolle, Brugère and Abel recommended using salts of PA, especially K, Amm or Na Picrates in mixts with K nitrate and charcoal as propnts and fillers for projectiles (Ref 12, p 6)

1869. Fontaine proposed a mixt of K Picrate and K Chlorate for loading torpedoes (Ref 12, p 6)

1869. Brugère proposed a mixt of Amm Picrate 54 & K Nitrate 46% for loading shells, while Abel recommended Amm Perc 40 & K Nitrate 60% (Ref 12, p 6)

1869. Beilstein & Kuhlberg investigated nitration of toluene with mixed nitro-sulfuric acid of various concns and isolated for the first time several isomers of mono- and dinitrotoluene. They also prep’d in 1870 TNT, but did not prepare Trinitrobenzene. At about the same time, mono-, di-, tri- and tetra-nitrophenylbenzenes were prep’d (Ref 12, p 18). Trinitrobenzenes were prep’d in 1882 by P. Hepp

1870. J. Howden of San Francisco, Calif proposed replacing Guhrodynamite by a Dynamite consisting of NG 75 absorbed by 25 parts of a mixt consisting of pulverized sugar, K nitrate and Mg carbonate. This was known as Howden’s Dynamite (Vol 5 of Encycl, p D1588-L).

1870. Nobel bought the patent of Norbin & Ohlsson which covered the use of AN as an “active base”. The patent also included the
1873. Nobel obtd Brit P 1570 for AN Dyna- 
mitite, in which particles of AN were waterproofed by 
stearine, ozocerite, etc (Vol 5 of Encycl, p D1588-L)

1873. Sprengel reported that PA in conjunction 
with suitable oxidizer forms a powerful 
expl (Ref 31a, p 166). However, he did not 
apply his discovery to practice as did Turpin in 
1885, who invented modern High Explosive 
Shell (Ref 12, p 1886)

1874. Accdg to info given in Encycl, Vol 2, 
p B46-R, Dinitrobenzenes became known

1875. After disastrous expln at the explosives 
plant of Messrs Ludlow at Birmingham, killing 
53 persons, the Brit Govt, on recommendation 
of Col Sir V.D. Majendie, passed the “Explosives 
Act”, which established the Inspectors of 
Explosives who were given power to inspect 
all magazines and factories in order to see that 
operations are carried out in a reasonably safe 
manner. As a result, the number of deaths in 
expls factories was very greatly reduced. The 
provisions of the English Explosives Act of 1875 
have been largely adopted by foreign countries, 
British Colonies and India (Ref 11, pp 46–47)

1875. Nobel obtd Brit P 4179 which covered 
Gelatins and Blasting Gelatin (Vol 5, p D1588-L)

1876. Nobel proposed using as antifreeze 
in Dynamites: methyl-, ethyl-, amyl-, or isoamyl- 
nitrates, but these substances were too volatile 
(Vol 5, p D1589-L)

1876. Accdg to Gorst (Ref 71, p 10), it was 
decided in Russia to load artillery projectiles 
with compressed Pyroxylin contg 18–20% 
water. As the original method of loading proved 
unsatisfactory, it was necessary to develop 
a new method which was not completed until 
the beginning of 1890

1877. Nitroguanidine (NGu) was first prepd by 
Jousselin, but only beginning in 1900 did it 
start to be used in prophts. During WWI NGu 
was used by the Germans as an ingredient of 
bursting charges (Ref 70, p 241 and Vol 6 of 
Encycl, p G158). It was used during WWI 
and later in triple-base propht Gudol (Vol 5, 
pp D1537 & D1538)

1877. A Commission was appointed in France 
to investigate the problem of ignition of fire-
damp. In the report made in 1880, it ad-
mitted that there was then no explosive known that would not ignite fire-damp (called “coal-
damp" by Marshall (Ref 11, p 45)  
1877. Wohlenberg & Sandström proposed o-MNT as an antifreeze, which proved to be better than those proposed by Nobel in 1876 (Vol 5, p D1589-L)  
1877-1878. Accdg to Dutton (Ref 58, p 153), the Turks used, during Russo-Turkish War, the propellant prep by treating NC with volatile solvent  
1878. In order to deal adequately with the many new inventions relating to exps, the French Govt created the "Commission des Substances Explosives" and Prof M. Berthelot (1827-1907) was appointed as its President (Ref 11, p 45 and Vol 2 of Encycl, p B104-R)  
1879. Tetryl or 2,4,6-Trinitrophenylmethylnitramine was first prepd by Michler & Meyer and studied in 1883 by van Romburg, in 1886 by Mertens. It was not used as an exp until WWI (Ref 31a, p 175 & Ref 70, p 339)  
1879. Hellhoff took out several Brit patents for Sprengel expls contg nitric acid either enclosed in glass tubes or absorbed in fossil flour or similar material (Ref 11, p 43)  
1879. Compression (or Crusher) Test of Hess is described in Vol 3 of Encycl, p C492  
1879. Nobel obtd patent covering Ammonium Nitrate Gelatins, known as Ammogelatins. This was the last patent granted to Nobel (Vol 5, p D1588-L)  
1880. TNT (Trinitrotoluene), also called Trotyl, was made in very pure state by P. Hepp (Ref 11, p 265) (See also under the years 1891 & 1902 and Ref 12, p 18)  
1880. The first du Pont NC & Dynamite plant, known as Repauno Works, was constructed at Gibbstown, New Jersey (Vol 5 of Encycl, p D1587-L)  

Note: Accdg to Dutton (Ref 58, pp 141-43), the Repauno Plant manuf Dynamite until 1954, when it was converted to chemicals. There were in 1960 other du Pont Dynamite plants, the largest located on the Potomac River near Martinsburg, W. Virginia. It was operated by remote control robots  
1880. S.R. Divine took out patents for Sprengel expls consisting of K chlorate & liquid combustibles. One of such expl, Rackarock, was used in 1885 for blasting of Hell Gate in New York Harbor (Ref 11, p 43)  
1880. Tetranitrocabazoles were first reported by C. Graebe, and they were studied at Picatinny Arsenal in 1951 by S. Livingston and in 1952 by D.B. Murphy et al. The results of their studies were confidential at that time (Vol 2 of Encycl, pp C48-C50)  
1880. Coal Mining Explosives, Testing for Permissability. It seems that the earliest tests were conducted in France, judging by the paper of E. Audibert, AnaMines 12, 278-98 (1937), but there seems to be no testing gallery until one was erected in 1907 at Liévin which was destroyed during WWI. The earliest gallery seems to have been erected in 1890 in England at Hebburn-upon-Tyne. This was followed by the gallery erected in 1891 in Austria at Mährisch-Ostrau. A brief description of galleries known to about 1965 is given in Vol 3 of Encycl, pp C370-C378  
1881. Prismatic Gunpowder was manuf by molding hexagonal prisms of pdr and pressing them in a special press (Ref 11, p 29)  
1881. Schultze's Smokeless Powder became very popular with sportsmen (See 1868), and it was decided to construct another factory. This factory at Hetzbach in Hesse-Darmstadt was completed in 1883 (Ref 11, p 48)  
1882. The next successful smokeless propnt was developed at the Explosives Co, England and became known as "EC Powder" (Ref 11, p 48). Its compn and props are described in Vol 5, pp E6-R to E8-R  
1882. Brown or Cocoa prismatic powder was made by the Germans, and in spite of attempts to keep the method of manuf secret, it was being made in England at Waltham Abbey two years later (Ref 11, pp 29-30) (See also Vol 2 of Encycl, p B173-L)  
1882. The size of cannons greatly increased and at the bombardment of Alexandria, the British had 16-inch cannons weighing 80 tons each. Such weapons required very dense and very large grain powder (Ref 11, p 30)  
1882. W.F. Read & D. Johnson patented in Germany a sporting propnt prep by agglomerating and hardening the NC grains by moistening their surface with eth-alc. This propnt was adopted in England by the Explosives Company at Stowmarket, England and manuf under the name of "EC Powder". It proved to be successful and was used for many years not only in shotguns but also as EC Blank Fire Powder.
1882. P. Hepp prep'd 1,2,4-Trinitrobenzene and 1,3,5-Trinitrobenzene, while 1,2,3-Trinitrobenzene was prep'd in 1914 by G. Köerner & A. Contardi (See Vol 2 of Encycl, pp B48 & B49).

1882. After introduction of rifled cannons by French and British during Crimean War (1853-1856), it was found by the Germans that ordinary BkPdr is too quick for larger caliber weapons. After prolonged research, the prismatic brown powder, C/82, made from partly carbonized charcoal, was introduced (See also Vol 5 of Encycl, p D1580-R, under Duttenhoefer). Then the Russians introduced in 1886 a brown powder with 19.6% of partly carbonized wood, 78.4% K nitrate and 2% sulfur (Vol 2 of Encycl, p B173-L).

Note: The above description was only partly reported by P. Tavernier in Ref 44, pp 243-44.

1883. Prep'n of Tetryl (or Tetranite) was first described by van Romburg in RecTravChim 2, p 108. Quoted from Marshall 1 (Ref 11, p 274).

1883-1884. M. von Duttenhoefer (1843-1903) nitrated brown charcoal (used for prep'n of C/82) by Schultze's method to obtain colloidal rifle powder RCP (Rottweiller Cellulose Pulver) which was adopted by the German Army in 1884, but its compn was kept secret until 1887 (See also Vol 5 of Encycl, p D1581, under "Duttenhoefer's Smokeless Propellant") (Ref 44, p 244).

1884. Ammonites are expts consisting of AN and nitrate derivatives of naphthalene. The original expt was known in England as "Miner's Safety Explosive". Some Ammonites cont'd TNT and were used for military purposes. A detailed description is given in Vol 1 of Encycl, pp A307-R to A310-L. Some Ammonites used by Germans during WWII as Substitute (Ersatz) Explosives cont'd RDX. They are listed in PATR 2510 (1958), p Ger 44 and on p E122 of Vol 5 of Encycl.

1884–1896. Nearly simultaneously and independently of Max von Duttenhoefer, Paul Vieille of France (1854–1934) invented in 1884 a completely colloided single-base NC propint. Originally called "poudre V" (V stands for Vieille). It was renamed "poudre B" in honor of General Boulanger, then Minister of War. It was adopted in 1886 for military rifles as BF (poudre B à fusil), in 1888 for field cannons as BC (poudre B à canon de campagne) and in 1896 for marine cannons as BM (poudre B marine). These propellants and their modifications were used as late as WWII (See also Vol 2 of Encycl, pp B1 & B2 and C32-L) (See also Ref 31a, pp 292-94; Ref 44, p 244 & Ref 11, p 49).

1884. 1,3-Diamino-1,3-trinitrobenzene first prep'd by Noëlting & Collin, was also prep'd in 1888 by Barr and in 1902 by Blinksma (Ref 70, pp 95-98 and Vol 5 of Encycl, p D1130-R).

1885. Prussian Govt established at Neunkirchen the first testing station for investigating mining explosives, which included a testing gallery. The method was soon adopted by other countries (Ref 11, p 45).

Note: Such galleries and method of testing are described in Vol 3 of Encycl, pp C368-R to C378-L. See also Naoûm (Ref 16, p 396).

1885. Accdg to Colver (Ref 12, pp 5, 7, 11 & 12), the discovery by French scientist E. Turpin (1848–1927) of detonating properties of PA (Picric Acid), was followed by its adoption for filling HE shell of his design (See next item).

Note: Accdg to Marshall (Ref 11, p 44), H. Sprengel was the first to draw attention to the fact that PA by itself could be initiated by a powerful detonator and was a very powerful expl, but no practical use was made of his discovery (See year 1873).

1886. In this year, the French Govt adopted the HE Shell designed by E. Turpin, using as a filler Picric Acid, designated as Mélinite. Its construction was essentially the same as modern High Explosive Shell. It is described on p 11 of Ref 12 and illustrated there, but not reproduced here.

Following is a historical development of projectiles given by Marshall 1 (Ref 11, pp 30-1):
The first projectiles used were made more or less like arrows with metal "feathers" and metal arrow-heads. They were soon found unsuitable and replaced by round shot made of stone, lead, iron or bronze. All these materials remained in use for several centuries, but stone was mostly used for large cannons. This was because it is less expensive than metal and also because the barrels of early cannons would not stand the strain of discharging heavy materials. Lead and iron bullets were usually employed in small arms. Attempts were made very early to throw from cannons incendiary missiles such as had been previously discharged from catapults, etc. In order to prevent the extinguishing of flame during flight, the projectiles (mostly stones), smaller than the bore of the cannon, were smeared over with incendiary matter and wrapped in cloth soaked in the same material. This method was used in 1469 at the siege of Weissenburg. Another incendiary, the "red hot shot", was dangerous to use because it could ignite prematurely the powder charge in the gun. This difficulty was overcome in 1579 when Stephen Batory, King of Poland, placed a thick wet pad on top of powder charge prior to introducing the hot-shot. Hot-shots were used with great effect by the English in the defense of Gibraltar in 1782.

Hollow projectiles (shells) filled with BkPdr could not be made until casting of metal was developed. But a sort of weak shell was made of earthenware, or by joining two metal hemispheres. These were filled with a slow-burning powder well rammed, and provided with an igniter, which was lighted by the flame of propelling charge, and burned for a few minutes before reaching the charge in the shell. Better results were obid with an igniter, invented in 1588 (qy), or with fuse invented by Bickford in 1831 (See Vol 2 of Encycl, p C112-L). The shells for early muzzle-loading rifled cannons were provided with studs to fit into rifling and with copper plates (gas-checks) over the base to prevent the escape of gases past the shell. For some of the early rifled breech-loading guns, the shells were coated with lead, but now they are provided with copper bands near the base to take the rifling and prevent escape of gases. Originally shells were filled with BkPdr, but now they are filled with HE's, such as TNT, Comp B, etc.

1885–1886. Accdg to Colver (Ref 12, p 14), three varieties of PA expls, invented by E. Turpin, were introduced in France. The first variety known as Explosif de Turpin consisted of powdered PA either pressed alone, pressed with the addn of sofn of a binder (such as gum arabic or collodion), or by pouring the molten material into containers of suitable form. Almost as soon as PA was introduced for military purposes, a mixt of 70 parts PA & 30 of Guncotton dissolved in 45 parts of acetone (or ether-alcohol) was used under the name of Mélinite. This was 2nd variety. Then later Guncotton was discarded and straight PA was heated at 130–135° and poured directly into shells. This was the 3rd variety, and the name was Mélinite (called Lyddite in England). The tests conducted in 1892 at LeBouchet showed that shells loaded with cast PA were safe and very powerful.

Mélinite (PA) was used in France for more than 25 years, and only during WWI was it replaced by TNT (Trotyl, in Fr)

Note 1: Since the mp of PA is relatively high (122.5°), the pouring process required to melt PA to that high temp was rather dangerous. Various additives have been suggested to lower the mp. Several low mp mixts were suggested by Girard, and four of them are listed in Colver on p 15. Most interesting of them is: PA/TN-Cresol in molecular proportion 1:1. Its mp is 78°

Note 2: PA is very poisonous and it dyes skin yellow, and this is hard to wash out. These were probably the reasons why PA was eventually replaced by other HE's, such as TNT, PETN, RDX, etc.

1886. Brown or Cocoa Powder was manufd at Okhta Plant, near StPetersburg, Russia (Vol 2 of Encycl, p B173-R)

1886. Accdg to Gorst (Ref 71, p 14), I.M. Chek'tsov proposed to load shells and mines with a mixt of AN 72.5 & AmmPicate 27.5%, named by him Gromoboy

1886. The British had 16.25-inch cannons weighing 110 tons which required very dense, very large grain BkPdr, but it was soon replaced by smokeless powder (called now smokeless propellant). With this propInt it was possible to propulse a shell weighing a ton a distance of 20 miles (Ref 11, p 30)
1887. P. Vieille devised in France the stability test known as “l’épreuve à 110°C de Vieille” (l’épreuve au première rouge), but it proved to be of no practical value until it was modified by Commandant Lepidi and became known as “l’épreuve à la résistance totalisée” (Test for Total Resistance). Both of these tests, of which the 2nd test became French official, are described in Vol 1 of Encycl, p XXI, under “Resistance to Heat Test”

1887. Another Commission was appointed in France to inquire into coal-mine explosions. Influenced by Berthelot’s work and theories, its attention was directed to the question of the heat evolved by an explosive and the resulting temperature of the products (Ref 11, p 45)

Note 1: As a result of the work of the Commission, a max temp of 1500°C for expls used in coal layers (explosifs grisou-couche) was established; and 1900°C for expls used for blasting rock in gaseous coal mines (explosifs grisou-roche) (Vol 3 of Encycl, p C369-L)

Note 2: Methods of calculating heat and temp of expls are described in Vol 3, pp C447–C449)

1888. In Germany and in England it was recognized that the temp of expls is only one of the factors in making an expl safe or dangerous to use in fiery mines. Consequently, reliance has been placed more upon testing in galleries, which are intended to imitate as nearly as possible conditions in the mines. The Committee was appointed in England in 1888 and a trial gallery at Hebbum Colliery completed in 1892 (1890?). After experimenting with various expls until 1895, the use of several expls, mostly based on AN (Ammonium Nitrate) was recommended. Based upon the results obt by this Committee, the Coal Mines Regulation Act of 1906 was founded. Following this Act, a gallery was constructed at Woolwich Arsenal and another more lately at Rotherham (Ref 11, p 46)

Note: More detailed description of this subject is given in Vol 3 of Encycl, pp C372 & C373)

1888. Cyclotrimethylene Trinitrosamine was discovered simultaneously by Griess & Harrow and by Mayer. It was studied in 1895 by Duden & Scharff and in 1896 by Delépine (Ref 70, pp 86–90). It was used during WWII in Germany under the name R-Salz (PATR 2510, p Ger 170-L). See also Vol 3 of Encycl, p C633

1888. Munroe-Neumann Effect or Shaped Charge Effect was accidentally discovered by C.E. Munroe of the US, but was not put to practical application until 1910 when C.E. Neumann of Germany proposed using the effect in the manuf of explosive ammunition (Vol 4 of Encycl, pp D442 to D454)

1888. PA (Picric Acid) was adopted by the German Govt, under the name Granattrüngung 88 (GfG 88). It consisted of PA pressed in cardboard or metal containers. It was used for filling shells, land mines, depth- and demolition charges. It was used as late as WWII, under the name Füllpulver 88, listed in PATR 2510, p Ger 46-R (1958), as Filler No 2

1888. Triaminotrinitrobenzene was first prepd by Jackson & Wing. More recently (1928), it was prepd by Flürscheim & Holmes (Ref 70, pp 364 to 366). Its uses are not indicated in Ref 70

1888. PA was adopted in England under the name of Lyddite (Ref 11, p 50); then later in Italy as Pertite and in Japan as Shimose (Ref 21, p 386)

1888. Alfred P. Nobel (1833–1896), famous scientist (born in Sweden, raised and educated in Russia, and worked in Sweden, Germany & USA), prepd a double-base smokeless proplnt by replacing camphor of Celluloid (Vol 2 of Encycl, p C95-L) with NG. He called the proplnt Ballistite. In 1889 he prepd Ballistite directly from Collodion Cotton (Vol 2 of Encycl, p C103) and NG using the “solventless” method of Lundholm & Sayers. More detailed description is given in Vol 2 of Encycl under Ballistite on pp B8–B9 (See also Vol 3, p C400 and Ref 31a, pp 293–95).

1888. Dr W. Kellner prepd in the laboratory of F. Abel the double-base proplnt Cordite (See Vol 3 of Encycl, p C532-L)

1889. Cordite, the 2nd successful double-base propellant, was patented for the British Government by F. Abel & J. Dewar and adopted as a military proplnt. It was Cordite Mark 1 or CSP1 (Cordite Smokeless Powder) (See Vol 3 of Encycl, p C532-L; also Ref 31a, p 295 & Ref 44, p 246)

1889. A. Nobel proposed using DPNA (Diphenylamine) as stabilizer for his Dynamite (Ref 44, p 250)

1890. Accdg to Dutton (Ref 58, pp 144–46),
attempts to use Dynamite as bursting chge in artillery shells were unsuccessful, because it blew up inside the barrels, thus destroying them and wounding or killing personnel. In order to reduce the muzzle velocity, pneumatic cannons operated by compressed air were invented. Here the Dynamite shell was propelled with lower velocity than possible in using a proplnt. At that time Americans were still using a brown variety of BkPdr, while all European countries switched to smokeless propellants. In 1890, the US Navy built the ship USS Vesuvius which had mounted on her bow three monster air cannons which hurled shells weighing a thousand lbs each and contg 600 lbs of Dynamite with some NC. An electric storage battery inside each shell ignited, on impact, NC which de-tonated the Dynamite. Great things were expected of Vesuvius when in 1898 it attacked Santiago, Cuba, but they were not fulfilled because the range of air-cannons, 1 mile, was shorter than the range of battery at the fort of Santiago. After this the US Congress decided to remove the air-cannons.

1890. Hydrogen Azide and Hydrazoic Acid were first prepd by T. Curtius and then investigated by D. Mendeleeff (Vol 1 of Encycl, pp A537-R to A542-R).

1890. Tetranitrocarbazole, first prepd by C. Graebe. A better method was developed in 1912 by Casella Co, and this method is still used. TNCHBz is suitable as component of igniter and pyrotechnic compns (Ref 70, pp 328–30).

1890. Cuprous Azide was first mentioned by T. Curtius in Ber 23, 3023, but was not properly investigated. History of later work is given in Vol 1 of Encycl, p A534.

1890. A. Wohl proposed polymerized glycerin as a gelatinizer for Dynamites, but it was not very satisfactory because it diminished strength & brisance of Dynamite, made it too difficult to detonate and affected adversely its oxygen balance (Vol 5 of Encycl, p D1589-L).

1890–1891. Mercurous Azide was prepd by T. Curtius (Vol 1 of Encycl, p A591).

1890–1891. Silver Azide was prepd by T. Curtius (Vol 1 of Encycl, p A598).

1890–1891. Sodium Azide was prepd by T. Curtius (Vol 1 of Encycl, pp A601 & A608).

1891. Hydrazine Azide was prepd by T. Curtius (Vol 1 of Encycl, p A536).

1891. Lead Azide was first prepd by T. Curtius (Vol 1 of Encycl, p A545).

1891. Ch.E. Munroe of the US prepd a double-base proplnt, Indurite, by gelatinizing Gun-cotton with about an equal amt of Nitrobenzene. Indurite was not found suitable for use in cannons (See Vol 2 of Encycl, p C32-L and Ref 31a, p 296 & Ref 44, p 246).

1891. C. Häusermann with the Griesheim Chemical Plant began the manuf of TNT, formerly (in 1880) prepd by Hepp in the laboratory. In 1901 the manuf was begun by the Carbonite Co and some expl firms soon followed. The first country to adopt it for military purposes was Germany (1902), as Sprengmunition 02. Other countries soon followed, such as Italy (1907), under the name Tritolo and Russia (1909), under the name of Trotil. In the English Service it was formerly known as T.N.T., but later as Tetryl. In the US it is called TNT and was adopted for military purposes in 1912 (Ref 11, p 265; Ref 18, p 20; Ref 64, p 24 (L) and Ref 71, p 12). It started to replace Mélinite in France as a bursting charge just before WWI.

1891. D.I. Mendeleeff (Mendeleev) of Russia (1834–1907) invented, quite independently of Vieille (who kept secret his poudre B), a completely colloided smokeless proplnt prepd by gelatinizing with eth-alc the NC of N 12.44%, which contained sufficient oxygen for complete combustion to CO & water. He called the proplnt Pyrokolodion (Pyrocollodion). It was adopted as “service” proplnt in newly developed, by Mosisin and Nagan, cal .30 (7.62mm), 5-cartridc rifle called “Trekshininyaya Vintovka” (Three-Line Rifle) (See Ref 51, pp Rus 22-R & Rus 23-L and Vol 2 of Encycl, p C32-L where the N content of Pyrocollodion is erroneously given as 13.44 instead of 12.44%)

Note: P. Tavernier did not say in his description in Ref 44, p 246, that Mendeleeff prepd, independently from Vieille, a NC contg 12.44% N and named it Pyrokolodion (Pyrocollodion), but said further that Lt Bernadou patented a single-base proplnt contg at first CP2 with 12.45% N and later pyrocollodion with N= 12.60%. All this is wrong. Lt Bernadou did not know the compn of French CP2, but he knew the compn of Mendeleeff’s Pyrocollodion,
which was only similar to CP₂.

Accdg to the books of: Lt John B. Bernadou, US Navy (1858–1908) (Ref 2a) and of Van Gelder & Schlatter (Ref 14, p 812–21), also listed in Vol 2 of Encycl, p B104 and described on p C32-L. Lt Bernadou, while serving as US Naval Attaché in StPetersbourg, learned in 1895 the method of prep of Mendeleff’s prop (He did not know the secret of Vieille’s “poudre B”) and decided to introduce it in Amer Navy in preference to Indurite. After investigating ballistic properties of Pyrocollodion, Bernadou found that better ballistic props as a cannon propint could be obtb if N content of NC were raised from 12.44% to about 12.6%. He gelatinized NC with eth-alc and called the resulting propint “Pyrocellulose Powder” or simply “Pyro”. It was adopted in 1897 by the US Navy and used in 1898 during Spanish American War. US Army adopted “Pyro” in 1899. Large scale manuf began in 1900 at a newly built “US Naval Powder Plant”, Indian Head, Md (now known as US Naval Ordnance Station). Picatinny Arsenal, Dover, NJ, started its manuf in 1907 (Ref 31a, p 297)

Note: Dutton (Ref 58, p 154) stated that Russia rearmed its Army in 1896 with Mendeleff’s propellant and that this propint modified by Lt Bernadou was adopted in 1901 by the US Govt 1891. It is the year of introduction by the Russian Armed Forces of 5-cartridge, cal 7.62mm rifle invented by Mossin and Nagan. It is known as trëkhlinyevaya vintovka (three-line rifle). It is briefly described in ArmyOrdn 30, 83–85 (1946) and in PATR 2145 (1955), pp Rus 22-R & Rus 23-L

1892. Tetranitroxyanilide was prepd by A.G. Perkin and used as a component of BPDr type expis and of pyrotechnic comprts (Ref 70, pp 331–34) (See also Hexanitroxyanilide)

1892. Hexanitroxyanilide was first prepd in England by A.G. Perkin. Used in pyrotechnic comprts (Ref 70, pp 170–72)

1892. Aminoguanidine and Derivatives were first prepd by J. Thiele (Vol 1 of Encycl, p A210) (See year 1928 for Nitraminoguanidine and Its Salts)

1892. Zinc Diazide was prepd by W. Willicnecus and in 1898 by T. Curtius & J. Risson (Vol 1 of Encycl, p A624)

1892. Ecrasite (Ekrasit) started to be used by Austro-Hungarian Army and this lasted until 1918, when its entire supply was blown up in Kiév, Russia. Its compn was never officially revealed, but it was suggested that it was either straight Ammonium Trinitro-m-cresylate (Cresylite) or its mixt with PA. More information on this subject is given in Ref 11, p 17 and in Vol 5 of Encycl, pp E8-R to E9-L

1892. 5-Amino-1H-tetrazole was first prepd by J. Thiele and examined in 1954 by L.F. Auridieth & J.F. Currier at Univ of Illinois and by D.B. Murphy, J.P. Picard, P. Rochlin & S. Helf at Picatinnny Arsenal. Was recommended as cooling agent in propints. Its salts and nitrated derivatives are explosive (Vol 1 of Encycl, pp A258-L to A260-R) [See also 5-Nitroaminotetrazole of Lieber, p A260-L]

1894. Stypnic Acid (2,4,6-Trinitrosorcinol) was first prepd by Hauff (Ref 11, p 281)

1894. Tetryl (or Trinitrophenylmethylaniline), first described in 1883 by van Romburgh was commercially manufd in Germany and tested as to its usefulness by military authorities (Ref 12, p 27)

1894. Hydrazine, Anhydroz was prepd in 1894 by Lobry de Bruyn, but its salts were prepd in 1875 by Emil Fischer, who coined the name “hydrazin”. Its derivs were investigated during WWII by Col L.F. Auridieth. More information is given in this Volume under Hydrazine and Derivatives

1894. Mercuric Azide was prepd by M. Berthelot & P. Vieille (Vol 1 of Encycl, p A590)

1894. Accdg to Gorst (Ref 71, p 11), attempts to use Melinit (Russ name for PA) in shells of cannon type 1877 were unsuccessful and the idea was abandoned

1894. Butanetriel Trinitrate was first prepd by Wagner & Ginsberg. A new method of prep was devised in 1948 by the US Rubber Co, and its props were evaluated in 1948 by the US Naval Powder Factory. Used as expl plasticizer for NC (Ref 70, pp 40–42)

1895. The Ammonium salt of Hexanitrodiphenylamine, known as yellow dye “Aurantia”, manufd by the Chemische Fabrik Griesheim, proved to be a valuable explosive (Ref 12, p 25)

1895. Adoption by the Russian Govt of modified revolver Nagan, invented in 1880's by Nagan. It is a 7.5mm (0.295-inch) weapon using 5 cartridges with bullets hidden in them.
It is described, with illustration of its cartridge, in PATR 2145 (1955), p Rus 11-R

1896. The AB BoFors, Nobelkrut of Sweden patented a smokeless propellant which was prep’d by gluing and compressing together several layers of different components. The outer layers were slow burning, while the speed increased progressively towards the inner layers (Ref 44, p 246)

1896. Introduction by P. Vieille of 2% amyl alcohol as stabilizer for smokeless propellant did not produce satisfactory results, especially in cannons of large caliber, and the amount was increased in 1906 to 8%. Nevertheless, two disastrous expls took place on battleships Iena in 1907 and Liberté in 1911. Both expls were ascribed to spontaneous ignition of poudre BAm stored in their powder magazines (Ref 44, p 248 & Ref 31a, p 308)

1897–1898. It is presumed that Research Establishment at Spandau, Germany, proposed to treat grains of small arms’ prop1nts on the outside with slow-burning substances like camphor. Such prop1nts were progressive to a certain extent. The GerP 120021 of 1898 proposed treating grains of NC with aromatic nitrocompds, such as DNT (Ref 44, p 246)

1897–1898. Chedites were invented in 1897 by E. Street of England, and several varieties of original compn were patented in 1898. A detailed description with several tables is given in Vol 2 of Encycl, p C155 to C164

1898. Cadmium Diazide was prep’d by T. Curtius & J. Risom (Vol 1 of Encycl, p A526)

1898. Calcium Diazide was prep’d by T. Curtius & J. Risom (Vol 1 of Encycl, p A527)

1898. Chromium Triazide was prep’d by T. Curtius & J. Risom (Vol 1 of Encycl, p A530)

1898. Cobalt Triazide was prep’d by T. Curtius & J. Risom (Vol 1 of Encycl, p A531)

1898. Cupric Azide was prep’d by T. Curtius & J. Risom (Vol 1 of Encycl, p A532)

1898. Iron Azide or Ferric Triazide was prep’d in soln by T. Curtius & J. Risom, but it was not until 1917 when L. Wöhler & F. Martin prep’d it again and isolated it (Vol 1 of Encycl, pp A543-L to A544-L)

1898. Lithium Azide was prep’d by T. Curtius & J. Risom (Vol 1 of Encycl, p A588)

1898. Manganese Diazide, Basic was prep’d by T. Curtius & J. Risom, and then in 1900 T. Curtius & A. Darapsky prep’d Mn(N₃)₂ (Vol 1 of Encycl, p A590)

1898. Nickel Diazide, Basic was prep’d by T. Curtius & J. Risom, and then in 1900 T. Curtius & A. Darapsky prep’d Ni(N₃)₂ (Vol 1 of Encycl, p A593)

1898. Potassium Azide was claimed to be prep’d by L.M. Dennis & C.H. Benedict, and in the same year by T. Curtius & J. Risom (Vol 1 of Encycl, p A595)

1898. Rubidium Azide was first prep’d by L.M. Dennis & C.H. Benedict and also by T. Curtius & J. Risom (Vol 1 of Encycl, p A596)

1898. Strontium Diazide was prep’d by the same persons as above (Vol 1 of Encycl, p A620)

1898. Thallium Azide was prep’d by the same persons as above (Vol 1 of Encycl, p A622)

1898. Azotetrazole was prep’d by J. Thiele, and its expl salts were recommended by E. von Herz and by H. Rathsburg for use in detonators. Its derivatives were described by L.F. Auwšteth & J.F. Currier in Univ of Illinois Rept, “Derivatives of 5-Aminotetrazole”, pp 22–23 (1954) (See Vol 1 of Encycl, pp A659 & A660)

1898. Simon Thomas developed stability test, which is described in Ref 28a, p 80. It was modified in 1920 and became the “Dutch Test” or “Loss of Weight Test”, described in Vol 5 of Encycl, p D1580-R

1899. Potassium Dinitrobenzofuroxan, first prep’d by P. von Drost, was examined at PicArsn in 1954 and 1955. It is in use in priming comps (Ref 70, pp 302 to 305)

1899. Cyclotrimethylenetetranitramine called RDX or Cyclonite, was first prep’d by Henning of Germany and he named it Hexogen. He used it for medical purposes. Then Brunswig prep’d it in 1916, but it was E. von Herz who recognized in 1920 its value as an expl. The first one who prep’d it in fairly good yield was G.C. Hale of Picatinny Arsenal (in 1925). Then nothing was done until 1940 when J. Meissner of Germany developed a continuous method of manuf. In 1940, Drs Ross & Schiessler of Canada developed a process which did not require the use of hexamine as a starting material, but the most important work on manuf and investigation of RDX was done by W.E. Bachmann. Four methods of manuf of RDX were developed in Germany (Vol 3 of Encycl, pp
C611 to C626). Expl mixtures based on RDX are described on pp C626 to C630. See also Ref 70, pp 69–75

1899. Oxylulit, invented by Linde, consisted of liquid air (or oxygen) absorbed in wadding, charcoal or other org material. As these mixts were hard to detonate, kieselguhr was substi-
tuted as absorbent with an addn of petroleum. It was inconvenient to use, as the cartridge had to be fired within 5 to 15 mins of its prep. It was tried in Austria and used in construction of the Simpon tunnel (Ref 11, p 44)

1899. Aluminized Explosives. The addn of Al to increase the performance of expls was first proposed by R. Escales of Germany and patented in 1900 by G. Roth. For more details see Vol 1 of Encyl, pp A146 to A151

1899. Ammonial, originally contg AN, charcoal & Al was proposed by R. Escales and patented in 1900 by G. Roth (Vol 1 of Encyl, p A287) (See also year 1917 for Austrian Ammonial T)

1899. Tritonal 80/20 (TNT 80 % & Al 20%) was developed in the US as filler for bombs. The addn of Al to expls was proposed in 1899 by Escales and patented in 1900 by Roth, but it was not properly investigated until 1943 and 1944 at PicArsn by W.R. Tomlinson Jr and proposed as filler for bombs of high blast ef-
fect (Ref 64, pp 7-68 & 7-69 and Ref 70, pp 386–90)

1900. It was proven during South African War (1899–1902) that Brit proplnt Cordite Mk 1, which contd 5% NG, caused very excessive erosion in gun barrels. As a result of this, the amt of NG was reduced to 30%, NC was increased to 65, and mineral jelly was 5% (Ref 44, p 247 & Vol 3 of Encyl, p C532-L)

1900. It was also shown during South African War that shells loaded with Lyddite (Eng l name for PA) were less satisfactory than those loaded with Black Powder. This was caused by incomplete deton of PA, which was attrib-
tuted to the faulty construction of detonators (Ref 71, p 11)

1900 is the year of discovery by the Badische Anilin- und Soda Fabrik of “contact process” for manuf of strong sulfuric acid and oleum, which was less expensive than the chamber method. As such acid was required for mixed nitr-sulfuric acid to nitrate aromatic hydro-
compds to tri- or higher states, Trinitrobenzene and Trinitrotoluene could be inexpensively manufd (Ref 12, p 17)

1900. Germans started to use in their NC rifle proplnts either 0.5 or 1.0% DPhA as a sta-
bilizer (Ref 44, p 250)

1900. Barium Diazide was prepd by T. Curtius (Vol 1 of Encyl, p A523-L)

1900. Konovalov of Russia prepd some expls by laboratory nitration of aliphatic derivs of petroleum, using weak nitric acid under pressure, but the yields were poor (Ref 12, p 29)

1900. Accdg to Gorst (Ref 71, p 15) it was proposed in Russia to replace MF (Mercuric Fulminate) detonators with those in which 3/4 of MF was replaced by Tetryl or Trotol (TNT)

1900. Chromatography, a physical method of separation in which the components are partitioned between two phases, was invented simultaneously and independently by American geologist D.T. Day and Russian botanist M.S. Tsvet (Tsweet). Their methods and later modifications are described in Vol 3 of Encyl, pp C289 to C298. Qualitative method of separation of ingredients by adsorption was known for centuries. This is described on p C289-R

1901. Hexanitrotribiphenyl was first prepd by F. Ullmann & J. Bielecki (Vol 1 of Encyl, p B124)

1901. Accdg to Colver (Ref 12, p 21), manuf of TNT by the method of P. Hepp, as improved by C. Häussermann, was conducted in 3 stages: 1) Nitration of toluene to MNT, using relatively weak spent mixed acid; 2) Nitration of MNT to DNT using stronger acid and 3) Nitration of DNT to TNT using strong acid with oleum. This method, which proved to be more economical than direct nitration of Toluene to TNT, was used in the US until WWII, when Dr I.A. Grageroff introduced the so-called “reverse process”. This method was twice as rapid.

Many plants switched to Continuous Methods (See Vol 3 of Encyl, p C505)

1901. Accdg to Colver (Ref 12, pp 20 & 24), German expls industry unreservedly adopted the manuf of TNT, mostly by the method of Hepp as improved by Häussermann (See above). Some factories used their own modifications and used names, such as Trotyl, Trymol, Tolril, etc. The names Triplastic and Plastortyol were used for TNT worked-up with DNT, MNT and a small amt of NC
1902. Accdg to Colver (Ref 12, p 23), Germany was the first country to adopt TNT for military purposes, mostly as filler of HE shells. Its official name was Füllpulver 02 (Fp 02). It is listed in PATR 2510 (1958), p Ger 46-R as Filler No 1

1902. Dr Edeleanu & G. Filip of Rumania took out a patent for manifold of nitrocompounds from petroleum, but they found no practical application (Ref 12, p 27)

1902. Germans proposed using cyanamide, di-cyanamide or tricyandiamine as auxiliary stabilizers in their propellants (Ref 44, p 251)

1902. Introduction of Bergmann-Junk Stability Test. It is described in Vol 2 of Encycl, pp B102-R to B103-L (See also Ref 44, p 251)

1902. Accdg to Colver (Ref 12, p 23), Germans also used TNT for demolition purposes, under the name Sprengmunition 02. It replaced Sprengmunition 88, which was PA (Pieric Acid)

1902. Cordeau détonant, developed in France, was improved in 1907 by L. Lheure. This type of cordeau was introduced in 1913 in the US by the Ensign-Bickford Co (Vol 3 of Encycl, pp C530 & C531). Addnl info is given on pp D103 to D106, under Detonating Cords or Detonating Fuses

1903. Accdg to Tavernier (Ref 44, p 252), Rottweil Fabrik of Germany proposed vaseline with 1–2% Na bicarbonate as muzzle-flash suppressor. This was replaced in 1907 by soap or rosin

1903. Until this year no satisfactory antifreeze for NG was found, and many industrialists preferred not to use any of those previously proposed, but let ordinary Dynamite be defrosted in winter. The disastrous expn of 1902 in Greisenau Mine, Germany, took place during defrosting of Dynamite and encouraged the renewal of research on antifreeze. In 1903 the SA de Poudres et Dynamites of France introduced DNT and TNT as antifreeze. They did not lower the fr p of NG sufficiently. The Glycerin Dinitrate (GDN) proposed in 1903 by A. Mikolajczak in Gelatin Dynamite was not very satisfactory because it lowered the strength of Dynamites too much (Vol 5 of Encycl, p D1589-L)

1904. The Dynamit AG of Germany proposed DNCH (Dinitrochlorohydrin) as an antifreeze. This compd was satisfactory, except that it produced very noxious fumes contg chlorine

1904. 2,3,5,6-Tetranitroanisole was prepbd by J. Blanksma (Vol 1 of Encycl, p A454-R)

1904. Introduction of Obermüller Stability Test which consists of heating 1 to 2g NC in a small tube under vacuum at 135–140° and measuring the pressure of evolved gas by vacuum manometer (Ref 38a, pp 87–88)

1904–1905. 2,4,6-Trinitroanisole or Methyl Picrate was patented for use in smokeless propellants in mixt with an equal amt of Pyrocellulose (Vol 1 of Encycl, p A452-R)

1906. The problem of producing quite satisfactory antifreeze remained unsolved until S. Nauckhoff of Sweden published his work in ZAngewChemie 18, pp 11 & 55 (1905), in which he formulated the requirements for a satisfactory antifreeze and gave a list of compds which might prove to be suitable

1906. Centralites were first investigated in Germany at the Zentralstelle für wissenschaftlich-technische Untersuchungen zu Neubabelsberg (Central Laboratory for Scientific & Technical Research at Neubabelsberg) as possible stabilizers for smokeless propellants. Several formulations were developed which proved to be very successful. A detailed description is given in Vol 2 of Encycl, pp C126 to C140

1906. Accdg to Gorst (Ref 71, pp 15–16, it was proposed in Russia to use LA (Lead Azide) in lieu of MF (Mercuric Fulminate) in compound detonators with Tetryl and later with Ten (PETN). Much work on improvement of LA detonators was done by Col A.A. Solonina. He also worked on detonators contg Lead Trinitroresorcinate, called in the US Lead Styphnate (LSt)

1906. After publication of Nauckhoff's paper (See under 1905), W. Will proposed TeNDG (Tetranitrodiglycerin) as an antifreeze, but it never was used in Germany. It was used in the US since 1912 (Vol 5, p D1589-R)

1906. Accdg to Tavernier (Ref 44, p 252), WASAG patented as muzzle flash suppressors, organic salts (such as tartrate, oxalate or citrate of sodium)

1906. Dr C. Claessen patented in Germany the use of totally substituted ureas as stabilizers in smokeless propellants. These substances can be subdivided into derivs of sym-N,N'-diphenyl-
ureas and asym-N,N-diphenylureas. Some of them served as gelatinizers. Sym-N,N-Diphenylureas were investigated beginning in 1906 at the Central Laboratory for Scientific and Technical Research at Neubabelsberg, near Berlin and named Centralites. There were developed Nos 1, 2, 3, 4 and Butyl Centralites, all good stabilizers for NC's, especially No 1. More detailed description is given in Vol 2 of Encycl, pp C126-R to C140-R asym-N,N-Diphenylureas, known now as Acardites were described before WWI by J.P. Reudler in Rec 33, 49–55 (1914). We were able to describe them in detail because Dr Hans Walter, who worked with them during WWII in Germany, gave us some info which is not in the literature (See Vol 1 of Encycl, pp A7-R to A9-L). Especially useful was Drs H & B Walter's help in description of Analytical Procedures (See Vol 1, pp A9-L to A10-R). P. Tavernier described Acardites in MP 38, pp 307–08 & 329 (1956).


1906. Adoption by French Navy for large caliber cannons of poudre B Marine with 8% amyl alcohol as stabilizer (poudre BAm₈) in lieu of previously used BAm₂. (Ref 44, p 249) 1907. Adoption in 1906 by the French Navy of proplnt with 8% amyl alcohol stabilizer did not prevent the disastrous expl on 1907 of battleship Ilena. As a result of this, previously tested DPhA (diphenylamine) was approved as stabilizer of Naval cannon proplnt, which became known as "poudre B(Bo)" (Ref 44, p 249) (See also Vol 1 of Encycl, p A395-L). 1907. Vezio Vender of Italy patented as an antifreeze for Dynamites, mixed glycerol esters, such as Dinitroacetin with Dinitroformin [Ref: SS 2, 21 & 195 (1907)]

1909. Ammonium Picrate (AmmPicr), Dunnite or Explosive D, was standardized in the US as a bursting charge for AP (Armor-Piercing) shells (Ref 64, p 2-3). Its prep'n and properties are described in Ref 70, pp 136–39, where it is stated that AmmPicr was first prepd in 1841 by Marchand and in 1869 used by Brugère in admixture with K nitrate as a proplnt. Used as an HE after 1900 (See also 1841)

1909. Accdg to Colver (Ref 12, p 18), low-nitrated aromatic hydrocarbons, such as MN-Naphthalene, MNbZ & MNT started to be used in coal mining expls based on AN. This was because of their stability and insensitivity to shock & to ignition 1909. Accdg to Gorst (Ref 71, p 12), Russia started to use Trotil (TNT) for loading shells 1909. Silvered Vessel Test, devised by Sir Robertson for detn of stability of Cordite, was described by F.L. Nathan in JSocChemInd 28, 443 (1909), and briefly described by Reilly (Ref 28a, p 81) and in Vol 1 of Encycl, p XXIV. It is also known as "Waltham Abbey Test.

1909. Accdg to P. Tavernier (Ref 44, p 252), one of the greatest German discoveries was Pulver ohne Lösung (POL), called in Fr Poudre sans dissolvant (psd) and in Engl Solventless Powder (or Propellant). Its discovery was attributed to Brunswig & Thieme, although many other scientists contributed to development of such proplns

1910. B. Flürschein patented the prep'n of 2,3,4,6-Tetranitroaniline, one of the most powerful expls. It was used during WWI by the Germans and Japanese (Vol 1 of Encycl, pp A411 to A413)

1910. Beginning in Oct, DPhA became the only stabilizer used in French proplns, but there were supplies contg amyl alcohol and one of the battleships, Liberté, had in 1911 a fire resulting in expln of poudre B(AM₈) of 1906 vintage. The ship was partially destroyed (Ref 44, p 249 & Ref 31a, p 308)

1910. Tetrazene, described in Vol 6, p G169 as Guanylnitrosoaminoguanyl Tetrazene was first prepd by K.A. Hofmann et al. The most extensive studies were done in 1931 by Rinenbach & Burton 1910–1911 & 1913. Dr C. Claessen developed a procedure for prep'n of solventless proplns contg about 30% NG. In his process the slightly wetted blend of NC–NG, known as galette (cake), with some additives was fed between two rolls (heated to 85–95°) by hot water circulated in them) in order to obtain the mass in the form of a sheet. This is called "lamination". Then the sheet passed thru a press to be drawn in the desired form. In order to facilitate the prep'n of such proplns,
some low N content NC was included in the cake or about 5% of NH₄₉a—Na oxalate was incorporated. Aromatic nitrocrops with mp's below 100°, such as DNT or TNT were also recommended. The 1st Ger solventless proplnt, RPC/12, contd 70% of blend of NC's, resulting in a N content of 11.7%, NG 25 and 5% of Centralite. It was made in large grains and, since it was solventless, the time consuming operation of solvent proplnts was not required. The proplnt was used in large caliber cannons (Ref 44, pp 252–53)

After 1911. As a result of disastrous explosions of battleships lëna (1907) and Liberté (1911), it was decided to investigate various stabilizers besides DPhA, such as naphthalene, MNaphthalene, carbazole and DPHNitosamine. None of them proved to be better than DPhA (Ref 31a, p 309 & Ref 44, p 250)

1912. Carbonit AI patented the following expls: Hexanitrodiphenylamine, Hexanitrodiphenylsulfide, Hexanitrosulfobenzide, Hexanitrodiphenyloxide and Hexanitrodiphenyl (Ref 12, pp 25 & 26)

1912. Acoday to E.M. Symmes (translator of Naoum's book), DGTen (Diglycerin Tetranitrate) introduced in 1906 by W. Will as an antifreeze was not used in Germany, but it was used in the USA to a very large extent until about 1926 (See footnote 5 on p 202 of Ref 16 and in Vol 5 of Encycl, under Diglycerol, p D1261-L)

1912. Modified British Cordite MD used in rifles, changed its form from cords to small tubes (Ref 44, p 253)

1912. TNT, first pred in 1891 (qv) by G. Häussermann and adopted in 1902 by the Germans, in 1907 by the Italians and in 1909 by the Russians. It began in 1902 in the US to replace PA, and then in 1912 became the standard bursting charge for HE shell for the mobile artillery of the US Army (Ref 64, p 2-4 and Ref 70, pp 350–58)

1912. Nitroisobutylylglycerol Trinitrate or Trimethylolnitromethane Trinitrate was first pred by Hofwimmer. Used as expl gelatinizer for NC (Ref 70, pp 244 & 245)

1912 & 1914. Although Nitroglycerol (NGe) or Ethylene glycol Dinitrate (EGcDN) was known since 1904, it was not properly described until before WWI [See MP 16, 73 (1912) and 17, 175 (1914)]. It proved to be the best antifreeze for Dynamites and was found to be suitable as a substitute for NG. Large quantities of NGe were used in Germany during WWI. As an antifreeze it was used in mixts of 20/80—NGe/NG (Vol 5 of Encycl, p D1590-L & R). Prepn and properties of EGcDN are described in Vol 6, pp E259-R to E278-L, under ETHYLENEGLYCOL DI-NITRATE

1912–1914. Russian proplnts were based on Pyrocoollodion of Mendeleeff. They contd 1% of DPhA stabilizer and were in the form of tubes

US proplnts were based on Pyrocellulose introduced by L. Bernadou. They contd 1% of DPhA and were usually multitudular in form

Italian proplnt, manufd from 1896 at the Royal Powder Factory at Liri was Solenite, which was similar to Brit Cordite contg about 35% of NG. It consisted of 1/3 Guncotton, 1/3 Collodion Cotton and 1/3 NG with added vaseline (replaced later by Centralite). Beginning in 1910, the Société Nobéle of Avigliana manufd C₂, which was a variety of Brit Cordites but in the form of tubes. Italians manufd also Ballistite of high NG content which contd 0.5% of aniline as stabilizer (Ref 44, p 253)

1913. Compression (or Crusher) Test of Kast is described in Vol 3 of Encycl, p C493

1913. A.S. Flexer of Vienna patented a process for manuf of nitrocrops from petroleum and tar but they found no practical application at that time (Ref 12, pp 27–28)

1913. Westfälisch-Anhaltische Sprengstoff AG (WASAG) devised solventless proplnts with low NG content in which stabilizers-plasticizers were anilides of organ acids in which hydrogen of imide group was replaced by a radical: ethyl or methyl acetanilide. The same Society proposed using esters of carbonic acid in which two hydrogens were replaced by radicals, such as diphenylurethane or methylphenylurethane. This permitted reducing the temp of lamination to 82° (Ref 44, pp 251–52)

1914. Lead Stypnate (LSt) or Nitratoresorcinol was first pred by E. von Herz (Ref 70, pp 193–96)

1914. G. Spica of Italy proposed using phenanthrene as stabilizer in solventless proplnts
with low NG content (Ref 44, p 254)

1914. **Cordite SC, Solventless** was developed at Waltham Abbey, England (Vol 3 of Encycl, p C532-R)

1914. **Trotyn (TNT)** started to replace in France Mûnitude (PA) as bursting charge

1914–1918. **World War I.** Prior and during the War many explosives, serving as substitutes or supplements of TNT, were developed in Germany, Austria, Russia, Italy and Great Britain. The most important were mixtures of HE's (such as TNT, TNX, TNZ and others) with AN and Al were: **Amatols** (British) (TNT with AN in various proprts) (Vol 1 of Encycl, pp A158 to A164); **Ammonals** (Austrian & US) (TNT or other HE with AN & Al) (Vol 1 of Encycl, pp A287 to A292) and **Alumatol** (British) consisting of AN 77, TNT 20 & Al 3%, used for filling bombs and trench mortar shells (Vol 1, pp A141-R & A142-L). In Russia, Col A.A. Solonina proposed using mixt of Trinitro-m-xylene (Ksili, in Rus) with AN & Al for loading hand grenades (Ref 71, p 14)

WWI. Incendiary bombs developed at that period were not very effective and comparatively easy to extinguish (Vol 2, p B243-R)

1915, April 22. **Modern Chemical Warfare** began with the German attack at Ypres, France, when 5700 barrels of chlorine gas were released against French and Canadian trenches. The attack was most effective because it was unexpected and many casualties resulted. This “gas-cloud method” was soon replaced by “chemical shells” which were widely used during WWI. Although “chemical warfare” was outlawed after WWI by the Hague Convention, all great nations continued development of new agents. The most efficient was Germany who developed before WWII several “nerve gases”, such as Trilon described in PATR 2510 (1958), p Ger 204-L and in Vol 2 of Encycl as GA & GB on p C167-R. They are also described in Vol 5, pp D1308-R to D1309-L, under Dimethylaminocyanophosphoric Acid, Monomethyl Ether. A comprehensive list of chemical warfare agents developed during WWI & WWII by various countries is given in Vol 2 of Encycl, pp C166 to C173, under “Chemical Warfare Agents”

A brief description of the use of poisonous and smoke gases in ancient times is given in this section under the years 431–404 BC

1915. **Dr C. Claessen patented in Germany a smokeless propellant containing volatile solvent, stabilized by introducing a large quantity of eutectic mixt DNT–TNT (Ref 44, p 251)

1915. **Amatols** (TNT+AN) were developed by the British in order to extend the available supply of TNT which was scarce (Vol 1 of Encycl, p A158)

1916. Dr C. Claessen patented solventless NC–NG proplnt contg acylaryl- or diaryl cyanamides as stabilizers (Ref 44, p 251)

1916. **Ballistite 50/50**, invented by Nobel, was manufd as propht for mortars. It was partially replaced by Attenuated Ballistite, which contd 25% NG and 15% DNT. The German name, **Würfelpulver** (WP), was assigned to Ballistite 50/50 used in mortars (Ref 44, p 252)

1916–1917. Single base NC proplnts were used in rifles and in the shape of tubes in some cannons. As gelatinizer for such propht an ether-alc mixture was often used, and as stabilizer diphenylamine (DPhA). Some NC–NG proplnts, gelatinized by acetone, contd small amts of organic nitrates. As stabilizer they used DPhA and later Centralite or Acardite. They were surface treated by Centralite or by aromatic nitrocomps. Such NC–NG proplnts were used in large caliber cannons (Ref 44, p 252)

1917. **Austrian Ammonal T** was developed by R. Fürig to be used in underwater expls, such as torpedoes and depth charges. It consisted of AN, Al & TNT (Vol 1 of Encycl, p A291-L & table on p A290)

1917. Accdg to Tavernier (Ref 44, p 252), WASAG patented as muzzle flash suppressor K chloride (0.5 to 5g)

1917. **Ardeer Cordite** was introduced (Vol 3 of Encycl, p C532-R)

After WWI. **Metril Trinitrate** or **Trimethylolmethane Trinitrate** was prepd in Italy and in 1927 in Germany. Used as an ingredient of rocket prophts (Ref 70, pp 206–08). Its prep and uses in Germany during WWII are described in PATR 2510 (1958), p Ger 113

1920. **Acidg to Davis (Ref 31a, pp 311–13), transformation of DPhA during aging was conducted beginning in 1920 by Desmaroux, Marqueyrol & Muraour and Marqueyrol & Lorrette; also by Davis & Ashdown (See also
Vol 5 of Encycl, pp D1417-R to D1418-L

1920. Cardox is a device for breaking coal in gaseous mines by the pressure produced on heating liquefied carbon dioxide. It is briefly described in Vol 2 of Encycl, p C67-R. Similar devices, Airdox and Armstrong Air Breaker, were developed in 1930 in the US. More successful were Hydrox developed in 1955 and Chemeeol developed by du Pont Co before 1958. They are also Hydraulic Coal Bursters (See Vol 3 of Encycl, p C434)

1921. It was known that some stabilizers act also as gelatinizers or colloidal agents

Davis (Ref 31a, pp 320–22), under the title "Gelatinizing Agents", describes results of investigation in France beginning in 1921 by Marqueyro & Florentin of a number of esters, amides, urea derivatives, halogen componds, ketones and alcohols as possible gelatinizers of soluble and insoluble NC. In 1922, Davis investigated many componds and gave in his book on p 322, a Table listing parts by wt required for complete gelatinization of Pyrocellulose (See also Vol 3 of Encycl, p C398-L, under the title "Colloiding Agents")

1923. O. Turek prep'd 1,3,5-Triazoido-2,4,6-trinitrobenzene and described its prep and properties in Chim&Ind(Paris) 26, 781 (1931) (See Vol 2 of Encycl, pp B43 & B44)

1923. W. Borschke, Ber 56B, 1942, described 1,3,4,5- & 1,2,4,5-Tetranitrobenzenes, both of them very expl (See also Vol 2 of Encycl, p B50 and Ref 70, pp 378–80)

1925. R.C. Moran prep'd 2(2',4',6'-Trinitro-N-nitranilino)-Ethanol Nitrate, also known as Pentryl. Not to be confused with Pentryl of A. Stettbacher, which designates mixts of PETN & TNT known in the US as Pentolites (See Vol 1 of Encycl, pp A425–29 and Ref 31a, p 229)

1927. Continuous Methods of Manufacture of Explosives. The first successful method was invented by Schmid of Austria, but some attempts were made as early as 1864. Description, with flow sheets is given in Vol 3 of Encycl, pp C501 to C510 for the methods of Schmid, Biauzzi, German and Bofors

1927. Diethyleneglycol Dinitrate was thoroughly examined by W.H. Rinkenbach and its current method of prep was patented in 1928 by A. Hough. A detailed description is given in the book of T. Urbanski, Vol 2, pp 149–54 (1965) and a brief description in Vol 5 of Encycl, pp D1232 & D1233-L (See also Ref 49, p 168)

1927. Trimethyleneglycol was first described by A. Rayner, but it was mentioned in 1895 by Noyes & Watkins as occurring in some glycerols (Ref 49, pp 206 & 209)

1928. B. Flörsheim & E. Holms prep'd 2,3,4,5,6-Penta nitrotrianiline in the pure state and found it to be a most powerful expl, comparable to RDX & PETN. It was stable while the same expl prep'd in 1910 in impure state was unstable (Vol 1 of Encycl, p A414)

1928. Nitraminoguanidine and Its Salts were prep'd by R. Phillips & J.W. Williams (Vol 1 of Encycl, pp A210-R to A212-L and A213-R)

1929. PVN (Polyvinyl Nitrate) was first prep'd in Germany and patented in 1938 in the US. Its uses are not indicated (Ref 70, pp 315–17)

1929. Acetyl to Dutton (Ref 58, p 174), the first cyclotron was built in the USA

1929. Copper Chlorotetrazole first prep'd by R. Stollé et al. More recently (1955), it was prep'd and investigated at PicArsn as possible ingredient of primary compns (Ref 70, pp 63 to 65)

1930. Dipentaerythritol Hexanitrate (DPEHN), obtd as impurity in PETN by W. Friederich & W. Brün was also prep'd in 1945 at PicArsn by S. Livingston and at Hercules Powder Co Lab at Radford, Va (Ref 70, pp 119–21 and Vol 5 of Encycl, p D1413)

1930–1931. Lead Azide, Dextrinated started to be used in the US, replacing gradually MF (Mercuric Fulminate) in priming compns (Vol 1 of Encycl, pp A545-R and A552-L). Plain, impure LA was first prep'd in 1891 by T. Curtius, but it was not until 1907 when T. Heonius of France succeeded in using it in explosives industry. It started to be used in 1920 in Europe, but in the US only dextrinated LA was allowed beginning 1930–31 (Ref 70, p 185)

1931. Propyleneglycol, although first prep'd in 1859 by Ch.A. Wurtz, it was not used until 1931 when it was produced on commercial scale by Carbide & Carbon Co (Ref 49, p 203)

1931. Glycerol Monolactate Trinitrate, patented in the US by Ch. Stine & Ch. Burke, was investigated in 1946 at PicArsn as possible expl plasticizer for NC (Ref 70, pp 140–42)

1931. 2,4-Dinitrotoluene (2,4-DNT) obtd
earlier as an impurity in crude TNT, was used in some expl mixtures (Ref 70, pp 116–18)

1933. Anilinotetrazole and Its Derivatives were first prep'd by R. Stolle et al (Vol 1 of Encycl, p A437-R)

1933. Pentrohit is a mixture of 20–50% of PETN with 80–50% NG. It was developed by A. Stettbacher (Ref 41, pp 83–4)

1933. Hexonit is a mixt of 20–50% RDX with 80–20% NG. It was developed by A. Stettbacher (Ref 46, p 111)

1934. Lead-2,4-dinitroresorcinale was prep'd and examined at PicArsn by J.D. Hopper and later (1944) by K.S. Warren (Ref 70, pp 187–89)

1934–1935. “G” Pulver. This proplnt was developed in Germany under the direction of Gen Uto Gallwitz. It was a “cool” double-base proplnt in which NG was replaced by DEGeDN or TEGeDN (Vol 3 of Encycl, p C400-R and pp CS11 & CS12)

1935. Ball Powder was developed by Fred Olsen of Olin Mathieson Chemical Corp (Vol 2 of Encycl, pp B11–B16)

1935. Accordg to Dutton (Ref 58, p 143), the World’s largest Dynamite Plant was constructed near Modderfontein, South Africa. It was operated by the African Explosive and Chemical Industries

1936. Penttaerythritol Trinitrate or PETRIN. Nitration products of PETN contg not more than 3-NO₂ groups were patented in Germany in 1936, but pure PETRIN was not prep'd until 1954 (Ref 70, pp 265–68)

1937. Baratol, which is the mixture of Bar nitrate and TNT (Tol), was developed by the British and used during WWII (Vol 2 of Encycl, pp B18 & B19)

1937. Gudolpulver was a “cool” proplnt developed in Germany by Dynamit AG Plant. It consisted of NC, DEGeDN & NGu (Nitroguanidene) (Vol 3 of Encycl, p C401-L)

1939. This year may be considered as the beginning of work on development in the US of atomic (or nuclear) energy. This work is briefly described by C.G. Dunkle in Vol 1 of Encycl, pp A500ff

1940. Composite Propellents consist of finely ground oxidizer (such as inorg perchlorate or nitrate) in a matrix of a resinous or elastometric material which serves as a fuel. The proplicants for use in JATOS were developed beginning 1940 (Vol 3 of Encycl, pp C464–C474)

1941. R.C. Elderfield prep'd N(2,4,6-Trinitro-N-nitranilino)-Trimethylmethane Trinitrate, designated as Heptryl, and called by G.B. Kistia-kowsky N-Nitro-N-picyl-trimethylmethylamine Trinitrate. It proved to be HE comparable in power and sensitivity to PETN (Vol 1 of Encycl, pp A441-R & A442-L)

1941. Lead Nitraminoguanidine was prep'd and examined at Picatinny Arsenal by A.J. Phillips. It was later recommended by L.R.V. Clark for use in primers and detonators (Vol 1 of Encycl, pp A212-R & A213)

Prior to WWII. Cartridge-Actuated Devices (CAD’s) or Propellant Actuated Devices. The first successful devices for pilot’s quick escape from a plane were developed by the Germans. They called them “Schleudersitzpatronen” and equipped them in many of their fighter planes. The British equipped their planes having speeds greater than 400mph with such devices beginning in 1945. The Americans started the study of emergency escape as early as 1940, but it was not until 1947 that the first device was standardized. It was named “M1 Personnel Catapult” (Vol 2 of Encycl, pp C70 to C72)

Prior to WWII. Composition B or Cyclotol, developed by the British, consisted of a mixture of RDX with TNT in various proportions. Some mixts contd wax. Its modifications as standardized during WWII in the US were: Compositions B–2; B–3; B–4 and B. Desensitized (Vol 3 of Encycl, pp C477-R to C484-L) (Ref 70, pp 46–50)

Prior to WWII. Under Dimethylaninoxyanophosphoric Acid, Monomethyl Ester are described in Vol 5 of Encycl, pp D1308-R to D1309-R

German Nerve Gases, such as Sarin, Tabun, Trilon and Soman. They are also described in PATR 2510 (1958), p Ger 204-L, under Trilons

Prior to WWII. Cyclotol, which are mixtures of Cyclonite (RDX) with TNT in various proportions were developed by the British and standardized in the US (Ref 70, pp 76–85) and Vol 3 of Encycl, pp C477-R to C484-L under Composition B Type Explosives and Cyclotol(

Prior to WWII were developed in Switzerland and Germany extremely toxic nerve gases, described under Trilons in Ref 56, p Ger 204-L

WWII. Items Developed or Standardized During
WWII. Ethylenedinitramine, EDNA or Haleite, first prep'd in 1887 by Franchimont & Klobbie and developed in 1935 as a military explosive by G.C. Hale, was standardized during WWII as a component of EDNATOL (EDNA+TNT) serving as a bursting charge for bombs and shells (Ref 70, pp 131-R and 154). See also Vol 6 of Encycl, p E238ff

**WWII. EDNATOL (EDNA 55 & TNT 45%).** See previous item and Ref 64, pp 7-74 & 7-75 and Ref 70, pp 130-32

**WWII. Cyclotetramethylene Tetranitramine or** Homocyclonite (Code named HMX, which means "High Melting Explosive"); was prep'd in 1943 by W.E. Bachmann (OSRD Rept 1981) and investigated at PicArsn by S. Livingston and O.E. Sheffield, et al (Ref 64, pp 7-64 to 7-66 and Ref 70, pp 173-77)

V-1 (Buzz Bomb) is described in Ref 56, p Ger 213-L

V-2 (Rocket) is described in Ref 56, p Ger 213-R

**WWII. Pentolites**, of which 50/50-PETN/TNT mixture is the most important, was standardized for use in shaped charges, bursting charges and demolition blocks (Ref 64, pp 7-69 to 7-71 and Ref 70, pp 272-75)

**WWII. Tetrytol**, of which 70/30-Tetryl/TNT, castable mixture is the most important in military applications. It was standardized for use in bursting type of chemical shells and in demolition blocks (Ref 64, p 7-72 and Ref 70, pp 341-49)

**WWII. Picratol (PA 52 & TNT 48%)** was developed as insensitive, castable fller for AP bombs and shells (Ref 64, pp 7-72 to 7-74 and Ref 70, pp 285 to 287)

**WWII. Torpex** (RDX 42, TNT 40 & Al 18%) developed by the British for use as a filler of warheads, mines and torpedoes (Ref 64, pp 7-77 & 7-78 and Ref 70, pp 259 to 263)

**WWII. HBX-1** (High Blast Explosive-1) is a modification of Torpex. Its compn & props are given on p 7-79 of Ref 64 and in this Vol of Encycl (See also Ref 70, pp 156 to 158)

**WWII. Demolition Explosives and Creteing Charge** are described in Vol 3 of Encycl, pp D54 to D61. See also Ref 64, pp 7-84 & 7-85

**WWII. Military Dynamites** are described in Ref 64 p 7-84

**WWII. DBX** (Depth Bomb Explosive) is an Amer expl developed to replace the Brit more sensitive Torpex (Vol 3 of Encycl, p D19-L and Ref 70, pp 91 to 94)

**WWII. Lead-4,6-dinitroresorcinol, Basic** was prep'd by the British for use in electric detonators (Ref 70, pp 190 to 192)

**WWII. Nipolitos** are types of NC-DEGDN-PETN propellants or expls patented by Dr E. von Holt, the inventor of Holtex (See under year 1958). Compnts and some props of Nipolitos are listed in PATR 2510 (1958), p Ger 117

**WWII. Bombs, Incendiary** were very efficient and hard to extinguish. They did more damage to the buildings than did other bombs. A brief description of such bombs and the damage inflicted by them during WWII is given in Vol 2 of Encycl, pp B235 to B237

**WWII. Composition A**, developed by the Brit, consisted of RDX 91 & beeswax 9%. When standardized in US, beeswax was replaced by synthetic wax and the expl became Composition A-2. Other modifications were Compositions A-3; A-4 and A-5 (Vol 3 of Encycl, pp C474 to C477)

**WWII. Composition C**, developed by the Brit, was a plastic RDX demolition expl. As standardized in the US, it consisted of RDX 88.3 & a nonexplosive plasticizer 11.7%. Its other modifications were Comp C-2, C-3 and C-4. The C-4, known as Harrisite, was developed before 1950 by K.G. Ottoson at Picatinny Arsenal. Their compns are given in Vol 3 of Encycl, pp C484 to C488

**WWII. Azon Guided Missile**, developed during the War, is briefly described in Vol 1 of Encycl, p A662-R

**WWII. Bangalore Torpedoes** were developed and are described in Vol 2 of Encycl, pp B16-B18

**WWII. Baronal**, consisting of Ba nitrate, TNT & Al, was described in 1942 in OSRD Rept 1035 (Vol 2 of Encycl, pp B21 & B22)

**WWII. Cast Double-Base Propellants for Rockets**, developed at the beginning of 1944 are described in Vol 2 of Encycl, pp C84 to C86

**WWII. Flashless Cordite contg NGu (Nitroguanidine) was adopted as the best, after investigation beginning in 1921, of various additives to Cordite (Vol 3 of Encycl, p C533 and Ref 30, p C536-L)
WWII. Cycloetramethyleneferrotetranitramine, called HMX by the British and also known as Octogen was described in OSRD Rept 652 (1942) and in Vol 3 of Encycl, pp C605-R to C610

WWII. RDX or Cyclonite. See 1899. Cyclo-trimethyleneferrotetranitramine

WWII. Demolition Explosives. Wide use was made in demolition practice of shaped charges, which utilize the Munroe- Neumann Effect, described in Vol 4 of Encycl, pp D442 to D443. US demolition shaped charges are described in Vol 3 of Encycl, p D58, while US plain demolition charges are on p D57. British, French, German, Japanese and Russian charges are on pp D58 to D60

WWII. Depth Charges are described in Vol 3, pp D86 & D87

WWII. Detonators are described in Vol 3, pp D92 to D96

WWII. Detonating Cords or Detonating Fuses are briefly described in Vol 3, pp D103 to D106. (See also 1902. Cordeau détonant)

WWII. Shaped Charge Effect, discovered in 1888 by C.E. Munro and applied in 1910 to manuf of expl items by E. Neumann, was extensively utilized during the war. The effect is described under Detonation, Munroe-Neumann Effect in Vol 4 of Encycl, pp D442 to D454

WWII. Triethylene glycol Dinitrate was prepd in Italy and then in Germany for use in “cool” double-base proplnts, “G” Pulvern, of Gen. U. Galliwitz intended for hot climates like N. Africa. It was also used as prop int in the “Nebelwerfer” (Ref 49, p 176 & Ref 70, pp 368-69)

WWII. H-6 is an HE consisting of RDX 45, TNT 30, Al 20 & D2 Wax 5% with 0.5% Ca chloride added (Ref 70, pp 146 to 148) (See also in this Vol of Encycl, p H1-L)

WWII. HMX (High Melting Explosive) is listed as cyclotrimethylene tetranitramine

WWII. Minol-2, Brit expl cong TNT 40, AN 40 & Al 20% developed for use in depth bombs (Ref 70, pp 209 to 212)

WWII. Tridevite. A castable mix of PA 88-90 & MNNaphthalene 12-10% developed by the British as an improvement over Trideite (PA 80 & DNPPhenol 20%). Used as TNT substitute in bombs and shells (Ref 70, pp 370 to 372)

WWII. PIPE is a mixture of PETN 81 and Gulf Crown E Oil (Ref 70, pp 294 to 295)

WWII. PTX-1 (Picatinny Ternary Explosive) is a eutectic mix of RDX 30, Tetryl 50 & TNT 20%. It was developed in 1943 at PicArtn and used in land mines and demolition charges (Ref 70, pp 306 to 308)

WWII. PTX-2 consisted of RDX 44-41, PETN 28-35 & TNT 28-35, developed in 1943 at PicArtn and used in shaped charges (Ref 70, pp 309 to 311)

WWII. PVA-4 (Polyvinyl Acetate) is a semi-plastic expl cong RDX 90, polyvinyl acetate 8 & DBuPh 2%. It was developed in Canada and used in demolition charges. (Ref 70, pp 312 to 315)

WWII. RIPE. Mixture of RDX 85 & Gulf Crown E Oil 15%; developed in the US as a plastic demolition expl (Ref 70, pp 318-19)

WWII. Hybrid Rocket Propellants are described in this Vol

WWII. PH-Salz (Ethylendiamine Dinitrate) was used by the Germans in mixts with AN and other ingredients for filling some shells (Ref 56, p Ger 131)

1945. This year there were completed the first three atomic (or nuclear) bombs. The first bomb was successfully exploded in the New Mexico desert on July 16, 1945, and a second over Hiroshima, Japan on Aug 6, 1945, and the third over Nagasaki on Aug 9, 1945. The bombs were of Fission Type and of tons of kilotons (thousands of tons of TNT equivalent) (Vol 1 of Encycl, p A499-L)

1945. PLX (Picatinny Liquid Explosive) consisting of Nitromethane 95 & Ethylene dinitramine 5% was developed at PicArtn by L.H. Eriksen & J.W. Rowen for use in minefield clearing (Ref 70, pp 298 to 301)

Note: Germans used during WWII similar liquid expls under the name of Myrol, which are described in PATR 2510 (1958), pp Ger 115 & Ger 116

1945. Tripeenthenylthiol Octanitrate was patented by J.A. Wyler of Trojan Powder Co, Allentown, Penna. It was used as an HE and as possible gelatinizer for NC (Ref 70, pp 382 to 384)

After WWII. Composition D-2 which is not an expl, but serves as an emulsifier and desensitizer of expls like HBX-1, consists of wax 84, lecithin 2 & NC 14% (Vol 3 of Encycl, p C488-R)

Compositions EL-387A and EL-387B. de-
veloped by the duPont Co., were slurry expls of compons given in Vol 3 of Encycl, pp C488-R & C489-L.

1948. The first Fusion Type Atomic Bomb (also known as Hydrogen Bomb, H—Bomb or Thermonuclear Bomb) was tested at Eniwetok and proved to be successful. The bomb was of several megatons (millions of tons of TNT equivalent) (Vol 1 of Encycl, p A499-L & R).

1950. MOX (Metal Oxidizer Explosives) were developed by National Northern, technical division of the National Fireworks Ordnance Corp, West Hanover, Massachusetts, for use mostly in small caliber antiaircraft shells. Their compons and props are described in Ref 70, pp 213 to 225.

1950. Trinitroethyl Trinitrobutyrate seems to have been first prepd at NavOrdLab and then at Hercules and US Rubber Labs. It is a high oxygen content expl (Ref 70, pp 375 to 377).

1951. Ethylene glycol Di(trinitrobutyrate) was prepd at the US Rubber Co Lab, Passaic, New Jersey (Ref 70, pp 133 to 135).

1952. Dynamite, Medium Velocity, Military was developed by W.R. Baldwin, Jr at Hercules Powder Co Lab (Ref 70, pp 125 to 127).

1952. 2,2-Dinitropropyl-4,4,4-trinitrobutyrate was prepd by M.E. Hill at NavOrdLab (Ref 70, pp 113 to 115).

1952. Bis-(2,2-Dinitropropyl)-fumarate was prepd by M.E. Hill at NavOrdLab and in 1954 by D.L. Koubal & H.D. McNiel of Hercules Powder Co (Ref 70, pp 107 to 109).

1952. PB—RDX (Plastic-Bonded RDX) was developed by Los Alamos Scientific Laboratory of the University of California for use as a mechanical strength expl. It consisted of RDX 90, poly- styrene 8.5 & diocetylphthalate 1.5% (Ref 70, pp 259 to 264).

1953. Bis-(2,2-Dinitropropyl)-succinate was prepd by M.E. Hill of NavOrdLab (Ref 70, pp 110 to 112).

1953. HEX (High Energy Explosives) developed at PicArsn are described in Ref 70, pp 164 to 169 and in this Vol, p H73-L.

1956. Pentacyrthritol Tetranitroacrylate or PETRIN Acrylate) was developed by Rohm and Haas Co, Redstone Arsenal Division (Ref 70, pp 269 to 271).

1957. Dynamite, Low Velocity, Military was developed at PicArsn by H.W. Voigt (Ref 70, pp 122 to 124).

1957. Veltex is the name given to a series of closely related NC compons prepd in 1957 at PicArsn by the solventless provess. They all contain a high percentage of solid HE. Hispano-Suiza Co investigated them to determine their suitability as Holtex type expls (See year 1958). Veltex expls are proposed for use as high mechanical strength machinable expls. Compn of Veltex No 448 is: NC (13.15% N) 15.0, HMX 70.0, NG 10.7, 2-NitroDPNA 1.3 & Triacetin 3.0% (Ref 70, pp 391 to 394).

1958. MASER and LASER. Acqg to "Time" magazine of July 12, 1968, pp 42—9, physicists A. Schawlon & C. Townes described in 1958 a device which was a variation of Townes earlier Nobel prize-winning invention named MASER. The first working model of this device was built in 1960 by physicist Maiman, who named it RUBY LASER. A brief description of these and later developed devices is given in Vol 4 of Encycl, pp D436 to D440, compiled by C.G. Dunkle, formerly of Picatinny Arsenal.

1958. HTA—3 (High Temperature Explosive), consisting of RDX 49, TNT 29 & Al 22%, was prepd at PicArsn by R. Brown & R. Velicky (Ref 70, pp 178 to 181). See also in this Vol.

1958. Octols 70/30 and 75/25 are mixts of HMX and TNT developed at Northern Corp as fillers for bombs and shells (Ref 70, pp 249 to 258).

1958. Holtex was patented by Dr E. von Hoit of Hispano-Suiza Co, who died in 1962 in an automobile accident. Holtex has several formulations which are listed in Vol 3 of Encycl, pp C396-L to C397-L.

Ref: Dr H. Freiwald of Saint Louis Laboratory, Germany; private communication, 12 Sept 1962
Note: See also in Vol 3 of Encycl, pp C396-L ff 1960. Detacead Process or Explosive Cladding, also called Explosive Bonding was developed by the duPont Co (See Vol 3 of Encycl, pp D96 & D97).

1960. Detacord, Detaflex Flexible Cord Explosive and Detashift Flexible Cord Explosives were developed by the duPont Co (See Vol 3 of Encycl, pp D97 to D101).

1967. Acqg to Nambo (Ref 66, p 39), the mining BkPdr of Nippon Kayaku Co, Ltd contd: saltpet 64, sulfur 18% & charcoal 18%, while its sporting BkPdr contd: saltpet 75.0, sulfur 12.5 & charcoal 12.5%.

1972. Acqg to Gorst (Ref 71, p 173), the
average compn of Russian Black Powder (called "Dymny Porochn", which means Smoke Powder) is saltpeter (mostly of K) 75, sulfur 10 & charcoal 15%

Refs for History of Explosives and Related Items:
1) M. Jähns, "Handbuch einer Geschichte des Kriegswesen von der Urzeit bis zur Renaissance", Leipzig (1880)
2) W.W. Greener, "The Gun and Its Development", Cassell, Petter & Co, 1st Edn, London (1881); 9th edn (1910) (See Ref 7)
2a) S.G. Romocki, "Geschichte der Explosivstoffe", Berlin & Hanover, Vol 1 (1895) & Vol 2 (1896)
2b) Lt J.B. Berndou, "Smokeless Powder, Nitrocellulose and Theory of Cellulose Molecules", J. Wiley, NY (1901) and 2nd Edn (1917)
5) Sir Andrew Noble, "Artillery and Explosives", Dutton Co, NY (1906)
8) P.F. Chalon, "Explosifs Modernes", Ch. Béranger, Paris (1911), 228
9) G.W. MacDonald, "Historical Papers on Modern Explosives", Whittaker, London (1912)
11) Marshall 1 (1917), 11-22 (Early History), 12-34 (Development of Gunpowder), 35-50 (Progress of Explosives in the Eighteenth and Nineteenth Centuries)
12) Capt E. de W.S. Colver, "High Explosives", Van Nostrand Co, NY (1918), pp 1-29 (History of High Explosives); 496-501 (History of HE Projectiles); 501-05 (History of Shell Fuzes); 563-64 (History of Fuses)
13) Anon, "History of Trench Warfare Materiel", Army Ordnance Pamphlet (1920) (Available at PicArsn Museum)
15) P. Charbonnier, MémArtIfran 7, 1227 (1927) (Artillery)
16) Ph. Naoum, "Nitroglycerine and Nitroglycerine Explosives" (Translated from Ger by E.M. Symmes), The Williams & Wilkins Co, Baltimore (1928), pp 1 to 22 (Discovery and Historical Review)
16a) H. Desvergnes, ArmyOrdn 10, 191-94 (1929) (History of development of military powders)
17) L. Vennin, E. Buret & H. Lecorché, "Les Poudres et Explosifs", Ch. Béranger, Paris (1932) (No history is described)
18) P. Charbonnier, MémArtIfran 11, 1er, 2ème, 3ème & 4ème Fascicules (1932) (Artillery)
20) Marshall 3 (1932), 1-2 (Early History); 3-5 (Development of Gunpowder); 6-8 (Progress of Explosives in the Eighteenth and Nineteenth Centuries)
21) A. Stettbacher, "Schiess- und Sprengstoffe", J.A. Barth, Leipzig (1933), pp 3-18 (Geschichtlicher Überblick)
22) T. Urbaski, MémArtIfran 13, 825-42 (1934) (Le centenaire de la nitrocellulose)
24) A. Basset, MémArtIfran 13, 529-78 (1934) (Bibliography on Artillery; more than 1000 refs are listed)
25) A. Basset, MémArtIfran 14, 881-1280 (1935) (Essais sur l’histoire des fabrications d’arme- ment en France)
(Geschichtliches)
28a) J. Reilly, “Explosives, Matches and Fireworks”, VanNostrand, NY (1938) (Describes various physical & expl tests)
29) V. Muthesius, “Zur Geschichte des Sprengstoffe und des Pulvers”, Hoppenstedt, Berlin (1941) [Reviewed by Dr Hassenstein in SS 37, 111–12 (1942)]
30) E.E. Sancho, “Química de los Explosivos”, A. Aguado, Madrid (1941), pp 263–61 (History of Black Powder and Ammunition & weapons using it)
33) J.R. Newman, “The Tools of War”, Double-day, Doran, NY (1943), pp 3–31 (Modern Warfare);
33) 33–46 (History of Small Arms); 69–108 (History of Field Artillery); 109–175 (History of Fortification and Siegecraft); 176–203 (History of Tanks); 204–67 (History of Tools of Sea War); 268–363 (History of Tools of Air War)
34) Gen U. Gallwitz, “Die Geschützladung” (Cannon Charge), Heereswaffenamt, Berlin (1944)
(Eng transnl available at PicArsn Library)
35) W. Hassenstein, SS 39, 1–9, 22–6 & 37–42 (1944) (The Chinese and the Discovery of Powder)
(Historical Review)
41) A. Stettbacher, “Spreng- und Schiesstoffe”, Rascher, Zürich (1948), 1–5 (Geschichtlicher Überblick)
& CA 47, 9014 (1953) (A Historic Development of Smokeless Powders) (The various stages are discussed that led to the discovery and formulation of smokeless powder. Accomplishments of various countries are given, with the more important dates up to the first World War)
45) LeRoux, RevIndustMinérale 33, 276–84 (1952) (History of development of expls Industry)
47) W. Gutmann, Sprengtechnik 3/4, p 39 (1952) (Chronological list of some expls, propyls and ammunition manufactured in Germany betw 1858 and 1937)
48) C. Belgrano, “Gli Esplosivi”, Hoepli, Milano (1952) (History of explosives and related items is not found) [See 2nd edition (1974)]
51) B.T. Fedoroff et al, “Dictionary of Russian Ammunition and Weapons”, PATR 2145 (1955); PB 159927, Available from NTIS
53) B.E. Schaar, “History of Accidental Scientific Discoveries”, Schaar & Co, Chicago 7, Ill (1955) (Discoveries of Dynamite, acetylene, petroleum jelly, etc are described)
54) Anon, “Principles of Artillery Weapons”, TM 9–3305–1 US Dept of Army, Washington,
DC (1956), 4–8 & 12–15
56) B.T. Fedoroff et al, “Dictionary of Explosives, Ammunition and Weapons” (German Section), PATR 2510 (1958); ASTIA Document AD 160636, Available from NTIS (National Technical Information Service), US Dept of Commerce, Springfield, Virginia 22151; pp Ger I & Ger II (Historical uses of expls in Germany during WWI and WWII)
57) M. Giua et al, “Trattato di Chimica Industriale”, UTET, Torino VI (1), 1959, 6–10 (Storia)
58) W.S. Dutton, “One Thousand Years of Explosives, from Wildfire to the H–Bomb”, Winston, Philadelphia (1960), pp indicated under the years in text
62) Coll, “Blasters’ Handbook”, Published by E.I. duPont de Nemours & Co (Inc), Wilmington, Del (1966), pp 1–6 (Early History of Explosives); 7–20 (DuPont’s Role Today) (See also Ref 68)
Gonnes, the earliest firearms, are listed on pp 29–33. (See also “Gonne” in Vol 6 of Encycl, p G120-R)
68) Coll, “Blasters’ Handbook”, Published by E.I. duPont & Co, Wilmington, Del (1969), pp 1–6 (Early History of Explosives); pp 7–24 (DuPont’s Role Today) (See also Ref 62)
71) A.G. Gorst, “Porokha i Vzryvchatyiye Veschehestva” (Powders and Explosive Substances), Izdat Mashinosstroeyntse, Moscow (1972), pp 5–17 (A brief historical description)

“Hive of Bees”: A Japanese pyrotechnic device of WWII, which consisted of a six by one inch cardboard (called Hive), mounted on a wooden block and provided with a fuse. The “Hive” was filled with miniature BkPdr charges (called “Bees”)

When ignited by a fuse the “Hive” injected a plume of smoke followed by some object shot at high speed and buzzed furiously away.
This display was followed by a machine-gun succession of "Bees", which may be called miniature unguided BkPdr rockets launched from a Roman candle


HMSO—Abbreviation for Her Majesty's Stationary Office (in London), which is the British Institution which corresponds to the US Government Printing Office (in Washington, DC)

HMT—See Hexamethylenetetramine in this Vol

HMTD—See Hexamethylenetriperoxidediamine in this Vol

HMX (High Melting Explosive or Her Majesty's Explosive). See CYCLOTETRAMETHYLENETETRANITRAMINE in Vol 3, pp C605-R to C610-R. It exists in four polymorphs of which beta-HMX is described on pp C606-R to C609-R and in AMCP 706-177 (1967), pp 173-77

HMX-Nylon Plastic Bonded Explosive. See Nylon-HMX Plastic Bonded Explosive

HNMI: Hexanitromannitol. See Mannitaol Hexanitrate in Vol 8

HNO. Abbrn for 2,4,6,2',4',6'-Hexanitroxanilide, described under Oxaanilide and Derivatives and in AMCP 706-177 (1967), pp 170-72

HNS. Abbrn for Hexanitrostilbene, also called Hexanitrodiphenylethylene. See Diphenylethylenehexanitro in Vol 5, p D1456-R

Hoch- und Niederuckkanone (High and Low Pressure Gun, abbreviated to H/L Gun) (Canon à tuére, in French). It has been known for a long time that the lower the peak pressure in a gun the thinner may be the walls of the projectile. This means that for a given total weight the projectile, used in a gun with lower peak pressure, can contain more explosive and do more damage to a target

This is of particular importance in the use of shaped charges because the penetration of targets does not depend upon the strength of the case (shell) but on the amount of the explosive charge. In order to achieve low pressure in a gun of conventional design, the barrel should be made longer and the chamber and cartridge case larger. Such guns were built but were found to be unsuitable because the propellant was difficult to ignite and it burned irregularly (due to the low pressure in the chamber). Also, the initial velocity of the projectile varied from round to round which means that no precision firing could be achieved

Better results were obtained in 1943 when Dr. Hermann and collaborators of the Rheinmetall-Borsig AG constructed the 8cm PWK 43 (80mm Antitank Gun). The description of this gun, called in French "canon antichar modèle 1943," was given by Travers and Touchard (Ref 3). They claimed that the "turbocanon Delamare-Maze," invented in France about 20 years earlier, may be considered as the predecessor of both the H/L and recoilless guns

The German gun, 8cm PWK 43, had a comparatively thin barrel with an inside diameter of 81mm and was 34 calibers long; the chamber had an enlarged diameter (105mm) and much thicker walls. The projectile (fintail type, 81mm in diameter, contained a shaped charge and weighed 3kg) was inserted first in the bore (as in separate-loading ammunition). This was followed by the cartridge (120mm long and 105 mm in diameter) which contained the propellant. The cartridge was closed by means of a disc provided with eight perforations (each 13mm in diameter). When the propellant burned the pressure of the gases developed inside the cartridge was about 850 kg/cm² but the pressure acting on the projectile was only 550 kg/cm² because the gases lost part of their velocity on passing through the holes in the disc

The relation between the high pressure inside the cartridge case and the lower pressure in the bore could be varied by increasing or decreasing the size or number of the openings in the separating disc. In order to protect the propellant in the container from spilling and from moisture, the perforated metallic disc was covered with a solid disc of paraffined cardboard

The ballistics for the H/L gun were worked out by Travers and Touchard in France and by
Corner in England

Note: Corner stated that towards the end of
WWII the Germans started to manufacture two
light antitank guns: the 8cm PAW 600 and the
10.5cm PAW 1000, but does not describe them.
He also mentioned the 8.8cm W71 gun, which
was built on the "three-pressure principle"
Refs: 1) J. Corner, JFranklinInst 246, 233 (1948)
2) J. Corner, "Theory of the Internal Ballistics
3) S. Travers & L. Touchard, MémArtilFr 26, 835-
58 (1952) 4) Ibid, 27, 219-36 & 245-78
(1953) 4) PATR 2510 (1958), pp Ger 90-R
& Ger 91-L

Hochdruckpumpe (HDP) oder V-3 (High Pressure
Pump, called also "Busy Lizzie" or "Münpede")
a German constant-pressure gun developed
during WW II by Condors, an engineer of the
firm Röchling, Saarbrücken, and intended to
fire the Arrow (Needle) Projectile (qv) across
the Channel to London. The barrel, caliber
150mm (5.9"), was of unalloyed crucible
cast steel made up of a great many Y-shaped
sections, each 12 to 16 ft long. With a gun
about 450 ft long containing about 28 prop-
ellant chambers (distributed along the bore),
a muzzle velocity of about 4500 ft/sec and
a range of about 130 km was expected to
be achieved (when using a projectile 8 ft long
and weighing 150 lb)

The gun could lie on the ground without
any carriage on wooden and concrete blocks
sloped at a 45° angle. The fin-stabilized, arrow
projectile was inserted in the barrel and the
base propellant charge electrically ignited. As
the projectile passed the separate Y-pieces,
additional propellant charges in the side arms
were electrically ignited one after another (in
pairs) thus accelerating the velocity of the
projectile as it progressed along the gun barrel

For servicing (reloading the Y-sections with
propellant charges between the rounds), the
gun required a great many soldiers. It was
planned to fire one round per gun every 5
minutes but this rate could not always be
achieved because the sections often exploded
and it was necessary to insert new Y-pieces
Refs: 1) L. E. Simon, German Research in
2) W. Dörnberger, "V-2," Viking, NY (1954),
3) A. I. Sprinz & H. H. Bullock of
Picatinny Arsenal; private communication
4) PATR 2510, p Ger 90-L & R

HDP Supergun
(Vergeltungswaffe 3)
(V-3)

Hochexplosivkörper. Ger for High Explosives
(HE's)

Hochstädtter Powder—patented in 1869 in
England was prep'd by blending in wet condition
KClO₃ (or Pb(Clo₃)2), KN0₃ (or NaNO₃),
charcoal and sulfur (or metallic sulfide) and
impregnating paper or dry vegetable materials
with this mixture
Refs: 1) Cundill (1889) in MP 6, 14 (1893)
2) Daniel (1902) 376 3) Giua, Trattato 6,
(1959), 392

Hohlladung (Ger for Shaped or Hollow Charge).
Considerable work was done in Germany before
and during WW II on the development of shaped charges. Among the most prominent contributors in this field were the personnel of Krümmel Fabrik, DAG. Among the shaped charge weapons developed at Krümmel may be mentioned:

a) Magnetic anti-tank shaped charge weighing 3 kg; blast penetration of armor was up to 250mm

b) Shaped charges for Faustpatrone, Panzerfaust, Panzerschreck, etc

Note: At Krümmel it was found that the best explosives for shaped charges were RDX-TNB and next, RDX-TNT mixtures. Substituting PETN for RDX led to a decrease in efficiency. The addition of aluminum powder was desirable but not in large quantity.

Krümmel was not the only place where work on shaped charges was conducted. Elsewhere the Germans developed a shaped charge shell which was shot from an 80mm mortar called "Panzerwurfkanone", and the warheads for several guided missiles.

Historical Discovery of the hollow (shaped) charge (HoC) effect is usually attributed to C. E. Munroe (USA) who described the effect in the AmerSci 36, 1888. It was claimed by H. Schardin that Max von Förster of Germany had in 1883 already shown that bare hollow charges gave an enhanced effect along the axis of the charge. The first practical application of the HoC effect for demolition charges, sea mines, torpedoes, projectiles etc, was patented in 1910 by E. Neumann & the Westfälisch-Anhaltische Sprengstoff AG (DRP Ann W36269). Neumann's work is described in SS 6, 356 (1911) and SS 9, 183 (1914). Important work on military applications of the HoC effect was done, prior to and during WW II, by H. Schardin et al in Berlin. Some work was also carried out by A. Stettbacher of Switzerland during this period.

Note: According to A. J. Dere, Ordnance Sergeant, October 1945, pp 3-13, hollow (shaped) charge ammunition was used by the Germans in many 75mm caliber weapons. There were at least four types of such projectiles: H1, H1/A, H1/B and H1/C. Most of these projectiles are briefly described in TM 9-1985-3 (1953). Some projectiles of calibers 88mm, 100mm, 105mm and 150mm also had shaped charges.

The following drawings represent some typical German hollow charges. (See next page)


Note: Gen description of Shaped Charge phenomenon is given in Vol 4, pp D442-R to D454-L under DETONATION, MUNROE-NEUMANN EFFECT AND LINED CAVITY EFFECT IN

Hoitsema's Stability Test: One of two interconnected U-tubes contains 1-2g of explosive, while the 2nd tube contains glass wool impregnated with a solution of 0.1g diphenylamine (DPhA) in 50ml of 5% sulfuric acid and 50ml of CP glycine. The first tube is heated for 15 minutes at 110° and while heating is continued a current of pure CO₂ is passed through the tubes. If the DPhA reagent turns blue, a new sample is taken and the test repeated at 10° lower and so on until a temp is reached at which the DPhA reagent no longer turns blue.

This test differs from Abel's test in using temperature of decomposition rather than time at constant temperature.

Ref: Daniel (1902), 679

Hoko. Ger abbrn for Hochkonzentriert (Highly Concentrated) Process for the manufacture of 98-99.5% nitric acid, developed during WW II, and used in several German plants. In this process, the concentration of the weak acid (50%) was effected by mixing it with liquid nitrogen tetroxide (N₂O₄) and adding the necessary extra oxygen under 50 atm pressure in an autoclave.

Description of this method as practiced by the IG Farbenind A-G subsidiary, the Wirtschaftliche Forschungsgesellschaft mbH (WIFO), Embrsen, Kr Lüneburg is given in the following BIOS Final Reports: 1232 (1947), pp 15-16 and 1442 (1947), pp 84-98

Hollings patented, in 1898 in England, the method of preparation of compressed charges of wet nitrocellulose in two stages: first by using a worm screw in a cylinder, provided with numerous small perforations, followed by hydraulic pressing. During the first operation air
and excess of water are driven off, while in the hydraulic press the NC assumes the desired shape, such as discs, blocks, etc
Ref: Daniel (1902), 376

Hollow Charge. See Hohlladung (Shaped Charge)

Hollow Charge Nose Attachment for AP Bombs.

SD 250

SAWDUST AND CEMENT

EXPLOSIVE

FUZE POCKET

Hollow Charge
Nose Attachment for SD 250

In order to permit greater penetrating power from low altitudes some German 250 kg AP bombs had a hollow charge (weighing about 4 kg) attached to the nose. This charge was detonated by its own nose fuze as soon as it hit the armor. The explosion of the HoC produced a hole in the armor (as deep as 7 cm) which permitted the AP bomb to enter inside the target. The AP bomb being provided with a short delay fuze did not explode until it was inside the target. In order to protect the bomb from premature detonation the space between the HoC and the nose of the bomb was filled with sawdust and cement

Holmgrens: An explosive used in Sweden between 1903 and 1906 for filling shells and claimed to be more suitable for that purpose than PA. No composition is given
Ref: D.H. Hone, SS 1, 109 (1906)

Holocellulose: Cellulosic material obtainable from wood after removal of lignin. The term therefore means total carbohydrates (cellulose acid hemicellulose) present in the wood (Refs 1 & 2)

Ritter (Ref 3) succeeded in isolating holocellulose from wood pulp by repeated chlorinations, followed by extraction with alcohol containing 3% of monoethanolamine. The resulting product was white but changed color on standing


Holtex. See Ref 7 thru Ref 11, pp C396-L to C397-L and Addnl Ref K, pp C402-R to C403-L in Vol 3 of Encycl

Holzmine 42. German Landmine developed during WW II
Ref: TM 9-1985-2 (1953), p 263

Homing Guidance Systems. A homing guidance system may be defined as a guidance system by which a missile steers itself toward a target by means of a self-contained mechanism which is activated by some distinguishing characteristic of the target. The homing guidance systems fall under one of the three general types: active homing, semiaactive homing, and passive homing. There are possible permutations and combinations of the three basic types, depending upon intended application. The basic homing types also may be employed in conjunction with other guidance techniques. For example, the control of a missile may involve several guidance phases each using a different guidance technique

An active homing guidance system is one in which both the source of energy to illuminate the target and the receiver of the energy reflected from the target are carried in the missile

The active homing guidance system, in its simplest form, consists of a transmitter and receiver of energy, which enable the missile to detect the presence of the target, a computer which predicts from the received energy the future position of the target, and missile control surfaces which respond to computed signals in such a fashion as to direct the missile to impact with the target. The energy used to illuminate the target may be in the form of radio, light, heat, or sound waves. A missile which uses active homing guidance is completely independent once homing starts; the missile does not require energy transmitted from an external source or externally derived guidance intelligence

A semiaactive homing guidance system is one wherein the receiver in the missile receives
energy reflected from the target, the energy having been transmitted from a point external to the missile. The semiactive homing guidance system consists of a receiver in the missile which detects the presence of the target, a computer (also in the missile) which predicts from the received energy the future position of the target, missile control surfaces which direct the missile along the correct flight path for impact with the target, and an externally located transmitter which illuminates the target. This external transmitter may be located at the missile launching station, or at a point separated from the missile launching station.

The transmitter may be a surface installation or it may be airborne. As in the case of active homing guidance, the transmitted energy may be in the form of radio, heat, or sound waves. The principal difference in the basic operation of an active and a semiactive guidance system is that the semiactive system is not independent of external sources. Its guidance intelligence is derived from energy transmitted from a point external to the missile.

A modified version of this semiactive technique has the transmitter located in the missile and the receiver of the reflected energy at some remote point. Computation of the desired flight path takes place at a remote point and suitable commands are sent to the missile. This system is known as a quasi-active homing guidance system.

A passive homing guidance system is a guidance system wherein the receiver in the missile utilizes energy emanating from the target. The basic difference between the passive homing technique and the two preceding techniques is that energy from which the guidance intelligence is derived in the passive homing system is generated in the target; thus no other transmitter is required. In this system, the receiver, computer, and missile control surfaces serve the same functions as described for the previous systems. The energy emanating from the target may be in the form of heat, light, sound, or radio waves.


Note: See also Guidance System of a Missile in Vol 8, p G175 to G178

Homocyclonite. One of the names for Cyclotetramethylene-tetrinitramine (HMX) described under Cyclotetramethylene-tetranitramine and Derivatives in Vol 3, p C605-R

Homogenization and Homogenizers (See also Emulsification and Emulsifiers). The term “homogenization,” according to Sloan (Ref 1), describes the process of putting incompatible or immiscible components into a stabilized, nearly homogenous suspension in a liquid medium. For instance, preparation of various cosmetic creams may be accomplished by homogenization of fats, tallow, and essential oils, coloring materials etc in water. Homogenization can produce particles of the order of 1 micron diameter, which are smaller than encountered in most natural emulsions, such as milk. As the smaller particles remain in suspension for a longer time than the larger ones, milk passed through a “homogenizer” (an apparatus used for “homogenization”) will not separate into cream and skimmed milk, particles of fat and casein are smaller than in ordinary milk, better digestibility is achieved.

Many types of homogenizers are on the market, but the most conventional types function by forcing the ingredients (at pressures of 500 to 3000 psi) past a spring-seated valve. Emulsification occurs not only while the components pass under the valve seat, but also when they impinge against the retaining wall that surrounds the valve. The high velocity combined with hydraulic shear, pressure release and impact, transforms the dispersed phase into a very fine state of subdivision.


Honest John. Popular name for the US Army 762mm rocket system, a surface-to-surface tactical missile, employing a solid propellant. It is launched from a rail-type launcher at an elevation which can be varied to obtain the desired point of impact. Honest John carries either a conventional HE warhead or a nuclear warhead.

Ref: OrdTechTerm (1962), p 156-L
Honey (Miel in French). A sweet, sticky substance made by the honey bee from the juices it collected from flowers. It can be nitrated to:

Nitrohoney (Nitromiel). Nitration is best conducted in mixtures of honey and glycerin, using the acids and procedure described under Nitroglycerin. The explosive properties of this mixed explosive are similar to those of Nitrohydrin (qv). “Nitrohoney” was patented for use in the so-called Thunder Powder (qv) Ref: Daniel (1902), p 564 under Nitromiel and p 767 under “Thunder Powder”

Hop Record of a Gun Carriage or Mount. The hop is defined as the motion of the carriage (or mount) from the instant of firing until equilibrium is attained. The “hop” record is a measure of the carriage (or mount) stability. The equipment and methods of testing are given in Ordnance Proof Manual No 40-78 (1943)

Hop-pao. A kind of incendiary charge resembling Greek fire, employed by the Mongols in the 13th century Ref: Daniel (1902) 377

Hope patented in 1884 in England—Black Powder in which part of charcoal is replaced by starch, flour, sugar, or other organic substances. Some bitumen, or other solid hydrocarbons was included in order to obtain more complete combustion Ref: Daniel (1902) 377

Hopkinson’s Pressure Bar Test. See under Physical Tests for Determining Explosive and Other Properties in Vol 1 of Encycl, p XVI

Horn-Seifert Stability Test. In the original test of Horn, 2g of smokeless proplnt was heated to 120° in a long test tube until brown fumes of nitrogen dioxide evolved. The degree of decomposition of the proplnt was judged approximately by the intensity of brown coloration as observed by looking down the length of the tube against a disc of white porcelain (Ref 2)

Seifert (Ref 1) modified the test by introducing a test paper (methyl violet or roseaniline) and observing the successive changes of color Refs: 1) Seifert, Voyenske-Technicke Zpravy (Prague), 4, 42 (1927) 2) Reilly (1938) p 79

Hornet Ball Cartridge. See under Cartridge, Ammunition in Vol 2, p C74-L

Horns. Metallic projections (lead, steel or copper) on top and/or sides of underwater mines, serving as initiators. Horns are of several types but the most common is the “chemial horn,” which contains glass vials filled with acid. When the ship hits one of these horns, it breaks the glass vial and the acid then flows between electrodes of a battery causing a current to be generated. The current fires an electric detonator, which causes detonation of the main charge of the mine Ref: Glossary of Ordn (1959), p 151-L; Ordn-TechTerm (1962), p 156-L

Horse Detonator. The name given to the American “M46” fuze because it can initiate much more effectively than standard fuzes, such as the Mark III. The “M46 fuze” can be used even with the worst exuding shell, where the standard fuzes nearly always fail. It compensates for both desensitized boosters and bursting charges. In all cases, it gives high-order detonations Ref: War Dept Tech Manual, TM9–1901 (1944), p 135

Horsedung Explosive. A blasting explosive composed of K nitrate 12 parts, horsedung dust 1, charcoal 1 & sulfur 3 parts Ref: Deutsche Pinkopst-Werke Karlsruhe, GerP 231598 (1907); CA 6, 2725 (1911)

Horsley Dynamites authorized in 1872 in England were prepdy by mixing at least 73 parts of NG with the following pre-pulverized mixtures: a) KClO₃ 75 & gallnut 25% (Ref 3) or b) KClO₃ 75, gallnut 12.5 & charcoal 12.5% (Ref 1)

According to Daniel (Ref 2), a mix of KClO₃ & NG, known as Seramine (qv) was proposed by Horsley. Another mixture, also known as Seramine was proposed by Björkman (Ref 2) Refs: 1) Cundill (1899) in MP 6 14 (1892) 2) Daniel (1902) 713 3) Gius, Trattato 6 (1959), 392

Hot Bar Test of High and Low Explosives. It is one of the tests used for the determination of ignition (deflagration or explosion) temper-
ature. The test originated in Spain and is described in Marshall (Ref 3) under the name "Temperature of Ignition." The apparatus shown in Fig 83, p 436 is reproduced here (qv)

![Image](image_url)

Fig 83. Apparatus for Determining the Temperature of Ignition

The copper bar (a) had a circular enlargement at one end in which there were five holes; the central one (F) contd mercury in which the bulb of a thermometer was immersed; a small copper tube, contg 0.1g of pulverized explosive to be tested, was placed in each of the other holes. The bar was supported on a stand and was heated by means of burners placed underneath it. The rate of heating could be regulated by moving the burners. The expl sample in tube (b) exploded first, then in (c) and (d) and finally (e). The mean of the thermometer readings, when explosion occured, was taken as the temperature of ignition. This method was used in Spain for testing NC propellants. If the ignition temp was 180° the sample was considered good; if it was between 178° & 180°, it was serviceable; if between 176° & 178° it had to be used immediately; and if below 176° the sample was destroyed.

**Note 1:** Instead of heating the bar by gas burners, it can be heated electrically, such as described by E. Berl & G. Rueff in Cellulosechemie 14, 43 (1933). The bar, known as *Bloc Maquenne*, generally used in France for determining melting points, could be adopted for use in determining the ignition (or explosion) point.

**Note 2:** Fisher Melting Point Apparatus, described by Fisher Scientific Co, Pittsburgh, Pa (1970), p 640, can also be used for determining ignition (or explosion) temperature.

**Note 3:** Hot bar and other methods for determining ignition (or explosion) temperature are described in the following refs:


**Hot Extrusion of Shells.** A modification of the Ugine-Sejournet hot extrusion process (using glass as lubricant) is used by Scaife Company of Oakmont, Pennsylvania. In this process a complete shell (such as 4.2 inch) can be produced in one piece from a simple billet. Important features of the development are in the substitution of readily available billet stock for seamless steel tubing, a critical material in times of war. Another feature of this process is that it requires about 25% less steel.

*Refs:* Ordnance, 38, 753 (1954) 2) Iron Age (April 1954) pp 98-102
Hot Spots

Introduction. A hot spot is a localized region of higher-than-average temperature. In explosion phenomena, the term "hot spot" is usually associated with the name of Bowden, although many other investigators have invoked the concept and contributed to its understanding.

There is a great deal of evidence that most explosive sensitivity phenomena are understandable in terms of the thermal decomposition of the explosive involved. Some of the early work on the interpretation of impact and friction sensitivity of explosives in terms of thermal effects is excellently summarized by Bowden & Yoffe (Ref 1). More recent studies which also include investigations of irradiation of explosives by nuclear particles and ionizing radiations are also reviewed by these authors (Ref 2), and by Bowden (Ref 7).

Cook (Ref 3) summarizes the adiabatic decomposition of explosives. Still more recently Bowden (Ref 7) reviewed energy localization effects in single crystals. (See Vol 4, pp D563-69). All these reviews as well as more studies (to be discussed below) emphasize the thermal nature of explosive initiation. Two simple examples will reveal that the thermal energy to produce initiation must be highly localized, ie unless the input energy, known to produce explosions, is concentrated into hot spots no explosions can occur.

1) In the B of M impact machine small samples of RDX explode at around 30 cm of a 2 kg weight. Assuming that all the drop energy goes into heating a 20 mg sample with no heat loss, then the maximum uniform temperature of the RDX is only ca 250°. At 250° the ignition lag for RDX is of the order of 1 sec, whereas it is found (Refs 1 & 3) that under impact measurement conditions the time lags are only several hundred microseconds or at least 1000-fold shorter. Clearly the impact energy must create hot spots that are at a considerably higher temp than 250°.

2) For PETN compacts of about 90% of crystal density, the 50% threshold for shock initiation is about 9 kbar. This threshold is very nearly the same for large diameter plane-wave tests (Ref 18) and for small diameter, divergent wave gap tests (Ref 17). From the Hugoniot data in Ref 17, the particle velocity, \( u_p \), corresponding to 9 kbar in the PETN is ca 0.30 mm/microsec. If all this Hugoniot energy (1/2 \( u_p^2 \)) goes into heat (and it certainly does not) the max uniform temp rise of the PETN is only ca 36° which is clearly insufficient to even start slow decomposition in the PETN, to say nothing of producing a detonation in the observed shock initiation times which are of the order of 1 microsec. Obviously the shock energy must be concentrated in hot spots.

Hot Spot Generation Mechanisms. Although there is general consensus on the existence of hot spots, there is far less agreement on their means of generation or the mechanisms by which they initiate explosions. Some of the possible modes of hot spot generation are:

a) an adiabatic heating of compressed gas spaces (Refs 1 & 2)

b) a frictional hot spot on the confining surface or on a grit particle (Ref 1 & 2)

c) intercrystalline friction of the explosive itself (Ref 1 & 2)

d) viscous heating of the explosive at high rates of shear (Ref 1)

e) heating of a sharp point where it is deformed plastically (Ref 2, p 76)

f) mutual reinforcement of relatively weak shock waves; probably at inhomogeneities in the shocked medium (Refs 7,8,10 & 16)

g) stagnation of particles spalled off a crystallite by the incoming shock and then stopped after flying across an air gap (the gap can be a void), by a neighboring crystallite (Ref 11)

h) Micro-Munro jets formed by shocking bubbles, cavities or voids whose interface (between solid or liquid and the cavity) is concave (Ref 7)

Modes a), b) & c) presumably operate in the usual impact and/or friction initiation of pressed solid explosives, and modes a) & b) in the impact and/or friction initiation of liquid explosives. Modes e), f), g) & h) operate in the shock initiation (and possibly the propagation) of detonation in solid explosive compacted or explosive liquids containing inhomogeneities. Mode d) is operative only at strong shock inputs and may be the main mode of initiation and propagation in homogeneous explosive liquids or defect-free explosive single crystals.

Theoretical Development. The foregoing shows
the complexity of the phenomena involved in generating hot spots. The mechanism by which hot spots initiate explosions is even more complicated since it involves:

1) dissipation of the externally applied energy leading to heat liberation in the materials during which the material is not heated uniformly, but at localized sites to form hot spots
2) thermal explosion of the hot spot
3) generation of a self-propagating process in the material, resulting in explosion

As full theoretical discussion of this problem involves enormous difficulties, of both physical and mathematical nature, the practice nowadays is to make use of calculations based on an idealized picture of the phenomenon. These idealizations are based on the following assumptions:

a) during hot spot formation an insignificant amount of chemical reaction takes place; i.e. the first and the second steps are clearly separated in time
b) the initial temperature profile in the hot spot is square wave
c) an exothermic first or zeroth order reaction not accompanied by any phase transformation or other physicochemical processes occurs in the hot spot and the medium

The real picture is naturally much more complicated. Some of the complications arising from the requirements of energy transfer from inhomogeneities to the explosive will be discussed in a later section.

The basic equation governing hot spot phenomena is:

\[ cp \frac{\partial T}{\partial t} = Q_k \exp\left(-\frac{E}{RT}\right) \]

\[ + \lambda \left( \frac{\partial^2 T}{\partial x^2} + \frac{n}{x} \frac{\partial T}{\partial x} \right) \]  \hspace{1cm} \text{[1]} \hspace{1cm} \text{(Eq 1)}

where

\[ 0 < x < \infty \text{ and } t \geq 0 \]

with conditions:

\[ t=0 \begin{cases} T=T_0, & x < r \\ T=T_1, & x > r \end{cases} \hspace{1cm} T_0 > T_1 \]

\[ t \geq 0, \hspace{0.1cm} x=0, \hspace{0.1cm} \frac{\partial T}{\partial x} = 0 \hspace{0.1cm} \hspace{0.1cm} x=\infty, \hspace{0.1cm} \frac{\partial T}{\partial x} = 0 \]

Symbols:

- \( T \) Absolute temperature
- \( x \) Space coordinate
- \( t \) Time
- \( T_0 \) Initial hot spot temperature
- \( T_1 \) Medium temperature
- \( r \) Hot spot radius
- \( Q \) Heat of reaction
- \( E \) Activation energy
- \( k \) Pre-exponential factor
- \( c \) Specific heat
- \( \lambda \) Thermal conductivity coefficient
- \( \rho \) Density
- \( n \) Hot spot symmetry factor

\( n=0 \) planar hot spot
\( n=1 \) cylindrical hot spot
\( n=2 \) spherical hot spot

The exact solution of Eq 1 is unknown. Only numerical or approximate analytical solutions are available. Solutions are obtained in terms of dimensionless variables and parameters, eg:

Variables:

\[ \theta = \frac{E}{RT^*_0} (T-T_0); \hspace{0.1cm} \xi = \frac{x}{r} \]

\[ \tau = \frac{t Q}{c \rho E} \frac{E}{RT^*_0} \kappa k \exp\left(-\frac{E}{RT_0}\right) \]

Parameters:

\[ \delta = \frac{Q}{\lambda} \frac{E}{RT^*_0} \exp\left(-\frac{E}{RT_0}\right); \hspace{0.1cm} \beta = \frac{RT^*_0}{E} \]

by Merzhanov (Ref 14) and

\[ \theta' = \frac{RT}{E}; \hspace{0.1cm} \xi' = \frac{x}{r} \left( \frac{RQk}{\lambda E} \right)^{1/2}; \hspace{0.1cm} \tau' = \frac{RQk}{c \rho E} \]

Parameters:

\[ \theta'_0 = \frac{RT^*_0}{E}; \hspace{0.1cm} \theta'_1 = \frac{RT^*_1}{E}; \hspace{0.1cm} \xi'_0 = \frac{x}{r} \left( \frac{RQk}{\lambda E} \right)^{1/2} \]

by Friedman (Ref 15)

In terms of the first (Merzhanov) set of dimensionless variables Eq 1 takes the form:

\[ \frac{\partial \theta}{\partial \tau} = \exp \frac{\theta}{\theta_0} + \frac{1}{\delta} \left( \frac{\partial^2 \theta}{\partial \xi^2} + \frac{n}{\xi} \frac{\partial \theta}{\partial \xi} \right) \]

\[ \tau = 0 \begin{cases} \theta = 0 & \xi < 1 \\ \theta = -\theta_0, & \xi > 1 \end{cases} \]

\[ \tau \geq 0 \hspace{0.1cm} \xi = 0 \hspace{0.1cm} \frac{\partial \theta}{\partial \xi} = 0 \hspace{0.1cm} \xi = \infty \hspace{0.1cm} \frac{\partial \theta}{\partial \xi} = 0 \]

(Eqs 4)
In Merzhanov's treatment the hot spot problem, as well as ordinary problems of thermal explosion, involves a critical value of the parameter δ which is a function of parameter $\theta_0$, namely

$$\delta_0 = f_0(\theta_0)$$  \hspace{1cm} (Eq 5)

Merzhanov states that the results of numerical calculations for spherical, cylindrical and planar hot spots fit the single empirical formula:

$$\delta_s = d_s (\ln \delta_s)^{m_s}$$

where $d_n$ & $m_n$ are (for $4 < \theta_0 < 25$):

- $n=0$$\quad d_0 = 2.66 \quad m_0 = 1.3$
- $n=1$$\quad d_1 = 7.39 \quad m_1 = 0.83$
- $n=2$$\quad d_2 = 12.1 \quad m_2 = 0.6$

Numerical solutions can be invalid outside the computational range for which they are made, and it is frequently cumbersome to use them in establishing the functional dependence of the solution on the variables of the process. Thus even approximate analytical solutions are often more instructive than the more accurate numerical solutions. However considerable caution must be used in this approach, since some of the approximations, employed to make the equations tractable, can lead to erroneous answers. A number of approximate solution for the hot spot system (Eq 1) are reviewed by Merzhanov and their shortcomings are pointed out (Ref 14). More recently, Friedman (Ref 15) has developed approximate analytical solutions for a planar (semi-infinite slab) hot spot. These were discussed in Sec 4 of Heat Effects on p H39-R of this Vol. To compare Friedman’s approximate solutions with the “exact” numerical solution of Merzhanov we computed $r_*$, the hot spot half-width, of a planar hot spot by both methods using the same thermal & kinetic parameters in both calculations. Over a wide range of input variables, the numerical solution gives values of $r_*$ which are 33 to 43% greater than the $r_*$'s of the approximate solution. Thus it appears that the approximate solution, from which the effect of the process variables are much easier to discern than from the numerical solution, gives answers that differ from the exact numerical solution by a nearly constant factor.

Other methods of making hot spot calculations are reviewed in Chap 10 of Ref 18. This Ref gives generalized, non-dimensioned curves for estimating hot spot temperatures and dimensions.

As an example of the usefulness of approximate solutions of Eq 1, let us consider how to compute the explosion time of a planar hot spot whose temperature and radius are given. A rough, but as will be shown, not too bad an estimate is obtained from solutions of the adiabatic explosion time equation:

$$r_*^2 = (\theta_0^2)^2 \exp(1/\theta_0^2)/a_c^2$$  \hspace{1cm} (Eq 6)

where $a_c^2 = \rho R Q k_r r^2 / \lambda E$, and the dimensionless $r_*$ is then converted into actual time via Eq 3. More accurately the explosion time, $r^*$, can be obtained from Friedman's results (Ref 15) namely:

$$\exp(-1/4r^*) = \pi^{1/2} (\theta_0^2 - \delta^2)/2(\theta_0^2 - \theta_1^2) \left(r^*ight)^{1/2}$$  \hspace{1cm} (Eq 7)

and

$$2(\theta_0^2 - 1.2 r^*) = -\delta(2\theta_0^2 + 1) + 2r^*$$

where the average explosion time $\bar{r}^*$ is taken to be 0.14. The following tabulation shows that Eq 7 underestimates $r^*$, but its values of $r^*$ that are closer to the “exact” (numerical solution of Eq 1) at small $\theta_0$ than those obtained from Eq 6. At large $\theta_0$ the $r^*$ from Eq 6 & 7 are indistinguishable and a little smaller than the exact $r^*$

$$\begin{array}{cccccccc}
\theta_0 & 0.03 & 0.04 & 0.06 & 0.08 & 0.10 \\
\theta_1^2 & 0.02 & 0.03 & 0.03 & 0.03 & 0.03 \\
\alpha_c^2 & 2.28 \times 10^{12} & 6.00 \times 10^8 & 4.36 \times 10^8 & 1.20 \times 10^8 & 1.73 \times 10^8 \\
r^* ("exact") & 0.114 & 0.179 & 0.141 & 0.150 & 0.138 \\
r^* (Eq 7) & 0.107 & 0.161 & 0.128 & 0.143 & 0.124 \\
r^* (Eq 6) & 0.150 & 0.200 & 0.143 & 0.144 & 0.124 \\
\end{array}$$

In contrast, the “exact” treatment of Merzhanov (Ref 14) does not give even approximate explosion times unless Eqs 4 are solved numerically.

Heat flow from a hot spot. One of the earliest recognized manifestations of hot spots was that of entrapped air or gas bubbles initiating explosion in an impacted liquid explosive (Ref 1). Originally Bowden's school suggested that heat from the adiabatic compression of such gas bubbles initiated explosion in the surrounding liquid. Johansson & coworkers (Ref 4), however, pointed out that heat flow from a compressed gas bubble to the surrounding liquid is much too slow to account for the observed phenomena, particularly at low impact energies. They have shown that to achieve explosion fine droplets
of liquid or the creation of foam are necessary within the compressed gas bubble. They also point out that too large a mass of droplets or foam within the bubble can result in too low a temperature of this mass. This immediately suggests a critical condition for explosion, namely, enough hot material within the compressed bubble to ignite the surrounding explosive after this material undergoes thermal explosion, but not so much material that its temperature rise (due to heat flow from the compressed gas) is too low for its thermal explosion. This criticality may account for the wide spread of results usually observed in the impact testing of explosives.

Hydrodynamic hot spots. For initiation via impact, characteristic times are of the order of several hundred microseconds and explosion usually starts with a deflagration which can turn into a detonation (Refs 1 & 5). For this process the heat flow described above appears to be adequate. However, for initiation of detonation, in granular explosive compacts or explosive liquids containing inhomogeneities, via shock the characteristic times are entirely too short for any appreciable heat flow from the hot spot to the surrounding explosive (Ref 6). Energy transfer from the hot spot to the surroundings is therefore presumed to occur entirely by shock and rarefaction waves. Mader (Ref 7) called this hot spot model the hydrodynamic hot spot and the development of this model is largely due to his work. In Ref 7 Mader considered the shock initiation of nitromethane containing inhomogeneities such as voids or grit particles. To quote Mader:

"In these calculations, the hot spots were idealized in initial geometry, since no method now exists for calculating the reactive dynamics of the actual two-dimensional configuration. In order to retain one-dimensionality in the computations, the hot spot was assumed to be a spherical disturbed region whose density and temperature was uniform throughout. The surrounding shocked nitromethane was likewise uniform in density and temperature. Initially, the fluid was everywhere at rest relative to the center of the hot spot, which was taken as the origin of the space coordinate.

Two types of hot spots were considered as representing the extreme possibilities. These are the "temperature hot spot," in which the density of the hot spot was the same as that of the surrounding nitromethane and the temperature was higher; and the "pressure hot spot," in which the density was also increased. In the pressure hot spot, the density and temperature chosen were values on the single shock Hugoniot. In both cases, the temperature was chosen such that decomposition of the hot spot would occur almost immediately relative to the surrounding nitromethane."

Based on these calculations, his conclusions were:

"The hydrodynamic hot spot is successful as a model for computing the critical sizes of hot spots in shocked nitromethane. The energy transfer in a hydrodynamic hot spot is effected by shock and rarefaction waves. The computed nitromethane hot spots, of reasonable size to result from interactions of a shock with a bubble, may initiate propagating detonation or not within the experimentally observed times of the order of 0.1 μsec. A hydro-dynamic hot spot may fail to explode if the rarefaction reaches the center of the hot spot before it can adiabatically explode. If the hot spot explodes, it sends a shock wave into the undetonated explosive, which heats the explosive. Whether or not it will initiate propagating detonation depends upon the initial strength of the shock wave and how rapidly the shock strength decreases at the interface."

More recently Mader developed computational procedures that enables him to examine 2-dimensional hot spots (Ref 10). We quote his conclusion of that study:

"The basic processes in the shock initiation of inhomogeneous explosives have been investigated theoretically using the model of a cylinder of nitromethane containing a void or an aluminum pellet. The interaction of a shock with the density discontinuities, the resulting formation of a hot spot, and the buildup to propagating detonation were computed using two-dimensional numerical hydro-dynamics of the "PIC" type with chemical reaction and accurate equations of state. The hot spots formed at aluminum pellets exhibit failure or propagation of detonation in approximately the same manner as the one-dimensional, hydrodynamic hot spots studied previously. The failure of hot spots formed at voids could be studied only in those cases in
which the failure mechanism did not depend on details of the structure of the reaction zone, as
this structure could not be reproduced in the calculation.”

Mader then reprogrammed his computations
for an Eulerian code and considered the interactions of 4 cylindrical voids rather than a single
void (Ref 16). He showed that shock interactions
with four holes lead to much greater & faster
computed nitromethane decomposition than the
shock interaction with a single hole for the same
initial conditions

He concludes:
“The basic two-dimensional processes involved
in the shock initiation of heterogeneous explo-
sives have now been numerically described.
The problem that remains is the study of the
interaction of a shock with a matrix of holes in
three-dimensional geometry. The basic twodimensional processes involved in the failure of
detonation, the failure diameter of explosives,
and the “spattering” initiation observed for
density discontinuities near the critical size have
been described. The three-dimensional study of
the interaction of numerous failures and re-
ignited detonations which is necessary for a
complete numerical description of these problems
must await new computing hardware.”

A novel method for estimating hot spot
temperatures. An interesting approach towards
estimating the hot spot temp during an impact
test has been presented by Bobolev & Bolkhov-
tinov (Ref 5). They suggest that in an impact
test in which the test sample is unconfined
some of the sample melts because of plastic
def ormation and the whole sample then behaves
as a very viscous liquid.* After considering the
heat flow in this system, and taking into account
that the melting point of the sample is raised
because the pressure increases during impact,
they assume a linear increase with a coefficient
a=0.02⁰/atm), they arrive at the following eqn:

\[ \frac{1 - \xi^*}{\xi^*} \frac{3\tau^* u^* (T^* - T_{\text{melt}})}{2ep_{gh}(T^* - T_0)\tau^*} \]

*Note: According to R. E. Winter of Cavendish
Labs, Cambridge: “Flat faces of Silver & Lead
Azide crystals were initiated when impacted with
spherical metal particles, small compared with the
sizes of crystals and above a critical impact
energy. Particles impacting below the critical
impact energy produced plastic indentations in
the crystals. On the basis of these results, it was
concluded that explosives can be initiated as a result
of rapid plastic flow. It is believed that in many
cases of impact and shock initiation plastic flow
can allow the localization of impact energy re-
quired for hot-spot formation” (Ref 20)
where \( \xi^* = \tau^*/\tau_0 \), \( \tau^* \) = distance from center of the
anvil to the spot where explosion starts (deter-
mined from the burning trace remaining on the
anvil after an impact that produces explosion);
\( \tau^* \) is the observed time from impact to explosion;
\( h \) is the test sample thickness; \( c \) & \( \rho \) have their
usual meaning; \( u^* \) is the critical impact velocity
to produce explosion; \( T^* \) is the hot spot tem-
perature. They obtain the following results with a
10 kg falling weight:

<table>
<thead>
<tr>
<th>Explosive</th>
<th>( u^* ) cm/sec</th>
<th>( \tau^* ) μsec</th>
<th>( \xi^* )</th>
<th>( T^* ) °C</th>
<th>( \tau_{\text{ad}} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>PETN</td>
<td>240</td>
<td>300</td>
<td>0.15</td>
<td>320</td>
<td>214</td>
</tr>
<tr>
<td>RDX</td>
<td>310</td>
<td>210</td>
<td>0.22</td>
<td>470</td>
<td>15</td>
</tr>
<tr>
<td>HMX</td>
<td>310</td>
<td>150</td>
<td>0.26</td>
<td>500</td>
<td></td>
</tr>
</tbody>
</table>

We have added the last column, \( \tau_{\text{ad}} \), the time
lag for an adiabatic thermal explosion at \( T^* \). In
our calculation \( T^* \) is the temp of a hot spot in
the condensed explosive. We used \( E=29.2 \text{ kcal/}
mole & \( k_e=1.2\times10^{12}\text{sec}^{-1} \) for PETN (J. Roth,
Addendum to Bulletin of 6th Army-Navy Solid
Propellant Group, p 41, I950) and \( E=41 \text{ kcal/}
mole & \( k_e=3\times10^{14}\text{sec}^{-1} \) for RDX (Ref 3). It is
clear that \( \tau^* \approx \tau_{\text{ad}} \) only for PETN. Why this is
so is uncertain at present. However the Bobolev &
Bolkhovitinov approach merits further study

As pointed out by Johansson (Ref 4) the
temperature of a compressed gas bubble within
a condensed explosive is usually much higher
than \( T^* \). This is emphasized by the findings
of Friedman (Ref 13) who used reflected shocks
in a shock tube to ignite explosive dusts and
sprays. Friedman claims that his results can
be interpreted to mean that bubble hot spot
 temperatures hot enough to ignite explosives
are between 500 & 800°C

A novel hot spot mechanism. Maycock and
Crabenstein have observed a piezoelectric effect in
single crystal HMX (Ref 12). This effect increases
linearly with static load up their max observed field
strength of 10V/cm at a load of 400g on a sample
area of 0.42cm². With the extreme assumption that the piezoelectric effect continues to increase linearly to much higher pressures, and that shock loading, which constitutes dynamic loading, is no different from static loading, fields of 10⁵ to 10⁶ v/cm might be generated at shock loadings of 10 to 100 kbars which are pressures typical of shock initiation processes. At such high field strengths electric breakdown and electron avalanche might be expected. This would create hot spots along the breakdown path. It is known (Ref 2) that Ag Azide breaks down and explodes at fields of 100 to 1000 v/cm, but breakdown strengths of secondary explosives, such as HMX, have not been measured.

Use of high energy radiation to create hot spots. Attempts have been made to initiate explosives by ionizing radiation such α-particles, high speed electrons, γ-rays, Pions etc (Refs 8 & 9). No initiations were observed. Cerny & Kaufman (Ref 9) take this absence of initiation to indicate failure of the hot spot model. However a crude preliminary calculation, based on the Friedman model (Ref 15), suggests that the dimensions of the Pion heated regions for Lead Azide (Fig 2 of Ref 9) and for RDX (Fig 3 of Ref 9) are smaller than the critical hot spot dimension at the corresponding temperatures.

Written by J. ROTH


Howard Powder. Same as Mercuric Fulminate

Howden Dynamite. A mixture compounded in 1870 in San Francisco by J. Howden and consisting of: NG 75 absorbed in a mixture of sugar, magnesium carbonate and potassium nitrate. This Dynamite was stronger and better than Nobel's Kieselguhr Dynamite. Ref: Blasters' Handbook (1952), p 5; (1958) p 5

Howittite. A mixt of PA, K chlorate and Na nitrate, which proved to be very sensitive and unstable. It was not authorized in England. Ref: 1) Cundill (1889) in MP 6, 15 (1893) 2) Daniel (1902), 378

Howitzer. See under Cannon, Gun, Howitzer, Mortar, Rocket and Trench mortar in Vol 2, p C27-L

HOX. Code name for Di- or Bis (2,2,2-Trinitro-ethyl) nitramine (BTNEN) described under Diethylamine and Derivatives in Vol 5 of Encycl, pp D1224-R to D1225-L

HS. Chemical warfare symbol (CWS) for "Mustard Gas". See under H in Vol 2, p C168-L

Hs (Henschel) Missiles. The following types were developed in Germany before and during WW II:

Hs117 (Henschel 117), also known as Schmutterling (Butterfly), was a rocket-propelled, radio-controlled missile for use against bomber formations. Some versions were for ground-to-air and some air-to-air. It used liq fuel called Tonka and an oxygen carrier called Salbei (qv) (Ref 1, pp 196–201 & Ref 2, p Ger 93-L)
Hs 293 (Henschel 293) was a radio-controlled missile released and directed to the target from an aircraft. The model fully developed and used was the Hs 293 A-1. Other models such as Hs 293 A-2, Hs 293 B, Hs 293 C, Hs 293 D, etc were not fully developed (Ref 1, pp 203-03 & Ref 2, p Ger 91-L)

Hs 298 (Henschel 298) was a rocket-propelled, radio-controlled missile designed primarily as an air-to-air weapon to be carried on fighter aircraft as well as the bomber types. There were several versions but the basic type was called Hs 298 V-2. It used a solid propellant (Ref 1, pp 203-05 & Ref 2, p Ger 91-L)

Note 1: Salbei was a code name for either 99.5% nitric acid or its mixt with 5-10% concd sulfuric acid, added to suppress corrosion (Ref 1, pp 216 & 231 & Ref 2, p Ger 170-R)

Note 2: Tonka was a liquid rocket fuel, such as a mixt of crude m-xylidine 57 and Triethylamine 43%(Ref 1, p 216 & Ref 2, p Ger 199-R)

Refs: 1) TM 9-1985-2 (1953) 2) PATR 2510 (1958)

HSC (Propellant). Same as Cordite HSC. See under CORDITE in Vol 3 of Encycl, p C535-L

HTA. Abbr for German mixture consisting of Hexogen (RDX), Trotyl (TNT) and Al (aluminum). One of such mixts contd RDX 40, TNT 40 and Al 20%

Refs: 1) G. Römöer, “Reports on Explosives (Germany)”, PBL Rept 85160 (1946), p 15 2) PATR 2510 (1958), p Ger 93-R

HTA—3 (USA). Designation of expl mixt consisting of HMX 49, TNT 29 & Al 22% (Type I); HTA—3, Type II contains HMX 49, TNT 28.65, Al 22 & Ca silicate 0.35%; mw 91, OB to CO₂—21%; cast, sp gr 1.90; Brisance by Sand Test 61.3g crushed in 200g Bomb; Detonation Rate for cast unconfined 1.0 inch diam chge 7866 m/sec; Explosion Temperature ca 370°C (flames erratically at 5 secs); Friction Pendulum Test—unaffected by fiber or steel shoe; Gas Volume Evolved on Explosion 680cc/g; Heat of Combustion 3687cal/g; Heat of Explosion 1190cal/g; Heat of Formation, no value reported; Heat, Specific 0.245cal/g°C; Impact Sensitivity, 2kg Wt: But Mines app for 20mg sample, no value reported; PicArsn App for 25mg sample 17 inches; Initiation, Sensitivity to—minimum detonating chge is 0.30g LA; Power by Ballistic Mortar 120% TNT; Rifle Bullet Impact Test in 3/16” steel—90% expls and 10% burnings; Ditto in 1/8” Al—50% expls and 50% unaffected; Sand Test—See Brisance by Sand Test; Sensitivity to Impact—See Impact Sensitivity; Sensitivity to Initiation—See Initiation, Sensitivity to; Stability Test—See Vacuum Stability Test; Vacuum Stability Test at 120°-0.37°C of gas evolved in 40 hrs from 1 g sample; Velocity of Detonation—See Detonation Rate; Viscosity, Efflux—24.8 Saybolt Seconds

Method of manuf of HTA—3 is similar to that used for Torpex, namely: purified TNT is melted by heating it to ca 100° in a steam-jacketed kettle, equipped with a stirrer. Water wet HMX is added slowly to molten TNT, while continuing to stir it and to heat, until all the water is evapd. Aluminum powder is added and the mixture continued to be stirred until uniform and then cooled, with stirring, to obtain a slurry suitable for pouring into shells or bombs to serve as their bursting charges


HTA—3, Analytical Procedures. Acdg to Pristera (Ref), up to 0.35% of Ca silicate may be used as an additive. The analysis of HTA—3 may be conducted as follows: TNT is extracted with CCl₄; HMX is extracted with acetone; the residue is Al. If Ca silicate is present, the residue is treated with 10% NaOH soin and washed with water, the remainder is Ca silicide


1. SCOPE
1.1 This specification covers an explosive com-
position having two types (Type I and Type II) designated as HTA-3 for use in the loading of warheads and other ammunition items.

1.2 Classification — Explosive Composition HTA-3 shall be of the following types as specified:
   Type I — See Table II and 3.3 to 3.5
   Type II — See Table II and 3.3 to 3.5

3. REQUIREMENTS

3.1 Material — The raw materials used in the manufacture of the composition shall be in accordance with the applicable specification in Table I

<table>
<thead>
<tr>
<th>TABLE I</th>
</tr>
</thead>
<tbody>
<tr>
<td>Material</td>
</tr>
<tr>
<td>----------</td>
</tr>
<tr>
<td>Octol</td>
</tr>
<tr>
<td>HMX</td>
</tr>
<tr>
<td>TNT</td>
</tr>
<tr>
<td>Aluminum</td>
</tr>
<tr>
<td>Ca Silicate</td>
</tr>
</tbody>
</table>

3.2 Composition — The composition of HTA-3, Type I and Type II shall conform to the applicable chemical requirements specified in Table II when tested as specified in the applicable paragraphs

<table>
<thead>
<tr>
<th>TABLE II</th>
</tr>
</thead>
<tbody>
<tr>
<td>Material</td>
</tr>
<tr>
<td>----------</td>
</tr>
<tr>
<td>TNT</td>
</tr>
<tr>
<td>HMX</td>
</tr>
<tr>
<td>Aluminum</td>
</tr>
<tr>
<td>Ca Silicate</td>
</tr>
</tbody>
</table>

3.3 Moisture content — The moisture content of Composition HTA-3, Type I and Type II shall be 0.10 percent maximum, when tested as specified in 4.3.2

3.4 Viscosity — The viscosity of Composition HTA-3, Type I and Type II, shall be 15 efflux seconds, maximum, when tested as specified in 4.3.3. See 6.6

3.5 Density — The density (specific gravity) of HTA-3, Type I and Type II shall be 1.85 grams per cubic centimeter minimum when tested as specified in 4.3.4. See 6.6

4. QUALITY ASSURANCE PROVISIONS. See in Specification

4.3 TEST METHODS AND PROCEDURES

4.3.1 Composition

4.3.1.1 TNT – Code No 04001 — Place an accurately weighed portion of approx 2.5g in a 200 ml beaker and add 75ml of benzene saturated with HMX. Cover the beaker with a watch glass and place beaker and contents on a steam bath. Break up the lumps with a glass stirring rod and agitate the solution occasionally by swirling. Remove the beaker and contents from the steam bath when all of the TNT has dissolved, as evidenced by the settling out of the other ingredients. Cool to RT and filter the solution quantitatively thru a tared medium porosity filtering crucible. Transfer any of the insoluble matter remaining in the beaker into the crucible by using an additional 150ml of benzene saturated with HMX. Draw air thru the crucible until the odor of benzene is no longer detectable. Dry the crucible and contents in an oven maintained at 100 ±5°C for one hour. Cool the crucible and contents in a desiccator and weigh. Retain the crucible and contents for the determinations which follow

Calculate the TNT content of the sample as follows:

\[
\text{Percent TNT} = \frac{100(W-A)}{W(MW)}
\]

Where:  
A = Weight of residue in crucible, g
W = Weight of sample, g
M = Percent moisture, expressed as a decimal (4.3.2)

4.3.1.2 HMX — Code No 05001 — Place the crucible and contents retained from the TNT determination on a filtering apparatus and wash with ten 20ml portions of acetone. Allow each portion of the acetone to remain in contact with the residue in the crucible for one minute before applying suction. Aspirate the crucible and contents until the odor of acetone is no longer detectable. Dry the crucible and contents in an oven maintained at 100 ±5°C for one hour. Cool in a desiccator and weigh. Wash the crucible and contents with an additional 20ml portion of acetone, and dry and weigh again. Repeat the washings and weighings until constant weight is obtained. Retain the crucible and contents for the determinations which follow

Calculate the HMX content of the sample as follows:

\[
\text{Percent HMX} = \frac{100(A-B)}{W(MW)}
\]

Where:  
A = Weight of residue in crucible retained from 4.3.1.1, g
B = Weight of residue in crucible, g
W = Weight of sample, g
M = Percent moisture, expressed as a decimal (4.3.2)

4.3.1.3 Calcium Silicate — Code No 06001 (applicable to Type II only) — Place the crucible and contents retained from the HMX determination on a filtering apparatus and wash with ten 10ml portions of the NaOH to remain in contact with the residue in the crucible for one minute before applying suction. Caution should be exercised when performing this procedure to preclude a too vigorous reaction between the aluminum and the NaOH. Aspirate the crucible and contents for a minimum of five minutes. Wash the crucible and contents with five 20ml portions of distilled water. Dry the crucible and contents in an oven maintained at 100±5°C for one hour or until constant weight is obtained. Cool in a desiccator and weigh.

Calculate the Calcium Silicate content of the sample as follows:

\[ \text{Percent Ca Silicate} = \frac{100C}{W-(100-MW)} \]

Where:  
C = Weight of residue in crucible, g
W = Weight of sample, g
M = Percent moisture, expressed as a decimal (4.3.2)

4.3.1.4 Aluminum — Code No 07001
Calculate the aluminum content as follows:

Percent Aluminum = 100—(A+B+C)
Where:
A = Percent TNT (4.3.1.1)
B = Percent HMX (4.3.1.2)
C = Percent Calcium Silicate (4.3.1.3)

4.3.2 Determination of moisture content

4.3.2.1 Special solvent — The special solvent shall be equal volumes of anhydrous methanol and benzene thoroughly mixed. If necessary the solvents shall be dried by distillation.

4.3.2.2 Method — Determine the moisture content in accordance with Method No 101.4 which is described in Specification MIL-STD-650, using solvent indicated in 4.3.2.1

4.3.3 Viscosity — Determine the viscosity (efflux) in accordance with Method No 212.1 which is described in Specification MIL-STD-650. See 6.6

4.3.4 Density — Determine the density in accordance with Method No 203.1 described in Specification MIL-STD-650

NOTE: Use a chunk of the sample to make this determination. See 6.6

5. Preparation for Delivery. See Specification

6. Notes:
6.3 Intended use of HTA-3. As a filler for warheads
6.4 HTA-3 can be made from Octol and the addn of TNT, Al & Ca silicate to conform to the desired compn
6.5 Calculation of control limits. See specification

6.6 The determination of density and viscosity is not necessary when the HTA-3 is manufactured at the time of loading

HTP (High Test Peroxide). See under Hydrogen Peroxide in this Vol

Hübner Powder. Smokeless powder prep'd by gelatinizing NC with a solution of potassium xanthogenate in alcoholic ether and adding 2-5% of nitronaphthal and nitromolasses or nitrosugar
Ref: Daniel (1902), 378

Hudson Explosive. Explosive mixture intended for use as a bursting charge in shells and prep'd in the USA in 1889 by mixing NG with NC, previously dissolved in acetone, ethyl acetate or ether-alcohol mixture
Ref: Daniel (1902) 378

Hudson Maxim Explosive. An explosive patented in the USA for loading HE shells. It was prep'd by blending NC 30-40 with NG 70-60. After the mass hardened it was pulverized and thoroughly mixed with 3-4 parts of wet, finely pulped Guncotton
Ref: Van Gelder & Schlatter (1927), 933

Huff-Duff (HFDF—High-frequency direction finder). An electronic long-distance device developed by Americans during WWII which permitted spotting within a split second any ship (such as submarine), or plane at distances much greater than radar's capability
Ref: Staff, ArmyOrdn 30, 280 (1956)
Hugoniots, more properly called Rankine-Hugoniots or R-H relations, are a series of equations and/or curves that describe conditions at a shock front. These are best illustrated by reference to an idealized plane shock, such as the one shown in Fig 1, and the following set of eqns that describe this shock:

<table>
<thead>
<tr>
<th>BEHIND</th>
<th>AHEAD</th>
</tr>
</thead>
<tbody>
<tr>
<td>PARTICLE VELOCITY</td>
<td>$u$</td>
</tr>
<tr>
<td>PRESSURE</td>
<td>$P$</td>
</tr>
<tr>
<td>DENSITY</td>
<td>$\rho \cdot NV$</td>
</tr>
<tr>
<td>INTERNAL ENERGY</td>
<td>$E$</td>
</tr>
<tr>
<td>TEMPERATURE</td>
<td>$T$</td>
</tr>
</tbody>
</table>

**Fig 1** Steady plane shock front propagating into undisturbed materials in laboratory coordinates

$$P-P_0 = \rho_0 uU$$  \hspace{1cm} (1)

$$U^2 = V_0^2(P-P_0)/V_0 - V$$  \hspace{1cm} (2)

$$u = (1-\rho_0/\rho)U$$  \hspace{1cm} (3)

$$E-E_0 = 1/2(V_0 - V)(P+P_0)$$  \hspace{1cm} (4)

Of these sets of equations it is Eq 4 that is commonly called the *Hugoniot equation*. Graphical representations of these equations (R-H curves), taken from Ref 20, are shown in Fig 2.

(a)  (b)  (c)

The curve (c) is a plot of Eq (1); curve (b) is a plot of a combination of Eqs 2 & 3; and curve (a) is empirical, although, in a way, it represents Eq 3, particularly for metals. Here $a_0$ is the bulk sound velocity in the medium ahead of the shock.

As indicated in Fig 2, curve (a) is useful in correlating experimental data, curve (b) is essentially a stress-strain curve, although it has other interesting features which will be discussed later, and curve (c) is most useful in considering shock and rarefaction effects across boundaries of different media.

Examination of Eqs 1–3, reveals that the simultaneous measurement of any two shock variables is sufficient to completely specify the remaining variables, provided that initial conditions in the unshocked medium are known. No satisfactory methods of measuring $V$ or $\rho$ are known for condensed media, therefore measurements must be restricted to $U$, $u$ & $P$. In Vol 4 of this Encyclopedia methods of measuring $U$, $u$ & $P$ are described on p D333, p D446 & D485. Theoretical Hugoniot curves for detonation products are shown on pp D706 & 707. More recent methods are reviewed in Ref 21 & 22. A novel electromagnetic method for measuring $u$ is described by Jacobs & Edwards (Ref 20).

An excellent compendium of shock Hugoniot data has been prepared by LRL (Ref 14). It includes some of the earlier Hugoniot data for explosives. These data as well as more recent data are summarized in Table 1 in the form of least square fits to the empirical equation in Fig 2a.

Most of the data in Table 1 is for granular explosive compacts. It must be pointed out that defining shock parameters for heterogeneous pellets consisting of explosive granules and air pockets present conceptual difficulties. At best the shock Hugoniots for such materials are useful only for describing the gross hydrodynamic behavior of granular explosives. It cannot be expected that the methods of continuum hydrodynamics applied to discontinuous media will yield any information on the fine structure of the state variables of the shocked explosive.
Table 1

"Unreacted" Hugoniot Data for Condensed Explosives

<table>
<thead>
<tr>
<th>HE</th>
<th>$\rho_0$ (g/cc)</th>
<th>$T_0$ (°C)</th>
<th>$a_0$ (mm/µsec)</th>
<th>$S$ (mm/µsec)</th>
<th>Range of $u$ for exp data (mm/µsec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AN</td>
<td>0.86</td>
<td>25</td>
<td>0.84 (a)</td>
<td>1.42 (a)</td>
<td>0.81-2.32</td>
</tr>
<tr>
<td>76/24 Baratol</td>
<td>2.63</td>
<td>25</td>
<td>2.79 ± 0.5 (a)</td>
<td>1.25 (a)</td>
<td>0.0-2.7</td>
</tr>
<tr>
<td>Comp B</td>
<td>1.70</td>
<td>25</td>
<td>3.0 ± 0.4</td>
<td>1.73 (a)</td>
<td>0.0-1.5</td>
</tr>
<tr>
<td></td>
<td>1.68</td>
<td>25</td>
<td>2.71 ± 0.05</td>
<td>1.86 ± 0.07</td>
<td>0.0-0.9</td>
</tr>
<tr>
<td>DATB</td>
<td>1.78</td>
<td>25</td>
<td>2.45 ± 0.04</td>
<td>1.89 ± 0.06</td>
<td>0.0-1.2</td>
</tr>
<tr>
<td>H-6</td>
<td>1.76</td>
<td>25</td>
<td>2.83 ± 0.07</td>
<td>1.70 ± 0.08</td>
<td>0.0-1.1</td>
</tr>
<tr>
<td>H-6</td>
<td>1.76</td>
<td>25</td>
<td>2.65 ± 0.10 (a)</td>
<td>1.98 ± 0.10 (a)</td>
<td>0.0-2.0</td>
</tr>
<tr>
<td>HBX-1</td>
<td>1.75</td>
<td>25</td>
<td>2.93 ± 0.08</td>
<td>1.65 ± 0.10</td>
<td>0.0-1.0</td>
</tr>
<tr>
<td>HBX-3</td>
<td>1.85</td>
<td>25</td>
<td>3.13 ± 0.02</td>
<td>1.61 ± 0.02</td>
<td>0.0-1.0</td>
</tr>
<tr>
<td>HNS</td>
<td>1.38</td>
<td>25</td>
<td>0.61 ± 0.21</td>
<td>2.77 ± 1.09</td>
<td>0.0-0.5</td>
</tr>
<tr>
<td>HNS</td>
<td>1.57</td>
<td>25</td>
<td>1.00 ± 0.05</td>
<td>3.21 ± 0.10</td>
<td>0.0-0.7</td>
</tr>
<tr>
<td>HNS</td>
<td>1.46</td>
<td>260</td>
<td></td>
<td></td>
<td>See Table 4 of Ref 18</td>
</tr>
<tr>
<td>LX-04</td>
<td>1.86</td>
<td>25</td>
<td>2.7 (a)</td>
<td>1.9 (a)</td>
<td>0.0-0.6</td>
</tr>
<tr>
<td></td>
<td>1.86</td>
<td>25</td>
<td>2.36 (a)</td>
<td>2.5 (a)</td>
<td>0.0-0.4</td>
</tr>
<tr>
<td>NM</td>
<td>1.14</td>
<td>25</td>
<td>2.00 (a)</td>
<td>1.38 (a)</td>
<td>0.0-1.8</td>
</tr>
<tr>
<td>NM</td>
<td>1.14</td>
<td>25</td>
<td>1.6 (a)</td>
<td>1.65 (a)</td>
<td>1.0-1.7</td>
</tr>
<tr>
<td>75/25 Octol</td>
<td>1.80</td>
<td>25</td>
<td>3.01 ± 0.4</td>
<td>1.72 (a)</td>
<td>0.0-1.2</td>
</tr>
<tr>
<td>PBX 9404</td>
<td>1.84</td>
<td>25</td>
<td>2.45 ± 0.21</td>
<td>2.48 ± 0.11</td>
<td>0.0-1.45</td>
</tr>
<tr>
<td>PBX 9404</td>
<td>1.84</td>
<td>25</td>
<td>2.31 ± 0.09 (b)</td>
<td>2.77 ± 0.11 (b)</td>
<td>0.0-2.00</td>
</tr>
<tr>
<td>PBX 9404</td>
<td>1.77</td>
<td>150</td>
<td>1.85 ± 0.54</td>
<td>3.47 ± 0.81</td>
<td>0.0-0.8</td>
</tr>
<tr>
<td>50/50 Pentolite</td>
<td>1.67</td>
<td>25</td>
<td>2.83 ± 0.4</td>
<td>1.91 (a)</td>
<td>0.0-1.2</td>
</tr>
<tr>
<td>PETN</td>
<td>0.82</td>
<td>25</td>
<td>0.47 (a)</td>
<td>1.73 (a)</td>
<td>0.76-3.50</td>
</tr>
<tr>
<td>PETN</td>
<td>1.0</td>
<td>25</td>
<td>0.76 (a)</td>
<td>0.66 (a)</td>
<td>0.28-0.42</td>
</tr>
<tr>
<td>PETN</td>
<td>1.59</td>
<td>25</td>
<td>1.33 ± 0.08</td>
<td>2.18 ± 0.27</td>
<td>0.03-0.37</td>
</tr>
<tr>
<td>PETN</td>
<td>1.60</td>
<td>25</td>
<td>1.32 (a)</td>
<td>2.58 (a)</td>
<td>0.2-0.4</td>
</tr>
<tr>
<td>PETN</td>
<td>1.72</td>
<td>25</td>
<td>1.83 (a)</td>
<td>3.45 (a)</td>
<td>0.2-0.6</td>
</tr>
<tr>
<td>PETN</td>
<td>1.55</td>
<td>110</td>
<td>-0.6 ± 0.5</td>
<td>8.7 ± 1.7</td>
<td>0.24-0.29</td>
</tr>
<tr>
<td>RDX</td>
<td>1.0</td>
<td>25</td>
<td>0.40 (a)</td>
<td>2.00 (a)</td>
<td>0.44-2.60</td>
</tr>
<tr>
<td>RDX</td>
<td>1.54</td>
<td>25</td>
<td>-0.7 (a)</td>
<td>3.2 (a)</td>
<td>0.25-0.6</td>
</tr>
<tr>
<td>RDX</td>
<td>1.58</td>
<td>180</td>
<td>0.71 ± 0.24</td>
<td>4.22 ± 0.42</td>
<td>0.25-0.32</td>
</tr>
<tr>
<td>RDX</td>
<td>1.64</td>
<td>25</td>
<td>0.70 ± 0.18</td>
<td>4.11 ± 0.37</td>
<td>0.35-0.47</td>
</tr>
<tr>
<td>RDX</td>
<td>1.80</td>
<td>25</td>
<td>2.87 (a)</td>
<td>1.61 (a)</td>
<td>0.75-1.6</td>
</tr>
<tr>
<td>RDX</td>
<td>1.64</td>
<td>25</td>
<td>1.93 ± 0.05</td>
<td>0.666 ± 0.168</td>
<td>0.11-0.35</td>
</tr>
<tr>
<td>RDX/EXON 94/6</td>
<td>1.60</td>
<td>25</td>
<td>1.33 (c)</td>
<td>1.99 (c)</td>
<td>0.35-0.93</td>
</tr>
<tr>
<td>TATB</td>
<td>1.85</td>
<td>25</td>
<td>2.34 ± 0.07</td>
<td>2.32 ± 0.08</td>
<td>0.3-1.4</td>
</tr>
</tbody>
</table>

(Continued)
Table 1
"Unreacted" Hugoniot Data for Condensed Explosives

<table>
<thead>
<tr>
<th>HE</th>
<th>( \rho_c ) (g/cc)</th>
<th>( T_c ) (°C)</th>
<th>( a_o ) (mm/μsec)</th>
<th>( s^+ ) (mm/μsec)</th>
<th>Range of ( u ) for exp data</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tetryl</td>
<td>1.00</td>
<td>25</td>
<td>0.35 (a)</td>
<td>1.35 (a)</td>
<td>0.76-2.80</td>
</tr>
<tr>
<td>Tetryl</td>
<td>1.30</td>
<td>25</td>
<td>2.16 (d)</td>
<td>1.438 (d)</td>
<td>0.3-1.1</td>
</tr>
<tr>
<td>Tetryl</td>
<td>1.40</td>
<td>25</td>
<td>1.61 (d)</td>
<td>1.978 (d)</td>
<td>0.3-1.25</td>
</tr>
<tr>
<td>Tetryl</td>
<td>1.50</td>
<td>25</td>
<td>2.17 (d)</td>
<td>1.628 (d)</td>
<td>0.3-1.0</td>
</tr>
<tr>
<td>Tetryl</td>
<td>1.60</td>
<td>25</td>
<td>2.36 (d)</td>
<td>1.538 (d)</td>
<td>0.3-1.25</td>
</tr>
<tr>
<td>Tetryl</td>
<td>1.70</td>
<td>25</td>
<td>2.48 (c)</td>
<td>1.42 (c)</td>
<td>0.4-1.2</td>
</tr>
<tr>
<td>TNB</td>
<td>1.64</td>
<td>25</td>
<td>2.32 ± 0.07</td>
<td>2.03 ± 0.12</td>
<td>0.0-0.75</td>
</tr>
<tr>
<td>TNT</td>
<td>1.00</td>
<td>25</td>
<td>See Ref 15</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TNT</td>
<td>1.62</td>
<td>25</td>
<td>2.493 (a)</td>
<td>1.61 (a)</td>
<td>0.75-1.6</td>
</tr>
<tr>
<td>TNT</td>
<td>1.63</td>
<td>25</td>
<td>2.57 (a)</td>
<td>1.88 (a)</td>
<td>0.0-1.25</td>
</tr>
<tr>
<td>TNT</td>
<td>1.62</td>
<td>25</td>
<td>2.27 ± 0.30</td>
<td>2.65 (a)</td>
<td>0.0-0.6</td>
</tr>
<tr>
<td>TNT</td>
<td>1.62</td>
<td>25</td>
<td>2.39 ± 0.03</td>
<td>1.36 (a)</td>
<td>1.0-1.5</td>
</tr>
<tr>
<td>TNT</td>
<td>1.61</td>
<td>25</td>
<td>2.08 ± 0.13</td>
<td>2.05 ± 0.03</td>
<td>0.0-1.4</td>
</tr>
<tr>
<td>TNT</td>
<td>1.64</td>
<td>25</td>
<td>2.4 (a)</td>
<td>2.1 (a)</td>
<td>0.2-1.4</td>
</tr>
<tr>
<td>TNT (Lq)</td>
<td>1.47</td>
<td>92</td>
<td>2.14 (a)</td>
<td>1.57 (a)</td>
<td>0.8-1.7</td>
</tr>
</tbody>
</table>

\( *U = \text{shock velocity, } a_o = \text{bulk sound velocity, } u = \text{particle velocity in the HE} \).

\( S \) is a constant.

† ± terms are twice std deviation.

(a) std deviation not given in Refs.

(b) based on widely divergent data.

(c) although std deviations are not given, data points agree very closely with this fit.

(d) \( u = a_o + Su \) - \( S^+ = \frac{a_o}{u} \); thus \( a_o \) column gives \( a_o \), & \( S^+ \) column gives \( S^+ \& S \).

Moreover, chemical reaction in or near the shock front can cause additional complications. Nevertheless, the data in Table 1 can be very useful as will be shown below, even though, aside from conceptual problems, there is some inconsistency in the data for some of the explosives.

A commonly encountered problem is that of a shock or rarefaction traveling from one medium into another. To solve such problems one converts the \( U-u \) data of Table 1 or Ref 14 into \( P-u \) data via Eq 1. Then the problem is readily solved by graphical means as illustrated in Figs 3 & 4 (taken from Ref 21).
Fig 3  Transmission of a plane shock wave into a material of lower impedance

Fig 4  Transmission of a plane shock wave into a material of higher impedance

Shock impedance is the product $\rho_A U$ and graphically it is represented by the slopes of the broken lines in Figs 3 & 4. Note that in Fig 3 the reflected wave is a rarefaction rather than a shock. Figure 5 (also from Ref 21) is a useful summary of shock effects in metals, rocks, plastics etc in contact with some common explosives.
Other interesting properties of Hugoniots are best viewed in the P-V plane. Figure 6 (taken from Ref 6) is an enlarged and more detailed view of Fig 2b.

Fig 6 The Rankine-Hugoniot curve defines states that can be induced in substance by shock compression in terms of pressure ($p$), specific volume ($V$), and internal energy ($E$). Shock compression from initial state B to shocked state C follows the R-H curve and dissipates energy shown by the hatched area. Thus shock compression is not a reversible process—unlike adiabatic compression, which is, at least ideally reversible.
Quoting Duvall (Ref 6): “Figure 6 gives the locus of all states \( (p_1, V_1, E_1) \), and so on) that can be reached from an initial state \( (p_0, V_0, E_0) \) by shock compression. In an analogous way, the ordinary adiabat or adiabatic curve may be defined as the locus of all states that can be reached by adiabatic compression.

At the point \( B \), which represents initial unshocked conditions in the material \( (p_0, V_0, E_0) \), the R-H curve and the adiabat have the same slope and curvature, but only at that point: at all higher pressures the R-H curve lies below the adiabat, because unlike adiabatic compression, shock compression dissipates energy, and is, therefore, irreversible.

As shown in Fig 6, the increase in internal energy in a shock whose pressure amplitude is \( p_1 \) is represented by area \( ABCD \). Loss of energy in a shock can be illustrated by comparing this area thermodynamically with that associated with a weaker shock, area \( ABCD' \), for example. It can also be shown by simple calculation that just as the internal energy increases or decreases as the shock is stronger or weaker, so the entropy of the final shocked state also increases with the shock strength. Although such calculations are valuable in computing the entropy of the shocked state, they are insufficient for calculating the total energy dissipation resulting from passage of the shock wave. However, referring again to Fig 6, the hatched area—bounded below by the Rankine-Hugoniot curve and above by the Rayleigh-Mikheil'son line, a straight line connecting the initial, unshocked point \( B \) with the final shocked state \( C \) — is a fair approximation to the energy dissipated in the shock cycle. But specifically this area is the waste heat of the cycle, not the energy dissipated. It is difficult to determine an exact expression for energy dissipated because thermal stresses are left behind in the material, even after the shock pressure has been relieved. Therefore, a precise calculation of the true energy dissipation in a decaying shock must account for hard-to-evaluate effects of thermally induced after-flow in the material. In practice we settle for the waste heat approximation.”

Duvall (Ref 5) also provides further discussion of P-V Hugoniot curves, eg the manifestations of multiple shocks, phase transitions, elastic limits etc. Cowperthwaite (Ref 11) considers the thermodynamic consequences of shock Hugoniot data and what additional information is required to specify an equation of state for the shocked medium.

Some very interesting information obtainable from Hugoniot data is dramatically illustrated in Fig 7 which is a plot of experimental P-V data (taken from Ref 15) for 1 g/cc TNT.

![P-V Hugoniot for 1 g/cc TNT](image)

Here the TNT sample is compressed at very low pressures from \( V=1 \) cc/g to \( V=0.62 \) cc/g (crystal density). Further compression (increase in pressure) then causes the sample to \textit{expand}! This can only mean that some \textit{heat} effect is overcoming this compression. Since it can be shown that uniform shock heating at pressures of the order of a few kbars is very small, this heat effect must be produced by exothermic chemical reaction at or very near the shock front. Thus shock Hugoniots for reactive materials can provide information on the presence or absence of chemical reaction at the shock front.

Written by J. ROTH
See also Hugoniot in Vol 4, pp D278-R to D281-R and pp D604-L to D607-R


Huile de Nobel. Fr for NG

Huile détonante, ou Huile explosive. Fr for one of the names for Nitroglycerin.

Hull. The outer covering or husk of any seed or fruit. Some of these substances, for instance, walnut or coconut hulls (shells) can be nitrated and probably would produce explosives resembling NC or NS

Hulls, Nitrated. Krüger studied the action of concd(1.52) nitric acid on a variety of plant hulls, such as those of chestnuts, walnuts, plums, etc. After 12 hours at -10º in the acid a considerable portion of the hulls dissolved leaving a residue (40-60% of the original weight) that contained 11-12%N primarily in the form of nitrate esters. When water was added the dissolved material, a slimy precipitate was formed. It also contained N but less than the residue. Moreover this N was only partly in the form of nitrates

Ref: W. Krüger, Ber 73B, 493 (1940) & CA 34, 6580 (1940)

Humidification and Humidity. The amount of water vapor in air is referred to as the humidity of air. The controlled increase in humidity is called humidification. The actual moisture content in air is called absolute humidity, and the ratio of the weight of water vapor to the weight of water vapor in moisture-saturated air at the same temp (expressed in %) is called relative humidity (RH)

Humidity can affect explosives and propellants both adversely and beneficently. If the humidity is very high, an explosive can pick up enough moisture to cause it to deteriorate or malfunction. A prime example of this is Ammonium Nitrate which must be stored and handled in special dehumidified areas to prevent caking. The humidity above which a substance deliquesces, or below which it ceases to be hygroscopic is called the critical relative humidity (CRH). It is equal to the humidity in equilibrium with a saturated solution of the substance at a given temperature

On the other hand, humidification can prevent buildup of dangerous electrostatic charges. This is especially important in the handling of such primary explosives as Pb Azide or Styphnate. General references on humidity control & humidification are Refs 2, 4, 5 & 6

CRH's can be used to determine whether a material will deliquesce or if it has a tendency to cake. Thus if the RH > CRH the substance will pick up moisture from the air (deliquesce) until the vapor pressure of water in the deliquescing material equals the partial pressure of water in the surrounding air. Conversely, if RH < CRH any moisture in the material will tend to escape
### Critical Relative Humidities for Some Explosives and Ingredients of Explosive and Pyrotechnic Compositions

<table>
<thead>
<tr>
<th>Substance</th>
<th>10</th>
<th>20</th>
<th>30</th>
<th>40</th>
<th>Other temperatures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ammonium chloride, NH₄Cl</td>
<td>75.3</td>
<td>69.8</td>
<td>59.4</td>
<td>52.5</td>
<td>48.4 at 50°C</td>
</tr>
<tr>
<td><strong>dichromate</strong> (NH₄)₂Cr₂O₇</td>
<td>98.1(c)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>nitrate</strong>, NH₄NO₃</td>
<td>75.0</td>
<td>66.0</td>
<td>60.0</td>
<td>53.0</td>
<td>48.6 at 50°C</td>
</tr>
<tr>
<td><strong>oxalate</strong>, (NH₄)₂C₂O₄</td>
<td></td>
<td></td>
<td></td>
<td>98.9(h)</td>
<td></td>
</tr>
<tr>
<td>** perchlorate**, NH₄ClO₄</td>
<td>95.6(c)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>permanganate</strong>, (NH₄)₂MnO₄</td>
<td>99.1(c)</td>
<td>97.0</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>sulfate</strong>, (NH₄)₂SO₄</td>
<td>79.4</td>
<td>80.0</td>
<td>81.0</td>
<td>79.0</td>
<td>75 at 108.2°C</td>
</tr>
<tr>
<td>Barium chloride, BaCl₂·2H₂O</td>
<td></td>
<td>88.0(g)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>nitrate</strong>, Ba(NO₃)₂</td>
<td>98.8</td>
<td>97.7</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Calcium chloride, CaCl₂·6H₂O</td>
<td>38.0</td>
<td>32.3</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>nitrate</strong>, Ca(NO₃)₂·4H₂O</td>
<td>66.0</td>
<td>56.0</td>
<td>51.0</td>
<td>46.0</td>
<td></td>
</tr>
<tr>
<td><strong>sulfate</strong>, CaSO₄·5H₂O</td>
<td>98.0</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cobalt chloride, CoCl₂·6H₂O</td>
<td>72.5(d)</td>
<td>67.3</td>
<td>61.9</td>
<td>56.6</td>
<td>49 at 55°C</td>
</tr>
<tr>
<td>Copper nitrate, Cu(NO₃)₂</td>
<td></td>
<td>75(d)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hexose, C₆H₁₂O₆·1/2H₂O</td>
<td>57(b)</td>
<td>56(f)</td>
<td>55(j)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lead nitrate, Pb(NO₃)₂</td>
<td>99.0</td>
<td>98.5</td>
<td>96.5</td>
<td>95.5</td>
<td>88.4 at 103.5°C</td>
</tr>
<tr>
<td>Magnesium chloride, MgCl₂·6H₂O</td>
<td>34.7</td>
<td>33.1</td>
<td>31.7</td>
<td>31.3</td>
<td>28 at 71°C</td>
</tr>
<tr>
<td><strong>nitrate</strong>, Mg(NO₃)₂·6H₂O</td>
<td>56(e)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mercuric Fulminate, Hg(ONC)₂</td>
<td>99.9(b)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oxalic acid, H₂C₂O₄·2H₂O</td>
<td></td>
<td>76.0</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Potassium chloride, KClO₃</td>
<td>98.0(h)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>chloride</strong>, KCl</td>
<td>87.4</td>
<td>86.3</td>
<td>84.5</td>
<td>82.8</td>
<td>74.7 at 100°C</td>
</tr>
<tr>
<td><strong>chromate</strong>, K₂CrO₄</td>
<td></td>
<td>88.0</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>** dichromate**, K₂Cr₂O₇</td>
<td></td>
<td>97.8</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>** nitrate**, KNO₃</td>
<td>95.1</td>
<td>94.2</td>
<td>92.5</td>
<td>89.4</td>
<td>61.2 at 110°C</td>
</tr>
<tr>
<td>** perchlorate**, KClO₄</td>
<td>99.8(c)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>sodium tartrate</strong>, KNaC₆H₄O₆·4H₂O</td>
<td>87.5</td>
<td></td>
<td>87.1</td>
<td>86.0</td>
<td></td>
</tr>
<tr>
<td><strong>sulfate</strong>, K₂SO₄</td>
<td>98.2</td>
<td>97.1</td>
<td>96.6</td>
<td>96.1</td>
<td>93 at 102.1°C</td>
</tr>
<tr>
<td>** sulfocyanate**, KCNS</td>
<td></td>
<td>47.0</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>** tartrate**, K₂C₆H₄O₆·1/2H₂O</td>
<td></td>
<td>75(c)</td>
<td></td>
<td>73.0</td>
<td></td>
</tr>
<tr>
<td>Sodium chloride, NaClO₃</td>
<td>98.8(b)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>chloride</strong>, NaCl</td>
<td>76.9</td>
<td>75.8</td>
<td>75.1</td>
<td>74.4</td>
<td>73.7 at 100°C</td>
</tr>
<tr>
<td>** dichromate**, Na₂Cr₂O₇</td>
<td></td>
<td>52.0</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>** nitrate**, NaNO₃</td>
<td>77.7</td>
<td>77.5</td>
<td>73.2</td>
<td>70.5</td>
<td>62 at 71°C</td>
</tr>
<tr>
<td>** sulfate**, Na₂SO₄</td>
<td>93(i)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>** tartrate**, Na₂C₆H₄O₆·H₂O</td>
<td></td>
<td>92.5</td>
<td>91.1</td>
<td>90.7</td>
<td></td>
</tr>
<tr>
<td>Strontium nitrate, Sr(NO₃)₂</td>
<td></td>
<td>86.0</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sugar, C₁₂H₂₂O₁₁</td>
<td></td>
<td>85.0(h)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Urea, NH₂CONH₂</td>
<td>81.5</td>
<td>80.5</td>
<td>73.3</td>
<td>68.7</td>
<td>62.7 at 50°C</td>
</tr>
<tr>
<td>Zinc chloride, ZnCl₂·xH₂O</td>
<td>10.0(b)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>** nitrate**, Zn(NO₃)₂·6H₂O</td>
<td></td>
<td>42.0</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>** sulfate**, ZnSO₄·7H₂O</td>
<td>94.7(a)</td>
<td>90.0</td>
<td>88.5(h)</td>
<td>84.0</td>
<td></td>
</tr>
</tbody>
</table>

*a - at 5°C  f - at 20°C*

*b - at 12°C  g - at 24.5°C*

*c - at 15°C  h - at 25°C*

*d - at 18°C  i - at 35°C*

*e - at 18.5°C  j - at room temp*
until equilibrium is established. Such evaporation can produce caking. Obviously caking will be strongly influenced by the original moisture content of the material.

The foregoing table (taken from Ref 1) gives CRH for some explosives and ingredients of explosive and pyrotechnic compositions.

Practical measures for controlling explosive dusts by humidification are discussed in Ref 3.

Ignition lag times of pyrophoric substances such as Al(BH₄)₃, AlMe₃, BE₅, etc were found to increase (except for BE₅) with decreasing humidity. In fact, Al(BH₄)₃ is not pyrophoric in dry air (Ref 4).

Suppression of firedamp and coal dust explosions, in high humidity coal mines, by stone dusting is affected by humidity. Agglomeration of moisture content of the dusting material influenced its effectiveness as explosion suppressor. Salt dusts become ineffective above 80% relative humidity because they dissolve (Ref 7).


**Hummel (Bumble Bee).** Nickname for a self-propelled mounting consisting of 150mm Medium Howitzer on the chassis of a PzKpfw III/IV tank. (See also under Panzer)

Ref: PATR 2510 (1958), p Ger 93-L

**Humus.** Decayed products of plant life, such as leafmolds, rotten wood, rotten straw, etc. It has been used as an ingredient of some explosive mixtures, as for instance of Bracket Sporting powder, manufactured in the USA, and which contained 18.9% of humus, according to the analysis by Munroe.

Ref: Daniel (1902) 82

**Hungarian Ammunition, Explosives and Weapons.** We have no information on post WWII Hungarian ammunition, exps & weapons

**Hunting Powders.** See Sporting Powders

**Hurst.** Patented in 1898 in the USA. The use of NG in frozen form as bursting charge for projectiles

Ref: Daniel (1902) 379

**Hybalines.** A family of novel metal hydride coordination compds, both liq & solid, trade name of Union Carbide Corp, New York. The company’s R & D program at its South Charleston, West Virginia technical center suggests Hybalines may find use as high-energy liq fuels and as additives in high-energy solid propellants.

Carbide is proposing a new concept of solid propelt grain construction. The process uses std ingredients, yet the grain burns more efficiently and can be fabricated faster than with conventional methods. There also exists a group of binders trademarked Carbitron which may find use in solid propelt because of their superior physical & ballistic props.


**Hybrid Rocket Propellants.** A special propelt combination of unlike materials, particularly of unlike physical characteristics. Typical hybrid propelt combinations are a solid fuel (or oxidizer) in combination with a liquid oxidizer (or fuel) in that order. Sometimes a grain of solid fuel is encased in the combustion chamber of a rocket engine and burned in combination with liq oxygen. Similarly a liq fuel may be injected into a combustion chamber in contact with a solid oxidizer. Another example is the use of concentrated hydrogen peroxide and a hydrocarbon fuel. In this case, the hydrogen peroxide is converted by decomp into a hot gas contg oxygen. The fuel is injected downstream of the first reaction, mixed with the hot oxidier-rich gas, and burns (Ref 1).

Hybrid rocket propelt & hybrid technology provide a number of propulsion-system capabilities in combinations not readily achievable with all-solid or all-liq systems. Foremost among
capabilities are throttling, safety, smokelessness, and low development cost (Ref 7).


**Hydantoins and Derivatives**

**Hydantoins, Glycoalyurea or Imidazoalideone** (called Lactam der Ureido-essigure in Ger), HN,CH₂CO,NH₂CO; mw 100.08, N 27.99%; colorless, odorless crystal, mp 220–21°, bp 7; sol in hot w & alk; sl sol in alc; nearly insol in eth. Was first prep'd by Franchimont & Kloebb (Refs 1 & 2). Wagner & Simons (Ref 3) prep'd it by two-step process from glycine, H₂N.CH₂.H. Used as intermediate in manuf of pharmaceuticals, synthetic resins, lubricants, etc, Givens on nitration an expl


**N-Nitrohydantoin or N-Nitroglycoalyurea**, O₂N,N,CH₂.CO,NH₂CO; mw 145.08, N 28.96%, OB to CO₂ –38.6%; colorless, sl hygroscopic plates; mp 170–71° (dec); bp–puffs off and sometimes ignites at 225°, but does not explode. Dissolves in w giving an acidic soln (pH 5). Can be prep'd either by nitrating hydantoin with mixed nitric-acetic acid or by dissolving hydantoin (15g) in white nitric (50ml), followed by evaporation of the soln to dryness on a steam bath.

It is a mild expl of satisfactory thermal stability and with sensitivity to impact comparable to that of TNT. Its heat of combstn is 295 kcal/mol.


No Azido- or Polyhynitro-derivs of Hydantoin are found in Beil or CA thru 1971

**Hydral-cellulose, (C₆H₁₀O₄)₄·H₂O, mw 666.58.** May be prep'd by digesting cellulose, such as filter paper, with a solution of H₂O₂ until completely disintegrated. On evaporation, a white substance is obtained which resembles "oxycellulose" in its properties. When treated with 10% aqueous NaOH, 33% of it dissolves, leaving a residue resembling cellulose. When cellulose is treated with concentrated H₂O₂ (60%) the resulting mixture is explosive (see Hydrogen Peroxide Explosives).


**Hydra Programs.** HYDRA is a US Navy code name for all its studies & tests of sea-launched missile concepts & systems. To simulate nuclear expls, Hydra programs have used chem HEs (Pentolite) which form noncondensible gas bubbles. In an effort to more accurately simulate nuclear detonations, which form steam bubbles, two steam producing expls were developed. These were an equal mixt by wt of RDX & alun [AlNH₄(SO₄)₂·12H₂O] and a pressurized stoichiometric mixt of hydrogen & oxygen. Using Friedman's formula for the bubble oscillation period and the experimentally determined bubble periods from Hydra studies, the energy partition (i.e., fraction of chge energy left for bubble oscillation after shock passage) of the three expls were found to be:
Since the min error variation in energy partition is much greater than the variation due to type of expl, it makes little difference which expl is used to simulate the nuclear case (Ref 6).

See also Refs 1, 2, 3, 4, 5 & 7 for further
detais.


Hydrated Cellulose, C₁₂H₂₀O₁₀·H₂O, mw 342.30; name proposed by Cross and Bevan for substances recovered from cellulose that has been mercerized (by treatment with aqueous NaOH) and then washed out, or that which has been converted into viscose and then regenerated.

The name “hydrated cellulose” is now considered to be obsolete, and the term “dispersed” cellulose has become more general (Refs 1 & 2). Hydrated cellulose gives off 1 mol of water at 120-125° while hydrocellulose retains it obstinately (Ref 2). See also Amyloid in Vol 1, p A398-L.


Hydrated Explosives. Permissible explosives (ie permissible for use in gassy mines) in which salts containing water of crystallization, such as MgSO₄·7H₂O with 51.22% H₂O, K₂Al₃(SO₄)₆·24H₂O with 45.57% H₂O, Al₂(SO₄)₃·18H₂O with 48.81% H₂O, CaSO₄·2H₂O with 20.92% H₂O etc., are characteristic ingredients. Explosives of this class are somewhat similar in composition to the ordinary low-grade Dynamites (all of which contain NG as the principle explosive ingredient and many contain AmmoniumNitrate), except that one or more salts containing water of crystallization are added to reduce flame temperature. These explosives are usually easily detonated, produce only small quantities of poisonous gases and generally can be used successfully in damp working places. A typical hydrated explosive has the following composition: NG42, hydrated Mg or Na sulfate 46, woodmeal 12%. Its strength compared to blasting gelatin: 200ml lead block expansion and 11mm crushing compared to 560cc and 24mm for 92/8 blasting gelatin.

See also Permissible Dynamites in Vol 5, p D1604-Rff


Hydrazide is a deriv of Hydrazine (qv) of general formula MHN.H₂N₂, where M means metal. One such compd, Sodium Hydrazide, NaHN.H₂N₂, mw 54.04, N 51.85%, is extremely explosive. It can be prepd by the action of Na on anhydrous hydrazine or from NaNH₂ and hydrazine

Hydrazides are compounds of the type RCONHNH₂. It is claimed that during recovery of pure anhydrous hydrazine from its mixtures with NH₄Cl & NH₃, by treating the mixture with a stoichiometric excess of alkali or alkaline earth metals in dilq ammonia, a large excess of the metal is to be avoided to prevent formation of explosive hydrazides
Refs: 1) F. T. Neth USP 2,735,752 (1956) & CA 80 8148 (1958) 2) No further refs to explosive hydrazides was found in CA 1957-71

Hydrazidicarboxylic Acid, Mercuric Ester.
C₈H₁₀O₄N₂Hg; this powder is claimed to explode on heating (total abstract)
Ref: R. Stolle, Ber 45, 273 (1912) & CA 6, 1005 (1912)

Hydrazidoxalic Acid—Diazide (called Hydrazidoxalsülze—diazid in Ger) N₈ COCONHNCOCO₃, mw 226.12, N 49.56%, OB −35.4; wh powd (from eth). Prep'd by treating hydrazidioxalic acid-dihydrazide with nitric acid. Explodes on heating
Ref: Beil 2, (244)

Hydrazinate of Diammoniodecaborane or Diammoniodecaborane (1,2) hydrazinate.
NH₃(B₁₀H₁₂)NH₃·N₂H₄; was prep'd (Ref 1) by adding hydrazine to a suspension of B₁₀H₁₂·2MeCN in benzene and refluxing the mixt. The benzene layer was decanted, the N₂H₄ layer was heated, EtOH was added & the mixt was cooled to 0° to ppt the salt. A suggested use is as a propellant component. No properties given

HYDRAZINE, ITS SALTS AND DERIVATIVES
Hydrazine, Anhydrous (Diamidogen or Diamide), H₂N.NH₂; mw 32.05, N 87.43%, OB to CO₂ −100%; colorless, corrosive, fuming hygroscopic liquid; sp gr 1.011 at 15°/4°(Lange) or 1.004 at 25°/4° (Ref 18a); fr p 2.0, bp 113.5°, fl p (open cup) 126°F; Refr Index 1.46979 at 22.3°. Vapor pressure can be called by the equation given in Ref 11; viscosity by the eqn given in Ref 8, thermodynamic data is in Ref 12, heat capacity of liquid is in Ref 2, heat capacity of gas is in Ref 11 and entropy is in Ref 11. Heat of combustion to water liq is 148.635 kcal/mole (Ref 9). Sol in polar liquids such as water, alcohols, ammonia & amines; insol in nonpolar liquids such as ether, chl, hydrocarbons and halogenated hydrocarbons. It is strong reducing agent and diamine base. Its vapor is explosive and toxic and especially dangerous to the eye (Refs 18a, 27c and 32a) (See also under Toxicity)
Preparation.

Anhydrous hydrazine was first prep'd in 1894 by Lobry de Bruyn, but its salts and derivs were prep'd beginning in 1875 by Emil Fischer, who coined the name "hydrazin". The "hydrazine hydrate" was first prep'd by Th. Curtius in 1887 (Ref 14a, p 1)
Aecdg to Clark (Ref 14a), only a small scale operation was producing hydrazine prior to WWII
In WWII the Germans discovered that N₂H₄ was an excellent rocket fuel when used with hydrogen peroxide, oxygen or fuming nitric acid. They used hydrazine as the propellant of their first rocket plane
The synthesis developed in 1907 by Raschig is the only preparative method that is now used commercially to produce N₂H₄ (Refs 3, 4 & 5). Sodium hypochlorite and NH₃ are first reacted to form chloramine according to:

\[ \text{NaOCl} + \text{NH}_3 + \text{NH}_4\text{Cl} + \text{NaOH} \]

Then hydrazine is formed by the action of excess ammonia upon chloramine:

\[ \text{NH}_4\text{Cl} + \text{NH}_3 + \text{NaOH} \rightarrow \text{N}_2\text{H}_4 + \text{NaCl} + \text{H}_2\text{O} \]

The Raschig synthesis yields a crude solution containing about 2% N₂H₄. After evaporating to precipitate sodium chloride, the purified hydrazine hydrate solution is distilled to yield an azetrop containing 58.5 mole% N₂H₄.
Commercial anhydrous hydrazine is produced by an azetric distillation using aniline process (Ref 18). In the US, Mathieson Chem Co began in 1947 research on the production of anhyd hydrazine with emphasis directed to its use as a rocket fuel. After constructing a pilot plant, an installation for commercial production of hydrazine and its derivs was erected at Lake Charles, Louisiana
A modification of the Raschig synthesis uses urea in place of ammonia to synthesize N₂H₄ according to (Ref 12):

\[ \text{NH}_3\text{CONH}_2 + \text{NaOCl} + 2\text{NaOH} + \text{N}_2\text{H}_4 + \text{H}_2\text{O} + \text{NaCl} + \text{Na}_2\text{CO}_3 \]
Audrieth summarizes many other lab prep methods based upon hyponitrous acid \((H_2N_2O_2)\), nitramide \((H_2NO_2)\), nitrosopolyhydroxylamine \([NH(NO)OH]\) derivatives, nitrosoamine \((H_2NO)\) derivatives, nitroso ketones, azo compounds, hydrazoic acid and azides, nitrogen and the decomposition of \(NH_3\) (Ref 12). In more recent work Anderson converted \(NH_3\) to \(N_2H_4\) by passing \(NH_3\) through a discharge tube using high frequency radio waves (Ref 17). Yields in terms of energy input are discouragingly low. Aerojet General has also studied a process based upon thermal decomposition of \(NH_3\) in a nuclear reactor but without achieving commercial production (Ref 26).

Accdg to Ref 38, although the classical Raschig Process was improved by West Germany's Bayer, many of its drawbacks still exist. These include the environmental problem of disposing of the large quantity of by-product chloride, the current tight chlorine situation and sizable consumption of primary energy.

Now chemists at the Central Research Laboratories of Produits Chimiques UGINE Kuhlmann (PCUK) at Lyons, France, developed beginning in 1970, under the direction of Dr F. Weiss, a novel continuous process for manufg hydrazine. The process is based on the discovery that ammonia, hydroperoxide, and a carbonyl compd, such as methyl ethyl ketone, react in presence of an amide and catalytic quantities of a phosphate to form an azine intermediate. This intermediate hydrolyzes quantitatively to hydrazine and to carbonyl compd which is recycled. The yield is claimed to be better than 75%.

The reaction proceeds as follows:

\[
\begin{align*}
\text{Ketone} & \quad \text{NH}_3 & \quad \text{NH}_2 \quad \text{OH} & \quad \text{C} = \text{NH} & \quad \text{H}_2\text{O} & \quad \text{C} = \text{NH} \\
\text{H}_2\text{N} = \text{NH}_2 & \quad \text{H}_2\text{O} & \quad \text{C} = \text{N} - \text{N} = \text{C} & \quad \text{C} = \text{O} & \quad \text{C} = \text{N} - \text{NH}_2
\end{align*}
\]

Accdg to PCUK's Dr J.P. Schirmann, the new method in addn to having a decided economic edge over existing methods, has the important advantage that chlorine compds are not involved in the reaction.

Encouraged by the operating success so far of a pilot plant in Lyons, PCUK is building a small-scale commercial unit there due on stream in 1975. A full scale plant is planned for 1977.

Hydrazine demand is rising continuously. Apart from being a constituent of rocket fuels, it is used to remove oxygen from boiler water to prevent the corrosion of the vessels. If, as some believe, hydrazine-based fuel cells eventually come into commercial use, demand will be enormous. Hydrazine hydrate production in non-communist countries now stands at about 25000 metric tons per year, of which 17000 metric tons are used by the US with about 7000 metric tons taken by the space program. US hydrazine capacity will be greatly increased by the end of 1975 when Mobay's 10000mt/yr unit constructed at Baytown, Texas will be ready.

The reaction of the PCUK process is described more fully by Dr Schirmann on p 19-L of Ref 38, while the additional advantages of the process are listed by Dr S. Delavarenne on p 19-M.

**Toxicology**

If spilled on the skin or in the eyes liquid \(N_2H_4\) can cause severe local damage or burns and can cause dermatitis. In addition it can penetrate skin to cause systemic effects similar to those produced when the compound is swallowed or inhaled. Inhalation of the vapor causes local irritation of the respiratory tract and eyes. On short exposure systemic effects involve the central nervous system. Resultant symptoms include tremors & on exposure to higher concentrations, convulsions and possibly death follow. Repeated exposures may cause toxic damage to the liver (fatty liver) and kidney (interstitial nephritis), as well as anemia. The threshold limit value of hydrazine is 1 ppm (1.3 mg/m³)(Ref 20) (See also Refs 27c and 33a)

**Uses**

The use of \(N_2H_4\) as a hypergolic propellant fuel for rocket propulsion is discussed in detail in this Vol under Hypergolic Propellants. It has also been used extensively as a monopropellant fuel or in combination with hydrazine nitrate and/or water as a thruster for maneuvering space vehicles (Ref 31).

Applications using hydrazine as a gas generator include tank pressurization, inflating pneumatic-power engine starters, turbines and positive displacement drives. The development of many of these uses was brought about by the introduction of decomposition catalysts discussed below in the section on combustion properties.
Hydrazine is used for the preparation of hydrazoic acid according to:

\[ \text{N}_2\text{H}_4 + \text{HNO}_2 \rightarrow \text{HN}_3 + 2\text{H}_2\text{O} \]

or for the preparation of sodium azide (Ref 14). Hydrazine is used also as a scavenger for oxygen in preventing corrosion in boilers (Ref 16) and as an oxygen scavenger to prevent external corrosion of oilwell casings (Ref 21). It has been found to be an excellent fuel for fuel cells (Refs 23, 25 & 27). Hydrazine is also used in a family of liquid exps developed & marketed under the trade name “Astrolite” by the Explosive Corporation of America, Issaquah, Wash (Refs 27b, 33a & 37)

Analytical

The water and ammonia content of anhydrous \( \text{N}_2\text{H}_4 \) are determined by a gas chromatography (Refs 37 & 38) method, and the analysis of the aniline in the mixture by ultraviolet spectroscopy. The total \( \text{N}_2\text{H}_4 \) content can be then determined by difference. Other methods are given at the end of section on Hydrazine

Chemistry

\( \text{N}_2\text{H}_4 \) has been characterized by Audrieth (Ref 7) as a saturated hydronitrogen belonging to the type given by the formula \( \text{N}_n\text{H}_{n+2} \). The members of this series of compounds are as follows:

- NH\(_3\) ammonia
- \( \text{N}_2\text{H}_4 \) hydrazine
- \( \text{N}_3\text{H}_8 \) triazane
- \( \text{N}_4\text{H}_6 \) tetrazane

Hydrazine may be looked upon as the nitrogen analog of hydrogen peroxide. It is a powerful reducing agent which makes it an attractive fuel for reaction with oxidizers such as \( \text{H}_2\text{O}_2 \), \( \text{O}_2 \), and fuming nitric acids and nitrogen tetroxide. Aqueous solutions of \( \text{N}_2\text{H}_4 \) have been employed to reduce various metallic ions such as copper, silver, gold and the platinum metals to the metallic state. It also reduces strong oxidizing agents such as permanganate, iodate, hypoiodate, iodine, cerate and the like. Many of these reactions have been used for the quantitative determination of hydrazine but side reactions giving off \( \text{NH}_3 \) and in some instances hydrazoic acid may present problems.

Anhydrous \( \text{N}_2\text{H}_4 \) is a thermodynamically unstable compound and tends to undergo decomposition to \( \text{N}_2 \), \( \text{H}_2 \) and \( \text{NH}_3 \) spontaneously. This is also true in aqueous solutions and is enhanced by the presence of catalytically active metals and oxides, such as Cu, Zn, Fe\(_2\)O\(_3\) etc (Ref 7)

Generally hydrazine reactions are similar to those of ammonia and amines except that as a diamine, further reaction takes place. The reaction of \( \text{N}_2\text{H}_4 \) with CO at 20-50\(^\circ\)C and high pressure produces semicarbazide (Ref 1). At higher temperatures and pressures the products are 4-amino-1,2,4-triazol-3-one and 4-amino-1,2,4-triazole. With cyanamide, hydrazine forms aminoguanidine which is treated with nitrous acid to form tetracene, a primary explosive (Ref 15, 19). Carboxylic acids are neutralized by hydrazine to form salts which decompose upon heating to give mono- or dicyclohydrizides (Ref 2). The hydrazine salts of oxidizing acids, such as nitric acid or perchloric acid, are sensitive to shock and decompose violently. Hydrazinium Selenate, \( \text{N}_2\text{H}_4\text{HSeO}_4 \), is also explosive, in fact a very sensitive explosive (Ref 27a).

Salts of very powerful oxidizing acids (eg chromic or permanganic) are unknown, and are unlikely to exist. Hydrazonium iodate may exist in solution at low temp (Ref 27a). Alkali metals, amides and hydrides react with hydrazine to give the corresponding alkali hydrazide. Sodium hydrazide explodes violently in the presence of \( \text{O}_2 \) or when heated above 100\(^\circ\)C—a typical behavior of the alkali hydrazides. For other reactions, see Ref 24

Explosive and Combustion Properties

Liquid anhydrous \( \text{N}_2\text{H}_4 \) is a non-explosive even though it is thermodynamically unstable. In the absence of decomposition catalysts, it has been heated above 500\(^\circ\)F with very little decomposition. It is completely insensitive to shock, friction or electrical discharge. Hydrazine vapor has a lower limit of inflammability of 4.67% by volume in air. The upper limit is 100% since the vapor may be exploded without the addition of air (Ref 10)

The use of \( \text{N}_2\text{H}_4 \) as a hypergolic rocket fuel is described under the topic Hypergolic Propellants in this Vol. Another major research effort has seen the development of hydrazine thruster rockets based upon the catalytic de-
composition of hydrazine. Typical thruster rockets give thrusts ranging from 0.1 ft-lb to 5 ft-lb with positive thrust pulses ranging to 1 million cycles (Refs 13 & 35). Catalysts studied for the heterogeneous decomposition of \( \text{N}_2\text{H}_4 \) include ruthenium, iridium and rhodium. Decomposition kinetics are detailed for ruthenium (Ref 34) and rhodium (Ref 36).

Due to the relatively high freezing point of pure anhydrous \( \text{N}_2\text{H}_4 \), simple binary and ternary solutions of \( \text{N}_2\text{H}_4 \) with water, ammonia and hydrazine nitrate form low freezing eutectics which provide a wide range of exhaust gas temperatures and compositions when used in gas generators. Typical examples are given in Ref 31. Ternary diagrams for the system \( \text{N}_2\text{H}_4-\text{N}_2\text{H}_4\text{NO}_3-\text{H}_2\text{O} \) give mixture freezing points, detonable limits and vacuum specific impulse for 20%, 40% and 60% \( \text{NH}_3 \) in the equilibrium mixture.

**Explosive Compositions**

Mixtures of hydrazine and other compounds form liquid explosives which have been described. A study of the detonablility of binary and ternary mixtures of nitromethane-hydrazine-methanol showed that \( \text{N}_2\text{H}_4 \) sensitizes nitromethane and nitromethane-methanol mixtures to detonation (Ref 29). From 2-10% \( \text{N}_2\text{H}_4 \) and 90-98% lower molecular weight nitroaraffins are claimed to be a water insoluble liquid explosive (Ref 22). Mixtures of hydrazine, ammonium nitrate and aluminum have been patented as explosive compositions with high air blast effects (Ref 32).

High performance explosive mixtures with densities \( \geq 1.4 \) have been obtained with a sensitizer containing an oxidizing acid and a base such as hydrazine (Ref 30). High detonation rate, low freezing point, low viscosity and low impact sensitivity are provided by mixture of \( \text{N}_2\text{H}_4\text{NO}_3 \) (and its mixtures with \( \text{N}_2\text{H}_4\text{ClO}_4 \) in a 4:1 ratio for increased sensitivity) and including up to 25% of metal salts such as Na, K, Ca and Al nitrates and perchlorates, 6-15% \( \text{N}_2\text{H}_4 \) and 2-15% \( \text{NH}_3 \) and water (Ref 28). The water content is varied to control the detonation velocity. For example a solution containing \( \text{N}_2\text{H}_4\text{NO}_3 \) 70%, \( \text{N}_2\text{H}_4 \) 7% and \( \text{NH}_3 \) 14% has a density of 1.31, detonation rate 8100 m/sec, fp -8°F and a viscosity of 12 cp. Variable amounts of perchlorates give impact sensitivities of 45-85 Kg/cm (nitroglycerine = 2 Kg/cm on the particular impact machine used).

Hydrazine forms numerous salts and derivatives, some of which are expl or used as ingredients of expls or for their prep. Several of them are described below


Hydrazine and Derivatives, Analytical Procedures.

See at the end of this section

Hydrazine Azide, Hydrazine Azoxime or Hydrazonium Azide. See list of Inorganic Azides in Vol 1, pp A536-537, and additional information below.

Apin et al (Ref 4), in a calorimetric investigation of mixtures of Hydrazine Azide with several metallic elements found that an explosion resulted with the formation of NH₃. The amount of NH₃ depended on the explosion conditions. The limiting conditions were either N₂H₄·HN₃ + 2%H₂ + 2%N₂ or + 1/2/3/N₃H₄ + 1/2/3/N₂

Calcium Azide Hydrazinates. Ca(N₃)₂·N₂H₄ and Ca(N₃)₂·2N₂H₄; the addn of alc to a soln of Ca(N₃)₂ in anhyd N₂H₄, or evapn of such a soln in a vacuum desiccator over sulfuric acid leads to the formation of a 2-hydrazinate. This solvate dissociates rapidly under reduced pressure and/or at higher temps to give the 1-hydrazinate, Ca(N₃)₂·N₂H₄ (Ref 1)


Hydrazine Chlorate, Hydrazinium Chlorate, N₂H₄·HClO₃ or N₂H₅ClO₃; mw 116.51, N 24.05%, OB to Cl₂ and H₂O +6.9%; crystals, mp 80°C; v sol in H₂O, diff sol in alc, insol in ether, benzene and chloroform; hygroscopic; may be prep'd by mixing eq solns of N₂H₄ and HClO₃ and evap at room temp; a powerful oxidizing agent and a very powerful explosive. Its power, as detd by Trauzl test is
about 278% of Mercury Fulminate or 100% TNT. It is less stable than Hydrazine Perchlorate. It has been proposed for use in primary and initiating mixtures (Ref 1)


Hydrazine Chlorate and Perchlorate Complexes. See Hydrazine Perchlorate and Chlorate Complexes*

Hydrazine (or Hydrazinium Chloride (or Monochloride) (or Hydrochloride), N₂H₄.HCl; mw 68.52, N 40.89%; colorless crystals, mp 92.6°, decomp at 240°; v sol in w; diff sol in absol alc; easily sol in liq NH₃ — Forms a number of double salts. Can be prepd by treating hydrazine hydrate with 1 mole of HCl


Hydrazine (or Hydrazinium) Chlorite, N₂H₄.HClO₄; mw 100.52, N 27.88%, mp—ignites spontaneously when dry. Was prepd in small amts from hydrazine hydrosulfate (N₂H₄.HSO₄), Ba hydroxide and Ba chloride [Ba(ClO₄)₂].

Refs: 1) Gmelins Hdb (1936), Syst 33, p 555 2) Clark, Hydrazine (1953), p 13

Hydrazine (or Hydrazonium)-N,N’-dicarboic Acid Diazone. See under HYDRAZINE SALTS OF ORGANIC ACIDS

Hydrazine (or Hydrazinium) Dichloride (or Dihydrochloride), N₂H₄.2HCl, mw 69.53, N 40.30%; wh crysts, sp gr 1.4226 at 20°/4°; mp 198° with dissociation; detonates on ignition, or when heated rapidly; hygroscopic; v sol in w; almost insol in hot absol alc. Can be prepd from hydrazine hydrate and excess of HCl or from hydrazine hydrosulfate and Ba chloride

Refs: 1) Gmelin’s Hdb (1936), Syst 23, p 553


Hydrazine Dihydrochloride. See Hydrazine Dichloride

Hydrazine Dinitrate. See under HYDRAZINE NITRATES

Hydrazine (or Hydrazinium) Hydrate. NH₂NH₂·H₂O; mw 50.06, N 55.97%; exists only in the solid state, contains 64% N₂H₄, bp 118.5° at 739.5mm (Ref 2a, p 100); fr p —51.7°C; density 25°C 1.031g/ml; fl p (open cup 163°F; refractive index 1.4284 at 20° (Ref 4); vapor pressure for the hydrate given in Ref 1. Misc with w & alc; insol in eth & chlif; strong reducing agent, weak base. Its aq soln was first prepd in 1887 by Th. Curtius. In the Raschig process (prior to 1924), NaOH, chlorine and ammonia react in aq soln to form dl soln of hydrazine, with Na chloride as a by-product; also by oxidation of urea by Na hypochlorite. For its purification can be used: fractional distillation, flash distillation or conversion to the slightly sol sulfate, followed by treatment of the latter with concd NaOH soln. Its vapor is expi and toxic; especially dangerous to the eyes

Analytical. Hydrazine content in aqueous solutions is determined by reacting with picryl chloride to form the yellow hexanitro hydrazobenzene, which, with alkali forms red or violet salts that can be measured colorimetrically (Ref 3). More concentrated solution of hydrazine can be titrated with either standard iodine or iodate solutions according to the following equations (Ref 4):

\[ \text{NH}_2\text{NH}_2 + 2\text{I}_2 + 4\text{NaHCO}_3 + 4\text{NaI} + \text{N}_2 + 4\text{H}_2\text{O} + 4\text{CO}_2 \]

or

\[ \text{NH}_2\text{NH}_2 + \text{KIO}_3 + 2\text{HCl} + \text{KCl} + \text{IC} + \text{N}_2 + 3\text{H}_2\text{O} \]

Heat of formation -10.3kcal/mole; heat of combustion +146.94kcal/mole with water liq (Ref 2a, p 28)

Thermal, Combustion and Explosion Properties. Ign temperatures of 85% hydrazine hydrate in pyrex and air was found to be 292°C. Oxygen lowered the ignition point to 218°C. On platinum foil an air atmosphere gave an ignition
temperature of 338°C and oxygen gave 132°C (Ref 1). The hydrate was not found to be sensitive to the impact of a 5kg weight dropped from a height of 1 meter. No explosion, burning or local reaction occurred when the 85% hydrate was subjected to the standard friction sensitivity test in the pendulum-friction machine. No explosion was obtained when hydrazine hydrate was subjected to a detonator in the ballistic mortar. No noticeable decomposition was observed in unconfined samples when a static spark of 12.5 joules was discharged through the liquid (Ref 1).

Used for prepn of anhydrous hydrazine and for the same purposes as listed under Hydrazine, Anhydrous. Also used as gas absorbent (Ref 2a, p 100)

Explosive Compositions. See hydrazine for explosive compositions containing water

Hydrazine Hydrate and Hydrazine Reactions with Explosive Violence.

The following reactions with inorganic substances are taken from the book of Clark, Hydrazine (1953), pp 4 to 18: Potassium reacts with N₂H₄ expl violently (p 4); Mercuric oxide reacts with N₂H₄.H₂O with expl violence (p 5); Chlorine, bromine and iodine react violently (p 8) with N₂H₄ and spontaneous ignition of N₂H₄ takes place in a chlorine atm. Exphs of N₂H₄ and iodine have been reported (p 12).

In Chap III of Clark's book entitled "Organic Chemistry of Hydrazine", it is stated on pp 83–4, that all indications are that anhydrous hydrazine reacts so vigorously with org substances that products will not be the same as with hydrazine hydrate or the salts. Anhyd N H is not used for making org hydrazides because it attacks other groups

Hydrazine (or Hydrazinium) Hydrochloride. See Hydrazine Chloride

Hydrazine (or Hydrazinium) Hydrosulfate
(Hydrazinium Acid Sulfate — so called by Clark (Ref 2) and its formula is given as N₂H₄.HSO₄). The same compd is called Hydrazine Sulfate in Lange (Ref 3) and in CondChemDict (Ref 4), where its formula is given as N₂H₄.H₂SO₄, which amounts to the same mw for both formulas equal to 130.13 and the same props: wh crysts, sp gr 1.378, mp 254°C (with evolution of gas); stable in storage, but contact with alkalies and oxidizing agents must be avoided; explodes with NaNH₂; v sol in hot w; diffic sol in cold w; insol in alc. Can be prepd by mixing hydrazine hydrate soln with sulfuric acid. It is a strong reducing agent. Used for prepn of other hydrazine salts and derivs; also in analysis of minerals and ores; in tests of blood; for separation of polonium from tellurium; as fungicide & germicide; and in adhesives

Hydrazine Mononitrate. See under HYDRAZINE NITRATES

Hydrazine (or Hydrazinium) Nitrite. N₂H₄.HNO₂; mw 79.06, N 53.15%; decomp or explodes on rapid heating; colorless to yellowish hygr solid; sol in w & alc; insol in eth; may be prepd by mixing solns of barium nitrite and neutral hydrazine sulfate, as described in Mellor (Ref 1). Explodes violently on impact and less so when rapidly heated. When heated slowly it decomposes according to the equation:
N₂H₄•HNO₂ → NH₃ + N₂O + H₂O and this decomp is greatly accelerated by nitrous acid

HYDRAZINE (OR HYDRAZINIUM) NITRATES
Hydrazine Mononitrate, HN, H₂N, hydrazinium nitrate. N₂H₄.HNO₃ or N₂H₄NO₃; mw 95.07; N 44.20%, OB to N₂ and H₂O +8.4%; white monoclinic tablets and rods existing in 2 forms,
α & β. β form is unstable, mp 62°; α form, stable, mp 70°; crystal axial ratio a:b:c = 0.957:1.0:0.492; interfacial angle (polar) 88°26'; crystal angle 90° (Ref 12); d 1.661. Mêdard (Ref 8) determined density at various pressures as follows: pressure in kg/sq cm, density: 34, 1.08; 88, 1.18; 136, 1.28; 204, 1.43; 306, 1.49; 425, 1.58; 680, 1.61; 1020, 1.635; 1700+, 1.64. If a conc'd aq soln is cooled from 100°, silky needles of β-form separate, but they undergo monotropic transition to α-form with evolution of heat at room temp. Soln in g/100g H₂O at 10°, 174.9; 20°, 266.3; 30°, 402.2; 40°, 607.2; 50°, 1034.0; 60°, 2127.0 (Ref 6)

Thermodynamic data. ΔH combustion (const vol) 112.1 kcal/kg; ΔH comb (const press) 111.1 kcal/kg; ΔH_f (const vol) 56.7 kcal/kg; ΔH_f (const press) 59.8 kcal/kg (Ref 10); ΔH transition β to α form 2.0 kcal/mole; ΔH_soln (H₂O) -8.72 kcal/mole; ΔH_soln (N₂H₄) +3.7 kcal/mole (Ref 21). Refractive ind at 5983Å & 25°C, α 1.605±0.004 and 1.620±0.005, β 1.458±0.003 (Ref 12). Density and viscosity curves of HN solns with hydrate and water are given in Ref 21

Preparations. Add HNO₃ by drops (to anhyd N₂H₄ dissolved in MeOH) to pH of 5.5. HN pts and is filtered off (Ref 10). May also be prepared from hydrazine carbonate and HNO₃, from N₂H₄·OH and HNO₃ or from N₂H₂·H₂SO₄ and Ba(NO₃)₂ (Ref 6). Weiss (Ref 16) reacted N₂H₄ and N₂O₃ at -133°, forming HN

Toxicology. Similar to hydrazine (Ref 24)

Analytical. Infrared absorption and x-ray diffraction spectra are given in Ref 21

Thermal, combustion and explosion properties. Decomposes on heating at 1 atm to 200-300° as follows: 4N₂H₂NO₃ +2NO + 5N₂ + 10H₂O (Ref 9). Mêdard (Ref 8) measured wt loss on heating at 100° as compared with Ammonium Nitrate: HN lost 0.005%/hr of its weight at a const rate as compared to 0.016%/hr for AN. Shidlovskii et al (Ref 14) in studying thermal decomp of function of temp found the decomposition started at 180°, increased rapidly above 240°, at 270° became explosive. The ignition temp detd by the “Bruce ton up-and-down” method which gave a 50% probability of ignition was 307° (Ref 21). Whittaker and Barham (Ref 15), using high-speed photography, measured average surface temp and melting point for NH₄NO₃: 303° ± 12° and 166° ± 10°; for HN: 195° ± 9° and 74° ± 5°; and for an NH₄NO₃—HN eutectic: 307° ± 11° and 39° ± 6°. Equilibrium flame temp of HN detonations have been reported to be 2400°C (Ref 21)

In the open air, HN supports combustion, but the flame was extinguished upon removal of the ignition source. To achieve stable burning Shidlovskii (Ref 14) found that addition of 10 wt-% of K₂Cr₂O₇ was necessary. HN ignites with permanganate, chromate-or peroxides. At slightly above its mp it will ignite with metallic Zn, or Cu, or their oxides, sulfides, nitrates or carbides (Ref 9)

Sensitivity to initiation: non-compressed HN with 0.5% moisture content, detonated with a cup contg 0.25g Mercury Fulminate; compressed to d 1.60, 1.5g MF was required for detonation (Ref 8)

Sensitivity to impact: values of 175 kg/cm and 200 to 150 kg/cm for 50% probability are reported for the “Bruce ton up-and-down” method; values of 32 kg/cm and 50 kg/cm are reported for the ERL type 12 tool test (Ref 21)

Card gap sensitivity: 6.25cm; brisance 82 (TNT = 100); Trauzl lead-block test 120.4 ± 0.5 (picric acid = 100); TNT equivalence = 1.42

Detonation velocity: Bureau of Mines detd det velocity of molten HN at 75° in thin-film expts to be 8500 m/sec (Ref 21)

Jacobs (Ref 4), with a rotating mirror camera, measured a detonation velocity of 5200 m/s with a 1" diam cartridge, 5600 m/s with a 1 5/8" diam cartridge, and 8500 m/s with a 2½" diam cartridge (d of 1.6 for pressed material)

Mêdard (Ref 8) using a 30mm diam and 170mm long cartridge found a maximum velocity of 5640 m/sec at a d of 1.25 g/cm³. Price et al (Ref 17) reported a det velocity of 8510 m/sec at d 1.59 g/cm³ for a 6.3 cm diam charge of pressed HN. Price and co-workers found that the infinite charge diam detonation velocity, D (in m/sec) for HN can be expressed as D = 5390 (ρHN-100) in which ρHN is the d of HN in g/cm³.
Kurbangalina et al (Ref 22) found that critical diam increases with increasing water content, and that molten HN & its aq solns can detonate with different velocities (2-8 km/se). Low-velocity detonations are less dependent on critical diam than high-velocity detonations.

**Explosive compositions.** Audrieth (Ref 11) patented an explosive compn of NH₄NO₃ 78.5 - 83.5%, HN 5-10%, TNT 7.5%; Al dust 0.5%; coal 3% and chalk 0.5%. Increasing HN content to 10% increases sensitivity and detonation rate. Bridgeforth et al (Ref 20) patented explosive compns with high detonation rate and high air blasts (up to 2-fold TNT) which contain N₂H₄ and HN with or without NH₃ + NH₄NO₃ + finely divided Al and a thickening agent. These compns have good cratering performance, comparable to TNT. Méardin (Ref 8) examined properties of HN mixtures with NH₄NO₃, Guanidine Nitrates, Nitroguanidine, and Ethylenediamine Nitrates. Also examined (Ref 8) was a mixt of 6% HN in a commercial explosive of NH₄NO₃ 87.5% and 12.5% wood flour. The original mixt of d = 0.80 required 0.80g MF for complete detonation, while the one with HN, d = 0.90 required only 0.30g MF for detonation. The gap test was 1.3cm for orig mixt and 4cm for mixt contg 6% HN. Incorporation of HN renders NH₄NO₃ explosives less hydroscopic. Thermal stability of HN at temps 100-200° is satisfactory and it can be incorporated in explosives loaded by casting. The low mp of HN lowers the casting temp of NH₄NO₃ mixtures (Ref 8).


**Hydrazine (or Hydrazinium) Dinitrate.**

N₂H₄·2HNO₃, mw 158.08, N 35.45%, OB to N₂ and H₂O 30.4%; crystals, mp 103–104° when heated rapidly. When slowly heated it decomps at 80-85° without melting, yielding among other decom products hydrazinide. Even in a desiccator over sulfd acid and at ord temp it decomps with evolution of hydrog azide, leaving a residue of N₂H₄·HNO₃ and NH₄NO₃. Readily sol in w, butaq solns contg more than 30% decom on heating, d 1.64. Prepd by treating 1 mole of hydrazine hydrate with 2 moles of nitric acid. It was examined in US during WWII and found to be a more powerful explosive than Tetryl but less powerful than PETN. It is slightly less sensitive to impact than Tetryl.

**Ref:** 1) Sabannejeff, ZAnorgChem 20, 21 (1899); not in CA 2) W.R.E. Hodgkinson, JSoChemInd 32, 519 (1913); not in CA 3) F. Sommer, ZAnorgChem 86, 71 (1914) & CA 8, 1932 (1914) 3a) Gmelins Hdb (1936), Syst 23, p 551 3b) Blatt, ORSD 2014 (1944) 4) Mellor 8, 327 (1946) 4a) ADL,
Hydrazine Nitrate Complexes. Salts of bivalent metals (Ni, Co, Zn, Cd and Mn) of complex hydrazine nitrates may be represented by the following formulas: $[\text{M(N}_2\text{H}_4\text{)}_2\text{]}(\text{NO}_3\text{)}_2$, or by $[\text{M(N}_2\text{H}_4\text{)}_3\text{]}(\text{NO}_3\text{)}_2$. In these complexes, called nitrohydrazinates, each $\text{N}_2\text{H}_4$ group plays the same role as two NH$_3$ groups in the corresponding ammonium complexes.

For Cu and Hg the only known nitrohydrazinates contained one $\text{N}_2\text{H}_4$ group.

Most of these complexes are stable in the dry state and remain so on heating to temperatures of at least 110°. At higher temperatures the complexes explode, some of them violently. They are nearly insol in water but are more or less hydrolyzed by it.

The complex of Ni was first prepared by Franzen and Mayer (Ref 1) and the formula $[\text{Ni(N}_2\text{H}_4\text{)}_2\text{]}(\text{NO}_3\text{)}_2$ was assigned to it. The same investigators prepared the nitrohydrazinates of Cd, Cu and Co.

Most of these complexes were prepared by the action of aqueous soln of hydrazine on ammoniacal complexes of the nitrates of Co, Ni, etc (contg six NH$_3$ groups). Mâder and Barlot (Ref 3) proposed the following procedure:

A soln of 20-30pts of a nitrate in 100pts of water (or preferably in ethanol) was gradually added to a mechanically agitated soln of 50pts of hydrazine hydrate in 100pts of ethanol, cooled to ca 10°. The amount of reagents should be calculated in such a manner that the final mixture contains 4 to 5% of excess of hydrazine.

As soon as all the nitrate was added the mixture was poured into a Büchner funnel connected to a suction flask. The resulting precipitate was washed with methanol or ethanol until the elimination of hydrazine was complete and dried first in air and then in a steam oven at 100°.

Note: During the investigation of various methods of preparation, it was observed by Barlot (Ref 2) that if solns of equimolecular amounts of nickel nitrate and hydrazine nitrate are mixed no precipitation is observed. If, however, some hydrazine, or ammonia is added, an immediate precipitation of pink complex is observed.

Cadmium Nitrohydrazinate. Cd(NO$_3$)$_2$·3N$_2$H$_4$.
Calcd: Cd 33.7% and N$_2$H$_4$ 28.9; found: Cd 33.1% and N$_2$H$_4$ 28.5%. White cryst powder, explodes on strong impact (such as of 2kg wt falling from the height of 3m) or on rapid heating to ca 245°; when spread in a thin layer it burns while melting and evolving brown fumes (Ref 3)

Cobalt Nitrohydrazinate. Co(NO$_3$)$_2$·3N$_2$H$_4$.
Calcd: Cd 21.0% and N$_2$H$_4$ 34.5%; found: Cd 22.1% and N$_2$H$_4$ 32.5%. Brownish (bister color) microcryst powder; explodes violently on impact (such as of 2kg wt falling from the height of 1.75m) or on heating to ca 210°. Addn of 25% water decreases sensitivity to drop test. Explodes also on contact with concd acid. It is unstable and decomposes slowly even in the presence of traces of water changing to a greenish color (Ref 3)

Manganese Nitrohydrazinate. Mn(NO$_3$)$_2$·2N$_2$H$_4$.
Calcd: Mn 22.8% and N$_2$H$_4$ 33.6%; found: Mn 23.6% and N$_2$H$_4$ 33.5%. Reddish (ochre) colored hexagonal crystals; ignites easily ca 150°, produces copious fumes of Mn oxide; is insensitive to shock (Ref 3)

Nickel Nitrohydrazinate. Ni(NO$_3$)$_2$·3N$_2$H$_4$.
Calcd: Ni 21.0% and N$_2$H$_4$ 34.5%; found: Ni 21.3% and N$_2$H$_4$ 34.0%. Rose-violet colored microcrystals; explodes violently on impact (such as of 2kg wt falling from the height of 1.50m) or on heating to ca 215°; the explosion is accompanied by a loud noise and a brilliant flash. Addn of 25% water decreases sensitivity to drop test. Power in the French lead block test (cup) ca 85 (PA 100) which is higher than for MF or LA. In cellophane tubes 200mm long at d = 0.62 (MF detonator), the velocity of detonation by the Dautriche method for various tube diams were: 2600 m/sec at 6mm diam; 2900 and 3100 at 8mm; 2700 at 10mm; 2900 at 12mm; and 3500 at 15mm (Ref 3)
Ellern and Olander (Ref 4) report a spontaneous explosion of a thoroughly washed and dried nickel nitrohydrinate \( [\text{Ni}(\text{N}_2\text{H}_4)_2 \cdot \text{(NO}_3)_2] \) after exposure to the atm for 10 min.

Barlot and Médard (Refs 2 & 3) describe an analytical procedure for nickel nitrohydrinate: decompose the complex cautiously (to avoid an explosion) by means of concd \( \text{H}_2\text{SO}_4 \) and determine Ni content by means of dimethylglyoxime. Determine the hydrazine content by means of an iodate soln.

Uses: The possibility of using Ni nitrohydrinate in commercial detonators was investigated by loading Cu caps of 5mm diam with 2.5g of Ni complex and trying to detonate them by means of a fuse, or an electric igniter. No detonations were produced unless the substance was primed by ca 1.5g of Black Powder.

In attempts to increase the detonability of Ni hydrazinate, it was mixed with equal parts of an easily ignitable initiating substance, such as Pb-styphnate or Pb-picrate, and then ignited. The best that could be obtained was a deflagration but no detonation. Mixtures of the Ni complex with RDX deflagrated and only in one case detonated (Ref 3).

**Zinc Nitrohydrinate.** \( \text{Zn(NO}_3)_2 \cdot 3\text{N}_2\text{H}_4 \).
Calcd: Zn 22.8% and \( \text{N}_2\text{H}_4 \) 33.6%; found Zn 23.0% and \( \text{N}_2\text{H}_4 \) 33.5%. White crystalline powder (apparently hexagonal); explodes occasionally under very strong impact (less sensitive than any of the above complexes with the exception of Mn); ignites with crackling and greenish sparks on heating to ca 310°C (Ref 3).

**Conclusions.** The following conclusions about the explosive properties of complex metallic nitrohydrinates were drawn by Médard and Barlot (Ref 3):

a) the complexes of Ni, Co and Cd possess properties similar to primary and initiating compounds, such as Mercury Fulminate.

b) they are much less sensitive to shock than Mercury Fulminate or Lead Azide and are more powerful.

c) they are much more difficult to ignite by means of a fuse, or an electrical igniter, than Mercury Fulminate and even when ignited they do not detonate but only deflagrate. This is also true, in a lesser degree, for mixtures of above complexes with more ignitable initiating compound, such as Lead Styphnate.

d) The complex Styphnate of Ni hydrazinate also deflagrated & did not detonate.

**Refs:**

**Hydrazine (or Hydrazinium) Perchlorate,** Hydrazinium Perchlorate, \( \text{N}_2\text{H}_4\text{HClO}_4 \) or \( \text{N}_2\text{H}_4\text{ClO}_4 \); mp 132.51; N 21.13%, OB to \( \text{Cl}_2 \) and \( \text{H}_2\text{O} \) +18.1%; orthorombic crystals; for a study of crystal structure see Conant and Rooj (Ref 16); mp 131–132°C; d 1.939 (dipercarbonate d 2.21, mp 191°C); solv in \( \text{H}_2\text{O} \): temp, g/100g satd soln: 0°C, 23.68; 40°C, 68.9; 60°C, 87.4; 75°C, 93.1 (Ref 3). Carleton & Lewis (Ref 11) studied the phase equilibria of Hydrazine Perchlorate—water system and observed: the limiting melting temp of the salt, 142.4°C; the occurrence of a solid hydrate decomposing at 64°C with a eutectic point at -4.1°C and 80% \( \text{H}_2\text{O} \); hydrate identified as \( \text{N}_2\text{H}_4\text{ClO}_4 \cdot \frac{1}{2}\text{H}_2\text{O} \); quadrupole pt 64°C and 88mm Hg. \( \Delta \text{H} \) of addn of \( \text{N}_2\text{H}_4 \) to \( \text{HClO}_4 \) = -44.4 kcal/mole; \( \Delta \text{H} \) of addn of \( \text{HClO}_4 \) to \( \text{N}_2\text{H}_4 \cdot \text{HClO}_4 \) = -18.8 kcal/mole; \( \Delta \text{H}^2 = 40.69 \pm 0.36 \text{ kcal/mole (Ref 15)} \) or \( -42.9 \pm 0.2 \text{ kcal/mole (Ref 5)} \); \( \Delta \text{H}^2 \) for dipercarbonate = -70.1 ± 1 (Ref 7); \( \Delta \text{H}_{\text{soln}} \) = 33.11 kcal/mole; \( \Delta \text{H} \) diln 1:1000 = ±9.77 kcal/mole.

**Preps:** May be prep’d by addn of concd \( \text{HClO}_4 \) to aq soln of \( \text{N}_2\text{H}_4 \) neutralizing and cooling to 0°C (Ref 3). Byrne (Ref 14) prep’d anhyd perchlorate by metathesis of a hydrazine salt with an inorganic perchlorate dissolved in a lower aliphatic alcohol. Stern (Ref 6) prep’d the hemihydrate by adding \( \text{N}_2\text{H}_4 \) to an aq soln of \( \text{NH}_4\text{ClO}_4 \), heating to remove \( \text{H}_2\text{O} \) and \( \text{NH}_3 \), and cooling below transition temp to crystallize.

**Analytical:** Jacobs & Russell-Jones (Ref 12) detd the IR spectrum and assigned absorption bands.

**Toxicology:** Unknown. Probably tox. (Ref 18)
Use: Rocket propellant (Ref 17)

**Thermal, combustion and explosion properties:**

Badot and Marsoule (Ref 3) reported the salt melting to a colorless liquid at 137.8°, with decompn beginning at 145°, becoming complete at 230°. Grelechi and Cruice (Ref 9) using a manometric technique to study thermal decompn, found the reaction rate to be proportional to the amt of free HClO₄ present. Shidlovskii et al (Ref 5) det the rate of thermal decompn at different temps. They also measured the vol of gases evolved in decompn to be 864 l/kg.

Jacobs and Russell-Jones (Ref 12) studied the thermal decompn of the hemihydrate. In vacuo it will partially dehydrate even at room temp; when heated, the remainder of the 0.5 mole H₂O is lost at 61°. At low temps, decompn is represented by the equation:

\[ 2 \text{NH}_4\text{ClO}_4 \cdot 0.5 \text{H}_2\text{O} + 0.8 \text{NH}_3 + 0.8 \text{HClO}_4 + 0.7 \text{O}_2 + 1.6 \text{N}_2 + 0.6 \text{Cl}_2 + 4.4 \text{H}_2\text{O} \]

but at high temp the rate of decompn of NH₄ClO₄ formed becomes appreciable. When ignited, hydradine perchlorate burns without exploding. The flame term is 2200°

Friedman and Levy (Ref 8) and Levy et al (Ref 13) investigated deflagration behavior.

Levy et al (Ref 10) studied deflagration of hydradine perchlorate both pure and with fuel and catalyst additives (Cu chromite, K₂Cr₂O₇ and MgO). Deflagration rates were measured photographically from 0.26 to 7.7 atm.

Shidlovskii et al (Ref 5) studied ignition behavior. Ignition temp, 277-80°, was lowered to 254-9° by addn of 5% MnO₂, and addn of 5% CuCl₂ resulted in explosion at 170°. In 1.05cm diam tubes hydradine perchlorate did not ignite, but with 5% MnO₂, CoO or Cu₂Cl₂ it maintained self combustion at room temp. The rate of burning in the presence of additives decreased in the order Cu₂Cl₂ > CoO > MnO₂. Self combustion was not maintained in the presence of 5% CuCl₂·2H₂O, but in the presence of 2.5% CuCl₂·2H₂O and 2.5% CuCl₂ about 2/3 of the perchlorate burned out before the residue exploded. The rate of burning of the perchlorate was 2-3 times as high as that of NH₄ClO₄. Hydradine perchlorate is an explosive comparable in power to the chlorate but is more stable. Salvadori (Ref 1) proposed using it in primary and initiating mixtures. Blatt (Ref 1b) gives its power by Trauzl Test as 113% PA, while ADL (Ref 1c) gives 122% TNT. Impact sensitivity FI 12% PA (Ref 1b).

In air it forms a semi-hydrate, mp 85°, which is insensitive to impact (Ref 1b).

**Refs:**
1. R. Salvadori, GazzChimItal 37 (2)

**Hydradine Perchlorate and Chlorate Complexes.**

Salts of hydradine chlorate and perchlorate complexes, such as [Co(N₂H₄)₂]

<sub>3</sub>ClO₃<sub>2</sub>, yellow cryst, and [Co(N₂H₄)₂](ClO₃)₂, were first prepd by Salvadori (Ref 1). Friederic and Vervoort (Ref 2) describe a number of complexes (Cn, Cd, Ni, Co & Zn) prepd by adding a soln of hydradine hydrate in H₂O or EtOH to a soln of a metallic chlorate or perchlorate with cooking and agitation, and washing and drying the ppt at low temperatures. For example, [Ni(N₂H₄)₂](ClO₃)₂, also Cd
and Cu chlorates and perchlorates, were prepd and their explosive characteristics were studied. They were found to be powerful and sensitive explosives, claimed to compare favorably with standard initiating explosives. Violent decomp occurs, presumably in accordance with the following hypothetical equations:

\[ \text{Me(N}_2\text{H}_4\text{)}_2 (\text{ClO}_3)_2 \rightarrow \text{MeCl}_2 + 2\text{N}_2 + 4\text{H}_2\text{O} + \text{O}_2 \]
\[ \text{Me(N}_2\text{H}_4\text{)}_2 (\text{ClO}_4)_2 \rightarrow \text{MeCl}_2 + 3\text{N}_2 + 6\text{H}_2\text{O} \]
\[ \text{Me(N}_2\text{H}_4\text{)}_2 (\text{ClO}_4)_2 \rightarrow \text{MeCl}_2 + 2\text{N}_2 + 4\text{H}_2\text{O} + 2\text{O}_2 \]
\[ \text{Me(N}_2\text{H}_4\text{)}_3 (\text{ClO}_4)_2 \rightarrow \text{MeCl}_2 + 3\text{N}_2 + 6\text{H}_2\text{O} + \text{O}_2 \]

The chlorates are more sensitive than the perchlorates (Ref 2); the Cu-chlorate complex detonates even on drying at room temperature. Maissen & Schwarzenbach (Ref 3) attempted to prepare \([\text{Ni(N}_2\text{H}_4\text{)}_2 (\text{ClO}_4)_2 \] by mixing Ni(ClO4)2 and N2H4 in water. After 5 days a blue ppt formed. When a glass stirring rod was introduced into the suspension, a violent explosion resulted.

Refs: 1) R. Salvadori, Gazz. 40, II, 9 (1910) & CA 5, 1568 (1911); Gazz 42, 448 (1912); JCS 98, III, 959 (1910) 2) W. Friederich & P. Vervoorst, SS 21, 49 (1926) & CA 21, 1184 (1927) 3) B. Maissen & G. Schwarzenbach, HelvChimActa 34, 2084 (1951) & CA 46, 3280 (1952)

**Hydrazine Picrate.** See under HYDRAZINE SALTS OF ORGANIC ACID

**HYDRAZINE, REACTIONS WITH METALS AND THEIR SALTS.**

Lithium dissolves slowly in anhydrous hydrazine with evolution of hydrogen and nitrogen. In the presence of N2H5.HSO4 the reduction is much more rapid.

Ref: Clark, Hydrazine (1953), p 4

**Mercuric Oxide.** Explosive action betw HgO and hydrazine hydrate was observed by Curtius & Schrader.


**Potassium reacts with anhydrous hydrazine with explosive violence.**

Refs: 1) T.W.B. Welsh & H.J. Broderson, JACS 37, 816–24 (1915) 2) Clark, Hydrazine (1953), p 4

**Sodium reacts with aqueous or anhydrous hydrazine with formation of NaNH2.NH2.** When dry the salt is extremely explosive.

**HYDRAZINE SALTS OF ORGANIC ACIDS**

**Hydrazine-N,N'-dicarboxylic Acid Diazide**

(Hydrazin-N,N'-dicarboxystyrene, in Ger)

N3C.CH.NH.C.N3; mw 170.10, N 65.88, OB

\[ \text{O} \]

\[ \text{O} \]

to CO2 –28.2%; crysts, mp 150–52 (decomp) and explodes if heated rapidly. Was obtd in 20% yield as by-product of the action of nitrous acid on the hydrazide of dicarboxylic acid. It is primary type expl more sensitive to impact than Lead or Silver Azides. One g of TNT can be initiated by 0.25g of this expl


**Hydrazine picrate, N2H4-C6H2(NO2)3.OH;**
mw 261.2, N 26.8%; OB -52.0%; sony detd by Gilbert & Huffman (Ref 1) in EtOH–H2O mixtures. Soly curve showed maxima & minima but no evidence of a stable alcohohle. Water of hydration was retained even in 99.8 % EtOH.

Forth (Ref 3) studied thermal stability of amine salts of Picric Acid; and rated, in order of increasing stability the following salts: Hydrazine Picrate; Aminoguanidine Picrate; N-Methylguanidine Picrate; Guanylurea Picrate; N-Ethylguanidine Picrate; and Guanidine Picrate.

Christensen & Gilbert (Ref 2) plotted the vapor pressure curve of the hydrated picrate, N2H4-C6H2(NO2)3.OH·2H2O; np to its melting point and found no break in the curve. It holds the water of hydration very tenaciously and begins to have an appreciable vapor pressure only at temps around 100°; calculated ΔH of dehydration is 16.855 kcal/mole and calculated ΔF298.1 is 2705 cal. No explosive props mentioned, but is evidently explosive.

Hydrazine Stypnate and Hydrazine Stypnate complex. Fauth (Ref 2) examined relative stability in terms of the temp of explosion of amine salts of stypnic acid, and rated, in order of increasing stability, the following salts: Hydrazine Stypnate; N-Methylguanidine Stypnate; N-Ethylguanidine Stypnate; Guanidine Stypnate; Guanyleurea Stypnate; and Aminoguanidine Stypnate. The impact sensitivity of the stypnates as indicated by the 5kg drop test, in order of increasing sensitivity was: Aminoguanidine Stypnate; Guanidine Stypnate; N-Ethylguanidine Stypnate; N-Methylguanidine Stypnate; Guanyleurea Stypnate and Hydrazine Stypnate

Medard & Barlot (Ref 1) prepd a small quantity of a Hydrazine Stypnate complex of Ni (no formula given) by dissolving freshly pptd Ni hydroxide in an aq soln of stypnic acid and treating the soln with aq hydrazine. The resulting ppt was dried and tested for following properties: Flammability: the complex, either alone or in mixts with RDX, was just as difficult to ignite by means of a fuse, or an electrical igniter as was the corresponding complex Ni-nitrohydrazinate. When ignited, the complex stypnate did not detonate and for this reason is considered to be unsuitable for use in detonators. Sensitivity to impact: explosions were obtained 20% of the time on dropping a 2kg hammer from the height of 1.5m, and 40% of the time from a height of 2.5m 


HYDRAZINE SUBSTITUTED DERIVATIVES:

Alkyl Substituted Hydrazines & Derivs

Of the many alkyl substituted hydrazines, only two are used as rocket fuels now: monomethyl hydrazine (MMH), CH₃NH₂NH₂, and unsymmetrical dimethylhydrazine (UDMH), (CH₃)₂NNH₂. (See Hypergolic Fuels in this Vol for combustion properties as rocket fuels). Their physical properties are summarized below

Monomethyl hydrazine, MMH, CH₃NH₂NH₂, mw 46.07; N 59.7%; fp -52.4⁰C, bp 87.5⁰C, d 0.871 at 26⁰C, heat of formation +12.7kcal/mole, heat of combstn-311.7kcal/m mole, explosive limits in air 2.5 to 98%, vapor pressure 49.63mm Hg at 25⁰C, viscosity 19.0cPs at -55⁰C. For other props see below. May be prepd by the alkylation of hydrazine:

CH₃Cl + 2N₂H₅ → CH₃NH₂NH₂ + N₂ + H₂Cl or by reaction of NH₂Cl with methylamine (Ref 3). Liquid MMH is not sensitive to impact or friction. It is hypergolic with some oxidants such as H₂O₂, N₂O₄, fluorine and nitric acid. When a film of MMH comes into contact with metallic oxides, such as those of iron, copper, lead and manganese, it may ignite owing to the chemical heat of decomposition (Ref 5)

Unsymmetrical Dimethyldihydrazine, UDMH, (CH₃)₂NNH₂, mw 60.10; N 46.62%, density 0.784 at 25⁰C; fr p -57⁰C; bp 63⁰C; Refr Index 1.4058 at ca 25⁰C; fl p (Tag CC) 34⁰F; vap press 157mm at 25⁰C; viscosity 0.51cPs at 25⁰C; flammability limits-15⁰ to ca 60⁰C; spontaneous ignition in air 250⁰C; heat of form (endothermic) 12.74kcal/mole at 25⁰C; heat of combstn-474.11kcal/mole; impact sensitivity—not detonable by 100g Tetryl chge; may be prepd by reduction of nitrosodimethylamine (Ref 3). Korabina et al (Ref 6) prepd the perchlorate salt of UDMH by reacting the base with HClO₄ at 50-60⁰C, cooling or evap to deposit crystals. Resulting salt is sol in H₂O and detonates easily.

UDMH is known to be miscible with the following: water, benzene, triethyl benzene, toluene, kerosene, ethyl alcohol, isobutyl alcohol, n-butyl ether, n-amyl ether, n-hexyl ether, diethyl ether, petroleum ether, petroleum naphtha, n-heptane, n-hexane, n-octane, n-decane, n-dodecane, n-hexadecane, cyclohexane, 1,2-dimethyl cyclohexane, phenyl cyclohexane, n-tetradecane, trichloroethylene, dichloroethylene, perchloroethylene, 1,1,1-trichloroethane, triethyl amine, ethylenediamine, diethylenetriamine, acetonitrile, aniline, cumene, tetrahydroxiphenyrene, tetraethylene pentamine, ethylene glycol and hydrazine (Ref 4)

UDMH should always be handled under a blanket of N (Ref 4)
Other alkyl hydrazines. Picard & Boivin (Ref 2) studied the action of 100% HNO₃, mixed at -30° with acetic anhydride, on various symmetrical disubstituted hydrazines. Westphal & Euhren (Ref 1) studied in great detail the properties and thermal decomposition of various tetra-alkyhydrazines


Hydrazine Substituted Aryl Derivatives. N,N′-dimtro-N,N′-dibenzoylhydrazine. PhCON(NO₂)₂-\( \text{N(NO}_2\text{)}\text{CO}_2\), mp 101.5°, may be prepared by reaction of nitric acid-acetic anhydride mixtures with N,N′-dibenzoylhydrazine; reacts with concd H₂SO₄, giving off nitrogen oxides (Ref 3)

2,4-dinitrophenyldihydrazine. 2,4-(NO₂)₂PhNH₂NH₂, may be prepared by reaction of 2,4-(NO₂)₂PhCl, AcOK and \( \text{N}_2\text{H}_4\cdot\text{H}_2\text{SO}_4\) (Ref 1); \( \Delta H_{\text{comb}}\) 3950 cal/g at const press, \( \Delta H_f\) 9.2 kcal/mole at const press (Ref 4)

2,6-dinitrophenyldihydrazine. 2,6-(NO₂)₂PhNH₂NH₂, may be prepared from 2,6-(NO₂)₂PhCl, AcOK and \( \text{N}_2\text{H}_4\cdot\text{H}_2\text{SO}_4\) (Ref 1)

2,4,6-trinitrophenyldihydrazine. 2,4,6-(NO₂)₃PhNH₂NH₂, may be prepared from picryl chloride, AcOK and \( \text{N}_2\text{H}_4\cdot\text{H}_2\text{SO}_4\) (Ref 1); \( \Delta H_{\text{comb}}\) 3070 cal/g at const press, \( \Delta H_f\) 5.8 kcal/mole (Ref 4)

\( \alpha\cdot\text{(2,4,6-trinitrophenyl)-6-methylhydrazine.} \)

2,4,6-(NO₂)₃PhN(CH₃)₂NH₂; crystals; mp 170°, may be prepared by reacting MeNH₂NH₂·H₂SO₄ with NaOH, filtering, and reacting filtrate with picryl chloride (Ref 2)

Written in collaboration with E. L. CAPENER R. M. WRIGHT


Hydrazine (or Hydrazinium) Sulfate. The name given in Clark (Ref 4) to the compound which is actually Dihydrazinium Sulfate (\( \text{N}_2\text{H}_4\cdot\text{SO}_4\cdot\text{H}_2\text{O} \)). It is prepared either by neutralization of hydrazine hydrate solution with sulfuric acid or by neutralization of hydrazine hydroxysulfate, followed by evaporation of the latter solution at 60° to obtain crystals of Dihydrazinium Sulfate, monohydrate, melting at 118.9°. Anhydrous salt can be obtained by slow cooling but in contact with soln it is stable only above the transition point, 47.3°; storage over \( \text{P}_2\text{O}_5\) gives anhydrous salt. The monohydrate is hygroscopic but insol in alc. Moist crystals have neutral pH. Can be used for preparation of other hydrazines (See also Hydrazine (or Hydrazinium) Hydrosulfate)

Refs: 1) Gmelins Hdb (1936), Syst 23, p 560
3) B.E. Christensen & E.C. Gilbert, JACS 56, 1897099 (1934)
4) Clark, Hydrazine (1953), 10

Hydrazine and Derivatives, Analytical Procedures.

The following procedures are described in the book of Clark, Hydrazine (Ref 5): p 114–Direct Iodate Method Using Solvent, originally described in Ref 4

pp 114–15–Direct Iodate Method with Internal Indicator, originally described in Ref 4

p 115–Sulfate in Hydrazine by the method developed by Mathieson Chem Corp

pp 115–16–Ammonia in Hydrazine by the method developed by Mathieson Chem Corp

pp 116–17–Iron in Hydrazine. Colorimetric method with \( \alpha\cdot\text{Dipyridyl, earlier described in Refs 1 & 2} \)

pp 117–19–Sulfides in Hydrazine, earlier described in Refs 2 & 3

p 119–Chloride in Hydrazine by the method developed by the Mathieson Chem Corp

The method for estimation of hydrazine content in unknown materials is described in
3.3 Filter. A filter with a 10-micron nominal and 40 absolute rating shall be installed between the manufacturer's plant system and the container to be filled for delivery.

3.4 Qualitative. The propellant shall be a colorless, homogeneous liquid when examined visually by transmitted light.

4. QUALITY ASSURANCE PROVISIONS.
See in Specification

4.5 Test Methods

4.5.1 Examination of product. The propellant shall be visually examined while performing test specified in 4.5.3 to determine compliance with the requirement as specified herein. Examination to ensure that the material conforms to 3.4 shall be conducted after the sample has been transferred to the 500-ml calibrated cylinder

4.5.2 Hydrazine assay and water. The propellant and water content of the sample shall be determined by the following method

4.5.2.1 Gas chromatographic method

4.5.2.1.1 Suggested column preparation. Weigh 10 grams of polyethylene glycol 400 and 20 grams of Anakrom B support material into separate beakers. Dissolve the polyethylene glycol 400 in reagent grade methylene chloride (dichloromethane). The final volume of the solution should be approximately that of the support material. Pour the support material into the polyethylene glycol 400 (stationary phase) solution with gentle stirring. Evaporate the solvent by spreading the mixture in a tray. Occasionally turn the mixture gently during the drying process. The column packing material is dry when it becomes a free-flowing powder. Fill a 1/4 inch x 1 meter (39.4 inches) stainless steel tube by pouring the prepared material thru a small funnel attached to one end. The bottom of the tubing is plugged with a small wad of glass wool. Tap gently or use a mechanical vibrator to facilitate packing. When the column is full, plug the inlet with glass wool and bend the tubing to the configuration required by the chromatograph oven

4.5.2.1.2 Procedure. Install the prepared column into the gas chromatograph but do not connect the column to the detector inlet. Condition the column for at least 4 hours by heating at 302°F (150°C) with the carrier gas (helium) flow set at approximately 50 milli-
liters per minute. After conditioning the column, set the column oven temperature at 230°F (110°C) and connect the column to the detector inlet. Adjust the carrier gas flow to 80–120 milliliters per minute. If the gas chromatograph is equipped with separate injector and detector temperature controls, set the detector at 302°F (150°C) and the injector at 230°F (110°C). Saturate the column by injecting two or three 10-microliter samples of propellant. Saturate the column prior to each series of analyses. When more than 30 minutes elapse between the elution of the hydrazine and the injection of a new sample, resaturate the column with one 10-microliter injection of propellant sample. Use a clean, dry 10-microliter hypodermic syringe and draw up 8 microliters of sample. Invert the syringe and expel the gas bubbles. Carefully set the syringe plunger to the 3-microliter mark and wipe the tip with a piece of tissue without touching the open end of the needle. Quickly inject the sample into the instrument injection port and then withdraw the syringe immediately. Measure the areas of all peaks in the chromatogram. The analyst may vary the temperature, flow rate, column size, and sample size to optimize the analysis.

4.5.2.1.3 Calculations

\[
\%N_2H_4 = \frac{A_{N_2H_4}}{\Sigma A_i} \times 100
\]
\[
\%H_2O = \frac{A_{H_2O}}{\Sigma A_i} \times 100
\]

Where:

- \(A_{N_2H_4}\) = the measured area of the \(N_2H_4\) peak multiplied by its signal attenuation factor
- \(A_{H_2O}\) = the measured area of the \(H_2O\) peak multiplied by its signal attenuation factor
- \(\Sigma A_i\) = the sum of all the measured peak areas in the signal attenuation factors

Assumption: The thermal conductivities of all components in the sample are equal.

4.5.2.1.4 Equipment and reagents. The following equipment and reagents shall apply as test conditions of 4.5.2

(a) Equipment

(1) Gas chromatograph: incorporating a thermal conductivity detector

(2) Recorder: potentiometric strip chart, 0–1 millivolt, 1 second F.S. response, with integrator

(3) Tubing: stainless steel, 1/4 inch O.D. x 1 meter (39.4 inches)

(4) Hypodermic syringe: 10 microliter, fixed needle

(5) Regulator: helium, to fit the cylinder

(b) Reagents

(1) Analrorm B: 90/100 mesh, Analabs, Inc, 9 Hobson Ave, Hamden, CT 06518, or equivalent

(2) Polyethylene glycol 400: Carbowax 400, or equivalent

(3) Methylene chloride: ACS reagent grade

(4) Helium gas: conforming to MIL-P-27407

4.5.3 Particulate. The propellant sample shall be tested for contamination in accordance with ASTM Designation D-2276-65T, Method A, with the following exceptions

4.5.3.1 Mix the sample thoroughly by shaking the sample container. Immediately pour 500ml of the sample into a clean 500-ml graduated cylinder. Use this 500ml of propellant for the particulate analysis.

4.5.3.2 Use a solvent resistant filter disc made from such materials as Millipore LSWP 04700, (Mitek-Teflon), Millipore URWP 04700, (Solvinert), or Gelman VF-6, (Fluoride-Metricel), plain, white, 10-3 microns, 47mm diameter instead of the filter specified in Method 2276-65T.

4.5.3.3 The drying oven temperature shall be 158°F (70°C) instead of the 194°F (90°C) specified in Method 2276-65T.

4.5.3.4 Filtered isopropyl alcohol shall be used for rinsing the sample bottle and filter holder instead of petroleum ether specified in Method 2276-65T.

6. Notes

6.1 Intended use of propellant described in this specification is as a fuel in rocket engines.

6.2 Particulate is the undisolved solid retained on a 10-micron filter membrane.

6.3 Density. For informational purposes, the range of density for material conforming to this Spec is 1.002–1.008g/ml at 25°C (77°F).
Hydrazinobenzene and Derivatives

Hydrazinobenzene or Phenylhydrazine (called Hydrazino-benzol in Ger), C₆H₄HN₂; mw 108.14, N 25.91%; pale-yellow monoclinic crystals or oily liq, mp 19.5°, bp 243.5° (with decomp), flash p 192°F (CC), auto ign temp 345°F, d 1.0978; sol in alc, eth, chl, benz, w and dil acids. Can be prep by reduction of diazotized aniline, followed by reaction with NaOH. Hydrazinobenzene is highly toxic by inhalation, ingestion & skin absorption. It forms a hydrate with 0.5 water. Used in anal chem as reagent for aldehydes & sugars, and in org syntheses. Also as fuel additive and gas absorbent


Hydrazinomononitrobenzene or Mononitrophenylnitrobenzene, O₃N.C₆H₄.NH.N₂; mw 153.14, N 27.44%. Three isomers are known: 2-Nitro-, brick-red ndls (from benz), mp 90° (Ref 1) 3-Nitro-, yel-orn ndls (from alc), mp 93° (Ref 2) 4-Nitro-, ornl-red lfts or ndls (from boiling alc), mp 157° (Ref 3)

Other props & methods of prepn are found in Beil


Hydrazinodinitrobenzene or Dinitrophenylnitrobenzene, (O₂N)₂C₆H₄.NH.N₂; mw 198.14, N 28.28%; OB to CO₂ -88.8%. Two isomers are known: 2,4-Dinitrophenylnitrobenzene, blsh-red crs (from alc), mp 194°, dec at 198°; insol in w; sol in alc & ethyl acetate, sl sol in ether, benz & chl; was prep'd in 1894 by Purgotti and in 1912 by Green & Rowe by heating a w bath 4-Chloro-1,3-dinitrobenzene with freshly prep'd hydrazine hydrate. Some of its salts are expl

Refs 1, 4, S & 6) 2,6-Dinitrophenylnitrobenzene, ornl-red ndls (from alc), mp 145°; insol in w; sol in hot w; sl sol in eth, benz & chl; was prep'd from 2-chloro-1,3-dinitrobenzene & hydrazine hydrate in dil alc


N-Nitroso-N-(2,4-dinitrophenyl)-hydrazine, (O₂N)₂C₆H₃.N(NO)NH₂; mw 227.14, N 30.84%; yel lfts or prisms (from eth), mp 72°, bp-decagrates; was prep'd from 2,4-dinitrophenylhydrazine & NaNO₂ in a soln of HCl, water & alc (Refs 1 & 2)


Hydrazinotrinitrobenzene or 2,4,6-Trinitrophenylnitrobenzene or Picrylhydrazine, (O₂N)₃C₆H₄.NH.N₂; mw 243.14, N 28.81%; red-brn prisms; nearly insol in w, eth, chl, cold alc or benz; sl sol in hot alc; sol in acetic acid; was prep'd from picrylchloride & hydrazine hydrate or acetate in alc. This compd is an expl less powerful & less brisant than PA


Hydrazinebenzene or Phenylhydrazine Reactions.

Several investigators have studied the reaction betwn Phenylhydrazine and various aromatic & aliphatic compds. The following Refs should be of interest

Refs: 1) F. Kehrmann & J. Messinger, Ber 25, 899 (1892) (Studies of reactions betwn Phenylhydrazine & dinitrochlorobenzene) 2) M. Giua, AttiAccadLincei 27 I, 379-82 & 247-52 (1918) & CA 13, 1450, 1586 (1919); GazzChim-Ital 49 II, 146-54 (1919) & CA 14, 1530 (1920) (Reactions betwn Phenylhydrazine and some aromatic nitrocompds contg a labile NO₂ group, such as TNT's, Trinitrobenzoic Acid, Trinitroxylene, Trinitrocreosol & Trinitroanisole were studied. In most cases the formation of nitrohydrazo compds was observed). 3) R.C. Elderfield, "Explosives from Hydroxy and Amino
Compounds", OSRD 158 (Oct 1941), pp 24-25 [a] On condensing Phenylhydrazine with 2,4-dinitrochlorobenzene, instead of the expected dinitrohydrazobenzene, 2-phenyl-5'-nitrobenzo-triazole,
\[ \text{HC} = \text{CH} - \text{C} = \text{N} \rightarrow \text{N.C6H5}, \text{was obtd.} \]
When nitrated, this compd yielded an expl,
\[ \text{HC} = \text{C(NO2)} - \text{C} = \text{N} \rightarrow \text{N.C6H3(NO2)2}, \text{called 2(2",4"-Dinitrophenyl)-1',5'-dinitrobenzo-triazole} \]
[b] Also on condensing Phenylhydrazine with 2,4-dinitrochlorobenzene, followed by refluxing in glacial acetic acid, 2-phenyl(5'-nitrobenzo-triazole)-1-oxide,
\[ \text{HC} = \text{CH} - \text{C} = \text{N} \rightarrow \text{N.C6H5}, \text{was obtd.} \]
When this compd was nitrated, an expl compd 2-(2",4"-Dinitrophenyl)-1',5'-dinitrobenzo-triazole-1-oxide,
\[ \text{HC} = \text{C(NO2)} - \text{C} = \text{N} \rightarrow \text{N.C6H3(NO2)2}, \text{was obtd} \]

**Hydrazino compounds, Analytical Methods for**

Three potentiometric methods of analysis of hydrazine compounds are described:

1) Dissolve sample in H2O, add coned HCl, cool & titrate until equilibrium is established between 700 & 800 mv. This takes place when the IO3 is reduced to ICl, losing 4 meq 2) Dissolve in H2O-H2SO & titrate to the formation of I 3) Dissolve in boiling H2O & keep under nitrogen while cooling. To cold soln add NaOH & titrate to 1° end pt. The 3rd method gives low results but is the only possible method for some hydrazino compounds


**Hydrazinodicarbonic Acid and Derivatives**

*Hydrazinodicarbonic Acid*, HOOC.HN.NH.COOH; mw 88.07, N 31.81%; may be considered as a parent compd of its deriv, although not used to prep it

*Hydrazinodicarbonic Acid Diazide or Hydrazodicarbonazide*, N3.OC.HN.NH.CO.N3; (called Hydrazindicarbonazide in Ger); mw 170.10, N 65.88%; OB to CO2 & N2 - 28.2%; ndls (from eth), mp 150-52° with evolution of gas (when heated slowly); explodes without melting, when heated rapidly. May be prep from hydrazindicarbonazide hydrochloride, H2NNHCONHNHC6H2.HCl, and an aqueous solution of NaNO2. Difficultly sol in cold w & bz; insol in chl & ligroin; sol in acetone, alcohol & hot ethylene bromide; very sol in eth

It is a primary explosive and is slightly more sensitive to impact than Lead and Silver Azides. It would require 0.25g of this explosive to detonate lg of TNT


**Hydrazinoethane.** See Ethyl Hydrazine in Vol 6, p E300-R

**Hydrazinoethane; 2,2-Dinitro.** See 2,2-Dinitro-ethyl Hydrazine in Vol 6, p E300-R

**Hydrazinoxyazobenzene and Derivatives**

*Hydrazinoxyazobenzene*, HOC6H4.N:NC6H4.HN.NH2; mw 228.25, N 24.55%; may be considered as parent compd of its dinitro deriv, although not used to prep it

*Ref: Beil, not found*

4,6-Dinitro-3'-hydrazino-4-hydrayxyzobenzene (called Phenol-(4azo3) - [4,6-dinitrophenylhydrazin] or 4,6-Dinitro-3'-hydrazino-4-oxo-azobenzol in Ger).

HOC6H4.N:NC6H4(NO2)2.NH.NH2; mw 318.25, N 26.41%; red-brown crystals; sl sol in boiling alc; sol in boiling nitro-benzene. Prep by reacting 5'-chloro-2',4'-dinitro-4-hydroxy-azobenzene & hydrazine hydrate in hot alcohol. Decomposes 178-80°

*Refs: 1) Beil 16, (262) 2) W. Borsche, Ber 54, 684 (1921)*
Hydrazinomethane or Methyl Hydrazine, CH₃.HN.NH₂; mw 46.07, N 60.81%; hygr flammable liq, sp gr 0.874 at 25°; fr p <80° (Lange), −52.4° (CondChemDict). It can be prepd by method of Thiele (Refs 1 & 2) consisting of condensing hydrazine with benzaldehyde to give azine, C₆H₅.CH:NH.NH₂, which is treated with dimethyl sulfate followed by water:

\[
\text{C}_6\text{H}_5.\text{CH}:\text{N} .\text{NH}_2 + (\text{CH}_3)_2\text{SO}_4 \rightarrow
\]

\[
\text{C}_6\text{H}_5.\text{CH}:\text{N} .\text{NH}_2 + \text{H}_2\text{O} 
\]

\[
\text{C}_6\text{H}_5.\text{CHO} + \text{H}_2\text{N} .\text{NH} .\text{CH}_3
\]

It is used in missile propels and as a solvent and intermediate (Ref )


Hydrazinophenol and Derivatives

4-Hydrazinophenol, C₆H₄(OH).HN.NH₂; mw 124.14, N 22.57%; crys, mp−unstable. It can be prepd as hydrochloride salt by heating N-[4-hydroxyphenyl]-hydrazine-N'-sulfonic acid K with alc & HCl; forms other salts

Refs: 1) Beil 15, 596 2) Altshul, JPrakt-Chem [2], 57, 202 (1898)

Hydrazinophenol Azide, N₃.C₆H₄(OH).HN.NH₂; not found in Beil

4,6-Dinitro-3-hydrazinophenol or 4,6-Dinitro-3-hydroxyphenylhydrazine, (O₂N)₂C₆H₄(OH).HN.NH₂; mw 214.14, N 26.17%, OB −74.6%; fine egg-yellow flocks, resembling yellow HgO when dry; mp 197° (decomp); sol in hot alc (with separation of red-brown spherical granules); sol in warm aq HCl; decomposed by dil NaOH with vigorous foaming. Prepd by heating 2,5-Dinitro-4-chlorophenol in alcohol with hydrazine hydrate & then acidifying with dil nitric acid

Refs: 1) Beil 15, [274] 2) W. Borsche, Ber 54B, 676 (1921) & CA 15, 2844 (1921)

Trinitrohydrazinophenol, C₆H₅N₅O₇, not found in Beil

Hydrazinopyridine and Derivatives

Hydrazinopyridine, H₂N.NH.C₅H₄N, mw 109.13, N 38.51%. Exists as 2-, 3- & 4-hydrazone isomers (Refs 1 & 2). May be considered as parent compd of nitrated derivs, although they were not prepd from it

Refs: 1) Beil 22, (688) & [487] 2) Beil 22, [488]

Hydrazinopyridine Azide, C₆H₅N₆O₇, not found in Beil

Nitrohydrazinopyridine, C₅H₄N₂O₄; mw 154.13, N 36.35%. Two isomers are known:

\[
\text{O}_2\text{N} \quad \text{N} \quad \text{NH} .\text{NH}_2;
\]

3-Nitro-4-hydrazinopyridine

\[
\text{N} \quad \text{NO}_2
\]

red ndls (from alc), mp 204−06° (dec); diff sol in common solv; sol in alkalis; can be prepd by treating 6-chloro-3-nitropyridine with hydrazine hydrate or alcoholic hydrazine soln under cooling (Ref 1); from 5-nitro-2-pyridine, solfonic acid & the appropriate amine (Ref 3), or by boiling 2-metroxy-5-nitropyridine with hydrazine (Ref 4)

\[
\text{N} \quad \text{NH}_2
\]

Dinitro, C₅H₅N₂O₄, and Trinitrohydrazinopyridine, C₅H₄N₆O₆, were not found in Beil

5-Hydrazino-tetrazole (Tetrazyl-5-hydrazine)

\[
\text{H}_2\text{N.NH} \quad \text{−N−N} \quad \text{or} \quad \text{H}_2\text{N.N}=\text{C} .\text{NH} .\text{N}
\]

\[
\text{N} \quad \text{N} \quad \text{NH} \quad \text{−N} ;
\]
mw 100.09, N 83.97%; crystals (from w), mp 199° (dec), bp—decomp explosively on heating above mp; sl sol in cold w; sol in alkalies & ammonia; insol in alc, eth & benz. It was first prepd by Thiele et al (Refs 1 & 2) by reduction of tetrazolediazonium chloride with stannous chloride in the presence of HCl. Other methods of prepn are given in Refs 1, 2 & 5


6) No more recent refs found in CA

Hydrazinotoluene and Derivatives

Hydrazinotoluene or Tolyldrazine,
H₃C.C₆H₄.HN.NH₂; mw 122.17, N 22.93%. There are three isomers, ortho-, meta-, & para-tolyldrazines

Refs: 1) Beil 15, 496, (147) & [222] (ortho)
2) Beil 15, 506, (152) & [229] (meta)
3) Beil 15, 510, (153) & [233] (para)

Hydrazinotoluene Azide, C₇H₉N₃S, not found in Beil

Mononitrohydrazinotoluene, C₇H₈N₃O₂; mw 167.17, N 25.14%. Two isomers are found in Beil:
5-Nitro-2-hydrazinotoluene, it golden-yel or orn-yel ndls with a violet tinge (from alc), mp 179—80°; diff sol in hot alc & xylene; was prepd by heating the K salt of N-[4-Nitro-2-methyl-phenyl]-hydrazine-N,N'-disulfonic acid with concd HCl (Refs 1 & 3)
3-Nitro-4-hydrazinotoluene, dk-red ndls (from eth), mp 110—11°; sl sol in alc; sol in eth, chl & benz; sl sol in petr eth; was prepd by diazotizing 2-nitro-4-methylaniline with Zn chloride & HCl. Its Hydrochloride salt, C₇H₈N₃O₂+HCl, orn-red ndls or plates (from w) dec at 190—91° (Refs 2 & 4)
Ref: 1) Beil 15, 505  2) Beil 15, 530

Dinitrohydrazinotoluene, C₇H₇N₄O₄; mw 212.17, N 26.41%. Five isomers are described in Beil:
3,5-Dinitro-2-hydrazinotoluene, orn ndls (from alc), mp 169° (dec); v sl sol in alc; was prepd from 2,3,5-Trinitrotoluene & hydrazine hydrate in alc (Refs 1 & 7)
2,4-Dinitro-3-hydrazinotoluene, red prisms (from alc or benz), mp 170°; sol in warm 10% HCl; was prepd by warming 2,3,4-Trinitrotoluene with hydrazine hydrate in dil alc and cooling the soln (Refs 2 & 7)
2,6-Dinitro-3-hydrazinotoluene, solid, mp—not reported; was prepd by warming 3-chloro-2,6-dinitrotoluene with hydrazine hydrate in dil alc on a water bath (Refs 3 & 7)
4,6-Dinitro-3-hydrazinotoluene, yel-red crysts (from alc), mp 194—95° (dec); sol in alc, benz & chl; was prepd by treating 2,4,5-Trinitrotoluene with hydrazine hydrate in alc (Refs 4 & 6)
3,5-Dinitro-4-hydrazinotoluene, copper-colored tablets (from alc), mp 139°; was prepd by heating an alcoholic soln of 3,4,5-Trinitrotoluene with alc hydrazine hydrate soln (Refs 5 & 7)

Other props are given in Refs

5) Beil 15, [244]  6) M. Giua, Gazz 49 II, 171 (1919); 53, 849 (1923)
7) O.L. Brady & J.H. Bowman, JChemSoc 179, 896, 899 & 901 (1921)

2,4,6-Trinitro-3-hydrazinotoluene,

mw 257.17, N 27.24%; golden-yel pltls (from alc), mp 176° (dec); sol in alc, benz, chl & acet; sol in alkalies giving a red-brn soln; sl sol in petr eth; was prepd by treating 2,4,6-Trinitro-3-methoxytoluene with hydrazine hydrate in alc (Refs 1 & 2)

Ref: 1) Beil 15, (153)  2) M. Giua, Gazz 49 II, 174 (1919)

3-Hydrazine-1,2,4-triazole,

HN—NH—C=N.NH₂ or HN—N=C—NH.NH₂
Hydrazobenzene and Derivatives. See N,N'-Di-phenylhydrazine and Derivatives in Vol 5 of Encycl, pp D1462-R to D1463-L and the following derivs not described:

2,4,2'-Trinitrohydrazobenzene,
\((O_2N)_2C_6H_3NNH.C_6H_4(NO_2)\); mw 319.23, N 21.94%; pale yellow needles (from HAc), mp 220°, puffs off on strong heating; sol in acetone, chll, ligroin & Et acetate. Prepd from 2-nitrophenylhydrazine & 4-chloro-1,3-dinitro-benzene

Refs: 1) Beil 15, 490 2) A. Werner & ? Stiasny, Ber 32, 3281 (1899)

2,4,4'-Trinitrohydrazobenzene,
\((O_2N)_2C_6H_3NNH.C_6H_4(NO_2)\); no props given. was prepd by heating 3-nitrophenylbenzene (2 mols) with 4-chloro-1,3-dinitro-benzene at 100–105°

Refs: 1) Beil 15, 490 2) A. Werner & ? Stiasny, Ber 32, 3280 (1899)

2,4,6-Trinitrohydrazobenzene (called N-Phenyl-N'[2,4,6-trinitrophenyl]-hydrazin in Ger.),
\(C_6H_3NNH.C_6H_4(NO_2)_3\); mw 319.2, N 21.9%; dark red prisms (from HAc or acetone); mp 181°; decomp on rapid heating & puffs off in a flame; sl sol in alc or chll; somewhat sol in acetone. Prepd from phenylhydrazine & picrylchloride (Ref 1 & 3) or trinitroanisole & phenylhydrazine in abs EtOH (Ref 2). It may be considered to be a mild explosive


4) No further refs were found in CA thru 1971

Hydrazobisformamidin e or Hydrazodicarbonamide. See Biguanidine in Vol 2, p B115-L

5,5'-Hydrazo-bis-1H-Tetrazole. See Bis (5-Tetrazolo)hydrazine in Vol 2 of Encycl, p B157-R and the following addnl refs

Refs: 1) F.R. Benson, ChemRevs 41, 8 (1947) 2) O.E. Sheffield, "Long Range Basic Research Leading to the Development of Ideal Propellants. Compounds of High Nitrogen Content in Propellant Powders," PATR 1694 (May 1948) 3) W.S. McEwan & M.W. Rigg, JACS 73, 4725 (1951) & CA 46, 4350 (1951) (give \(\Delta H^\circ\))

Hydrazodicarbonamide. See Hydrazinodicarbonic Acid Dizide

Hydrazoic Acid. See under Azides in Vol 1, p A537-42

Hydrazotates. See Azides

5,5'-Hydrazotetrazole. Compd investigated for use as a primary explosive. Impact sensitivity was found unsatisfactory. Some pertinent properties: Impact sensitivity 32 cms, Vac Stab, 1.7 ml of gas/48 hrs/100°C, Hot Bar Ign temp 239°C
Ref: Francis Taylor, NAVORD Rep 2800, Primary Explosives Research, Naval Ord Lab, White Oaks, Md (1953)

NOTE: See also 5,5'-Hydrazo-bis-tetrazole in Vol 2 of Encycl, p B157-R

Hydrazulmin. \(C_6H_4N_4\), mw 162.16, N 51.83%, soot-black shiny leaflets. Prepd from cyanogen & dry ammonia. Decomposes on heating to give paracyanogen; decoump in water
Ref: Beil 2, 553

Hydrides. Both ionic & covalent compounds of H with another element usually metallic. These will generally be described under the "other" element eg Al-hydride in Vol 1, p A145 & B-hydrides in Vol 2, pp B253-56. However certain
props common to the hydrides follow below:

Toxicity: Variable. The hydrides of phosphorus, arsenic, sulfur, selenium, tellurium and boron which are highly toxic, produce local irritation and destroy red blood cells. They are particularly dangerous because of their volatility and ease of entry into the body. The hydrides of the alkali metals, alkaline earths, aluminum, zirconium and titanium react with moisture to evolve hydrogen and leave behind the hydroxide of the metallic element. This hydroxide is usually caustic. See also sodium hydroxide.

Hydrides, metallic, primary type:

This group includes the hydrides of calcium, lithium, magnesium, potassium, sodium and strontium. In the presence of moisture they are readily converted to hydroxides which are highly irritating to the skin by caustic and thermal action. Similar effects can occur on contact with eyes and respiratory mucous membranes.

Fire Hazard: The volatile hydrides are flammable, some spontaneously so in air. All hydrides react violently on contact with powerful oxidizing agents. When heated or on contact with moisture or acids an exothermic reaction evolving hydrogen occurs. Often enough heat is evolved to cause ignition. Hydrides require special handling instructions which should be obtained from the manufacturers.

Explosion Hazard: The volatile hydrides (such as hydrides of boron, arsenic, phosphorus, selenium, tellurium) form explosive mixtures with air. The nonvolatile hydrides (such as sodium, lithium, calcium) readily liberate hydrogen when heated or on contact with moisture or acids. Furthermore, hydrides form dust clouds which can explode due to contact with flames, sparks, heat or oxidizers.

Disaster Hazard: Highly dangerous; when heated, they can ignite at once or liberate hydrogen; they react with moisture or acids to evolve heat and hydrogen; on contact with powerful oxidizers violent reactions can occur. (Ref 2). Metal hydrides (Al, Li, LiAlH₄ etc) in Me ether solns should not be evaporated because of the danger of explosions (Ref 1). Similarly the use of LiAl-hydride is not recommended for drying Me ethers (Ref 2).


Hydridin is the reaction product of a polyhydric alcohol and an inorganic salt, e.g., chlorhydrin is formed from HOCH₂CH(OH)CH₂OH + HCl = HOCH₂CH(OH)CH₂Cl + H₂O. Ref: Hacke's (1944), p 418-L

Hydridinwestalite. A German explosive containing DNT. Ref: Colver (1918), p 167

Hydrogel. Brit Expls used for Pulsed Infusion Shot firings: NG+NGc 37.9-40.9, NC 1.0-3.0, woodmeal 0.6-1.6, AN 19.0-21.0, NaCl 26.1-28.1, Barytes 8.7-10.7, chalk 0.1-0.5, diammonium phosphate 0.1-0.8, Acid Magenta 0.001-0.005, AS No 2 (optional) 0.001-0.01%, d 1.7; vel of deton 5000 m/sec; temp of expln—not given. To initiate hydrogel chges under high hydrostatic pressure, it is necessary to use a specially designed Cu tube & submarine electric detonators. Ref: McAdam & Westwater "Mining Explosives," Oliver & Boyd, Edinburgh (1958), p 109

NOTE: The compn of AS No 2 is not reported.

Hydrocarbon. C₂₀H₄₆; mw 264.39, crystals, mp 38°, bp 330°. Was prep'd by Puranen & Ehrnrooth from cymene by treating it at low temperature with a solution of nitrosylsulfuric acid in concd H₂SO₄, who proposed it as a starting material for preparation of various explosives. Ref: N. Puranen & E. Ehrnrooth, GerP 611461 (1935) & CA 29, 4020 (1935)

Hydrocarbons. Compounds consisting of carbon and hydrogen. The number of such compounds is immense and they may be classified into: "aliphatic" (acyclic) and "aromatic" (cyclic) compounds. In the former class, the principal carbon atoms are arranged in chains, while in the latter class they are arranged in one or several rings. Hydrocarbons may also be divided into "saturated"—in which all four valences of C are satisfied, and "unsaturated"—in which there are one or more double or triple bonds between carbon atoms.

Many hydrocarbons furnish explosive nitrates or nitrocompounds on nitrination. (See under individual compounds)
The literature on hydrocarbons is overwhelmingly voluminous. Only three standard references are listed below:


Hydrocarbons As Rocket Fuels or Explosives. Liquid methane & other hydrocarbons are considered as propellant fuels (Ref 1). Nitrogen tetroxide and hydrocarbons (kerosine) are claimed as a blasting explosive (Ref 2)


Hydrocellulose. C_{12}H_{22}O_{11}, mw 342.30. The term has been employed since about 1880 to designate a cellulose which has been weakened or tendered as the result of treatment with acids. It retains water strongly. Hydrocellulose is not a single substance but a mixture of hydrolyzed products less complex than cellulose itself and more or less related to it and to glucose (Reps 4 & 6)

Hydrocellulose differs from hydrated-cellulose in its properties (Ref 3). According to Stettbacher (Ref 5) hydrocellulose lies between the hydrated cellulose and oxyz cellulose. Its use as a flash reducer in a propellant has been claimed by C.R. Franklin in USP 1564549 (1925) & CA 20, 505 (1926). Acetol to CISO Rept 31-68 (1945), pp 6-7, hydrocellulose was used during WWII by Germans in some rocket propellants, presumably to improve their burning characteristics. For instance the so-called Ammon-pulver contained 5% hydrocellulose and the EP (Einheitspulver) contained about 3%. Hydrocellulose was also used in some rocket propellants to increase the rate of burning at low temperature

Note: Their Einheitspulver Standard (or Unit) Propellant was “G” Pulver (qv), which containd 3% hydrocellulose & 1.5% K-nitrate. Its props are given in PATR 2510 (1958), p Ger 190-R

Hydrocellulose can be nitrated, forming an explosive (Reps 2 & 5). Hydrocellulose is claimed to be C_{24}H_{40}O_{20} mw 648.56 by Tavernier (Ref 7), who gives (Q_f)_v = 1400 cal/g & (Q_f)_p = 1428 cal/g

Hydrocellulose Nitrate or Nitrohydrocellulose. A substance resembling NC in its properties (but slightly more sensitive to shock), was prep by Cross and Bevan and Beadle by nitration of 1 part of dry hydrocellulose by means of 3 parts of nitric acid and 9 parts of sulfuric acid (Reps 1, p 25, & 2, p 556); products corresponding to hexa- and heptanitrohydrocelluloses are formed. Its use in explosives was recommended by Luck, Durnford and Ungania (Ref 2, p. 557). Trauhl (Ref 2, p 199) proposed using detonators charged with a gel prep by impregnating 1 part of Nitrohydrocellulose with 1 to 3 parts of NG. Such detonators could be used underwater


Hydrochloric Acid or Muriatic Acid. See Hydrogen Chloride, Anhydrous and Hydrochloric Acid

Hydrocinnamic Acid and Derivatives

Hydrocinnamic Acid or 3-Phenylpropionic Acid, C_6H_5CH_2CH_2COOH; mw 150.17, crystals, mp 46°; sl sol in w; sol in alc or eth. Prep by the reduction of cinnamic acid with Na-amalgam

α-Azidohydrocinnamic Acid or α-Azido-β-phenylpropionic Acid (called α-Amido-Hydroximtsäure in Ger), C_6H_5CH_2CH(N_3)COOH; mw 191.19, N 21.98% plates, mp 24°-7°. Its Silver salt, AgC_6H_5N_3O_2, cryst (from w), explodes mildly on heating; the Ammonium salt, white cryst, burns quietly on contact with a flame

Reps: 1) Bell B, (205-6) 2) A. Darapsky & H. Berger, JPraktChem 96, 320-1 (1917); JCS 114 I, 507-8 (1918) & CA 13, 1304-5 (1919) 3) No further refs found in CA thru 1971
Hydrocinnamic Acid Azide (called Hydrozimiture-Azid in Ger). C₉H₈CH₂CH₂CON₂, mw 175.19, N 23.99%, powdery crystals (from eth); insol in w, sol in alc or eth. Puffs off on heating. Prepd by reacting hydrocinnamic acid hydrazide in HCl with Na-nitrite
Refs: 1) Beil 9, 513 2) No further refs found in CA thru 1971

Hydrocyanic Acid or Hydrogen Cyanide (Formonitrile or Prussic Acid). HCN, mw 27.03, N 51.82%, OB to CO₂ -14.9%; highly poisonous, colorless liquid or gas; d 0.697 at 18°, fr p -13.3°, bp 25.6°, fl p 0°F, auto ignition temp 100°F. Very sol in w, alc & eth. Exists in two isomers, H:C:N and H:N:C. This fact was first stated by Enklaar and then confirmed by Wöhler and Roth, who also examined the explosive properties of HCN (Ref 3). May be prepd by treating a cyanide, such as calcium cyanide, with dilute sulfuric acid; also catalytically from hydrocarbons and ammonia (Ref 6). HCN is present in the products of detonation of various explosives. The dissociation energy for H-CN is 129 ± 3 kcal/mole (Ref 9). It is an explosive which requires a stronger initiation than that supplied by a No 8 cap for its detonation. For instance, 3g of HCN, cooled to 0°, may be detonated with a cap containing 0.4g of Lead Azide and 1.25g of PETN. If the detonation takes place in a lead block 10 x 10 cm, the block is generally split in two. The explosion of HCN is favored by its preliminary polymerization by alkalies (Ref 3). It may even polymerize explosively (Ref 12). Presence of NaOH, but not water, and increasing temp were found to favor violent HCN polymerization. The heat of polym is 377 cal/g (Ref 8). In the presence of NO & argon, HCN explodes when heated in a reflected shock wave to give CN, NCN, NH & OH radicals during the pre-explosion induction period. The reaction NO+HCN= CN+HNO is claimed to be rate-controlling (Ref 13).

The combustion of HCN has been studied extensively. Ignition limits in air are ca 42.5 & 8 vol%. In the presence of CO₂ these limits coalesce for the mixture CO₂ 36 vol% & HCN 12 vol% (Ref 6). Burning velocity, flame temp & thickness of reaction zone for HCN/O₂/N₂ mixtures have been measured. The overall activation energy for HCN/O₂ flames is claimed to be 42-44 kcal/mole (Ref 11). HCN has also been considered as a rocket fuel. Its combustion enthalpy and specific impulse are high & even its toxicity is relatively low when compared with diborane & hydrazine (Ref 10)

Some of its salts are also explosive, for example, Hg(CN)₂ and Hg(CN)₂·HgO. Its methyl salt is explosive when in the CH₃NC form, while the CH₃CN form is not an explosive

HCN is used in the manufacture of polymers & fumigants (Ref 14)

NOTE: Accdg to Blatt, OSRD 2014 (1944), HCN is an expl of low sensitivity to initiation because it cannot be detonated by No 8 cap. In order to detn its power by Trauzl Test, a 3g sample in lead block 10x10cm, cooled to 0° was detonated with 0.4g LA+1.25g PETN initiator. It gave an expansion of 95–100cc and this usually split the block


Hydrodynamics. Flow systems are governed by the laws of hydrodynamics. How these laws relate to detonation was discussed in Vol 4, pp D602-22. Recent books and reviews on this subject are listed below:

3) S. Chandrasekhar, “Hydrodynamic & Hydro- 
magnetic Stability”, Clarendon Press, London 
(1961)
4) G.H.A. Cole, “Fluid Dynamics”, John Wiley, 
NY (1962)
5) V.G. Levich, “Physicochemical Hydrodyna-
metrics”, Prentice-Hall, Englewood Cliffs, NJ 
(1962)
6) C. Truesdell, “Fluid Dynamics I”, Vol VIII 
of Encyc of Physics, ed S. Fluegge, B.H. 
Blackwell, Oxford (1963)
Macmillan, NY (1964)
8) M. Weintraub “Chemical Engineering Aspects 
of Fluid Dynamics.” Review with 252 references. 
IndEngChem 56 (4), 43 (1964) & CA 60, 15453 
(1964)
9) B. Alder et al, Eds, “Fundamental Methods in 
Hydrodynamics” (Methods in Computational 
Physics, Vol 3), AcademicPress, NY (1964)
10) J. Happel & H. Brenner, “Low Reynolds 
Number Hydrodynamics”, Prentice-Hall, Engle-
wood Cliffs, NJ (1965)
11) R.J. Seeger & G. Temple, Eds, “Research 
Frontiers in Fluid Dynamics”, Interscience, NY 
(1965)
12) M.A.T. Cocquerel “Unit Operations, Heat 
Transfer, Fluid Flow.” Review with 73 refs. 
ReplProgrApplChem 50, 117 (1965) & CA 66, 
39243 (1967)
Theory and Plane Shock Waves in Metals. I. 
Theory.” Title of the paper in 4thONRSympDeton 
(1965), p 289 (Abstract) and of PATR 3464 (1966)
14) L. G. Lototskynsk “Mechanics of Liquids & 
15) Ya. B. Zel'dovich & Yu. P. Raizer “Physics 
of Shock Waves & High-Temperature Hydro-
dynamic Phenomena” 2nd Ed Moscow, Nauka (1966)
16) S. I. Pai et al, Eds, “Dynamics of Fluids & 
17) R.H.F. Pao, “Fluid Dynamics”, Merrill, 
Columbus (1967)
18) R. A. Strehlow “Detonation & the Hydro-
dynamics of Reactive Shock Waves.” Review of 
recent work. ACS, DivFuelChemPreprints, 11, 1 
(1967) & CA 70, 98391 (1969)
19) W. N. Gill et al “Fluid Dynamics.” Review 
with 1065 refs. IndEngChem 59, 69 (1967) & 
CA 68 51292 (1968)
20) W. N. Gill et al “Fluid Dynamics.” Critical 
review of the 1967 fluid dynamics literature. 
IndEngChem 61, 41 (1969) & CA 70, 69515 
(1969)
21) W. N. Gil et al “Fluid Dynamics.” Review of 
literature for 1968 with 791 refs. IndEngChem 
22) W. N. Gill et al “Fluid Dynamics.” Review 
74, 55600 (1971)

Hydrodynamic Detonation Velocity. Same as 
Ideal Detonation Velocity, briefly described in 
Vol 4 of Encyc, p D630-R

Hydrodynamic Theories of Detonation. See 
Vol 4 of Encyc, p D610-L

Hydrofluoric Acid or Fluothydric Acid. See 
Hydrogen Fluoride, Anhydrous and Hydro-
fluoric Acid in this Vol

Hydrofluosilic Acid. See Fluorosilic Acid 
in Vol 6 of Encyc, p F141-R

Hydrogen, H, at wt 1.008. The lightest element; 
usually exists as a molecule (H₂) of mw 2.016 
in the form of colorless, highly flammable gas; 
sp gr of gas 0.06948 (Air 1.0); and of liq 
0.0709 at −252.7⁰; fr p −259.18⁰; bp −252.77⁰; 
autoignition temp 1085⁰F; low expn limit 4.1% 
by vol, upper expn limit 74.2%; very sl sol in 
w, alc or eth. It can be prepd by the follow-
ing methods: a) By the action of steam on 
natural gas at high temperatures, and subsequent 
purification b) By treatment of water gas 
with steam and absorption of the carbon dioxide 
c) Dissociation of ammonia d) Thermal de-
composition of hydrocarbons e) Catalytic re-
forming of petroleum f) By reaction of iron 
and steam g) Catalytic reaction of methanol 
& steam and h) By electrolysis of water 
(Ref 2, p 587-R)

Accdg to Ref 2, p 588-L, the molecular hy-
drogen exists in two varieties: ortho and para, 
named accdg to their nuclear spin types. Ortho 
molecules have a parallel spin, para an anti-
parallel spin. By cooling to liquid air temp and using 
a catalyst, the normal equilibrium of three 
ortho to one para molecules is displaced and 
para-H₂ may be isolated. Para type of H₂ is 
preferred for liquid fuels
Liquid hydrogen is used in rocket fuels, while hydrogen gas can be used for the following purposes: production of synthetic ammonia and methanol; hydrogenation of organic materials such as oils, phenol or naphthalene; reducing agent for organic synthesis; reduction of metallic ores; reducing atmospheres to prevent oxidation; as oxyhydrogen flame for high temps; for atomic hydrogen welding; making hydrochloric and hydrobromic acids; for filling small balloons, etc. It was previously used in Europe for inflation of passenger-carrying balloons and for observation and barrage balloons (during WWII). Its use in dirigibles was discontinued after disastrous fire of Ger dirigible, "Hindenburg". Hydrogen has not been used for inflation of US dirigibles or balloons because the non-combustible, lighter than air helium is readily available.

**Note:** The explosion and combustion properties of hydrogen mixtures are described as a separate item.

**Refs:**
1) Gmelins Handb (1936), not found
3) Kirk & Other Vol 11 (1966), pp 338-79
   by R.M. Reed 4) Sax (1968), p 823

**Hydrogen Bomb.** See under **Atomic Bomb** in Vol 1, A499-L.

**Hydrogen bond.** An attractive force, or bridge occurring in polar compounds such as water, in which a hydrogen atom of one molecule is attracted to two unshared electrons of another. The hydrogen atom is the positive end of one polar molecule and forms a linkage with the electronegative end of another such molecule. In the formula below, the hydrogen atom in the center is the “bridge”:

\[
\begin{align*}
\text{H-O:O}&:\text{H-O:} \\
\text{H} & \quad \text{H}
\end{align*}
\]

Hydrogen bonds are only one-tenth to one-thirtieth as strong as covalent bonds, but they have pronounced effects on the properties of substances in which they occur, especially as regards melting and boiling points and crystal structure. They are found in compounds containing such strongly electronegative atoms as nitrogen, oxygen and fluorine. They play an important part in the bonding of cellulosic compounds, eg, in the paper industry.

**Refs:**
2) G. C. Pimentel & A. L. McClellan "The Hydrogen Bond" Reinhold (1960)
3) CondChemDict (8th Ed 1971), p 455

**Hydrogen Bonding Concept in Gelatinization of NC.** Tests of a number of hydrogen bonding agents dissolved in ether, alcohol or benzene showed that the concept of hydrogen bonding was certainly involved in the mechanism of gelatinization, but the results were modified by the character of the volatile solvent used.

Tests of the best hydrogen bonding agents in ether-alcohol showed the following compounds: trimethyl-, trimethylammonium, tributyl phosphates, isophorone, dimethyl acetamide, and dibutyl tartrate were superior to dibutyl phthalate or triacetin in the amounts required for gelatinization and in the viscosities of the resulting solutions.

When a major portion of Guncotton was dispersed with acetone-alcohol and the residual portion with one of the above compounds, equal weights of the compds were superior to dibutyl phthalate and to triacetin first, with respect to time required for complete dispersion and secondly, in producing solns of low viscosity.

**Ref:** A.J. Philips, "The Relation of the Hydrogen Bonding Concept to the Gelatinization of Nitrocellulose," PATR 1249 (March 1943)

**Hydrogen Chloride, Anhydrous and Hydrochloric Acid.**

**Anhydrous Hydrogen Chloride** is a colorless gas, which on liquefaction gives colorless (or sl yellowish) fuming, strongly corrosive liquid; mw 36.47, sp gr of gas at 0° 1.268 (Air); fr p -114.3°; bp -111° (Lange); bp -84.8°; sol in w, alc, eth & benz. It can be prepd by furnace combustn of hydrogen, methane or water-gas, in chlorine or by solvent extraction from hydrochloric acid (aqua HCl).

It is used in reactions where aqu HCl is not suitable, such as production of vinyl chloride from acetylene or of alkyl chlorides from olefins (Ref 2, p 256 & Ref 3, p 588-R)

**Hydrochloric Acid** (known in commerce as *Muriatic Acid*), is a conc, aqu soln of anhydrous HCl. It is colorless or sl yel, fuming, pungent, liquid, poisonous unless it is very diluted. The commercial product contains 38% HCl and has
sp gr of 1.19. A more concd product of 45.2% HCl listed in Lange (Ref 2), has sp gr 1.48 and freezes at \(-15.37^\circ\). The acid is sol in w & alc. It can be prepd by several methods such as:

a) By-product of chlorination of benz or other hydrocarbons  
b) By the action of sulfuric acid on common salt; or  
c) By Hargreaves process which is described in Ref 3, p 585-R.

The acid has many uses in chem & metallurgical industries, as indicated on p 586-R of Ref 3.

**Note:** Salts of HCl are called chlorides, many of which are described in Vol 3 of Encycl, pp C236ff.

**Refs:**  
1) GmelinsHdb, Syst No 5 (1927), p 86 & Teil B (1968), p 2  
2) Lange (1961), p 256  

#### Hydrogen Fluoride, Anhydrous and Hydrofluoric Acid.

**Anhydrous Hydrogen Fluoride**, HF, mw 20.01; colorless, corrosive gas which forms at temp below 19.4°, colorless, fuming, corrosive, poisonous liquid, producing painful burns on the skin. It freezes at \(-83^\circ\) and its sp gr is 0.988 at 3.6° (Lange). Both gas and liquid consist of associated molecules; the vapor density corresponds to HF only at high temps. Both gas and liquid are very sol in water. Can be prepd by distillation from the product of reaction of Ca fluoride (fluorspar) and sulfuric acid. Used as fluorinating agent in organic and inorganic reactions; production of fluorine and Al fluoride; prepn of hydrofluoric acid; production of fluoroborates, fluorosilicates, etc; as an additive in liquid rocket propellants; and for refining of uranium (Ref 3, p 588-R).

**Hydrofluoric Acid** is an aqueous soln of hydrogen fluoride of strength up to 70%. These solns are colorless, fuming, corrosive, mobile liquids, producing terrible sores when allowed to touch the skin. Only a moderately strong acid, but unlike other acids will attack glass and any silica-contg substance. Prep by absorbing in w, HF gas, which is distilled from a mixture of fluorspar (CaF₂) and sulfuric acid; also by dis-

solving HF liquid in w.

Lange (Ref 2, p 256) gives aor 35.35% acid, sp gr 1.15, fr p -35° & bp 120°.

Hydrofluoric acid is used for polishing, etching & frosting of glass; pickling Cu & other metals; electropolishing of metals; purification of filter paper and cleaning castings (Ref 3, p 587-L).

**Refs:**  
1) GmelinsHdb, Syst No 5 (1926), p 31 & 5 (1959), p 142  
2) Lange (1961), p 256  
5) Sax (1968), 823  
6) CondChemDict (1971), pp 454 & 455  
7) US Spec O-H-795(2) (July 1957) (Hydrofluoric Acid, Technical Grade)  

**Hydrogen Gun.** A gun that shoots bullets at a speed of 4 miles per sec (or even faster) was constructed by E. B. Mayfield of the Naval Ordnance Test Station, Inyokern, California. The propellant is liquid hydrogen touched off by an injection of oxygen and the projectiles are smaller than rifle bullets. The prime purpose of this gun is the study of what happens when a projectile moves through air at very high speeds.  

**Ref:** Anon, Ordn 36, (No 191), 780 (1952)

**Hydrogen-ion Concentration (pH) or Potential of Hydrogen.** The concentration of H⁺ per liter \([H^+]\) of an aqueous solution. It denotes the true acidity or alkalinity of such solutions, and is expressed by the pH value (potential of hydrogen). This value is the concentration of H⁺ in terms of the reciprocal logarithm of the number of gram ions of hydrogen per liter:

\[ pH = \log_{10} \frac{1}{C_H} = -\log_{10} C_H \]

where \(C_H\) is effective concentration of hydrogen (molar or normal). Pure water has pH value = 7.0, because one liter dissociates to contain only \(10^{-7}\) g of H⁺. As there is also an equal number of \([OH^-]\) present, its pOH is also 7, which makes a total \(pH + pOH = 14\). Accordingly pH values from 0 to 7 indicate an acid solution, \(pH = 7\) indicates neutrality and \(pH\) 7 to 14 an alkaline solution.

Hack's Dictionary, p 422 (Ref 13) gives a table of relation between pH, pOH, Hydrogen-ion concentration (normality) and hydroxyl ion concentration (normality).
pH values may easily be calculated if the value $C_H$ (also denoted as $A_{H^+}$) is known. For instance, pH of a solution which is 0.0002 molal in hydrogen ion ($C_H = 2 \times 10^{-4}$).

$$\text{pH} = \log_{10} \left( \frac{1}{C_H} \right) = \log_{10} \left( \frac{1}{2 \times 10^{-4}} \right) = \log_{10} 5 \times 10^3 = 3.699$$

Determination of pH values may be done either by electromotive force measurements, by colorimetric methods (indicator papers, or indicator solutions), or by the glass electrode method.

Since the pH value is of great importance in many manufacturing processes (including explosives), pH meters are installed to exercise control over the pH (see Refs 5 & 6).

There are also other methods for measuring hydrogen ion concentration, for instance, Sörensen's (pH) unit or the activity coefficient unit (pH), based on emf measurements.


**Hydrogen-ion Concentration Stability Test.** See Hansen's Stability Test in this Vol.

**Hydrogen Peroxide, Hydrogen Dioxide or T-stuff.** HOOH; mw 34.02; colorless or faint blue syrupy liquid, mp $-2^5$, bp 158°C (Ref 24), vap press 1mm at 15°C & 47mm at 80°C, d 1.46; sol in w, alc & eth; insol in pet eth. Thermody data: $\Delta H_f^{298}$ -44.84 & $\Delta F_f^{298}$ -28.2 kcal/mole for liq; $\Delta H_f^{298}$ -45.65 kcal/mole for aq H$_2$O$_2$ -31.83 kcal/mole for gaseous H$_2$O$_2$ (Ref 6). Bond strengths: H-OH 89.6 kcal/mole (Ref 18); HO-OH 47.1 kcal/mole (Ref 22)

**Prepn:** a) BaO$_2$ + H$_2$SO$_4$ (aq) = BaSO$_4$ + H$_2$O$_2$, the aq H$_2$O$_2$ is then concentrated by evaporating w under reduced press (Ref 2)

b) Electrolytic oxidation of concd H$_2$SO$_4$ with subsequent hydrolysis of the peroxydisulfuric acid (Ref 2)

c) Hydrogen reduction of 2-ethylanthraquinone followed by air oxidation to regenerate the quinone (Ref 25)

Pure H$_2$O$_2$ is a very unstable material which decomposes violently above 80°C. The decomp H$_2$O$_2$ = H$_2$O + 1/2O$_2$ is catalyzed by many subts such as Ag, MnO$_2$, HBr & salvia. Because of this instability, commercial H$_2$O$_2$ is usually a water soln—commonly 27.5, 35, 50 & 70%, although solns ranging from 3 to 90% are also available. H$_2$O$_2$ solns are stabilized with small amts of acetopheneditin inhibitor (Ref 25)

**Toxicology:** Pure H$_2$O$_2$, its solutions, vapors and mists are irritating to body tissue. This irritation can vary from mild to severe depending upon the concn of H$_2$O$_2$. For instance solutions of H$_2$O$_2$ of 35 wt% and over can easily cause blistering of the skin. Irritation caused by H$_2$O$_2$ which does
not subside upon flushing of the affected part
with water should be treated by a physician.

The eyes are particularly sensitive to irritation
by this material (Ref 24)  

Uses: Bleaching and deodorizing of textiles,
wood pulp, hair, fur, etc; source of organic and
inorganic peroxides: pulp and paper industry;
plasticizers; rocket fuel; foam rubber; manufacture
of glycerol; antichlor; dyeing; electroplating;
antiseptic; laboratory reagent; epoxidation;
hydroxylation; oxidation and reduction; viscosity
control for starch and cellulose derivatives;
refining and cleaning metals; bleaching and
oxidizing agent in foods; neutralizing agent in
wine distillation; seed disinfectant; cosmetics
(Ref 25)  

Chemistry: In some respects chemistry of \( \text{H}_2\text{O}_2 \)
is quite similar to that of \( \text{H}_2\text{NNH}_2 \) or \( \text{NH}_2\text{OH} \)
its ammonia system analogs. It is a powerful,
but usually slow, oxidizing agent in acid or alkaline
solsn. It can also act as a reducing agent, eg with
\( \text{MnO}_4^- \) or \( \text{Ag}_2\text{O} \) (Ref 2). On the other hand, \( \text{H}_2\text{O}_2 \)
does not tend to form salts or substitute compds
(replacement of H) like its ammonia analogs
(See Ref 17)  

Analytical: Delicate qual tests for \( \text{H}_2\text{O}_2 \) are its
reactions with titanic chromate or sulfate in
acid to produce the bright blue peroxychromic
or bright yellow peroxy titanic acid both of
which are sol in ether. Quantitatively \( \text{H}_2\text{O}_2 \)
may be detd by its oxidation of I⁻ and titration of
the \( I_2 \) (or \( I_3^- \)) thus formed (Ref 2)  

Explosive and Combustion Props: Pure \( \text{H}_2\text{O}_2 \)
is readily detonable and its heat of expln is given as
24.6 kcal/mole. It is claimed that aq solns of
less than 94% \( \text{H}_2\text{O}_2 \) will decomp explosively if
catalyzed, but will not detonate (Ref 4). More
recent studies, however, show that 86% \( \text{H}_2\text{O}_2 \),
at 50° or higher and contained in 1.61 inch
ID Al tubes, will detonate at 5600 m/sec when
strongly boostered, but not at smaller diameters.
A 90.7% soln detonates at 5500 to 6000 m/sec
even at 0.5 inch ID and 25°. The gap over
which the 90.7% soln will propagate detonation
increases with increasing temp (Ref 23)  

Mixtures of \( \text{H}_2\text{O}_2 \)gas/air will not detonate
at subatmospheric press. However, a 35 mol% mixt
will detonate at 6700 ft/sec at 1 atm. The
ignition limit under these conditions is about
30 mol% \( \text{H}_2\text{O}_2 \) vapor (Ref 21)  

In an earlier study the min concns of \( \text{H}_2\text{O}_2 \)
in a vapor mixt below which powerful external
ignition cannot initiate a self-propagating re-
action were determined over the pressure ranges
of 1 to 6.5 atm. These min concns were found
to be 25.6 mole% at 1 atm (or a little lower
than above) and 20.7 mole% in the 2-6 atm
region (Ref 13)  

The explosive limits of \( \text{H}_2\text{O}_2 \) vapor were also
determined by using hot wire heated surfaces &
catalytic surfaces for ignition. At 1 atm, vapors
contg 26.0 mole% or more \( \text{H}_2\text{O}_2 \) could be ex-
ploed but only when catalytic surfaces were
used. Non-catalytic surfaces had to be pre-heated
to 150° to obtain explosions. The lower limits
at 200 and at 40mm Hg were 33 and 55 mole% 
resp. Hydrogen Peroxide explosion is believed to
be thermal, involving straight chains only,
with the initiating reaction being thermal
rupture of the O=O bond (Ref 5)  

At lower pressures (200mm), presumably at
elevated temps or on catalytic surfaces, the
ign limit is 32.5 mol% \( \text{H}_2\text{O}_2 \). It is not affected by
diluting the air with oxygen, nitrogen or He
in amounts of 0 to 39 mol%. Decreasing the
system volume by packing it with Raschig rings
shifts the ign limit to higher Hydrogen Peroxide
concns (Ref 10)  

Quenching distances & minium ignition
ergies of \( \text{H}_2\text{O}_2 \) & \( \text{H}_2\text{O} \) vapor mixtures were
studied in a flow system in which liq \( \text{H}_2\text{O}_2 \)
& \( \text{H}_2\text{O} \) was fed to a boiler where it completely
vaporized and passed through the explosion
vessel contg the ignition source & then was
condensed (Ref 14). The quenching distances &
min spark ignition energies of mixts contg be-
tween 35 & 50 mol% \( \text{H}_2\text{O}_2 \) were detd at pressures
between 15 & 200 mg of Hg at a temp 9°
above the condensation temp. The quenching
distances varied between 0.51 & 1.63cm & the
ignition energies between 0.53 & 25.5 millijoules,
decreasing with increasing \( \text{H}_2\text{O}_2 \) content &
pressure. The observed min energy for ign (in
millijoules) is related to the quench distance
(in cm) by:

\[ E_{\text{min}} = 3.84d_0^{3.04} \]  

Hydrogen peroxide, pure or aq, is readily
detonable when mixed with organic materials.
Detonation rates for \( \text{H}_2\text{O}_2 \) mixts with MeOH,
EtOH & glycerol were measured by Haeuseler
(Ref 7). These rates could be as high as 6700
m/sec for H₂O₂/H₂O/EtOH. The amount of energy to sustain explosive reaction in H₂O₂/H₂O/organic mixes depends not only on the peroxide/organic ratio but also on the water in the mix, the water acting as an energy sink. To a lesser extent explosibility is also affected by sample vol, initial temp, and to a considerable degree by the type of initiation used. Chem changes that occur in a mix can make a powerful explosive out of an apparently non-explosive mixt. Consequently, careful experimentation should precede the mixing or use of mixtures containing Hydrogen Peroxide and organic substances (Ref 15).

In view of these many factors found to influence the explosibility of H₂O₂/H₂O/org subst mixes, the simplified approach of Shanley and Perrin (Ref 12), which is based primarily on the enthalpy contents of these mixes, should only be used as a first attempt at determining the explosion hazard of these mixes. Their suggested isenthalpy line of 0.8 kcal/g of mixt defines the experimentally observed expl limits fairly well for initiation by blasting cap. However, their value of 1.2 kcal/g for impact initiation suggests that some mixes are safe which in fact were found to be in the explosive range (Ref 15).

The work of Kuchta (Ref 16) confirms that the relative concns of Hydrogen Peroxide/org subst are more important in defining the explosibility of a mixt than the type of org subst used. He points out that detonation can be initiated in certain H₂O₂ purge liquors that may form in commercial production of Hydrogen Peroxide.

**Hydrogen Peroxide Explosives.** Attempts were made in Austria during WWI to use H₂O₂ for the preparation of explosives, but they were not successful. Later, Bamberger and Nussbaum (Ref 1) succeeded in preparing several Hydrogen Peroxide explosives, which were successfully tried in blasting operations. The following explosives were prepared by mixing organic compounds with strong aqueous solutions (about 60% or higher) of peroxide:

a) Mixture of H₂O₂ (60% solution) with parafinomalousaldehyde forms a brisant explosive, which may be detonated either by heat or by a blasting cap. This mixture reacts with lead forming a crystalline compound of high brisance and sensitiveness, melting at about 50°

b) A mixture was prep'd by treating cellulose with 83.4% H₂O₂ to form a gelatinous mass. This mixt is more powerful than TNT or Tetryl in the nitramine, but is insensitive to shock or friction and burns without detonation when dropped in a red hot crucible or ignited by flame. Its ignit temp is 194-208°. It cannot be stored for longer than 24-48 hours, as bubbles, resulting from decomposition of the H₂O₂, are evolved and the mass hardens. Moreover, its explosive power decreased after 48 hours of standing (Ref 1).

A Spanish patent claims an explosive compn prep'd by mixing hexamethylenetetramine (See p H79 in this Vol) with Hydrogen Peroxide and then adding HCl (Ref 9).

An explosive insensitive to mech and rifle bullet impact, but detonable by a blasting cap, is claimed in a US patent. It consists of Hydrogen Peroxide, water & glycerol (Ref 3).

Another patented high explosive composition contains powdered metal and Hydrogen Peroxide, e.g. 29.7% pow'd B & 70.3% H₂O₂ (Ref 19). **Hydrogen Peroxide in Propulsion.** German use of high-conc Hydrogen Peroxide in propulsion, assisted take-off, torpedoes, special submarine turbines, etc during WWII is described in Ref 8.

Use of high-conc Hydrogen Peroxide for rocket propulsion, details of its manufacture and safety are described by Cleaver (Ref 11).

A review of the fire and explosion hazards of flight vehicle fuels includes discussion of Hydrogen Peroxide. It gives vap press data (Ref 19).

Acctg to Ref 14a, Hydrogen Peroxide has been used as torpedo propnt and rocket fuel oxidizer.


Hydrogen Peroxide (Additional Information on German Manufacture and Uses During WWII). Accdg to Dr Hans Walter, communicated during his work at Picatinny Arsenal, the name Wasserstoffperoxyd or Wasserstoffsuperoxid applies to any strength compd, while the name T-Stoff (T-stuff in Engl) is applied to 80–85% H₂O₂ and 20–15% H₂O. It was a clear viscous liquid fairly stable at ordinary temp and pressure when stored with a small amt of phosphoric acid, serving as a stabilizer. Such a mixt was known as T-Stoff(S) (Ref 4, p 8 & Ref 13, p Ger 210-R)

When H₂O₂, contg ca 20% H₂O, was stabilized with oxyquinoline (400mg per liter), the mixt was known as T-Stoff(SS) (Ref 4, p 9 & Ref 13, pp Ger 210-R & 211-L)

However, even with the greatest care, it was not possible to prevent on prolonged storage, a slow decompn into water and oxygen

For rapid estimation of its strength, either a hydrometer or titration with K or Na permanganate was used

Methods of manuf of H₂O₂ in Germany are described in Refs 2, 3, 5 & 6

One of the most interesting applications of T-Stoff was as a source of power turbines driving submarines as proposed by Dr Helmuth Walter. (See U-Boot, Walter in Ref 13, pp Ger 211-R). Seven of such submarines (300 to 500 tons each) were accepted by the German Navy up to the end of WWII

Besides submarine, the following applications of T-Stoff are listed in Ref 13, p Ger 210 a) A 500kg ATO (Assisted Take-Off) T-Stoff monofuel unit b) A 300-kg thrust, rocket propulsion unit for guided missiles c) A bipropellant 1000 to 1500-kg ATO d) A catapult with T-Stoff propulsion unit (decompn only) for launching V-1, 2 hicch is described in Ref 13, p Ger 213-L e) Controllable propulsion of a 750-kg thrust unit for Messerschmitt 263 f) Rocket training airplane and a controllable power plant giving to 2000 kg thrust for the Messerschmitt 263B

Accdg to Ref 4, p 8, T-Stoff was used as an oxygen carrier in some guided missile propgts, such as in Hecht (Pike), listed in Ref 13, p Ger 132-L as Pike Missile and described in Ref 11, pp 116–17. In order to accomplish this, T-Stoff was mixed with Z-Stoff which was an aq sln of K or Na permanganate, described in Ref 13, p Ger 264-R. This mixt produces superheated steam, attaining temp of 180°. This steam was also suitable for driving rockets and ATO, but was not suitable for driving turbines, because it contd small particles of MnO₂ which might foul the blades of turbine. However, the superheated steam formed as result of mixing T-Stoff with MP-14 catalyst, described in Ref 13, p Ger 114-L can be used for driving steam turbines

When T-Stoff was mixed with B-Stoff (Hydrazine Hydrate), in the presence of K cuprocyanide, the resulting liquid spontaneously ignited

Accdg to Ref 4, p 8, T-Stoff was also known
as Ingolin. Accdg to Ref 7, p 23, the code name T-Stoff was used only for 82% H₂O₂, while the code names Auroil, Neuralin and Subsidol were used for any 80 to 86% Hydrogen Peroxides


Hydrogen Selenide, H₂Se, mw 80.98; colorless gas, fr p -64°, bp -41.4°, d 3.614g/liter (gas), 2.12 at -42° (liq); vap press 10 atm at 23.4°; can be formed in small ams on heating Se in hydrogen & by action of nascent H on selenious acid. Pure H₂Se is obtd by action of water on Al selenide (Refs 1 & 2)

The compd is highly toxic by inhalation, strongly irritating to the skin, and causes damage to the lungs & liver. It reacts violently with oxidizing materials and forms expl mixts with air (Refs 1, 3 & 4)


Hydrogen Sulfide. H₂S, mw 34.08, colorless gas, mp -83.8°, bp -60.2°, auto ign temp 500°F, d 1.54g/l at 0°. It is highly toxic by inhalation and is a strong irritant to eyes and mucous membranes. It is highly flammable and has wide explosive limits in air: lower limit 4.3 vol%; upper limit 46 vol%

Ref: CondChemDict (1971), p 456

Hydrogen Telluride. H₂Te, mw 129.63, colorless gas or yellow needles, mp -49.0°, bp -2°, d 5.81g/l. It is highly toxic and is an irritant but it does not present the fire & explosion hazard of H₂S or H₂Se

Ref: CondChemDict (1971), p 456

Hydrogen Tetraoxide. H₂O₄, mw 66.02. This substance exists only at low temps and is probably H₂O₄ rather than H₂O₃ (Ref 3). It is probably non-planar with a ΔH°₂⁻ = -27.9kcal/mole (Ref 1). It has been prepd by reacting H₂O₃ or H₂O and its reactions with NH₃, Cl & C₂H₄ have been examined (Ref 2). A review of the general subject of higher peroxides, including H₂O₄ is given in Ref 3


Hydrolysis. The chemical reaction of a substance with water to form one or more new substances. Examples of hydrolysis are: the catalytic conversion of starch into glucose; the catalytic or enzymatic conversion of sucrose into glucose and fructose; the conversion of natural fats into fatty acids and glycerin.

In general hydrolysis is a reaction of the type: AB+H₂O = AOH+HB, which in its ionic form: H₂O = H⁺+OH⁻ is the reverse of neutralization. As examples:

In inorganic chemistry:
CuSO₄ + 2H₂O = Cu(OH)₂ + H₂SO₄

In organic chemistry:
RCOO₉⁺ + H₂O = RCOOH + RH⁺

In biochemistry (by enzymes):
R.CNH₂R₂⁺ + H₂O = R.COONH₂ + NH₄⁺
C₁₂H₂₂O₁₁ + H₂O = 2C₆H₁₂O₆

Refs: 1) Hackh's Dict (1944), 423-R
2) Groggins (1952), 651-70

Hydrolysis of Plasticizers for NC by Water at 22° and 60°C. Following tests were conducted in 1945 at Picatinny Arsenal by P. F. Macy and A. A. Saffitz:

a) Place a sample of about 5g of material to test in a small flask containing 12.5 ml of CO₂-free neutral distilled water. Stopper and shake for 10 days at 22° in a rotary, end over end, shaker apparatus placed in a constant temperature room.

b) Determine the liberated during the exposure period by titrating the sample water mixture with either N/10, or N/50 sodium hydroxide solutions in presence of phenolphthalein indicator. Titrations should be conducted separately on the oil and water layers of the mixture.

If the organic acid liberated during the hydrolysis is more soluble in the oil layer than in the water layer, 5ml of cp acetone should be added to the oil to reduce possible errors arising due to low solubility of the standard alkali in the oil. Blank determination shall be made in all cases with water and acetone, and the necessary corrections applied.

c) Express the acidity as grams of acid produced (mineral acid in case of ester nitrates, and organic acid in case of phthalates, acetates, lactate-nitrates, acetate nitrates) per 100g of starting material at the specified temperature for the entire test period.

d) Hydrolysis at 60° ± 1°C was conducted by shaking for 5 days in a rocker device contained in Cenco Electric Oven, Model No 95, 105A.

Following are some results:

<table>
<thead>
<tr>
<th>Substance</th>
<th>Hydrolysis: % acidity</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>At 22°</td>
</tr>
<tr>
<td>Butanediol-1,3-dinitrate (BDN-1,3)</td>
<td>—</td>
</tr>
<tr>
<td>Butanediol-1,4-dinitrate (BDN-1,4)</td>
<td>—</td>
</tr>
<tr>
<td>Diacetyl mononitrate (DAM)</td>
<td>—</td>
</tr>
<tr>
<td>Dibutylphthalate (DBP)</td>
<td>0.002</td>
</tr>
<tr>
<td>Diethylene glycol dinitrate (DEGDN)</td>
<td>0.003</td>
</tr>
<tr>
<td>Dinitroethylbenzene (DNEB)</td>
<td>0.007</td>
</tr>
<tr>
<td>Dipropylene glycol dinitrate (DPGDN)</td>
<td>—</td>
</tr>
<tr>
<td>Dimethyl methylol nitromethanemononitrate</td>
<td>—</td>
</tr>
<tr>
<td>Ethyltrimethyl methanetrimonitrile (EMMET)</td>
<td>0.003</td>
</tr>
<tr>
<td>Glycerol mono lactate trinitrate (GLTN)</td>
<td>0.021</td>
</tr>
<tr>
<td>Glycol mono ethylether mononitrate (GEEN)</td>
<td>—</td>
</tr>
<tr>
<td>Glycol mono methyl ether mononitrate (GMEN)</td>
<td>—</td>
</tr>
<tr>
<td>Metroll trinitrate (MTN)</td>
<td>0.018</td>
</tr>
<tr>
<td>Monoacetinedinitrate (MAD)</td>
<td>—</td>
</tr>
<tr>
<td>Nitroglycerin (NG)</td>
<td>&lt;0.002</td>
</tr>
<tr>
<td>Triethylene glycol dinitrate (TEGDN)</td>
<td>0.032</td>
</tr>
<tr>
<td>Trinitrophenyl ethanol mononitrate (TNPEN)</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

Ref: P.F. Macy & A.A. Saffitz, PATR's 1616 (1946); 1638 (1947)

Hydrolysis Test for Nitrocelluloses. The following hydrolysis test was developed and is used at Picatinny Arsenal.

a) Weigh accurately a 5g sample of dry NC, place in a tared 250ml Pyrex flask provided with a ground connection to fit a Pyrex condenser.
b) Add 100ml of distilled neutral (conductivity) water and boil the water under reflux condenser during 240 hours.

c) Cool the flask and filter off the water into a 500ml volumetric flask. Rinse the NC with several portions of conductivity water, using sufficient amount to fill the flask to 500ml mark.

d) Stopper the flask, shake and take an aliquot. Determine its pH, using a standard procedure.

e) Take another aliquot and titrate it with a standard NaOH soln in presence of methyl red indicator.

Note: In one of the tests conducted at PicArsenal the Pyro (N=12.6%N) gave 1.22%, calc as HNO₂ and Guncotton (N=13.45%) gave 1.03%

Refs: 1) D.D. Sager, PATR 174 (1932)
2) W.R. Tomlinson & O.E. Sheffield, AMCP 706-177 (Jan 1971), p 9

Hydrolitic Reaction. Pertaining to Hydrolysis. Comp with Hydroxylation
Ref: Hackhi's Dict (1944), 423-R

Hydrometer. A device for measuring specific gravity of liquids. Its description with Fig is given in Vol 3 of Encycl, p D67, under DENSITY AND SPECIFIC GRAVITY

Hydronitrile Acid. Same as Hydrazoic Acid or Azoinide. See in Vol 1, p A537-R

Hydronitriles. Same as Azides

HYDRONITROGENS. Compounds containing only hydrogen and nitrogen. Some thirteen different hydronitrogens are known, mostly not in the free state, but as derivatives. There are saturated and unsaturated derivatives.

Saturated Hydronitrogens (Type formula NₓHᵧ). Ammonia NH₃, Hydrazine (diaza- mide) H₂N, NH₂; Triazane (proazane) H₃N, NH₂, NH₂, Tetrrazane (buzane, hydrotetrazane) H₄N, NH₂, NH₂, NH₂. Ammonia and Hydrazine are known in the free state; the Triazanes are rather poorly defined as a class, whereas Tetrrazanes exhibit a tendency towards instability by undergoing dissociation in solution to yield hydrayzyl radicals.

Unsaturated Hydronitrogens (Type formula NₓHᵧ). Diamide HN:NH; Triazene (diazoamine) HN:NH₂; Tetrazine (tetrazene, or 2-tetrazene) H₂N:NH₂; Isotetrazene (1-tetra- zene, diazohydrazene, buzylene) HN:NH₂; Ammonium Azide, NH₄N₃; Hydrazine Azide, N₃H₅.N₃.

Type formula NₓHᵧ-

Hydrogen Azide (hydrazoic acid, azoinide, hydronitric acid, triazoeic acid) HN:N; Diamino-
Hydrazine HN:NH₃.NH₃; Bisdiazooamine
HN:NH₂.NH₂; Hexadiaziodene HN:NH₂.NH₂; Hexadiazoidene HN:NH₂.NH₂.

Type formula NₓHᵧ-

Octazotriene (octazone) HN:NH₂.NH₂.NH₂.NH₂

Many of the derivatives of the above compounds are explosive and they are described separately under corresponding names. Some of the compounds included in these tables (as for instance ammonium azide and hydrazine azide), do not possess the structural formula of real hydronitrogens but they are included for the sake of comparison, because their empirical formulas correspond to the type NₓHᵧ-

2) C.C. Clark, “Hydrazine”, Mathieson Chem Corp, Baltimore, Md (1953)

Hydronitrous Acid or Nitroxylic Acid, H NO₂, mw 48.02. The hypothetical acid from which nitroxyls are derived. The nitroxy radical is -NO₂, when attached to a strongly electronegative group such as F or Cl or to metal forming compds such as NO₂F, NO₂Cl or Cu₂(NO₃)₂.

Ref: Hackhi's Dict (1944), 424-R & 580-R

Hydronium ion. The solvent hydrogen ion, H⁺(H₂O) or H₃O⁺, which is considered to be present in all acids:

HCl+H₂O → H₃O⁺+Cl⁻

Hydroperoxides, Organic. Are monosubstituted derivatives of hydrogen peroxide ROOH, where R is alkyl or aralkyl radical. They generally may be prep by the following methods: a) alklylation of H₂O₂ with alkyl halides, sulfates, or alcohols in the presence of strong acids; b) controlled oxidation of the hydrocarbon RH with O₂; c) addition of O₂ to Grignard reagents. A brief discussion of prepn and properties of hydroperoxides (including explosive: methyl-, ethyl-, isopropyl-, t-butyl-, tetralin-, decalin-, and phenylhydrazone- hydroperoxides) is given in Ref 2.
A table of the explosive nature of peroxides, taken from Ref 3, is given below:

**EXPLOSIVE NATURE OF PEROXIDES**

<table>
<thead>
<tr>
<th>Peroxide</th>
<th>Remarks Concerning Explosiveness</th>
<th>Ref</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>ALKYL HYDROPEROXIDES</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methyl Hydroperoxide</td>
<td>Explodes violently; sensitive to jarring especially at warm temps; greatest caution in making, Ba salt in dry state extremely explosive</td>
<td>A. Rieche &amp; F. Hitz, Ber 62, 2458 (1929)</td>
</tr>
<tr>
<td>Ethyl Hydroperoxide</td>
<td>Explodes on superheating quite violently, Ba salt, heat and percussion sensitive</td>
<td>A. Baeyer &amp; V. Villiger, Ber 34, 738 (1901)</td>
</tr>
<tr>
<td>Isopropyl Hydroperoxide</td>
<td>Explodes just above bp</td>
<td>S. Medvedev &amp; E.N. Alexejeva, Ber 65, 133 (1932)</td>
</tr>
<tr>
<td>t-Butyl Hydroperoxide</td>
<td>By distn under normal pressures explosions can result; otherwise relatively harmless</td>
<td>N.A. Milas &amp; D. Surgener, JACS 68, 205 (1946)</td>
</tr>
<tr>
<td>Tetralin Hydroperoxide</td>
<td>Detonated by superheating</td>
<td>H. Hock &amp; W. Susenichl, Ber 66, 22 (1944)</td>
</tr>
<tr>
<td>Decalin Hydroperoxide</td>
<td>Very stable; can be sublimed in small quantities at atm pressure</td>
<td>R. Criegee, Ber 77, 22 (1944)</td>
</tr>
<tr>
<td>Triphenylmethyl Hydroperoxide</td>
<td>Not explosive</td>
<td>H. Wieland &amp; J. Maier, Ber 64, 1205 (1931)</td>
</tr>
<tr>
<td>Phenylhydrazone Hydroperoxide</td>
<td>Detonates in dry state after a short time with no evident cause</td>
<td>W. Busch &amp; W. Dietz, Ber 47, 3277 (1914)</td>
</tr>
<tr>
<td><strong>DIALKYL PEROXIDES</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dimethyl Peroxide</td>
<td>Extremely explosive on heating and jarring; vapors also sensitive to shock</td>
<td>A. Rieche &amp; W. Brumshagen, Ber 61, 951 (1928)</td>
</tr>
<tr>
<td>Methylethyl Peroxide</td>
<td>Liquid and vapor forms shock-sensitive; explodes violently on superheating</td>
<td>A. Rieche, Ber 62, 218 (1929)</td>
</tr>
<tr>
<td><strong>DIACYL PEROXIDES</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diacetyl Peroxide</td>
<td>Highly explosive; to be handled only with extraordinary caution</td>
<td>L.P. Kuhn, C&amp;EN 26, 3197 (1948)</td>
</tr>
<tr>
<td>Bishexahydrobenzoyl Peroxide</td>
<td>Large quantities can explode without apparent reason</td>
<td>F. Fichter &amp; W. Siegrist, HelvChimActa 15, 1304 (1932)</td>
</tr>
<tr>
<td>Difuroyl Peroxide</td>
<td>Explodes violently by rubbing and heating</td>
<td>N.A. Miles &amp; A. McAlevy, JACS 56, 1219 (1934)</td>
</tr>
<tr>
<td>Dibenozyol Peroxide</td>
<td>Exploded by heat; carry out recrvstns of large quantities without heating</td>
<td>K. Nozaki &amp; P.D. Bartlett, JACS 68, 1686 (1946)</td>
</tr>
<tr>
<td>1-Naphthoyl Peroxide</td>
<td>Explodes by rubbing</td>
<td>S. Mededeco &amp; O. Bloch, ChemZhl 1935 I, 2670</td>
</tr>
</tbody>
</table>


## EXPLOSIVE NATURE OF PEROXIDES (Cont'n)

<table>
<thead>
<tr>
<th>Peroxide</th>
<th>Remarks Concerning Explosiveness</th>
<th>Ref</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydroxymethyl Peroxide</td>
<td>Explodes on heating; not sensitive to friction; higher homologs not explosive</td>
<td>A. Riche &amp; R. Meister, Ber 68, 1465 (1935)</td>
</tr>
<tr>
<td>Hydroperoxide</td>
<td></td>
<td>A. Riche, Ber 64, 2328 (1931)</td>
</tr>
<tr>
<td>Hydroxymethyl</td>
<td>Violently explosive, on heating becomes percussion-sensitive</td>
<td>A. Riche &amp; F. Hitz, Ber 62, 2458 (1929)</td>
</tr>
<tr>
<td>Methyl Peroxide</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1-Hydroxyethyl</td>
<td>Detonates on heating</td>
<td>A. Riche, Ber 63, 2642 (1930)</td>
</tr>
<tr>
<td>Ethyl Peroxide</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bis-hydroxymethyl Peroxide</td>
<td>Highly explosive; strongly friction-sensitive</td>
<td>H. Wieland &amp; H. Sutter, Ber 63, 66 (1930)</td>
</tr>
<tr>
<td>Bis(1-hydroxycyclohexyl) Peroxide</td>
<td>Harmless by itself; but explodes on attempted vacuum distn</td>
<td>M. Stoll &amp; W. Scherrer, HelvChimActa 13, 142 (1930)</td>
</tr>
<tr>
<td>Bis-hydroperoxydicyclohexyl Peroxide</td>
<td>Detonates very actively in a flame</td>
<td>R. Criegee, W. Schnorrenberg &amp; J. Becke, Ann 566, 7 (1949)</td>
</tr>
<tr>
<td>1-Hydroperoxy-1-acetoxy-cyclodecan-6-one</td>
<td>Detonates on removal from a freezing mixture</td>
<td>R. Criegee &amp; G. Wenner, Ann 564, 9 (1949)</td>
</tr>
<tr>
<td>Dimeric Ethyldene Peroxide</td>
<td>Explodes with extreme violence just by touching; <em>greatest caution!</em></td>
<td>A. Riche &amp; R. Meister, Ber 72, 1933 (1939)</td>
</tr>
<tr>
<td>Polymeric Ethyldene Peroxide (Ether Peroxide)</td>
<td>Extremely explosive, even below 100°C</td>
<td>Eg, A. Riche &amp; R. Meister, AngewChem 49, 101 (1936)</td>
</tr>
<tr>
<td>Trimeric Propyldene Peroxide</td>
<td>Extremely explosive; very friction-sensitive</td>
<td>A. Riche &amp; R. Meister, Ber 72, 1938 (1939)</td>
</tr>
<tr>
<td>Dimeric Acetone Peroxide</td>
<td>Explodes violently by percussion and rubbing</td>
<td>A. Baeyer &amp; V. Villiger, Ber 32, 3632 (1899)</td>
</tr>
<tr>
<td>Trimeric Acetone Peroxide</td>
<td>Very explosive; can penetrate a plate of iron when heated on it</td>
<td>A. Baeyer &amp; V. Villiger, Ber 32, 3632 (1899)</td>
</tr>
</tbody>
</table>
Properties of hydroxalkyl hydroperoxides and hydroxyalkyl peroxides are shown in the following table (from Ref 3):

**PEROXY DERIVATIVES OF ALDEHYDES AND KETONES**

<table>
<thead>
<tr>
<th>Name</th>
<th>Formula</th>
<th>( n_D^{20} )</th>
<th>mp, C</th>
<th>bp, C/mm Hg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydroxymethyl Hydroperoxide</td>
<td>HOCH OOH</td>
<td>1.4205 (16)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>1-Hydroxyethyl Hydroperoxide</td>
<td>CH CH(OH)OOH</td>
<td>1.4250 (24)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>1-Hydroxyhexyl Hydroperoxide</td>
<td>C H CH(OH)OOH</td>
<td>-</td>
<td>40</td>
<td>-</td>
</tr>
<tr>
<td>1-Hydroxydodecyl Hydroperoxide</td>
<td>C H CH(OH)OOH</td>
<td>-</td>
<td>65 to 67</td>
<td>-</td>
</tr>
<tr>
<td>1-Hydroxycyclohexyl</td>
<td>HOOHO</td>
<td>-</td>
<td>76 to 78</td>
<td>-</td>
</tr>
</tbody>
</table>

**B. Bis(1-hydroxyalkyl) Peroxides**

<table>
<thead>
<tr>
<th>Name</th>
<th>Formula</th>
<th>( n_D^{20} )</th>
<th>mp, C</th>
<th>bp, C/mm Hg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bis(hydroxymethyl Peroxide)</td>
<td>CH(OH)OOCH(OH)</td>
<td>-</td>
<td>62 to 64</td>
<td>-</td>
</tr>
<tr>
<td>Bis(1-hydroxyethyl) Peroxide</td>
<td>CH CH(OH)OOCH(OH)CH</td>
<td>1.4265 (16)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Bis(hydroxybenzyl) Peroxide</td>
<td>C H CH(OH)OOCH(OH)C H</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Bis(hydroxy-(\beta),(\beta),(\beta)) trihaloethyl) Peroxide</td>
<td>CCl CH(OH)OOCH(OH)CCl</td>
<td>-</td>
<td>122</td>
<td>-</td>
</tr>
<tr>
<td>Bis(1-hydroxycyclohexyl) Peroxide</td>
<td>HOHOHOHOHOHO</td>
<td>-</td>
<td>68 to 70</td>
<td>-</td>
</tr>
</tbody>
</table>

The reduction of hydroperoxides with LiAlH\(_4\) yields the corresponding alcohols probably via an LiAl(OR)\(_4\) intermediate. However this reaction with B\(_2\)O\(_2\) resulted in an explosion (Ref 1).

Cumene hydroperoxide (91-95\% pure) will not detonate even when strongly booster. However, it is easily ignitable & can burn with sufficient violence to rupture steel distillation equipment (Ref 4).


**Hydroquinone and Derivatives,**

Hydroquinone, Hydroquinol, 1,4-Benzenediol or \(p\)-Dihydroxybenzene (Hydrochinon in Ger). See 1,4-Dihydroxybenzene in Vol 5 of Encycl, p D1270-R

**Hydrous.** A compd contg water, as opposed to anhydrous. In case of salts it is water of crystallization

Ref: Hackh's Dict (1944), 425-R; (1969), 335-L
Azidohydroquinone, N₃C₆H₃(OH)₂; mw 151.12, N 27.81%, OB to CO₂ -132.3; leaflets, mp explodes violently on heating; sol in alc or eth, insol in petr eth. It can be prep'd from quinone and an excess of hydrazoic acid in benz.
Refs: 1) Beil 6, (419) 2) E. Oliveri-Mandala & E. Calderaro, Gazz. 45 I, 312 (1915); 45 II, 120 (1915) & CA 10, 596-7 & 1514-5 (1916)

Diazidodinitro hydroquinone. (OH)₂C₆H₃(N₃)₂ (NO₂)₂ (probably), mw 284.15, N 39.41%, OB 11.2, fine yellow needles (from mixed acid). It is claimed that the K salt is prep'd by reacting nitroaminoresorcinol with conc. mixed acid containing an excess of K-nitrate.
The 3,5-dihydroxyquinonediazide (sic) is claimed to be prep'd similarly from the mono or diaminophenol glucinols. These cmpds are claimed to be unstable in detonators in conjunction with PETN or other HE as the base charge.
Refs: E. von Herz, BritPat 207563 (1922) & CA 18, 1574 (1924)

Nitrohydroquinone. OH C₆H₃(NO₂)OH, mw 155.11, N 9.03%, red rhomboidal grains or pyramids (from w) mp 133-134°, sl sol in w; v sol in alc & eth. Prep'd from o-nitrophenol & aq NaOH + ammonium persulphate and then dil sulfuric acid (Ref 1); by nitrating monobenzensulfonate (Ref 4).
The effect of pH on the uv spectrum of nitrohydroquinone has been studied (Ref 2), also its dissociation constants, redox potentials & dipole moments (Ref 5), as well as its electronic spectra (Ref 6). Its thin layer chromatographic props were also investigated (Ref 3).

2-6 Dinitrohydroquinone, (OH)₂C₆H₃(NO₂)₂, mw 200.11, N 14.00%, golden platelets (from w) mp 135-136° (some color change), sol hot water, alc, eth. Prep'd by nitrating hydroquinonediacetate (Ref 1); nitration of monobenzensulfonate (Ref 2). Its Ba salt, BaC₆H₃(NO₂)₂, dk-blue crysts, when anhyd is very expl (Ref 1a)

Trinitrohydroquinonediethylether,
(O₂N)₂ C₆H₄(OC₂H₅)₂; mw 301.21, N 13.95%; yellow needles that turn yellow-orange on exposure to light, mp 130°, insol in w, somewhat sol in alc, eth or benz. Prep'd by nitrating hydroquinonediethylether with mixed acid.
Refs: 1) Beil 6, 859 2) Nietzki, Ann 215, 153 (1882)

Hydrosilicofluoric Acid. See Fluorosilicic Acid in Vol 6 of Encycl, p F141-R

Hydrosol. A colloidal suspension in water (Ref 1). Also Brand name of L.B. Holliday & Co, Ltd, Huddersfield, England for a proprietary product of the hydrosulfite class for wool bleaching (Ref 2).

Hydrostatic Fuze. See Vol 4, p D881-R

Hydrostatics. The science of physics which deals with liquids in equilibrium (Comp with Hydrodynamics)
Refs: 1) Hack's Dict (1944), 425-L; (1969), 335-L

Hydrox. Is a blasting device for underground mining primarily used in coal mines. Like Cardox (see Vol 2, p G67-R) it employs a gas generating chemical reaction to rupture a disk in a heavy-walled steel tube with the resulting flow of high pressure gas then doing mechanical work on the surroundings. The usual gas producing system is (Ref 1):

NaNO₂ + NH₄Cl = NaCl + N₂ + 2H₂O

This reaction is self-sustaining (once started) but
does not generate flames. Usually the rupture disk is selected to break at about 10 tons per sq inch (Ref 1).

A simple calcn shows that for this reaction the heat of explosion is 52.4 kcal/mole or 428 kcal/ kg; the explosion temp (the temp that can be reached by explosion within a closed vessel) is about 1430°, and the vol of the gas evolved at normal temp & pressure is 540/l. The ignition temps of mixts of various NaNO₃/ NH₄Cl ratios, r, was found to be the lowest for r=1:2. The mixts, when stored over CaCl₂, undergo spontaneous explosion after 30-50 days, but stabilizers such as 2% of: MgO, CaO, CaC₂, BaCl₂, MgCO₃, Na₂CO₃ were effective in preventing explosions for more than a year.

The max pressure developed by the explosion was 790 kg/cm² at a loading dens of 0.3; 1200 kg/cm² at a loading dens of 0.4; and 1700 kg/cm² at a loading dens of 0.5 (Ref 2).

Other reactions have been patented for use in Hydrox. Some additional ingredients claimed in these patents are AN, NH₄HCO₃, chromates and dichromates. These compositions and the use of Hydrox are discussed in detail in Refs 3 & 4.


Hydrox Fuel Cell. See under Fuel Cells in Vol 6 of Encyc, p F210-R

Ref: CondChemDict (1961), 517-L (Fuel Cells); (1971), 403-R

Hydroxides. The generic name for compounds of the type $M_a(OH)_b$, where M is a metal and a & b are integers. Most hydroxides are insol or sparingly sol in water but are sol in acq acid with the formation of salts. Most hydroxides have been used in neutralizing nitrated explosive made by mixed acid or nitric acid nitration although bicarbonates are more commonly employed. The more common hydroxides are listed below:

Aluminum hydroxide. $\text{Al(OH)}_3$, mw 77.99, white cryst powder, d 2.42, insol in w, sol in acids & NaOH. Derived from bauxite

Ref: CondChemDict (1971), p 33

Ammonium hydroxide or Ammonia, Aqueous. See under AMMONIA in Vol 1 of Encycl, p A296 to A305 and in CondChemDict (1971), p 52-R

Barium Hydroxide. Exists as Anhydrous, $\text{Ba(OH)}_2$, available commercially; Monohydrate, $\text{Ba(OH)}_2 \cdot \text{H}_2\text{O}$, wh powd, highly toxic, has many industrial uses; Pentahydrate, $\text{Ba(OH)}_2 \cdot 5\text{H}_2\text{O}$, translucent free-flowing wh flakes, d 65lbs/cu ft, highly toxic; and Octahydrate, wh crys, mp 78°, loses w of crystallization at 408°, d 2.18 g/cc; sol in water, alc & eth, highly toxic; prep by dissolving Ba oxide in w with subsequent crystallization and by other methods, used in org preps, Ba salts & in analytical chem


Calcium hydroxide. $\text{Ca(OH)}_2$, mw 74.10, rhombic colorless crystals, mp (loses water at) 580°, d 2.343, v sl sol in w, insol in alc, sol in acq acids; absorbs carbon dioxide from air. Prep by action of water on CaO


Lithium Hydroxide. LiOH, mw 23.95, white powder, mp 450°, d 1.43; sol in water & acids, sl sol in alc. Prep by action of water on Li or Li₂O


Magnesium Hydroxide. $\text{Mg(OH)}_2$, mw 58.34, colorless trigonal crystals, loses w at 380°, d 2.38, sl sol in hot water, sol in alc, sol in acq acids & in solns of ammonium salts. Obtainable from sea water by precipitation with CaO

Ref: CondChemDict (1971), p 535

Potassium Hydroxide. KOH; mw 56.11, white, rhombic, deliquescent crystals; mp 360°, bp 1320°, d 2.044; sol in w, alc or acids; insol in eth. Prep by electrolysis of concd KCl solns

Sodium Hydroxide. NaOH; mw 40.01; white deliquescent crystals, mp 318°, bp 1390°, d 2.13; sol in w, alc, glycerin & acids; insol in eth or acetone. Prep'd by electrolysis of NaCl solns
2) US Spec for Tech NaOH O-S-598A, IntAmel
(1) (July 1968)

Hydroxy (Hydroxyl). A prefix indicating the presence of the monovalent OH group in an org compd analogous to acids of the lactic series. It is preferable to oxy- used by the Germans
Ref: Hack's Dict (1944), 425-R; (1969), 335-R

Hydroxyacetanilide and Derivatives
Hydroxyacetanilide. Same as Acetamidophenol, Vol 1, p A20-R
2-Hydroxy-3,4,6-trinitroacetanilide. See 3,5,6-Trinitro-2-acetamidophenol in Vol 1, p A21-R

Hydroxyacetamide. See Glycolic Acid Azide, or Glycolyl Azide in Vol 6, p G115-R

Hydroxyacetophenone and Derivatives
Hydroxyacetophenone. OHC6H4COCH3, mw 136.14. Three isomers are known: ortho — greenish-yel oil, fr p 4—6°, bp 215° at 717mm, sp gr 1.1037 at 21°, ref. ind. 1.5580 at 20°; sol in w; sol in alc & eth (Ref 1, 4 & 5); meta — crysts, mp 95—6°, bp 296° at 756mm; sol in alc & eth and hot w (Ref 2); para — crysts, mp 109°, bp 147—48° at 3mm, sol in w, sol in alc & eth (Ref 3)

Hydroxyacetophenone Azide (called Azido-4-oxy-acetophenone in Ger) HOC6H4COCH2N3, mw 177.16, N 23.72%; yel leaflets (from w) mp 136°, sol in the usual solvents except benzene; decomposes in boiling w Prep'd by heating 4-hydroxy-chloroacetophenone with Na Azide in alc. No explosive props are mentioned
Ref: Beil 8 [87]

Hydroxy-trinitro-acetophenone (called Trinitro-4-oxy-acetophenone in Ger), HOC6H4(NO2)3COCH3; mw 271.14, N 15.50%, crysts; sl sol in hot w. Prep'd by heating 4-hydroxyacetophenone with 1.3 g/cc nitric acid; No explosive props mentioned
Refs: 1) Beil 8, 89 2) Nencki & Stoebel, Ber 30, 1770 (1897)

Hydroxyalkyl Alkylene Dinitramines and Their Nitrate Esters. Nitrate esters such as N,N′-bis-(2-nitroxyethyl) ethylene-nitramine.
O2NO-C6H4-N(NO2)2-C6H4-N(NO2)2-C6H4-ONO2 (qv); were proposed as non-volat plasticizers for NC in prepn of propellants. The method of prepn consists of the hydroxyalkylation of a nitramine (such as ethylene-nitramine) by means of an alkylene oxide (such as ethylene oxide), followed by nitration:
CH2-NH2NO2 + CH2 = CH2. NO2 = CH2 = CH2. NO2
HNO3 + CH2-NH2NO2 + CH2 = CH2. NO2 = CH2 = CH2. NO2
N,N′-bis(2-hydroxyethyl) ethylene-nitramine
Ref: J. R. Johnson et al, USP 2683165 (1954) & CA 49, 7590 (1955)

Hydroxyaminanthraquinones and Derivatives.
See Aminohydroxyanthraquinone and Derivatives in Vol 1 of Encycl, p A217-L. Its Tetranitro-
derivative (Aminochrysanemic Acid), described on p A217-L, forms expl salts of Ammonium, Barium and Potassium

Hydroxyaminobenzene. See Aminophenol in Vol 1, pp A241-L to A244-R. Its expl derivs: 1-Hydroxy-2-amin-4,6-dinitrobenzene or Picramic Acid and 1-Hydroxy-4-amin-2,6-
dinitrobenzene or Isopicramic Acid are on pp A241-R & A243-R

Hydroxyaminobenzoic Acid. See Aminosalicylic Acid in Vol 1, p A257
1-Hydroxy-6-aminobenzotriazole or 6-Aminobenzotriazidone (called oxy-6-aminobenzotriazole in Ger)

\[
\text{H}_2\text{NC}_6\text{H}_3\xrightarrow{\text{N}}\text{N}_\text{OH, mw 150.14, N 37.32\%, rose-}
\]

-colored powder, mp 220° (with carbonization), puffs off at 235-36° without melting; sol in hot water & hot aq; insol in eth or ligroin. Prep by treating 1-hydroxy-6-nitro-benzotriazole with Sn & concd HCl.


Hydroxyaminobutane. See Aminobutanol in Vol 1, p A192-R

Hydroxy and Amino Compounds, Explosives from. A number of commercially available amino alcohols have been condensed with 2,4-dichloro-benzene to give products which, when nitratated, are explosive. Products from the nitration of several glycerol derivs, guanidine derivs & phenylbenzotriazole derivs were also studied. Prepn of the following compds & their props are reported:

- Trinitrophenylnitroguanidine
- N-(beta-Hydroxyethyl)-N'-nitroguanidine Nitrate
- Tris-Hydroxymethylguanidine
- N,N'-Dinitropiperazine
- Tetryl

Of other expls produced, Heptyl, Ditetyl & Trinitrophenylbutylnitramine Nitrate appeared especially worthy of further study.


Hydroxyaminomethylbenzene or Hydroxyaminotoluene. See Aminocresol in Vol 1, p A193-R

Hydroxyaminomethylpropane. See Aminomethylpropanol in Vol 1, p A233-L

Hydroxyaminopropane. See Aminopropanol, Vol 1, p A253-L & R

Hydroxy- and Amino-methylnitramines. A review with 93 refs of the chemistry of nitramines is reported by Lamberton (Ref) Ref: A. H. Lamberton, Quart Revs 5, 92-95 (1951) & CA 46, 6081 (1952)

Hydroxyanilinobenzene-diazonium. See Anilino-benzenediazonium Hydroxide in Vol 1, p A421

Hydroxyanilinopropane. See Anilinopropanol in Vol 1, p A436-L

Hydroxyanthroquinone and Derivatives

1-Hydroxyanthraquinone, C_{6}H_{4}(CO)_{2}C_{6}H_{3}-OH; mw 224.20. Two isomers of this formula are known:

1-Hydroxy-, red- ornd ndls (alc), mp 194-95°, bp - subl; sol in alc & v sol in eth. Can be prep'd by heating phthalic acid anhydride with phenol in the presence of ZnCl_{2} & HCl at 120-25° (Refs 1 & 3)

2-Hydroxy-, yel ndls (alc), mp 306-08°, bp - subl; v sol in w; sol in alc & eth. Can be prep'd by diazotizing 2-a�noinoanthraquinone in HCl soln and boiling the diazonium sulfate soln (Refs 2 & 4)

Refs: 1) Beil 8, 338, (650) & [388]

1-Hydroxy-4, 5, 8-trinitroanthraquinone

(NO_{2})_{3}C_{6}(OH) COOC_{6}H_{4}, mw 359.20, N 11.0\%, cryst, mp 245°. Prep'd by treating the corresponding dihalogen or dinitro anthraquinone with a metal nitrite in an organic solvent.

Refs: 1) P. Belshaw et al, USP 2,587,093 (1952) & CA 46, 8679 (1952) 2) P. Belshaw et al, BritPat 670,720 (1952) & CA 46, 10205 (1952)

Hydroxyazapropane Nitroxy-Nitro Derivative. See NENA or N-(beta-Nitroxyethyl)-nitramine in Vol 1 of Encycel, p A201-L
1-Hydroxy-III-azidoiminomethyl-triazene or N'-
Hydroxy-N-azidoiminomethyl-triazene,
HON.NNHC.(NH)N₃, mw 129.09, N 75.97%.
Inasmuch as this compound was never isolated
in the free state but was prep'd only in solution,
it's structure has not been definitely established.
Hofmann et al (Refs 1 & 2) claimed that when
they treated guanyldiazoguanyltetrazene (called
by them simply "diazohydrate," but now known as "Tetrazene")
with KOH or NaOH in the
presence of water, the insoluble Tetrazene went
into solution accompanied by the evolution of ammonia.
When this solution was acidified, the
original "Tetrazene" was not recovered. When the
alkaline solution of Tetrazene was allowed
to evaporate in a vacuum over concd H₂SO₄,
an explosion took place while the mass was still
wet.

When an alkaline solution of Tetrazene was
acidified with dilute HNO₃ and then treated
with an excess of AgNO₃, a pale-yellow ppt was
obtained. This ppt was washed with a 3% solution
of HNO₃ and the residue dissolved in a
2.5% solution of ammonia. The resulting solution
with a pale yellow color gave, after evaporating
the ammonia in vacuum over H₂SO₄ or on
acidification by HNO₃, a white ppt which the
authors called "Tetrazylazoimidazilsilber,"
N₃CH₄,Ag. This compound exploded violently
when heated in water on a water bath. It also
exploded when the incompletely dried material
was touched with a feather.

The same authors claimed that they prep'd
a copper salt of the following composition:
\[
\text{CN₇HO.Cu, or N₃.C} = \frac{\text{Cu.O}}{\text{NH.N:N}} \quad \text{N 51.41%}, \text{light-}
\text{green plates}
\]

For preparation of this salt, they dissolved
Tetrazene in very dilute pure NaOH solution and
then added immediately, without acidification, an
excess of copper acetate solution. The resulting
brownish ppt was separated and washed with
2-3% solution of acetic acid followed with water.
On treating the ppt with 3% solution of ammonia, a blue solution was obtained,
leaving a brown residue of copper cyanimide.
When the blue filtrate was evaporated partly
in vacuum over concd H₂SO₄ until the
separation of a small amount of greenish plates
was obtained, they proved to be a mixture of
the copper salt, CN₇HO.Cu, with other com-
ounds, which Hofmann called "Amidotetrazot-
kupfer," but did not give a formula for it.
The filtrate remaining was evaporated to
dryness in vacuum over phosphorous pentoxide
which gave thin plates, appearing sky-blue by
reflected light and pale-green by transmitted
light. The analysis gave Cu 32.84% and 33.85%
and N 51.78%, which closely coincides with the
composition; Cu 33.37%, N 51.41% of the
formula CN₇HO.Cu

The dry copper salt is very sensitive to
friction or impact, but it is insensitive when
wet. It explodes violently on heating.

When a solution of the copper salt in very
dilute nitric or hydrochloric acid was evaporated
to dryness on a water bath, a greenish powder
remained. This residue is so sensitive that it
exploded on being touched with a glass rod
even when wet. It is presumed that this is
the copper salt of Tetrazyl-azoimide, since it
was converted to the corresponding silver salt,
N₃CN₄.Ag

Refs: 1) Beil, 3 (60) 2) K. A. Hofmann et al,
Ber 43, 1093 (1910)

Hydroxyazobenzene and Derivatives
Hydroxyazobenzene, HO.C₆H₄.N:N.C₆H₅; mw
198.22. Three isomers are known: ortho –
orn ndls, mp 82.5–83° (Ref 1); meta – yel
crysts, mp 114–15° (Ref 2); and para – orn
crysts, mp 155–56° (Ref 3); all three sl sol in
w; sol in alc & eth
Refs: 1) Beil 16, 90, (233) & [32] 2) Beil
16, 94 & [85] 3) Beil 16, 96, (233), [38]
& [86]

Mononitrohydroxyazobenzene, C₁₂H₉N₃O₃; mw
243.22, N 17.28%. Several isomers are found in
Beil:
5-Nitro-2-hydroxy-azobenzene,
HO.C₆H₄(NO₂)N:N.C₆H₅, orn-red pltls (from
dil acetic acid), mp 150–51° (Ref 1)
3-Nitro-4-hydroxy-azobenzene, l-yel pltls (from
benz), mp 128° (Ref 2)
2'-Nitro-4-hydroxy-azobenzene,
HO.C₆H₄.N:N.C₆H₄.NO₂, dk-red ndls (from
dil MeOH), mp 162–63° (Ref 3)
3'-Nitro-4-hydroxy-azobenzene, orn-yel crystals
(from acet ac), mp 147–64° (Ref 3)
4′-Nitro-4-hydroxy-azobenzene, crystals (from alc or xylol), mp 219–219.5° (Ref 5). Other props & methods of prepn are given in Refs

Dinitrohydroxyazobenzene, C H N O : mw 288.22, N 19.44%. Four isomers are known:
2,3′-Dinitro-4-hydroxy-azobenzene,
HOC6H3(NO2)2N=N:C6H4NO2; red crystals (from ace acid), mp 202° (Ref 4)
3,2′-Dinitro-4-hydroxy-azobenzene, brown-red crystals (from 1-ethoxy-2-ethanol), mp 190° (Ref 3)
5,4′-Dinitro-4-hydroxy-azobenzene, orangered crystals (from aq ace), mp 205–07° (Ref 1)
2′,4′-Dinitro-4-hydroxy-azobenzene,
HOC6H4N=N:C6H4(NO2)2; crystals (from MeOH), mp 185–08° (Ref 2)
Other props & methods of prepn are found in Beil
Refs: 1) Beil 16, (84)  2) Beil 16, (88)  3) Beil 16, (100)  4) Beil 16, (101)

Trinitrohydroxyazobenzene, C12H5N3O7; mw 333.22, N 21.02%. Two isomers are known:
4,6,4′-Trinitro-3-hydroxy-azobenzene,
HOC6H4(NO2)3N=N:C6H4NO2; red ndls (from ace acid + w), mp 179°; was prepd by heating benzoxquinone-1,4-oxime-4,6-trinitro-3-hydroxyphenylhydrazone with nitric acid (d 1.39) & ace acid (Refs 1 & 3). No expl props are reported
2′,4′,6′-Trinitro-4-hydroxy-azobenzene,
HOC6H4N=N:C6H4(NO2)3; dk-red, blue colored ndls (from dil ace acid), mp 194–95°; was prepd from 2,4,6-trinitrophenylhydrazone & benzoxquinone (1,4) in alc HCl (Refs 2 & 3). No expl props reported
Refs: 1) Beil 16, (37)  2) Beil 16, (40)  3) W. Borsche, Ber 54, 678 & 1287 (1921)
NOTE: No higher nitrated derivs of hydroxyazobenzene were found in the literature

Hydroxybenzaldehyde and Derivatives
Hydroxybenzaldehyde, HO:C6H4CHO; mw 122.12. Three isomers are known: ortho – or Salicylaldehyde, colorless liq or dark-red oil with bitter almond odor; combustible; fr p –7°, bp 196.5°, sp gr 1.153 at 25°/4; refr 1.5736 at 20°; v sl sol in w, v sol in alc & eth; can be prepd by the interaction of phenol & chlor in presence of aqueous alkali. It is used in anal chem & perfumery (Refs 1, 4 & 5); meta – or-pink crystals; mp 106–08° (Ref 1), bp ca 240°, sol in alc, benz, hot w & aromatic hydrocarbons. It is used as intermediate for dyes, plastics & pharmaceuticals and in anal chem (Refs 2, 3 & 4); para – colorless crystals, mp 116–17°; bp – subl; sol in alc, eth, ace, benz & w. It is used in pharmaceuticals (Refs 3, 4 & 5).

Ortho-Salicylaldehyde Azide or Salicylic Acid Azide, HO:C6H4CON; mw 163.13, N 25.76%; plates (from eth), mp 27°, extremely volatile; was obtd by action of ice cold furning HNO3 & NaN3 on salicylic acid hydrazide. No expl props recorded (Refs 1 & 2)

4,6-Dinitro-3-hydroxybenzaldehyde gives dark crimson needles of 2,4,6-Trinitro-hydroxybenzaldehyde-[4-nitropheny1hydrazone], O3NC6H4NN=CC6H4(NO2)3OH; which explode at 228–30°. Its p-Bromophenylhydrazone, deep olive green crystals with a metallic luster, also explodes violently at 218–20°. The p-nitrophenylhydrazone derived from 2,6-dinitrohydroxybenzaldehyde, deep terra cotta crystals, explodes at 240–42°, but the corresponding p-bromophenyl-
hydrzone is not explosive. Many other phenyl-hydrazone derivatives are known but although they are unstable to heat they do not appear to be explosive (Ref 2).


Many nitro derivs of hydroxybenzaldehyde are known (See Beil 8, 56-7, 62-3, 83 & [63]) but none of these is explosive, even the trinitro derivative, (NO₂)₃C₆H₃OHCHO, although it melts (161°) with decomposition. However reacting the latter with hydrazine sulfate produces the explosive:

**Azin of 2,4,6-trinitro-3-hydroxy-benzaldehyde**, also called 2,4,6,2',4',6'-Hexanitro-3,3'-dihydroxybenzadazin (also Azin des 2,4,6-Trinitrobenzaldoxyds in Ger). [HOC₆H₃(NO₂)₂CH:N]-; mw 510.28, N 21.96%, OB -53.0%; in the form of bright yellow needles that are sol in alc or 50% HAc, but insol in benz. Prepd by treating 2,4,6-Trinitro-3-hydroxybenzaldehyde in hot aq NaOH with hydrazine sulfate


**Hydroxybenzazide or Hydroxybenzoylazide.**

See under Hydroxybenzoic Acid and Derivatives in this Vol

**Hydroxybenzene.** See Phenol or Carbolic Acid

**Hydroxybenzenearsonic Acid and Derivatives**

**Hydroxybenzenearsonic Acid** (called Oxyphenylarson säure in Ger), HO.C₆H₄AsO(OH)₂; mw 218.03. Three isomers are known:

2-Hydroxybenzenearsonic Acid, ndls (from w), mp 190–91°; readily sol in water, MeOH & alc; insol in eth (Ref 1)

3-Hydroxybenzenearsonic Acid, crystals (from w), mp 159–73°; sol in w, MeOH, alc & gl acet acid; insol in chlf & benz (Ref 2)

4-Hydroxybenzenearsonic Acid, crystals or ndls (from hot acet), mp 177–78°; sol in w, alc & dil mineral acids; si sol in eth & eth acet (Ref 3)

Other props & methods of prepn are found in Beil


**Hydroxybenzenearsonic Acid and Derivatives**

**Hydroxybenzenearsonic Acid** (called Oxyphenylarson säure in Ger), HO.C₆H₄AsO(OH)₂; mw 218.03. Three isomers are known:

2-Hydroxybenzenearsonic Acid, ndls (from w), mp 190–91°; readily sol in water, MeOH & alc; insol in eth (Ref 1)

3-Hydroxybenzenearsonic Acid, crystals (from w), mp 159–73°; sol in w, MeOH, alc & gl acet acid; insol in chlf & benz (Ref 2)

4-Hydroxybenzenearsonic Acid, crystals or ndls (from hot acet), mp 177–78°; sol in w, alc & dil mineral acids; si sol in eth & eth acet (Ref 3)

Other props & methods of prepn are found in Beil


**Hydroxybenzenearsonic Acid and Derivatives**

**Hydroxybenzenearsonic Acid** (called Oxyphenylarson säure in Ger), HO.C₆H₄AsO(OH)₂; mw 218.03. Three isomers are known:

2-Hydroxybenzenearsonic Acid, ndls (from w), mp 190–91°; readily sol in water, MeOH & alc; insol in eth (Ref 1)

3-Hydroxybenzenearsonic Acid, crystals (from w), mp 159–73°; sol in w, MeOH, alc & gl acet acid; insol in chlf & benz (Ref 2)

4-Hydroxybenzenearsonic Acid, crystals or ndls (from hot acet), mp 177–78°; sol in w, alc & dil mineral acids; si sol in eth & eth acet (Ref 3)

Other props & methods of prepn are found in Beil


**Hydroxybenzenearsonic Acid and Derivatives**

**Hydroxybenzenearsonic Acid** (called Oxyphenylarson säure in Ger), HO.C₆H₄AsO(OH)₂; mw 218.03. Three isomers are known:

2-Hydroxybenzenearsonic Acid, ndls (from w), mp 190–91°; readily sol in water, MeOH & alc; insol in eth (Ref 1)

3-Hydroxybenzenearsonic Acid, crystals (from w), mp 159–73°; sol in w, MeOH, alc & gl acet acid; insol in chlf & benz (Ref 2)

4-Hydroxybenzenearsonic Acid, crystals or ndls (from hot acet), mp 177–78°; sol in w, alc & dil mineral acids; si sol in eth & eth acet (Ref 3)

Other props & methods of prepn are found in Beil


**Mononitrohydroxybenzenearsonic Acid,** C₆H₄NO₃As; mw 263.03, N 6.20%. Many mono nitro derivs are found in Beil

Ref: Beil 16, (454, 455, 456), [465, 466, 468, 469] & (1070, 1073)

**Dinitrohydroxybenzenearsonic Acid,** (O₂N)₂C₆H₄AsO(OH)₂; mw 308.03, N 9.16%. Two derivs are found in the literature:

3,5-Dinitro-2-hydroxybenzenearsonic Acid, yel ndls (from w), mp 244–46°; prepd by diazo-izing picramic acid with Na arsonite in alc soln; forms crystals & amorph salts (Ref 1)

3,5-Dinitro-4-hydroxybenzenearsonic Acid, yel crystals (from w), mp >275° w/o melting, d 2.0565 at 20°; dec on heating in a flame; an expl which liberates a poisonous gas (AsH₃). It can be prepd by nitrating 4-hydroxybenzenear-sonic Acid Na salt with 4 moles of nitric acid (d 1.52) in concd sulfuric acid at 15–20° (Ref 2)

A lab expln took place when a thick wet cake of the compd was heated in a flask (Ref 3). Some work done in Phillip’s Lab indicates that the acid as well as its salts might prove to be as expl as PA to which it is closely related.

NOTE: No higher nitrated derivs of Hydroxybenzeneazophonaphthol were found

Hydroxybenzeneazophonaphthol and Derivatives

Hydroxybenzeneazophonaphthol (called Phenolazo-naphthol in Ger), C₁₆H₁₂N₂O₂; mw 264.27, N 10.60%. Four isomers are known:

1-(2-hydroxybenzeneazo)-2-naphthol,

[Diagram]

OH

N:N

HO; dk-violet crysts (from alc or eth acet), mp 193° (Ref 1); forms numerous metal salts

1-(3-hydroxybenzeneazo)-2-naphthol,

[Diagram]

OH

N:N

HO; red-brn powd, mp 231–33°; forms a Cu salt (Ref 2)

1-(4-hydroxybenzeneazo)-4 azo 1-2-naphthol,

[Diagram]

OH

N:N

N:C₁₀H₆OH; red crysts, ndls (from MeOH), ptlls (from alc) or dk-violet ndls (from alc + benz), mp 194°; forms Cu & Ni cryst salts (Ref 3)

1-(4-hydroxybenzeneazo)-4 azo 4-1-naphthol,

[Diagram]

HO

N:N

N:C₁₀H₆OH; ptlls, mp - not reported; readily sol in alcalies giving a dk red color (Ref 4)

Other props & methods of prepn are found in Beil

Ref: 1) Beil 16, 169, (262) & (141) 2) Beil 16, (144) 3) Beil 16, 170, (264) & [74] 4) Beil 16, 158

Mononitrohydroxybenzeneazophonaphthol,

C₁₆H₁₀N₄O₄; mw 309.27, N 13.59%. Several isomers are known:

5-Nitro-1-hydroxybenzene-(2 azo 1)-2-naphthol,

[Diagram]

OH

O₂N

N:N:C₁₀H₆OH; purple-red crysts, mp 228°; sol in alc, glac acet ac & eth acet; sl sol in benz & chlf; forms a K salt, KC₁₆H₁₀N₅O₄, green + bronze luster crysts (from alc), which explodes on heating (Refs 1 & 5)

4-Nitro-1-hydroxybenzene-(3 azo 1)-2-naphthol,

[Diagram]

OH

O₂N

N:N:C₁₀H₆OH; red crysts, mp >260°; sol in H₂SO₄; v sl sol in org solvs (Ref 2)

2-Nitro-1-hydroxybenzene-(4 azo 1)-2-naphthol,

[Diagram]

OH

N:N:C₁₀H₆OH; no description reported (Ref 4)

Other props & methods of prepn are found in Beil


Dinitrohydroxybenzeneazophonaphthol, C₁₆H₁₀N₄O₆; mw 354.27, N 15.81%. Three isomers are found in Beil:

4,6-Dinitro-1-hydroxybenzene-(3 azo 1)-2-naphthol,

[Diagram]

OH

O₂N

N:N:C₁₀H₆OH; brn-red ndls (from benz), mp >260°; insol in alc; sol in concd H₂SO₄ giving a violet color (Ref 1)

2,6-Dinitro-1-hydroxybenzene-(4 azo 1)-2-naphthol,

[Diagram]

OH

O₂N

N:N:C₁₀H₆OH; red ndls (from Nitrobenz), mp 269–70°; sol in alk; sl sol in alc; v sl sol in glac acet acid (Ref 2)

5-Nitro-1-hydroxybenzene-(2 azo 1)-4-nitro-2-naphthol,

[Diagram]

OH

O₂N

N:N.
2,4,6-Trinitro-1-hydroxybenzene-(3 azo 1)-2-naphthol,

\[
\begin{align*}
\text{O}_2\text{N} & \quad \text{HO} \\
\text{NO}_2 & \quad \text{N}:\text{N}:\text{C}_1\text{H}_4\text{OH}; \text{mw} \ 399.27
\end{align*}
\]

N 17.54%; brownish crystals (from Nitrobenz), mp — not reported; sol in concd \(\text{H}_2\text{SO}_4\) with blue color; sol in dil \(\text{NaOH}\) with dk-red color; the alkaline soln decomp on warming gently. It was prepd by coupling the diazotized salt of 2,4,6-Trinitro-3-aminophenol with \(\beta\)-naphthol in glacial ac \& concd sulfuric acid (Refs 1 \& 2). No expls props are reported.

Refs: 1) Beil 16, [73] 2) E. Misslin, Helv 3, 637 (1920)

NOTE: No higher nitrated derivs of Hydroxybenzenazonaphthol were found in the literature.

Hydroxybenzoic Acid and Derivatives

Hydroxybenzoic Acid, \(\text{HO.C}_6\text{H}_4\text{COOH}\); mw 138.12. Three isomers are known: ortho or Salicylic Acid, colorless, combustible, monocry.

st formstd from phenol with \(\text{CO}_2\) and acidifying the Na salt thus formed. Used as org intermediate and for manuf of aspirin (See Vol 1, p A87) \& other medicinals. Its dust forms an expi mixt with air (Refs 1, 4 \& 5); meta, colorless crystals, sp gr 1.473, mp 200–01°C; sol in w, eth \& hot alc. Can be prepd by passing light thru a hot suspension of 3-hydroxybenzaldehyde in \(\text{NaOH}\). Used as intermediate for plasticizers, in resins and as petroleum additives (Refs 2 \& 4); para, colorless crystals, sp gr 1.468 at 4°C; mp ca 215°C (Lange); 210°C (Ref 4); sol in w, alc \& eth. Prepd by interaction of p-aminobenzoic acid with nitrous acid. Used as intermediate and for prepn of synthetic drugs (Refs 3 \& 4)


Hydroxybenzoic Acid Azide, Hydroxybenzoyl Azide or Hydroxybenzoyl Azide, \(\text{HO.C}_6\text{H}_4\text{CO}_3\text{N}_3\); mw 163.13, N 25.76%. Three isomers are known: 2-Hydroxybenzoyl Azide or Salicylic Acid Azide, plts (from eth), mp 27°C, extremely volatile; was prepd from salicylic acid hydrizide by action of an ice-cold soln of aq \(\text{H}_2\text{O}_3\) \& \(\text{NaNO}_2\) (Refs 1 \& 4)

3-Hydroxybenzoyl Azide, crystals (from alc \& w), mp 95°C; sol in alc, eth \& cly; sl sol in petr eth; prepd by nitratng 3-hydroxybenzoyl hydrizide with \(\text{H}_2\text{SO}_4\) \& \(\text{NaNO}_2\) (Refs 2 \& 4)

4-Hydroxybenzoyl Azide, nlds (from alc \& w), mp 132°C, was prepd by action of \(\text{NaOH}, \text{NaNO}_2\) \& acetic acid on 4-hydroxybenzoyl hydrizide (Refs 3 \& 4)

No expl props were reported for the above azides


5-Nitro-3-hydroxybenzoyl Azide (called 5-Nitro-3-oxy-benzazid in Ger), \(\text{HO.C}_6\text{H}_3(\text{NO}_2)\text{CO}_3\text{N}_3\); mw 208.15, N 26.92%, OB to \(\text{CO}_2\) 92.2%; red-yel flocculent ppt, mp — deigr on heating; insol in w; sol in alc or eth. Prepd from 5-nitro-3-aminobenzhydrazide in aced acet + aq \(\text{NaNO}_2\) (Refs 1 \& 2)


Hydroxybenzoic Acid, 3-Chloro-5-diazo or 5-Diazo-3-Chloro-Hydroxybenzoic Acid (called 3-chloro-5-diazo-salicylsäure in Ger).

\[
\begin{align*}
\text{HOOC} & \quad \text{Cl} \\
\text{N} & \quad \text{N}
\end{align*}
\]

mw 198.57, N 14.11%; yel scale-like crystals (from glac HAc); sl sol in water or alc.
Prepd from 3-chloro-5-amino-salicylic acid and Na-nitrite in dil HCl. Explodes at 193°C.


**Hydroxybenzoic Acid, 4-Diazo or 4-Diazo-3-chloro-hydroxybenzoic Acid** [called Anhydro-(3-oxo-benzoësäure-(4) in Ger),

```
COOH
\( \text{N}_2 \text{O} \)
```

mw 164.12, N 17.07%; yel powd, mp — very unstable; prepd by diazotizing 4-amino-3-hydroxybenzoic acid (Refs 1 & 2)

Refs: 1) Beil 16, (369) 2) R. Mellet, Chem-Ztg 34, 1073 (1910)

**Hydroxybenzoic Acid, 3-Diazo-5-nitro or 3-Diazo-5-Nitro-Hydroxybenzoic Acid** (called 5-Nitro-3-diazo-salicylsäure in Ger).

```
COOH
\( \text{O}_2\text{N} \)
```

mw 209.13, N 20.10%, OB to CO₂ – 80.3%; yel ndls (from alc); turns brown at 100°C, decomp at 145-150°C, explodes on rapid heating. Prepd by reacting 3-amino-salicylic acid with Na-nitrite in HCl. Infra-red spectra confirm diazonium structure shown above (Ref 2). Its lead salts also have this structure. They are light yellow, sparingly sol in org solvents, and are sensitive to impact, friction & flame


The mono & dinitro derivatives of hydroxybenzoic acid are stable, non-explosive compounds (Refs 1, 3 & 4)


Some of the salts of the dinitro derivative are explosive:

**2-Bromo-4,6-Dinitro-Hydroxybenzoic Acid, Calcium Salt.** Ca[OOC₆H₄(OH)(Br)(NO₃)₂]₂

It is prepd by nitrating 2,4,6-Tribromo-3-hydroxybenzoic acid with nitric acid and neutralizing with Ca-hydroxide (Ref 1)

**3,5-Dinitro-4-Hydroxybenzoic Acid, Copper Salt.** Cu[OOC₆H₄(H)(NO₃)₂]₂₃H₂O; green crystals, starts to decompose at 110°C when heated slowly (Ref 6), loses water at 180°C & explodes violently at 320°C (Ref 2). It also explodes on rapid heating (Ref 6). Prepd by action of CuAc on Dinitrohydroxybenzoic acid

**3,5-Dinitro-4-Hydroxybenzoic Acid, Lead Salt.** Pb[OOC₆H₄(H)(NO₃)₂]₂₃H₂O; mw 715.50, N 7.83%; yellow ndls (from water or dil HAc), sl sol in water. Prepd by reaction of Pb-nitrate and Dinitrohydroxybenzoic Acid (Refs 2 & 5). It explodes violently on rapid heating (Ref 6) and is claimed to be as shock-sensitive as Mercuric Fulminate (Ref 3). Its use as a primary explosive has been claimed in Ref 4


**2,4,6-Trinitro-3-Hydroxybenzoic Acid, (called Trinitro-ox-benzoësäure in Ger).**

HOC₆H₄(NO₃)₃COOH; mw 273.13, N 15.39%, OB -38.2%; shiny crystals (from conc nitric acid), platelets and prisms, containing 1 mole H₂O (from w), mp ca 105°C (loses water) of hydrated compd, 186°C of anhydrous, darkens at 200°C & puffs off on rapid heating; sol in water, alc or eth, sl sol in benz. Prepd by heating diazoaminobenzoic acid with concd nitric acid, or reacting 3-amino-benzoic acid with fuming nitric acid (Ref 1). It is quantitatively precipitated by a phenylacridine soln (Ref 3)

The Barium, Copper and Lead salts of Trinitrohydroxybenzoic acid are explosive:

**Barium Salt.** Ba[OOC₆H₄(H)(NO₃)₂](OH)₂·2H₂O; yellow needles; very explosive (Ref 1)

**Copper Salt.** Cu[OOC₆H₄(H)(NO₃)₂](OH)₂·2H₂O; green needles; explodes at 299°C or on rapid heating (Ref 1)

**Lead Salt.** Pb[OOC₆H₄(H)(NO₃)₂]²·H₂O. Schmitt determined the ignitability of pellets of this salt (Ref 2). On exposure to flame, pellets pressed at 500 kg/cm² detonated, while pellets
pressed at 1000, 2000 or 3000 kg/cm² merely burned
See Dead-Pressed Explosives in Vol 3, p D20-L. 
Ficheroule & Kovache claim that this salt is not useful as an explosive primer (Ref 4) 

**Hydroxybenzotriazole.** See Benzotriazolol in Vol 2, p B87-R. Additional explosive derivatives of Benzotriazolol are described below:

1-Bromobenzo-3,4-(3'azimidole), C₇H₅BrN₂O₂, mp 201.5-202.5°, sol in alkalies. Explodes on heating (Ref 1)

Vis (Ref 2) lists some other explosive derivatives of 1-hydroxy-1,2,3-benzotriazol, for example:

4,6-Dichlor-1-hydroxy-1,2,3-benzotriazol, mp 193°; powerful explosive 

Hydrazine salt of the above compound, mp 186-93°; explodes violently on heating

4,6-Dibrom-1-hydroxy-1,2,3-benzotriazol, mp 222°; powerful explosive 

Hydrazine salt of the above compound, mp about 222°; very explosive

Hydrazine salt of 4-nitro-6-chlor-1-hydroxy-1,2,3-benzotriazol, orange-yellow amorphous solid; very powerful explosive 

Goldstein & Voegeli (Ref 3) describe:

3-Hydroxybenzotriazolol-5-carboxylic acid, decamp 225°; defrages at 245-7° 
Ref: 1) A. Mangini, Gazz Chim Ital, 66, 675-84 (1936) & CA 31, 4961 (1937) 2) B. Vis, Rec Trav Chim, 58, 847-55 (1939) & CA 33, 8612 (1939) 3) H. Goldstein & R. Voegeli, Helv Chim Acta, 46, 475-81 (1943) & CA 37, 5709 (1943)

**Hydroxy-6-nitro-1,2,3-benzotriazolol-5-acetic Acid** 
See under Benzotriazolol in Vol 2, p B88-R

**Hydroxybenzylaniline, Tetranitro Derivative** 
xxxx(NO₂)₄(OH)C₆H₂CH₂NHC₆H₄; mw 379.27, N 18.47%; yellow ndls (from benz), mp 66° (decomp); insol in water; sol in alc, benz or ligroin. Prepd by mixed acid nitration of 2-hydroxybenzylaniline

Ref: Beil, 13, 580

**Hydroxybenzyltoluidine, Tetranitro Derivative** 
(called Tetranitro [(2-oxy-benzyl)-p-toluidin] in Ger). 
xxxx(NO₂)₄(OH)C₆H₂CH₂CH₂NH₂C₆H₄(CH₃);* mw 393.30, N 17.81%; yellow ndls (from benz), mp 168°; sol in alc or benz; insol in water. Prepd by mixed acid nitration of 2-hydroxy-benzyl-p-toluidine. Explodes on rapid heating

*Note: Position of NO₂ groups has not been determined. Consequently, the number of H groups on each benzene ring is also uncertain 
Ref: 1) Beil, 13, 581 2) Not found in CA, 1957-71

**α-Hydroxybibenzyl or α,β-Diphenyl-β-hydroxyethane.** Its pentanitro deriv is:

2,4,2',4',5'-Pentanitro-α-hydroxybibenzyl or α-2,4,5-Trinitrophenyl-β-2',4'-dinitrophenyl-β-hydroxyethane. 

\[
\begin{align*}
\text{CH}_2 & \quad \text{CH}_2 \\
O & \quad \text{NO}_2 \\
\text{NO}_2 & \quad \text{NO}_2
\end{align*}
\]

mw 423.25, N 16.55%; pale cream needles (from toluene), mp 187.3°, v sl sol in alc or eth; appreciably sol in benz, toluene, acetone, ethylene dichloride & acetylene tetrachloride. It was prepd by adding tetranitro-diphenylamine to fuming sulfuric acid at 85° for 4 days, after cooling fuming nitric acid (95%) was added and the temp was raised slowly and maintained at 85° for two days. The separated solid was collected, washed with alc & recrystallized from toluene

The compd exploded in 5 secs when heated to 360°, and exploded at 18 inches when impacted with a 2 kg wt. It could not be detonated with 0.4g of MF, but when initiated with 0.25g of Tetryl & 0.25g MF in the sand test, it crushed 21g of sand (Ref)

Ref: W.H. Rinkenbach & H.A. Aaronson, JACS 52, 5042-44 (1930) & CA 25, 508 (1931)
Hydroxybutyl and Derivatives. See Butanol and Derivatives in Vol 2, p B372-R

Hydroxychlorophenyl and Derivatives. See Chlorophenyl and Derivatives in Vol 3, p C362-L

3-Hydroxycumarone-2-azide. See 2-Azide-3-coumaranone in Vol 3, p C548-L

1-Hydroxycyclohexyl-1-hydroperoxide and 1-Hydroxy-1'-hydroperoxidecyclohexylperoxide. See under Cyclohexanone Peroxide in Vol 3, p C598-L

2-Hydroxy-1,1,3,3-Cyclopentanetetramethanol-pentanitrate.

\[
\begin{align*}
\text{O}_2\text{NOCH} & \quad \text{CH}_2\text{OON} \\
\text{H}_2\text{C} & \quad \text{HCON} \\
\text{H}_2\text{C} & \quad \text{CH}_2\text{OON} \\
\text{H}_2\text{C} & \quad \text{CH}_2\text{OON} \\
\end{align*}
\]

mw 431.23, N 26.24%, OB to CO\textsubscript{2} –35.2; mp 90.5. Prepared by mixed acid nitration of 2-hydroxy-1,1,3,3-cyclopentanetetramethanol. It is a powerful explosive only slightly weaker than RDX: ballistic mortar 151% TNT vs 161% TNT for RDX. It is slightly less impact & friction sensitive than RDX and has an unsatisfactory vacuum stability (12-15ml at 100\textsuperscript{o} in 12 hrs) (Ref 1). Its lowest explosion temp is 200\textsuperscript{o} (Ref 2)


Hydroxydialkylamines, Nitrated Derivatives. Compounds of the general formula

\[
\text{O}_2\text{N}N\text{R}_1 \quad \text{R}_2
\]

where \text{R}_1 is a nitroxyalkyl group and \text{R}_2 is either a nitroxyalkyl or an alkyl group (alicyclic, cyclic or substituted) were proposed as explosive, essentially non-volatile, plasticizers for NC used in prep of double-base propellants

Typical examples of this type of nitramine are DINA, EtNENA, MeNENA and Me\textsubscript{3}NENA. They are described separately


7-Hydroxy-2,5-diazaheptane. See N-Ethanol-N'-methyl-ethylenediamine in Vol 6, p E183-L

Hydroxydichlorobenzene and Derivatives. See Dichlorophenol and Derivatives in Vol 5, p D1213-R

Hydroxydiethylperoxide. See Ethylethanolperoxide in Vol 6, p E294-L

Hydroxydimethylperoxide. (Called methyl-oxymethyl-peroxide in Ger). CH\textsubscript{3}OOCH\textsubscript{2}OH, mw 78.07, OB –82%; colorless oil of penetrating odor, will not freeze in ice-salt mixture, bp 45\textsuperscript{o} at 17 mm; d 1.11 at 15\textsuperscript{o}, refract index 1.398 at 15\textsuperscript{o}; sol in water, alc, eth, benz & pet eth. Prepd by reacting methylhydroperoxide with formaldehyde in ether. Explodes on heating above 50\textsuperscript{o} & decomposes on long storage. Claimed to become impact sensitive on heating (Ref 3)

Refs: 1) Beil 1, [641] 2) A. Riche & F. Hitz, Ber 62B, 2458 (1929) & CA 24, 1079 (1930) 3) See table under Hydroperoxides in this Vol

2-Hydroxy-3,5-dinitrophenyl Azide or 2-Azido-4,6-dinitrophenol.

\[
\text{O}_2\text{N}_2\text{O}_2\text{N}_2\text{OH}
\]

mw 225.12, N 31.11%; prep'd from diazodinitrophenol & Na Azide. Its alkali or alkaline earth metal salts are used with various admixtures, such as basic Pb Trinitroresorcinate, bari-um nitrate, Sb sulfide, Pb sulfocyanate & ground glass, for use as priming comps to initiate expls (Refs)

Refs: 1) J.D. McNutt, USP 1906394 (1933) & CA 27, 3612 (1933); USP 1930653 (1934) & CA 28, 328 (1934); & USP 2005197 (1935) & CA 29, 5274 (1935) 2) A. H. Blatt & F. C. Whitmore, OSRD 1085 (1942), p 54 3) Blatt, OSRD 2014 (1944) (Under Azides)
Hydroxydiphenylamine. See Anilinophenol in Vol 1, p A433-R and additional entries below:

Hydroxy-tetranitro-diphenylamine (called 3,5,2',4'.
Tetranitro-oxy-diphenylamine in Ger).
(O₂N₄)₂C₆H₄NHCH₂H₂(NO₂)₂(OH), mw 365.22, N 19.18%, yellow powder, mp 236°, almost insol in all usual solvents. Prepd by condensation reaction of 4-chloro-1,3-dinitro-4 aminophenol in alc containing NaAc. No explosive props mentioned

Ref: 1) Beil 13, 528 2) Not found in CA thru 1971

4-Hydroxy-3,5,2',4',6'-Pentanitrodiphenylamine
(Also called 2,6-Dinitro-4-Picrylaminophenol in Ger).
(O₂N₄)₂C₆H₄NHCH₂H₂(NO₂)₂OH; mw 410.22, N 20.49%, OB to CO₂ -62.4%; ochre-yel crystals (from HAc), mp 248°; insol in water, sl sol in alc; sol in alkalis with formation of a deep brown color. Prepd by heating picryl chloride, isopicarminic acid & NaAc inaq alc. No explosive props are mentioned for it nor for the tetranitro derivatives (Ref 2). The Na salt, of the pentanitro derivative NaC₁₂H₄O₁₁N₆, red platelets, explodes above 300°. The 3'-Hydroxy-4-methyl-pentanitro diphenylamine is not explosive (Ref 3)

Ref: 1) Beil 13, (191) 2) Beil 13, 396, 528 & 531 3) Beil 13, 412

N-Hydroxyethenylamidoxime. See under Ethenylamidoxime in Vol 6, p E185-L

Hydroxyethyl. See Ethanol and Derivatives in Vol 6, p E154-L

Hydroxyethylamine and Derivatives. See under Amino-ethanol and Derivatives in Vol 1, pp A200-202

β-Hydroxyethyaminobenzene or β-Hydroxy-
ethylineline. See under 2-Anilinoethanol and
Derivatives in Vol 1, pp A424-31. This includes Pentryl (Trinitroanilinoethanol Nitrate) & various halogen derivs of anilinoethanol

N-(2-Hydroxyethyl)-butylamine. Its nitrated
deriv N-(2-Nitroxyethyl)-butylnitramine (BuNENA),
O₂NOCH₂CH₂.N(NO₂)₂.C₆H₄; mw 207.19, N 20.28%; pale yel oil, frp -9.9°, RI 1.4750 at 20°, was prepd & described by Blomquist & Fiedorek (Ref)


Hydroxyethylcelluloses. Title of a doctorate thesis submitted to the University of Paris, France, 9 June 1960, 89pp & 144 refs

Ref: J. Quincon, “Contribution à l'étude des hydroxyethylcelluloses,” MP 42, (1960)

N-(2-Hydroxyethyl)-cyclohexylamine. Its nitrated
deriv N-(2-Nitroxyethyl)-cyclohexylnitramine
(Cyclohexyl-NENA),

CH₂-CH₂-CH₂
CH₂-CH₂
O₂NO.CH₂.CH₂.N(NO₂)₂.CH

mw 233.22, N 18.02%; is described in Vol 3, p C600-L

N-(2-Hydroxyethyl)-ethylamine. Its nitrated
deriv N-(2-Nitroxyethyl)-ethylnitramine (EtNENA),
O₂NO.CH₂.CH₂.N(NO₂)₂.C₂H₄; mw 179.14, N 23.46%; pale yel oil, frp 4-5.5°, d 1.32 at 25°, RI 1.479 at 25°; prepd by addg ethylethanol-
amine dropwise to a stirred 98% nitric acid soln at 10° and adding the resulting mixt to a soln of acetic anhydride & acetyl chloride at 35° (Ref)


N-(2-Hydroxyethyl)-glucosamine Hexanitrate.
C₅H₅(OONO₂)₃.CO.NHCH₂CH₂ONO₂; mw 520.26, N 18.85%, OB to CO₂ -12.3%; prepd by nitric acid nitration N-2-hydroxyethylamidogluconamide, C₅H₅(OH)₂.CO.NHCH₂CH₂OH. Claimed to be useful in blasting cap charges

Ref: W.F. Filbert, USP 2443903 (1948) & CA 43, 1796 (1949)

N-(2-Hydroxyethyl)-glycolamide Dinitrate.
O₂NOC₂H₅.CO.NHCH₂CH₂ONO₂; mw 209.12, N 20.10%, OB -34.4%; prepd by nitric acid nitration of N-2-hydroxyethylglycolamide, HOCH₂.CO.NHCH₂H₂O. Claimed to be useful as a blasting cap charge

Ref: W.F. Filbert, USP 2443903 (1948) & CA 43, 1796 (1949)

Hydroxyethylglycolurethane. N-Nitro-Dinitrate.

ONO₂CH₂CH₂OCN(NO₂)₂CH₂CH₂ONO₂; mw
284.14, N 19.72%, OB -22.4%; white crystals. Prepd by nitrating N-hydroxy-ethylglycolurethane with 98% nitric acid. No properties given
Ref: G. Desseigne, FrP 1094959 (1955) & CA 53, 11159 (1959)

Hydroxyethylguanidine and Derivatives

Hydroxyethylguanidine [called β-Guanidino-ethylalkohol or (2-Hydroxy-ethyl)-guanidin in Ger] NH₂CH₂CH₂·OH
C·NH

NH₂

mw 103.13, N 40.75%; exists in the form of its salts. May be considered as the parent compd of its nitrated derivs, although not used to prep them
Ref: Beil, 4, (730) & (710)

N-(β-Hydroxyethyl)-N'-nitroguanidine or 1-(2-Hydroxyethyl)-3-nitroguanidine.
HO·CH₂CH₂·NH·C:(NH)NH·NO₂; mw 148.13, N 37.83%; wh crystals (from alc), mp 118° (dec); was obtd when methylnitrosoguanidine was added portionwise to a soln of ethanolamine in water cooled to 5-10°
Refs: 1) Beil, not found 2) A.F. McKay & J.E. Milks, JACS 72, 1617-18 (1950) & CA 44, 10661 (1950)

N-(β-Nitroxyethyl)-N'-nitroguanidine or 1-(2-Nitroxyethyl)-3-nitroguanidine.
O₂NO·CH₂CH₂·NH·C:(NH)NH·NO₂; mw 238.13, N 35.30%; wh crystals (from MeOH), mp 84.5 to 85.3°; was prepd by addg β-hydroxyethylguanidine to a nitration mixt of 99.8% nitric acid & acetic anhydride at 0° and after 30 mins pouring the mixt onto ice. This compd is useful in proplnts (Ref 3)
Refs: 1) Beil, not found 2) A.F. McKay & J.E. Milks, JACS 72, 1619 (1950) & CA 44, 10661 (1950) 3) A.F. McKay, USP 2538907 (1951) & CA 45, 4264 (1951); BritP 673709 (1952) & CA 48, 9123 (1952)

1-Hydroxyethyl Hydroperoxide.
CH₃CH(OH)OOH, or its higher homologs are not explosive. See Table under Hydroperoxides in this Vol

N-(β-Hydroxyethyl)-hydroxyacetamidine.
HOCH₂CONHCH₂CH₂OH; mw 119.12, crystals, mp 72.5-73.5°. May be prepd by treating methyl glycolate with monoethanolamine

A colorless oil was obtained when 2g of the compound were dissolved in 10ml of white nitric acid and then heated at 50-60° for one-half hour and poured on ice. This oil was not further investigated, but it might have been a liquid explosive, suitable as a plasticizer for NC etc
Refs: 1) Beil, not found 2) R. Adams & C.S. Marvel, ODRD No 86 (1941), pp 12 & 36-37 3) CA not found thru 1971

N-(β-Hydroxyethyl)-methylamine. Its nitraterd deriv N-(β-Nitroxyethyl)-methyltrimine (MeNe-NA). O₂NO·CH₂CH₂·N(NO₂).CH₂; mw 165.11,
N(β-Hydroxyethyl)-N'-methyl-ethylenediamine.

Its nitrated deriv N-(β-Nitroxyethyl)-N'-methyl-ethylenedinitrime.

\[ \text{O}_2\text{N}_2\text{O}_2\text{CH}_2\text{CH}_2\text{N}(_2\text{N}_2\text{O}_2)_2\text{CH}_2\text{CH}_2\text{N}(_2\text{N}_2\text{O}_2)_2\text{CH}_3; \text{mw} 253.18, \text{N} 22.64%; \text{glistening crystalline (from 50% acetonitrile), mp 88.5-90°, was prep'd by cleavage with 98% nitric acid of the compd iso-N}_{1}:_{N}_2\text{-dimethyl-triethylenetetramine,}\]

\[ \text{H}_2\text{C}-\text{N}(_2\text{N}_2\text{O}_2)_2\text{CH}_2\text{-CH}_2\text{N}(_2\text{N}_2\text{O}_2)_2\text{CH}_2\text{-CH}_2\text{O}:\text{N}:
\text{N}_2\text{CH}_2\text{-CH}_2\text{N}(_2\text{N}_2\text{O}_2)_2\text{CH}_3, \text{at 10°} \]

Ref: A.T. Blomquist, OSRD 4134 (1944), pp 9 & 81

α-Hydroxyethyl methylperoxide or Methylhydroxyethylperoxide \[ (α\text{-Oxyethyl-methyl-peroxy in Ger.).}\]

\[ \text{CH}_3\text{CH(OH)}\text{OOCH}_3; \text{mw} 92.09, \text{easily flowing oil, fairly stable; bp 29-31° at 22mm; d 1.029 at 15°, RI 1.3930 at 15°. May be prep'd from methylhydroperoxide, CH}_3\text{OOH, and acetaldehyde, CH}_3\text{CHO; diff sol in water, miscible with alc and eth; easily sol in benz.}\]

Puffs off weakly on heating

Refs: 1) Beil, not found 2) A. Rieche & F. Hitz, Ber 63, 2642 (1930) & CA 25, 911 (1931)

N,N'-bis(2-Hydroxyethyl)-oxamide. See Diethylolamide and Derivatives in Vol 5, p D1243-R

Hydroxyethyl-3-oxo-1,2-propanediol trinitrate or 3(β-Hydroxyethoxy)-1,2-propanediol trinitrate.

\[ \text{ONO}_2\text{CH}_2\text{CH}_2\text{OCH}_3\text{CH(ONO}_2\text{)}\text{CH}_2\text{ONO}_2, \text{mw 271.18, N 15.50%, OB -26.5%. Prepd by mixed acid or straight nitric acid nitration of 3(β-Hydroxyethoxy)-1,2-propanediol (Ref 1) or nitrating the reaction product of epichlorohydrin & glycol (Ref 2). It is a liquid which is less volatile than NG. Desseigne has claimed its use as an explosive and as a gelatinizing agent.}\]

Desseigne also prepd in a similar manner

Hydroxypropyl-3-oxo,1,2-propanediol Trinitrate.

\[ \text{CH}_3\text{CH(ONO}_2\text{)}\text{CH}_2\text{OCH}_3\text{CH(ONO}_2\text{)}\text{CH}_2\text{ONO}_2\]

and also claimed its use as an explosive and gelatinizing agent


Hydroxyethyl picramide, or 2,4,6-Trinitrophenylaminoethanol. See under Anilino-ethanol and its Derivatives in Vol 1, p A425-L

N(β-Hydroxyethyl)-N'-phenyl-1,2-diaminoethane and Derivatives. See Anilinopropylaminoethanol and Derivatives in Vol 1, p A431-L

Hydroxyheptylperoxide. [CH3(CH2)3CH(OH)]2O2; mw 262.38

The commercial product of the Lucidol Division contains a minimum of 95% of peroxide; active oxygen 5.8% (min). It is a white fine powder; mp 60-65°.

It is insol in w and moderately sol in organic solvents or monomers. Unstable at ordinary room temp and should not be stored at above 70°F (in order to avoid loss of active oxygen) and away from all sources of heat. It should not be subjected to frictional heat or grinding. It has been used as a room or low temp catalyst for polymerization of polyester resins

Ref: Lucidol Division, Novadel-Agen Company, Buffalo, NY, Organic Peroxides, Data Sheets No 18 (1948) and No 32 (1950)

1-Hydroxy-1'-hydroperoxy-dicyclohexyl peroxide.

See under Cyclohexane Peroxide in Vol 3, p C598-R

1-Hydroxy-5-hydroxymethyltetrazole and Derivatives.

1-Hydroxy-5-hydroxymethyltetrazole (called 1-Oxy-5-hydroxymethyltetrazol in Ger).

\[ \text{N}—\text{N} \text{OH} \]

\( \text{C}_{\text{CH}_2\text{OH}}; \text{mw} 116.08, \text{N} 48.27%, \text{OB} \]

-55.1%. Prepd by treating 1-hydroxytetrazol with formaldehyde. It is a volatile material which is an explosive less powerful & brisant than PA

Refs: 1) Beil 26 (109) 2) Blatt OSRD 2014 (1944)
1-Hydroxy-5-nitrooxymethyltetrazole.

\[
\begin{align*}
\text{N—N—OH} & \quad \text{C—CH}_2\text{ONO}_2; \text{mw 161.08, N 43.48%;} \\
\text{N—N} & \quad \text{OB -14.9%. Prep by nitrating above compd. It is} \\
& \text{less brisant and powerful than PA}
\end{align*}
\]

Ref: Blatt OSRD 2014 (1944)

6-Hydroxyindazole (called 6-Oxyindazol in Ger).

\[
\text{MW 134.13, N 20.89%; leaflets, mp 215-16\degree;} \\
\text{sol in dil acid or alkali, sl sol in water or eth.} \\
\text{Prepd by diazotizing 6-aminoindazole. When nitrat-}
\]

\[
\text{ed it forms:}
\]

5,7-Dinitro-6-hydroxyindazole,

\[
\text{MW 224.13, N 25.00%; OB to CO\textsubscript{2} -78.5%, pale-yel} \\
\text{leaflets (from glacial HAc); mp 232-33\degree; sol in} \\
\text{water. Puffs off above mp}
\]

Ref: 1) Beil 23, 377 2) Not found in CA thru 1971

Hydroxymethylbenzene, Hydroxyltoluen or Methylphenol. See Cresol and Derivatives in Vol 3, p C556-L

2-Hydroxymethyl-1,3-propanediol-2-aniline. See Anilinotrimethylolmethane and Derivatives in Vol 1, p A441. Its highly expl deriv, designated as Hemtryl is on pp A441-R to A442-R

1-Hydroxy-2-propanone. See Acetol and Derivatives, in Vol 1, p A33-R

HYDROXYLAMINE AND DERIVATIVES

Hydroxylamine, Oxammonium (HA), NH\textsubscript{2}OH, 

\[
\text{mw 33.03, N 42.3%, colorless, deliq low melt} \\
\text{needles, mp 33\degree, bp 56.5\degree at 22mm, 110\degree at} \\
\text{760mm (Ref 8), flp explodes at 265\degree F, d 1.23,} \\
\text{for liq RI 1.440 at 23.5\degree, sol in cold w, alc & acet; v sl sol in eth, chl & benz; decomp in} \\
\text{hot w. } \Delta H^f = 25.5, \text{ for cryst, -21.7, for soln,}
\]

kcal/mole (Ref 7). Dissoc const for dil soln

\[
1.1 \times 10^{-4}; \Delta H \text{ subl 15.3 kcal/mole} \quad \text{(Ref 9)}
\]

It is an acute local iritiant, and systemically it can cause methemoglobinemia (Ref 10)

Preparation

Since free NH\textsubscript{2}OH is unstable it is usual to prepare it in the form of the hydroxylammonium salts, which resemble ammonium salts. A number of methods were proposed but only 5 offer practical methods

1) Reduction of HNO\textsubscript{3} or its Salts. When a soln of NaNO\textsubscript{3} is reduced by NaHSO\textsubscript{3}, sodium hydroxylamine-N,N-disulfate is formed:

\[
\text{NaNO}_3 + \text{NaHSO}_3 + H_2SO_4 \rightarrow (\text{SO}_3 \text{Na})_2 \text{OH+H}_2\text{O.}
\]

Heating the resulting soln produces hydroxyl- 
aminon hydrogen sulfate:

\[
(\text{SO}_3 \text{Na})_2 \text{OH+H}_2\text{O} \rightarrow \text{Na}_2\text{SO}_4 + \text{NH}_3\text{OH} + \text{H}_2\text{SO}_4
\]

The crude hydroxylammonium salt may be obtained from the soln by fractional crystallization. Several modifications of the method exist (Ref 3)

2) Reduction of Nitric Acid. Electrolytic reduction of HNO\textsubscript{3} in the presence of HCl produces hydroxylammonia chloride:

\[
\text{HNO}_3 + 6\text{H} + \text{HCl+6e} \rightarrow (\text{NH}_3\text{OH})\text{Cl}+2\text{H}_2\text{O}
\]

Yield ab 80% (Ref 2)

3) Hydrolysis of primary nitroparaffins. A primary nitroparaffin gives on hydrolysis with a strong acid, the corresponding fatty acid and the hydroxylammonium salt of the acid. For instance: CH\textsubscript{3}CH\textsubscript{2}NO\textsubscript{2}+H\textsubscript{2}O+HCl → (NH\textsubscript{3}OH)\text{Cl}+ CH\textsubscript{3}COOH. This reaction is used for producing hydroxylammonium salts industrially

Free NH\textsubscript{3}OH may be liberated from any of the above salts, by a base stronger than hydroxy- 
aminol itself, eg K, Na, hydroxides or carbonates in aq or alc solns would liberate the base (Ref 4)

4) In the prepn of crystalline NH\textsubscript{3}OH it is impractical to use a water solvent since it is difficult to remove it. It is better to use the anhydrous alcoholver of the corresponding solvent eg:

\[
\text{NH}_2\text{OH+HCl+BuONa in BuOH} \rightarrow \text{NH}_2\text{OH+NaCl} \\
\text{+BuOH. MeOH or EtOH and their alcoholver may also be used} \quad \text{(Ref 5)}
\]

5) Similarly:

\[
\text{NH}_2\text{OH+HCl+NH}_3\text{(liq)} \rightarrow \text{NH}_2\text{OH+NH}_4\text{Cl}
\]

The excess NH\textsubscript{3} is evaporated and the HA is extracted with abs alc (Ref 6)

Removal of HA from the solvent alc is
achieved by cooling (the HA crystallizes) and/or addn of ether. Volatile alics may be removed by vac distillation

**Chemical Props**

There is similarity in chem props between NH$_2$OH and hydrazine. HA may be regarded as hydroxylated product of ammonia, thus representing the first member of such series, which includes dihydroxyammonia and ortho nitrous acid. It may be regarded as ammonolized HOOH (Hydrogen Peroxide). Hydroxylamine, like NH$_2$OH, NH$_2$.NH$_3$, forms a large group of salts, but only strong acids form stable salts, of which chlorides and sulfates are the best known. However, all of these salts decompose at about 150$^\circ$.

Like H$_2$O$_2$ and N$_2$H$_4$, solns of the free base, NH$_2$OH, are susceptible to thermal and catalytic decomposition. It is not improbable that auto-oxidation and catalytic decomp reactions are catalyzed by traces of metallic ions, as has been noted in case of H$_2$O$_2$ and of N$_2$H$_4$. The stability of HA is intermediate between hydrazine & hydrogen peroxide. It is less stable than N$_2$H$_4$ and somewhat more stable than H$_2$O$_2$. Hydroxylamine is a highly hydrogen bonded subst in both the solid and liquid forms, but the vapor is a single molecule at low press (Ref 11). The latter studies give the bond strength of NH$_2$-OH as 61.3 kcal/mole.

NH$_2$OH very seldom exhibits acidic properties. It is claimed, however, that it forms unstable hydroxylamites, NaONH$_2$, Ca(ONH$_2$)$_2$ (Ref 6). As a base NH$_2$OH resembles ammonia and other amines. Although its basic ionization constant is considerably lower than that of NH$_3$ and N$_2$H$_4$, it forms a series of ammonia-like inorg and org salts. In general, these salts are more stable than the parent base. Therefore, hydroxylamine is usually prepd and shipped in the free form of its salts.

NH$_2$OH also forms few double salts and coordination compds analogous to those formed by ammonia. Some of these compds are very stable to heat.

Soln of HA, especially acidic solns are strong reducing agents. They will reduce FeCl$_3$, ammonical AgNO$_3$, CuSO$_4$ etc. In some instances, however, particularly in alkaline solns, HA is capable of oxidizing such subs as chromous chloride & CuOH.

Liquid anhydrous NH$_2$OH will dissolve many substances such as KI, KCN, KBr, NaN$_3$, Ba(NO$_3$)$_2$, NaCl, KCl, NaOH, Ba(OH)$_2$, NH$_3$ (Ref 6).

**Explosive & Combustion Props.** NH$_2$OH, like N$_2$H$_4$, is a strong reducing agent. The base ignites in a stream of chlorine and even reacts violently if air is blown through it

If it is heated in air, it explodes between 60 and 100$^\circ$. It will explode upon contact with an open flame. On oxidation HA liberates a considerable amount of energy and may therefore be considered as a specialty fuel.

A rough calculation of the overall heat of the reaction according to:

$$2\text{NH}_2\text{OH} (\text{gas}) + \frac{1}{2} \text{O}_2 \rightarrow 3\text{H}_2\text{O} (\text{gas}) + \text{N}_2$$

indicates that the heat of combustion of HA should liberate about 59.2 kcal/mol.

The combustion vel of a decomposition flame of a 74% HA in nitric acid soln is 0.1 cm/sec. The flame is orange and ammonia is one of the flame products (Ref 8).

**Stability.** Some authors claim that NH$_2$OH decomposes merely on standing, but L. L. deBruyn (Ber 27, 967 (1894)) claimed that one sample decomposed only very slightly after a period of several years. In the opinion of Audrieth et al (Ref 6), some of the impurities cause catalytic decomposition and removal of these impurities improves the stability of product. On standing NH$_2$OH presumably decomposes, either as 3NH$_2$OH - NH$_3$ + N$_2$ + 3H$_2$O, or 4NH$_2$OH - 2NH$_3$+N$_2$O+3H$_2$O depending on conditions (Ref 6).

**Analytical.** A number of analytical procédures for determining and determining of NH$_2$OH are available (Ref 6). The best method is to add an excess of some oxidizing agent such as ferrie ammonium sulfate to a soln of free base or one of its salts, and then determine the amt of oxidizing agent which has been reduced by standard permanganate or ceric sulfate solns.

An alternate method consists of titrating aq NH$_2$OH with a std acid using a methyl orange indicator.

Qualitatively, amounts as low as 1 part/100,000, may be detected by treating aq NH$_2$OH with ammonium CuSO$_4$ to get a red ppt of Cu$_2$O.

**Uses:** Besides its uses as a reducing agent and in org synth, HA is claimed as an ingredient in several propellant formulations, in combination with AN & AP.
**Hydroxylamine Salts & Complexes**

Most strong acids form acid salts with HA. The following are fairly stable but some of them are explosive:

**Hydroxylamine acid sulfate.** NH$_3$OH.H$_2$SO$_4$; mw 131.12, N 10.68%, white to brown crystals; v hygroscopic, mp indefinite, bp decomp; sol w, MeOH; sl sol EtOH. Toxic and may explode on heating. It is used as a reducing agent, photog developer & in synths of dyes, pharmaceuticals & rubber chars.

*Ref: CondChemDict, 8th Ed (1971), p 460*

**Hydroxylamine hydrochloride.** NH$_3$OH.HCl; mw 69.50, N 20.16%; colorless, hygr crystals (monoclinic), mp 152°, bp decomp. d 1.67; sol w, alc, glycol; insol eth. It is an irritant. It is toxic (causes prolonged methemoglobinemia [Ref 3]) and it may react violently when heated. It is used in org synths, in photography & in medicine (Ref 3).

Prepd by hydrolyzing MeNO$_2$ with HCl (Ref 1); from acetooxime, water & Me$_2$CO with product crystallized at 20° (Ref 2).


**Hydroxylamine nitrate.** NH$_3$OH.HNO$_3$; mw 96.05, N 29.17%; white crystals, mp 48°, bp decomp at about 100°. Sol. cold w, alc; decomp in 70°. Prepd by reacting NH$_3$OH.HCl with AgNO$_3$ or (NH$_3$OH)$_2$H$_2$SO$_4$ with Ba(NO$_3$)$_2$ (Ref 1). It is claimed as a propell ingred in combination with AN, AP & a curable liq binder (Ref 2).

*Refs: 1) G. Bourjoul, MP 34, 63 (1952) & CA 46, 9236 (1953) 2) CondChemDict (1971), 1971, p 460-L*

**Hydroxylamine Perchlorate (HAP).**

NH$_3$OH.HClO$_4$, mw 135.50, N 10.50%; hygr crystals, mp 87.5-90°, decomp 120°, sol eth. ΔH° = 66.2 kcal/mole (Ref 3). Prepd by reacting NH$_3$OH.HCl or (NH$_3$OH)$_2$H$_2$SO$_4$ with BaClO$_4$. Used to increase burning rate in propellants. Impact sens 15cm with 2 kg falling wt. A solid propell formulation based on HAP has been patented (Ref 2). It contains about 80% HAP (or LiClO$_4$, 2NH$_3$OH or MgClO$_4$, 4NH$_3$OH) and 20% epoxy binder.

The self deflagration of HAP & hydrazine nitroform was studied (Ref 4). It was found that HAP exhibits a low pressure deflagration limit of 146 atm analogous to the 20 atm limit of AP. Combust temps were determined.


**Hydroxylamine sulfate.** (NH$_3$OH)$_2$H$_2$SO$_4$, mw 164.14, N 17.07%; colorless monoclinic crystals, mp 177°, bp decomp. Sol w, eth, sl sol alc (Ref 2).

Prepd by hydrolysis of MeNO$_2$ with H$_2$SO$_4$. This reaction yields HA in an anhydrous medium: CH$_3$NO$_2$+H$_2$SO$_4$→HOSO$_2$CC(CH$_3$)(OH)(I)

(II)+2H$_2$O+HCOOH+NH$_3$OH or H$_2$SO$_4$+CO+NH$_3$OH (Ref 1). Hydroxylamine sulfate is highly toxic. Its uses are similar to those for the acid sulfate.

*Refs: 1) G. Bourjoul, MP 34, 63 (1952) & CA 46, 9236 (1953) 2) CondChemDict (1971), 1971, p 460-L*

**Hydroxylamine complexes** of some inorganic salts. The prepn & characterization of some NH$_3$OH complexes of various inorg salts are described. (NH$_3$OH)ClO$_4$ forms complexes with both 1 and 2 moles of NH$_3$OH (see HAP) while the nitrate forms only a single complex. These complexes are, in general, less hygroscopic than the salt itself.

Hydroxylamine Derivatives. Many $\text{NH}_2\text{OH}$ derivatives may be obtained by substituting for the $\text{NH}_2$ or $\text{OH}$ hydrogens. Replacement of the $\text{NH}_2$ hydrogens gives hydroxylamates, and $N$-hydroxy-derivatives (also called $\beta$-hydroxylamines) of various degrees of substitution. Replacement of the $\text{OH}$ hydrogens gives $\alpha$-hydroxylamines (also called $\alpha$-hydroxylamines) and hydroxylamine-acids. Inorganic and organic derivatives produced by both types of hydrogen replacement are listed below:

$\text{NH}_2$ Hydrogen Substitution

**Calcium Hydroxylamate.** $\text{Ca(NHOH)}_2$, mw 104.13, N 26.90%. HA reacts with Ca at 5° to form a very explosive salt

Ref: E. Ebner & E. Schott, JPraktChem 78, 289 (1908) & CA 3, 754 (1909)

**Sodium Hydroxylamate.** $\text{Na(NHOH)}$, mw 54.0, N 26.6%. When an ether soln of $\text{NH}_2\text{OH}$ is treated with metallic Na an extremely unstable explosive compd is formed

Ref: L. F. Audrieth et al “Compounds of High Nitrogen Content,” 1st Quart Rept, U of Illinois, Jan 1951

**Zinc Hydroxylamate.** HA reacts with Zn at 5° to give an explosive salt of indeterminate composition

Ref: E. Ebner & E. Schott, JPraktChem 78, 289 (no year) & CA 3, 754 (1908)

Many $N$-hydroxyamine derivatives are known. They appear to be non-explosive, stable compounds eg $\text{CH}_3\text{HNOH}$ (Ref 1) or $\text{HOOCCH}_2\text{NH(OH)}$ (Ref 2)

Ref: 1) Beil 4, 534, [1952] & 1715 2) Beil 4, 542

However some of $N$-hydroxylamino-alkanoic acids form nitroso compounds and some of the metal salts of such compounds are explosive as shown below:

**Hydroxylaminoacetic Acid or 2-(Hydroxylamino)-ethanoic Acid.** (Hydroxylaminooessigsäure or Hydroxylaminäthersäure in Ger), $\text{HO,NH,CH}_2,\text{COOH}$ is described in Beil 4, 542. It is the parent compd of its nitroso deriv described below

**Nitrosohydroxylaminocetic Acid, Isonitramino-acetic Acid or 2-(Nitrosohydroxylamino)-ethanoic Acid.** (Isonitraminoessigsäure in Ger), $\text{HO,N(NO)},\text{CH}_2,\text{COOH}$, mw 120.07, N 23.33%. Needles (from ether + ligroin), mp-decomp 103-4°; stable only in solns. Prep'd by passing NO thru an alc soln of ethylsoda-acetateacetate and then reacting the acid to form the lead salt.

It's silver, calcium & ammonium salts are explosive.

Ref: 1) Beil 4, 575 2) W. Traube, Ber 28, 1791-3 (1895) & JCS 68i, 503 (1895)

2-(Hydroxylamino)-1-butanoic Acid or $\alpha$-Hydroxylaminobutyric Acid. (2-Hydroxyamino-butanäsäure in Ger). $\text{HO,NH,CH}_2,\text{CH}_2,\text{COOH}$ is described in Beil 4, 543. It may be considered as the parent compd of its nitroso-deriv described below

2-(Nitrosohydroxylamino)-1-butanoic Acid or $\alpha$-Isonitraminobutyric Acid. ($\alpha$-[Nitrosohydroxylamino]-buttersäure or $\alpha$-Isonitraminobuttersäure in Ger). $\text{H}_3\text{CO,CH(NH,OH),COOCCH}_2,\text{H}_2$ is not found in Beil 4, but its nitroso deriv (see below) is described

2-(Nitrosohydroxylamino)-3-butanoic-1-acid, Ethylester or $\alpha$-(Nitrosohydroxylamino)-ethylesterr. $\text{CH}_3,\text{CO,CH(NH,OH),COOCCH}_2,\text{H}_2$ is not found in Beil 4, but its nitroso deriv (see below) is described

2-(Nitrosohydroxylamino)-3-butanoic-1-acid, Ethylester: $\alpha$-(Nitrosohydroxylamino)-ethylesterracetate or $\alpha$-Isonitraminonactocetoxy Acid, Ethylester. (Ger names: Athylester der 2-Nitrosohydroxylamino-butanon-(3)-säure, $\beta$-Oxo-$\alpha$-[Nitrosohydroxylamino]-propan-1-carbonsäureäthylester, $\alpha$-[Nitrosohydroxylamino]-acetessigsäureäthylester or $\alpha$-Isonitraminoacetessigsäureäthylester) $\text{CH}_3,\text{CO,CH(NO),OH,COOCCH}_2,\text{H}_2$ Its disodium salt, $\text{C}_6\text{H}_4\text{N}_2,\text{O}_5,\text{Na}_2,\text{H}_2,\text{O}$ is explosive and was obtained when dry nitric oxide was passed into a 10 per cent alc soln of ethylic sodaacetate as described in Ref 2

Ref: 1) Beil 4, 577 2) W. Traube, Ber 27, 1507 (1894) & JCS 66i, 400 (1894)

2-(Hydroxylamino)-1-propanoic Acid or $\alpha$-Hydroxylaminopropionic Acid. $\text{HO,NH,CH(CH)},\text{COOH}$, is not found in Beil 4, but its nitroso deriv (see below) is described
2-{Nitrosohydroxylaminol}-1-propanoic Acid or α-Isonitrinmomproponic Acid \(\{\alpha-[\text{Nitroso-}
\text{hydroxylamino}]\}\)-propanioäure or \(\alpha\)-Isonitrino-
momproponiäure} \(\text{HO.N(NO)O.CH(CH}_3\text{).COOH;}
\text{mw}\ 134.09, N\ 20.89\%.\ \text{It\ is\ stable\ only\ in\ solns.\ May\ be\ prepd\ by\ warming\ with\ soda\ the\ crude\ soln\ obtained\ by\ the\ action\ of\ nitric\ oxide\ on\ ethylic\ methylacetoacetet,\ until\ the\ alcohol\ is\ driven\ off;\ the\ acid\ is\ then\ precipitated\ as\ the \textbf{Lead Salt, PbC}_3\text{H}_4\text{N}_2\text{O}_4,\ \text{which\ is\ explosive}}\).\ \textit{Refs:}\ 1) Beil 4, 576 2) W. Traube, Ber 28, 1793 (1895) & JCS 68, 503 (1895)

\textbf{Aryl derivatives of N-hydroxylamine} form azo and nitro compounds which are unstable towards heat but not explosive

\textbf{Azo-Derivatives of N-phenylhydroxylamine} crystalline comp of the type

\[
\begin{array}{c}
\text{Y} \\
\text{NO}_2 \\
\text{N:N} \\
\text{NHOOH}
\end{array}
\]

are known where X is Cl or OH, and Y is NO₂ or NO. They decom in the range of 210-230°. The dinitro Cl deriv is prepd by reacting benzoquinone-1,4-monoxyne with 5-chloro-2,4-dinitrophényhylidrazone, and 4,6-Dinitro-3-hydraxino-phenol to get the OH compd. Similarly, the cyano deriv is prepd from the above oxime and 2-nitro-4-phenoxylidynehydrazine

\textit{Ref:}\ Beil 16, [260]

\begin{itemize}
  \item \textbf{Nitro-Derivatives of Ayl-N-hydroxylamine}
  \item 6-Nitro-2,4-Dihydroxyaminophloroglucinol (called -phloroglucin in Ger). \((\text{HO})_3\text{C}_6(\text{NO}_2)_2(\text{NNOH})),\ \text{mw}\ 223.14, \text{N}\ 18.03\%, \text{OB}\ -58.5\%;\ \text{yellowish needles, mp}\ 166°\ \text{(decomp)}\ \text{has similar props} \text{and same meth of prep as the dinitro derivative described below}
  \item 2,4,6-Trihydroxyaminophloroglucinol. \((\text{OH})_3\text{C}_6(\text{NNOH})_3;\ \text{alum colorod needles (from eth); sol in eth} \ \text{dil alk solns. Very unstable. Other props and method of prep same as above}
  \item 4,6-Dinitro-2-hydroxyaminophloroglucinol (called 4,6-Dinitro-2-hydroxyaminophloroglucin in Ger). \((\text{HO})_3\text{C}_6(\text{NO}_2)_2(\text{NNOH}));\ \text{mw}\ 247.12, \text{N}\ 17.01\%, \text{OB}\ -42.3\%;\ \text{yel needles (from eth), mp}\ 146-148°\ \text{(decomp); readily sol in hot w, org solns & dil numeral acids. ALC or eth soln oxidizes readily in air. Formed (in addn to other products, see above) by heating, in air, 2,4,6,2',4',6'-Hexanitro 3,3'-dioxo-azobenzene in a concd KOH soln, is converted to the trinitrophloroglucin in boiling w}
\end{itemize}

\textit{Ref:}\ Beil 15 [27]

2,4,6-Trinitro-3-hydroxylaminotoluene, N-[2,4,6-Trinitro-3-methylphenyl] hydroxylamine. \(\text{H}_3\text{CC}_6\text{H(NO}_2)_3(\text{NNOH});\ \text{mw}\ 258.15, \text{N}\ 21.71\%, \text{OB}\ -62\%;\ \text{dark yellow crystes (from aq alc), mp}\ 99°\ \text{(decomp); sol in alkalis or aq ammonia. Prep by heating 2,4,6-Trinitro-3-methoxy-
toluene with an alc soln of hydroxylamine. With coned nitric acid it forms 2,3,4,6-
tetranitrotoluene}

The 4,6-dinitro derivative has also been prepared

\textit{Ref:}\ Beil 15 [15]

\textbf{OH Hydrogen Substitution}

\textbf{Hydroxylamine-o-Sulfonic acid, NH}_2\text{OSO}_3\text{H;}\ \text{mw}\ 113.10, \text{N}\ 12.40\%;\ \text{white, hygr, microcrystalline solid, decom 210°. Stable if stored in moisture-free atmos. Prep by: the room temp reaction of (NH}_3\text{OH).H}_2\text{SO}_4 + 2\text{H}_2\text{SO}_4 + 3\text{SO}_3 = 2\text{NH}_3\text{OSO}_3\text{H + 3H}_2\text{SO}_4;\ by reacting a hydroxylamine salt with an excess of chlorosulphonic acid; by bubbling HN}_3\text{ thru fuming H}_2\text{SO}_4 at 60-80°}

\(\text{NH}_2\text{OSO}_3\text{H} \text{Reacts with ammonia and amines to give, respectively, hydrazine and substbd hydrazines}


\textbf{o-Methylhydroxylamine, α-Methylhydroxylamine, Methoxylnamine. CH}_3\text{ONH}_2;\ \text{mw}\ 47.06, \text{N}\ 29.77\%, \text{OB}\ -153\%;\ \text{fluid liq, bp}\ 49-50°;\ \text{sol in w, alc & eth. Prep by reaction of C}_6\text{H}_4\text{C(OC}_2\text{H}_5)_2: NOCH}_3 & \text{HCl; by heating the di-K salt of o-methylhydroxylamine-N,N-disulfonic acid in dilute acid soln. Explodes at 300°. Atomic constants and IR spectra have been measured (Ref 1)}

It can be stabilized against explosion by shock (adiabatic compression by a piston) by adding 75 parts by weight of ether, MeOH, N₂H₄ or MeNH₂ (Ref 2)

\textit{Ref:s:1) Beil 1, 288 (143) [275] & (1212) 2) S.A. Greene, USP 3117415 (1964) & CA 60, 9093 (1964)}

\text{Written by J. ROTH}
Hydroxymethyl alkanes. Compounds such as CH₃C(CH₂OH)₃ or C₂H₅C(CH₂OH)₃, called also “polymethyloalkanes.” See under “Polyhydroxy alcohols.”

Hydroxymethylbenzoic Acid.
5-Hydroxy, 2,4,6-Trinitro-3-methylbenzoic acid (also called 5-Oxy-2,4,6-trinitro-m-toluic acid). HOC₆(NO₂)₃(CH₃)COOH; mw 287.16, N 16.64%, OB to CO₂ 52.9%, crystals containing 1 mole H₂O, starts to melt (with decomp) between 170 & 180°, pulls off above 180°. Prepd by heating 5-hydroxy-3-methyl benzoic acid with 1.38 g/cc nitric acid.

Ref: Beil 10, 227 & (98)

2,2-Bis(hydroxymethyl)butanol (1) trinitrate.
C₆H₁₂C(CH₂ONO₂)₂; mw 269.17, N 15.61%, OB to CO₂ 50.5%; crystals (from acetone), mp 51.2°. Prepd by mixed acid nitration of 2,2-Bis(hydroxymethyl)butanol (1). It was used in explosive compositions by the Dupont Co.

Ref: Beil 1, (2350) 2) Not found in CA 1957-1971

N-Hydroxymethyl ethylenedinitramine or N-[Hydroxymethyl]-N,N'-dinitro ethylenediamine.
O₂N,N-HN.CH₂.CH₂.N(NO₂)CH₂OH; mw 196.13, N 31.1%, OB -40.7%; prisms, mp 127-130° (decomp on prolonged heating in water/bath)

May be prepd by adding 40% formaldehyde to ethylenedinitramine in boiling water.

Ref: F. Chapman et al, JCS 1949, 1635-36 & CA 44, 1411 (1950)

Hydroxyethyl hydroperoxide, Methylhydroperoxide or Methylhydroperoxide. (Called Oxyethylperoxy or Mono-oxymethylperoxy- in Ger). HOCH₂OH; mw 64.01; Oil of medium consistency; fairly stable in storage; RI 1.4205 at 16°. May be prepd by treating anhydrous formaldehyde in dry ether with anhydrous H₂O₂;

CH₂O + H₂O₂ = HOCH₂OOH

Easily sol in w, alc, eth, acetone & dioxane; sl sol in chlf & petroleum ether; insol in benz.

It is an explosive which is not sensitive to friction but explodes with extreme violence when heated in a flame. Its higher homologs are not explosive (Ref 4)

Its IR absorption spectrum has been studied (Ref 5)


Hydroxyethyl nitramines. See under Hydroxy- and Amino-methyl-nitramines

3-Hydroxy-6-nitro-2-phenylindazolone.

H N
\( \text{C}::\text{N}::\text{C}_6\text{H}_5 \)

mw 242.23, N 11.57%; crystals, mp > 260°. Prepd by refluxing 2,4-Dinitrobenzylidene aniline, \((O_2N)_2C_6H_5CH:NPh\), with Na-carbonate in EtOH and then treating with HAc. The Na salt of this product explodes on heating.

Ref: S. Sacareanu & I. Lupas, Bull soc chim (5) 1, 373 (1934) & CA 28, 5445 (1934)

1-Hydroxy-2-propanone. See Acetol in Vol 1 of Encycl, p A33-R

Hydroxypropylamine and N-(2-Nitroxypropyl)-nitramine, designated as Iso-Me-NENA, are described under Aminopropanols and Derivatives in Vol 1, p A253

α-Hydroxypropylethylperoxide (α-Oxypropyl- α-thyl-peroxy- in Ger). C₆H₅CH(OH)OOC₂H₅; mw 120.15, OB -173%; non-viscous oil; d 0.974 at 21°; bp 50-52° at 50mm; RI 1.4021 at 21.4°. May be prepd by treating propionaldehyde with ethylhydroperoxide, C₂H₅OOH. Explodes on heating.

Refs: 1) Beil, not found 2) A. Riche & F. Hitz, Ber 63, 2648 (1930); CA ref not found

Hydroxypropyl-3-oxyl-1,2-propanedioltrinitrate.
See under Hydroxyethyl-3-oxyl-1,2-propanedioltrinitrate in this Vol
2-Hydroxypyridine, 3,5-Dinitro (Called 3,5-Dinitro-oxy-pyridin in Ger).

\[
\begin{array}{c}
\text{O} \hspace{1cm} \text{N} \\
\text{N} \hspace{1cm} \text{NO}_2 \\
\text{OH}
\end{array}
\]

\[\text{mw 185.10, N 22.70\%, OB -56.2\%; yellow ndls (from eth), mp 133^\circ, sol in benz, alc or warm watter. Prepd by mixed acid nitration of 2-hydroxy-pyridine. Explodes weakly on heating above mp. Its alkali-salts are very explosive.}\]

The 3,5 Dinitro-4-hydroxy-pyridine, mp 325^\circ, on the other hand, or its alkali salts are not explosive. \(\text{Ref: Beil 21, 48 [33 & 35]}\)

**Hydroxyopyruvic Acid (Oxyypyruvic Acid).**

\[\text{C}_6\text{H}_4\text{O}_4; \text{mw 104.06. One of the products of the alkaline saponification of NC and nitro-oxycellulose (Ref 1). Berl and Smith (Ref 3) obtained it, calling it oxypyruvic acid, by the alkaline hydrolysis, not only of NC, but also of nitrates of glucose and levulose.}\]

This substance may be present as an impurity in commercial NC. \(\text{Ref: 1) W. Will, Ber, 24, 400 \& 3831 (1891)}\) \[\text{2) Marshall, T., (1917), 192} \]

\[\text{3) E. Berl \& W. Smith, JSCI, 27, 534 (1908)}\]

**HYDROXYQUINOLINE and DERIVATIVES**

Many hydroxquinoline isomers are known. Only the 2-hydroxy or 8-hydroxy quinolines appear to have nitro derivatives that are near explosive.

2-Hydroxyquinoline or Quinolone (Called 2-Oxy-chinolin in Ger).

\[
\begin{array}{c}
\text{N} \\
\text{OH}
\end{array}
\]

\[\text{mw 145.15, N 9.65\%; prisms (from alc), mp 199–200^\circ, v sl sol in w; sol in alc. Prepd by heating 2-Acetaminobenzaldehyde with NaOH in aq alcohol.}\]

\(\text{Ref: Beil 21, 77}\)

4-Hydroxyquinoline (Called 4-Oxy-chinolin in Ger).

\[
\begin{array}{c}
\text{O} \hspace{1cm} \text{N} \\
\text{OH}
\end{array}
\]

\[\text{mw 145.15, N 9.65\%; monoclinic ndls, mp 201^\circ, difficult to sublime; sol in warm water \& alc; sl sol in eth or benz. Prepd by heating 2-formaminoacetophenone in aq NaOH.}\]

\(\text{Ref: Beil 21, 83}\)

8-Hydroxyquinoline or 8-Quinolinol, oxyquinoline or Oxine (Called 8-Oxy-chinolin or Chinophenol in Ger).

\[
\begin{array}{c}
\text{HO} \\
\text{N}
\end{array}
\]

\[\text{mw 145.15, N 9.65\%; prisms (from alc), mp 75–76^\circ, bp 267^\circ, sublimes readily. Prepd by heating acidified 2-Amino-phenol with 2-Nitrophenol, glycerin \& concd sulfuric acid.}\]

\(\text{Ref: Beil 21, 91}\)

The mononitro derivatives of the 2-hydroxy & 8-hydroxy quinolines are stable (Beil 21, 99-100). The di \& tri-nitro derivatives are less stable but not really explosive. These are described below.

5,7 Dinitro-2-hydroxyquinoline.

\[
\begin{array}{c}
\text{NO}_2 \\
\text{N} \\
\text{OH}
\end{array}
\]

\[\text{mw 235.15, N 17.87\%, OB -104.2\%; golden-yel leaflets, mp 267^\circ (with decomposition); insol in most solvents. Prepd by nitrating 2-hydroxyquinoline with conc nitric acid.}\]

\(\text{Ref: Beil 21, 100, (220) \& [58]}\)

3,6-Dinitro-4-hydroxyquinoline.

\[
\begin{array}{c}
\text{O}_2\text{N} \\
\text{N} \\
\text{OH} \\
\text{NO}_2
\end{array}
\]

\[\text{mw 235.15, N 17.87\%, OB -104.2\%; mp 352–4^\circ (decomp). Prepd by mixed acid nitration of 4-hydroxyquinoline.}\]

5,7-Dinitro-8-hydroxyquinoline,

\[
\begin{array}{c}
\text{O} \\
\text{N} \\
\text{O}_2^+ \\
\text{OH} \\
\text{N}
\end{array}
\]

mw 235.15, N 17.87%, OB -104.2%; yellow crystals (from alc), mp 276° (Ref 1); mp 276-79° (from ethyleneglycol) with decomposition (Ref 3), mp 320° (decomp) (Ref 2). Prepd by concd nitric acid nitration of 8-hydroxyquinoline.


3,5,8 Trinitro-2-hydroxyquinoline,

\[
\begin{array}{c}
\text{O}_2 \text{N} \\
\text{NO}_2 \\
\text{OH}
\end{array}
\]

mw 280.15, N 20.00%, OB -76.8%; bright yellow needles (from xylene), mp 182° (decomp). Prepd by concd nitric acid nitration of 2-hydroxyquinoline.

Ref: Beil 21, (220)

3,6,8 Trinitro-4-hydroxyquinoline or 3,6,8-Trinitro-4-quinolinol,

\[
\begin{array}{c}
\text{O}_2 \text{N} \\
\text{NO}_2 \\
\text{OH}
\end{array}
\]

mw 280.15, N 20.00%, OB -76.8%; yel prisms, mp 296° with decomposition. Prepd by mixed acid nitration of 3-nitro-4-hydroxyquinoline (Ref 1); or reaction of AcONO₂ with quinoline N-oxide in Ac₂O (Ref 2).


Hydroxyquinone Diazides, Dinitro Derivatives of.

Percussion caps or detonators are loaded with one or more of the ortho- or para-nitroated quinone diazides of the polymeric phenols or their metallic salts. The K salt of dinitro-m-hydroxyquinone diazide is obtained from nitraminoresorcinol by treatment of the hot strong H₂SO₄ & HNO₃ soln with excess of KNO₃ & long boiling. On cooling, the salt crystallizes out in fine yellow needles. Dinitro-3,5-dihydroxyquinone diazide is prepd in the same way from mono- or di-nitroaminophloroglucinol.

Ref: E. von Herz, BritP 207563 (1922) & CA 18, 1573 (1924)

m-Hydroxytetrol. Same as 2,4,6-Trinitro-3-methylnitroaminophenol described under.

Methylaminophenol and Derivatives.

α-Hydroxytoluene. See Benzyl Alcohol in Vol 2 of Encycl, p B91-L

2-Hydroxytoluene. See α-Cresol in Vol 3 of Encycl, p C556-L

3-Hydroxytoluene. See m-Cresol in Vol 3 of Encycl, p C556-L

4-Hydroxytoluene. See p-Cresol in Vol 3 of Encycl, p C556-L

Hydroxy-tetrazole. See Tetrazolol and Derivatives. These include:

1-Hydroxytetrazole
1-Hydroxy-5-azidotetrazole
5-Hydroxytetrazole
Hydroxytetrazolylmethanol
Hydroxytetrazolylmethanol Nitrate

Hydroxy-triazole Derivatives. See Triazole Derivatives. These include:

3-Hydroxy-asym-triazole
Hydroxybenzotriazole, Chloro Derivatives
1-Hydroxy-4-benzoyl-5-methyl-sym-triazole
1-Hydroxy-4,5-dicarboxylic acid-sym-triazole
[2'-Hydroxynaphthyl]-1'-azo-3'-5'-azido-sym-triazole]
4-Hydroxymethyl-5-hydroxy-sym-benzotriazole
5-Hydroxy-1-(p-nitrophenyl)-3-carbonyl azide-asym-triazole
1-Hydroxy-5-phenyl-tetrazole
1-Hydroxy-sym-triazolopyrimidine

Hygiene-Industrial, in War Plants. The reader is referred to the following lectures & reviews:

"Industrial Hygiene at work in the national defense program" (Ref 1); "Mobilization of
Industrial hygiene for national defense" (Ref 2); "Industrial hygiene at work in defense industries" (Ref 3); "Industrial hygiene support in a missile program" (Ref 4).


Hygrometric Tests: Tests designed to determine hygroscopicity (see under Hygroscopicity Tests)

Hygroscopicity. Hygroscopicity is the property possessed by some substances of readily absorbing atmospheric moisture. Some explosives, as well as substances serving as ingredients of explosive mixtures, are hygroscopic. Most nitratated organic compounds per se are only slightly hygroscopic, but the materials added as oxidizers, such as NaNO₃, NH₄NO₃ etc., or as explosion temperatures suppressors, such as NaCl etc., are very hygroscopic.

Hygroscopicity is a property which is very undesirable in explosives, because moisture lowers their power, brisance and sensitivity. Moisture also lowers the ballistic potential of propellants. For this reason, it is advisable to keep explosive mixtures as non-hygroscopic as possible. This may be done either by excluding hygroscopic ingredients or by coating these ingredients with substances such as vaselin, paraffin, wax, oils, rosin etc. Nitrocellulose is made non-hygroscopic by gelatinization with solvents. Hygroscopic dynamites are protected from atmospheric moisture by enclosing them in paraffin coated paper cartridges.

Hygroscopicity Tests. Hygroscopicity is the affinity of a substance for water vapor. It is a complex phenomenon which is controlled by the rate of diffusion of water across the vapor-liquid interface. This rate depends on temperature, surface area, liquid depth, and liquid and vapor film coefficients. Inasmuch as it is impractical to measure the effect of all these variables, simplified empirical tests have been designed to determine the relative hygroscopicity of various substances.

Some of these tests applicable to explosives and propellants are as follows:

1) Desiccator test.

a) Using an analytical balance, weigh a large flat weighing dish provided with a glass stopper (W₁).

b) Weigh separately in a scoop, on a rough balance, 5-10g of a test sample (of known moisture content) and introduce it into the weighing dish. Stopper and reweigh (W₂).

c) Remove the stopper and place the dish in a desiccator containing a saturated solution of potassium nitrate containing a few crystals.

d) After leaving it in the desiccator for 24 or 48 hours, reweigh the dish (W₃).

% Moisture absorbed = \( \frac{W_3 - W_2}{W_2 - W_1} \) \times 100

Note: Instead of using the potassium nitrate solution, Wilson (Ref 1) recommends the use of aqueous solutions of sulfuric acid. The following table gives the strength of H₂SO₄ required to give different relative humidities (RH) at various temperatures.

<table>
<thead>
<tr>
<th>% Sulfuric Acid Required</th>
<th>0°</th>
<th>25°</th>
<th>50°</th>
<th>75°C</th>
</tr>
</thead>
<tbody>
<tr>
<td>RH</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>63.1</td>
<td>64.8</td>
<td>66.6</td>
<td>68.3</td>
</tr>
<tr>
<td>25</td>
<td>54.3</td>
<td>55.9</td>
<td>57.5</td>
<td>59.0</td>
</tr>
<tr>
<td>35</td>
<td>49.4</td>
<td>50.9</td>
<td>52.5</td>
<td>54.0</td>
</tr>
<tr>
<td>50</td>
<td>42.1</td>
<td>43.4</td>
<td>44.8</td>
<td>46.2</td>
</tr>
<tr>
<td>65</td>
<td>34.8</td>
<td>36.0</td>
<td>37.1</td>
<td>38.3</td>
</tr>
<tr>
<td>75</td>
<td>29.4</td>
<td>30.4</td>
<td>31.4</td>
<td>32.5</td>
</tr>
<tr>
<td>90</td>
<td>17.8</td>
<td>18.5</td>
<td>19.2</td>
<td>20.0</td>
</tr>
</tbody>
</table>

A chart giving a detailed relation between relative vapor pressures, %H₂SO₄ and density of solutions is given on p 328 of Ref 1.

2) British Hygroscopicity Test for Black Powder.

Put about 65g of powder (of known moisture content) into a tared small tray with a bottom of fine gauze and reweigh. Place it inside a box over a tray containing a saturated solution of KNO₃. Allow it to remain there either 24 hours for small grain powders, or for 48 hours for large grain powders. Reweigh the tray and determine the percentage of weight gain. Add to this the percentage of moisture originally present to obtain the hygroscopicity of the powder (Ref 2).
3) **Hygroscopicity Test at 15° in a moisture-saturated atmosphere.** A sample of 10-100g of dried material is placed in a constant temperature room at 15° saturated with moisture. The sample is reweighed periodically until constant weight is attained (Ref 4)

4) **US Armed Forces Test at 30° and 90% RH**

   a) Transfer a sample of about 100g to a tared (W₁) wide glass weighing dish with a ground glass cover. Weigh tare & sample on an analytical balance (W₂).

   b) Remove the cover and place the dish in a humidor containing 1 liter of a sulfuric-water mixture (18.6 ± 0.5% H₂SO₄), which will maintain a relative humidity of 90 ± 0.25% at 30°. The humidor shall have a total capacity of about 10 liters and should be capable of being sealed hermetically.

   c) Place the humidor in an oven maintained at 30 ± 2° and leave it there for 3 days.

   d) On the fourth day, remove the dish, stopper it and weigh (W₃).

   e) Repeat the weighings every day thereafter until the sample ceases to gain weight, indicating that the powder has reached equilibrium with 90% RH (W₄).

   \[
   \% \text{Gain} = \frac{W₄ - W₂}{W₃ - W₁} \times 100 = \%G, \text{where}
   \]

   W₂ - W₁ is usually 100g

   This figure has to be added to the total moisture content (%M) as determined on another sample of 100-200g and the sum gives the % hygroscopicity (%H) = %G + %M (Ref 3).

Total moisture content is determined on a separate 100 to 200g sample by distilling it in a 500ml flask with 200ml of dry CCl₄.

5) **Humidity Equilibrium Method** (Ref 1)

   a) Set up an apparatus consisting of 3 large bubbler bottles, containing aqueous solutions of H₂SO₄ of the desired humidity at the temperature of the test connected to a wide tube containing glass wool or cotton and to a tared (W₁) straight or U-tube containing 20-60g of loose powder without any cotton or glass wool. The weight of the tube with the sample is designated as W₂, and both weighings should be done on an analytical balance.

   b) Pass a slow current of compressed air (50 to 500cc per min, using the higher rates at lower temperatures) through the bubbler bottles and then through the glass wool tube into the tube containing the sample.

   c) Weigh the tube containing the sample every few hours until constant weight is reached (W₃).

   d) Disconnect the tube with the sample and pass a stream of warm air (50 to 125°) through it. This air stream is predried by passing it thru a U-tube containing P₂O₅, until constant weight in the sample tube is reached (W₄).

   The % "equilibrium moisture content" of the sample at the temperature and humidity of the test, and calculated on the dry weight of the sample is equal to: \(W₃ - W₄\), where (W₁) is the weight of the empty tube, (W₃) the weight of the tube plus humid sample and (W₄) the weight of tube plus dry sample.

   In order to make certain that equilibrium has been reached in any given case, it is always desirable to approach it from both the dry and the moist sides. The simplest and quickest way to accomplish this with a single sample is to pass dry air through the sample at the start and then determine the equilibrium weights at 10, 25, 50, 75 and 90% humidity. After this, saturated air (100% humidity, obtained by bubbling air through water) is passed for a short time thru the sample and the same points redetermined in the reverse order. Finally, dry hot air is passed thru in order to obtain the dry weight.

   Alternate means of controlling humidity by using saturated solutions of salts have been proposed (Ref 7). For instance, at 30° a saturated solution of KNO₃ gives about 92% RH; KCl = 84.5%; NaCl = 75.5%; Ca(NO₃)₂.4H₂O = about 50% etc.

6) **French test (LeRoux)**

   Fifty grams of a previously dried explosive sample is placed in a closed container above 21° Be'sulfuric acid. The % moisture pick-up of the explosive is determined daily and plotted on a chart (Ref 9).

   In addition to the general tests listed above a number of studies & tests for specific ingredients have been published. These data for propellants, NC and for ammonium nitrate (AN) are summarized below:

   The hygroscopicity of a propellant is determined rapidly in an apparatus consisting of 2 identical gas washing bottles provided with fritted glass plates near the bottom. The outlets at the bottom are connected through a diaphragm pump.
having a capacity of 400 l/hr. The top outlets are connected to complete the path for air circulation. In one flask is placed 50 g of the propellant in the form of shavings 0.1 mm thick. In the other is a soln of CaCl₂ of such concn that the vapor pressure of water over it at 20° is 0.75 of that of pure water. The entire apparatus is immersed in a thermostat at 20°. The moist air is circulated until equil is attained, which requires ~1½ hrs, as measured by increase in wt of the sample. The method may be used for materials other than propellants (Ref 15).

Various attempts have been made to relate the hygroscopicity of NC to its nitrogen content. For uncolloided NC at 30 to 70% RH the % moisture ≈ 14.6% N. Thus for "pyro cotton" (12.6% N) calc % moist = obs moist = 2% & for guncotton calc moist = 1.2 = obs moist = 1.3% (Ref 6). For colloided NC the above formula has been modified to % moist = 14.6-%N-0.4, but the calc moistures based on this modification do not always agree with experiment (Ref 6).

The linear (inverse) relation between NC moisture content and NC nitrogen content was also noted in Ref 5.

Urbanski (Ref 12) gives the moisture contents of 13% N & 12.5% N as 1 to 1.5% & 1.5 to 2% respectively.

Tavernier (Ref 10) discusses the effects of hygroscopicity and several other factors on the ballistic props of colloided double base propellants and incompletely gelatinized oxidizer-containing propellants. The studies of Ficherouille & Kovache (Ref 8) on effects of hygroscopicity (and several other factors) on the potential usefulness of substances as priming explosives should also be mentioned, although NC is not one of the substances examined.

Because of its importance in commercial explosives, and because it is highly hygroscopic, the hygroscopicity of AN has been studied intensively.

Equilibrium moistures (hygroscopic points) in AN determined by static or flow methods do not agree. In general, flow methods are recommended. To increase the rate of moisture exchange (at a given RH) the temp of system should be raised (Ref 11).

The hygroscopicity of AN can be decreased by the addition of calcium carbonate. The reduction in hygroscopicity is proportional to the calcium carbonate content if the latter is similar in grain size to the AN. Kaolin can be used almost as effectively as calcium carbonate (Ref 14).

AN starts to pick up moisture at 55% RH. With the addition of 1.12% P₂O₅ or 1.73% Ca(NO₃)₂ these humidities decrease to 45 & 33% RH respectively. AN begins to dissolve at 63-67% RH (Ref 13).

Effect of surface-active agents & mineral additions on the hygroscopic nature & caking tendency of ammonium nitrate was examined in the following lab experiments in order to improve the storage properties of AN: (a) AN was crystd from satd solns contg 0-1.0% of cationic, anionic or nonionic surfactants; (b) granules of AN (0.5-1.0mm) were contacted with a satd NH₄NO₃ sln contg varying amts of surfactants; (c) granules of NH₄NO₃ were wetted with mineral oil, excess of oil removed, & the granules coated with talc or clay. The effect of the concn of several surfactants on caking tendency & hygroscopicity, under varying pressures & relative humidity, was detd & the results related to those obtained with untreated AN. According to process (a), caking tendency decreased 2.5 fold & hygroscopicity decreased 2.0 fold; according to (b), these decreases were 4 & 5 times respectively when 0.1% of Dispergator NF or 0.2% of carboxymethylcellulose was used; and according to (c) these decreases were 4 & 1.5 times respectively (Ref 17).

To extend the shelf life of explosives contg NH₄NO₃, it is important to minimize its hygroscopicity. Samples were tested for hygroscopicity at relative humidity levels of 60, 70, 80 & 90% & between 25-30°. Most of the samples were coated; some were hand cut by a process producing prills; clay material was added in 2 cases. The results indicated that the NH₄NO₃ which contained the least amt of impurities or additives was also the least hygroscopic (Ref 18)

Laboratory investigations of Igdenit (ammonium nitrate-fuel-oil explosives). Flaked & prilled Igdenit, differing in particle-size distribution, d, & amt & kind of coating agent or additive were obtained from 8 sources. Hygroscopicities, relative rates of drying & ease of fracture of the dried particles after wetting were detd (Ref 16).
Hypercompression of Explosives. See Vol 3, p D20-R under Dead-Pressed Explosives

Hypergolic Propellants. Hypergolic propellant systems are those which ignite spontaneously on mixing. Those which do not are called non-hypergolic. Of course, the real difference between hypergolic and non-hypergolic systems lies in the rates of chemical reactions at the mixing temperatures. Hypergolic propellants react quickly enough at ambient temperatures to liberate heat so that they therefore ignite spontaneously. Ignition delay measurements normally are indicative of rates of reaction; and so much work has been done in this area.

Ignition Delay. Ignition delays in the range of 39-80 msec were measured by the impingement method for 80% H2O2 and mixtures of N2H4, CH3OH and H2O. Delay varied with impingement angle, mixture ratio and temp (Ref 1). For combinations of HNO3 with aniline, furfuryl alc, and mixts of aniline and furfuryl alc, ignition lags ranged from 10 to 400 msec depending upon temp, acid compn, fuel compn and metalic additives (Ref 2). Wetting agents significantly decrease ignition lag of HNO3 combinations with furfuryl alcohol dicyclopentadiene. Oleamide, triethanolamine oleate, glyceryl oleate, sodium sulfocinate and alkyl-aryl sodium sulfate were most effective (Ref 3). Ignitibility and delay times were also clearly related to chemical structure of the fuel. Comparisons have been made between hypergolic characteristics of primary, secondary, and tertiary amines as well as the effect of substituent groups such as methyl, hydroxy, and phenyl on the a- and β-carbon atoms of various amines with white fuming nitric acid oxidizer (Ref 4). With highly reactive combinations such as N2H4-H2O2, N2H4 and N2H4-NH3 mixtures with RFNA (red fuming nitric acid) and WFNA (white fuming nitric acid) the mixing time is the rate-controlling step in the process and ignition delays in the order of a millisecond are observed (Ref 5). The National Advisory Committee for Aeronautics extensively studied ignition delays of nitric acid oxidizers. Their noteworthy conclusions are summarized in Ref 6.

The ignition delay phenomena have been extensively studied for hypergolic combinations of N2O4 with various fuels because of the almost exclusive use of this oxidizer in the upper stages of the US military and civilian space program. Propellant combinations used in specific launch vehicles are given in Ref 34. A summary of a literature and industry survey of the hypergolic ignition spike phenomenon is given in Ref 35. The physics of spray formation, chemistry of combustion, physico-chemistry, gas dynamics, and transport are dealt with. Ignition limits have been detd for various combinations of N2O4 and N2H4 under press and temp conditions of space chambers of varying sizes. Vapor phase and condensed phases were investigated (Ref 36). The reactivity of N2O4 with Aeroxide-50 (50/50 hydrazine/unsym dimethylylhydrazine) was detd at pressures from 10^-5 to 1 atm. At pressures
below 0.01 atm explosions and detonations occurred under certain specified conditions (Ref 37). Various physical phenomena influencing ignition delay of HNO₃ and N₂O₄ oxidizers and mixts include diameter of injection tube, stream velocity, mixing ratio, chamber pressure, fuel & oxidizer temps, water content of fuel & oxidizer. Fuel mixts investigated include furfuryl alc & primary amines, UDMH (unsym dimethylhydrazine) and phenylacetylene (Ref 38). Ignition delay data are given in Ref 39 for fuels made hypergolic by addition of UDMH. Included are alkanes, aldehydes, alcohols, esters, C₆H₆ and derivatives, olefins, naphthalene & derivatives, furan & derivatives, cyclohexane & derivs, and methylcyclopentane & derivs. The amount of UDMH required decreased with increasing chain length. Oxidizers included HNO₃, N₂O₄, H₂SO₄ and H₂O. A mathematical model for the hypergolic ignition in space for N₂O₄/MMMH (monomethylhydrazine) and N₂O₄/UDMH has been developed. It predicts ignition delay times for space rocket engines and ignition pressure spikes (Ref 40).

The vacuum ignition characteristics of flox/diborane (flox is a mixt of F₂ & liq O₂) and oxygen difluoride/diborane were investigated in 100-lbf rocket engines for possible use in space engines. Variables tested included hardware temp, propellant temp, oxidizer leading or lagging fuel, chamber press and injector configuration (Ref 41).

**Hypergolic Oxidizer and Fuel Mixtures.** A list of oxidizers and fuels which are hypergolic follows: RFNA with: aniline (Ref 7); N₂H₄ (Refs 8 & 9); H₂ (Ref 10); toluene (Ref 11); diethylenetriamine (Ref 8); MMH (Ref 8); UDMH 50% & N₂H₄ 50% (Ref 8); ethylenediamine (Ref 8); UDMH 60% & diethylenetriamine 40% (Ref 8); pentaborane (Ref 8); triethylaluminum (Ref 12); diethylzinc (Ref 12); trimethylaluminum (Ref 12); diborane (Ref 12).

WFNA with: aniline (Ref 2); N₂H₄ (Ref 8); MMH (Ref 8); UDMH (Ref 8); UDMH 50% & N₂H₄ 50% (Ref 8); diethylenetriamine (Ref 8); UDMH 60% & diethylenetriamine 40% (Ref 8); pentaborane (Ref 8); amines (Ref 4); JP-4 & dicyclopentadienyliron (Ref 13).

N₂O₄ with: H₂ (Ref 10, 14); pentaborane (Ref 8, 10); N₂H₄ (Ref 8, 10, 15, 16); MMH (Ref 8, 10, 14, 17); ethylenediamine (Ref 8); diethylenetriamine (Ref 8); UDMH (Ref 8, 14); UDMH 50% & N₂H₄ 50% (Ref 8, 10, 14, 18).

Oxygen with: N₂H₄ (Ref 10); UDMH 50% & N₂H₄ 50% (Ref 10); pentaborane (Ref 10, 20); triethylaluminum (Ref 21); trimethylaluminum (Ref 22); boranes & derivatives (Ref 19).

Ozone with: diethylenetriamine (Ref 10); N₂H₄ (Ref 10); hydrogen (Ref 10); pentaborane (Ref 10); UDMH (Ref 10); trimethylaluminum (Ref 19); morpholines (Ref 24); polyamines (Ref 23).

H₂O₂ with: UDMH (Ref 10); H₂ (Ref 10, 14); MMH (Ref 10, 17); pentaborane (Ref 10); triethylaluminum (Ref 14); triethylboron & other borane derivatives (Ref 19).

Fluorine with: N₂H₄ (Ref 10); H₂ (Ref 10, 14); MMH (Ref 10, 17); pentaborane (Ref 10); triethylaluminum (Ref 14); triethylboron & other borane derivatives (Ref 19).

Oxygen/Fluorine (FLOX) with: ethylene (Ref 26); methane (Ref 26); propane (Ref 26); propylene (Ref 26); diborane (Ref 27).

ClF₃ with: N₂H₄ (Ref 8, 10, 16, 28); H₂ (Ref 10); MMH (Ref 8, 17); pentaborane (Ref 8, 10); UDMH (Ref 8); diethylenetriamine (Ref 8); ethylenediamine (Ref 8); UDMH 60% & diethylenetriamine 40% (Ref 8); kerosene (Ref 19); triethylaluminum (Ref 19); triethylboron (Ref 19); diborane (Ref 19); UDMH 50% & N₂H₄ 50% (Ref 8, 10); PBAA (polybutadiene/acrylic acid/acrylonitrile) + NH₄ClO₄ + Al (Ref 29); polysulfide + NH₄ClO₄ + Al (Ref 29); polyurethane + NH₄ClO₄ + Al (Ref 29); double-base propellant (Ref 29).

ClO₃F with: UDMH 50% & N₂H₄ 50% (Ref 8); diethylenetriamine (Ref 8); ethylenediamine (Ref 8); pentaborane (Ref 8); N₂H₄ (Ref 8, 10); H₂ (Ref 10); MMH (Ref 8, 10); UDMH (Ref 8, 10); UDMH 60% & diethylenetriamine (DETA) 40% (Ref 8, 10).

N₂F₄ with: NH₃ 36% & N₂H₄ 64% (Ref 10); N₂H₄ (Ref 10); H₂ (Ref 10).

Oxygen difluoride with: diborane (Ref 27); H₂ (Ref 10); UDMH 60% & diethylenetriamine 40% (Ref 10); MMH (Ref 10); RP-1 fuel (Ref 10).
BrF$_3$ with: N$_2$H$_4$ (Ref 8); MMH (Ref 8); UDMH (Ref 8); UDMH 50% & N$_2$H$_4$ 50% (Ref 8); diethylenetriamine (Ref 8); ethylenediamine (Ref 8); UDMH 60% & diethylenetriamine 40% (Ref 8); pentaborane (Ref 8)  
ClF$_3$ with: N$_2$H$_4$ (Ref 30, 31); Li, Ca, Na, NaK and K (Ref 32)  

**Physical Properties.** Physical properties and handling hazards for the following hypergolic propellant fuels and oxidizers are given in Ref 42: alkyl borane, aniline, ClF$_3$, F$_2$, N$_2$H$_4$, H$_2$, H$_2$O$_2$, MMH, RFNA, N$_2$O$_4$, O$_2$, pentaborane, ClO$_3$F, and UDMH  

**Thermochemical and Thermodynamic Properties.** The heat release rate for the liq phase reaction of N$_2$O$_4$ & N$_2$H$_4$ was measured and found to be equal to that for HNO$_3$/N$_2$H$_4$ determined by Somogyi & Feiler (Ref 43). Both heat & gas release were detd for liq H$_2$O$_4$ with N$_2$H$_4$, MMH, and UDMH which were mixed by free jets impinging and quenched. Calorimetric and PVT measurements established total heat and gas release. Reactivity increases in order of: N$_2$H$_4$, UDMH, and MMH (Ref 44). Enthalpy, entropy, phase compn, press, temp, and sp vol for Aerozine 50 (50% N$_2$H$_4$ & 50% UDMH) have been detd and charted (Ref 45). Significant chem species and reactions in propellant exhausts containing C, H, O, N, F, Cl, and either Al, Be, B or Li were detd for use in nonequil performance calcs. Propellant systems included hypergolic liquid propellants and hybrid propellants (Ref 46). Important chem reactions in the following propellant mixts were used to describe the kinetics of nonequil expansion and calculated theoretical sp imp: O$_2$/H$_2$; F$_2$/H$_2$; N$_2$O$_4$/ASO (49% UDMH & 51% N$_2$H$_4$); ClF$_3$/N$_2$H$_4$; OF$_2$/B$_2$H$_6$; F$_2$/N$_2$H$_4$; FLOX (69.3% F$_2$ & 30.7% O$_2$)/RP-1; liq O$_2$/RP-1 (Ref 47). Theoretical performance calcs have been made for O$_2$, F$_2$/beryllium; lithium hydride/H$_2$ (Ref 48); aviation gasoline/fuming nitric acid (Ref 49); and oxygen difluoride/diborane propellants (Ref 50)  

**Explosion Properties.** Occurence of detonation within rocket engines employing certain aminated fuels and nitric acid propellants are related to the formation of amine or hydrazine nitrates which tend to decompose explosively under the influence of sudden temp or pressure rises (Ref 51). Tangential-mode rocket motor instabilities were analyzed using 1-dimensional, 2-phase detonation wave as a reaction model. Theoretical analysis compared favorably with exptl data from the literature on N$_2$O$_4$–HNO$_3$/N$_2$H$_4$ rocket motor firings.  

TNT equivalent blast yields and fireball dimensions were detd experimentally for the hypergolic propellant N$_2$O$_4$/Aerozine 50 (Ref 53, 54). Explosive reactions have been reported for ClF$_3$ with methane and propane (Ref 55). Violent reactions occur between N$_2$O$_4$ and many halogenated hydrocarbon solvents (Ref 56). Most reactive are the partly chlorinated cmpds of C$_2$H$_6$, C$_2$H$_4$, & CH$_4$. Explosive sensitivity decreases in solvents: 1) with greater ratios of Cl:H atoms; 2) that are symmetrical; 3) with no double bonds and 4) with more F than Cl atoms  

**Propellant Performance Data.** Specific impulse and chamber temperature for a number of more common hypergolic propellant combinations are in the following table. The values are based on shifting equilibrium conditions with a chamber pressure of 1000 psia. Data are from Ref 33  

**Chlorine Trifluoride (ClF$_3$)**  
Heat of formation, liquid, $\Delta H_f^\circ = -44.4$ kcal/mole  
Density of liquid, 1.81 g/ml at 298°K  
Melting point, -82.6°C  
Boiling point, 11.3°C  

**Maximum Specific Impulse (1000 + 14.7 psia)**  

<table>
<thead>
<tr>
<th>Fuel</th>
<th>Oxidizer</th>
<th>$T_c$, °K</th>
<th>$\rho$ (g/cc)</th>
<th>Isp (Sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>H$_2$</td>
<td>92</td>
<td>3403</td>
<td>0.612</td>
<td>318</td>
</tr>
<tr>
<td>N$_2$H$_4$</td>
<td>73</td>
<td>3882</td>
<td>1.48</td>
<td>293</td>
</tr>
<tr>
<td>B$_2$H$_6$</td>
<td>88</td>
<td>4487</td>
<td>1.47</td>
<td>290</td>
</tr>
<tr>
<td>UDMH</td>
<td>75</td>
<td>3794</td>
<td>1.37</td>
<td>280</td>
</tr>
<tr>
<td>Polyethylene</td>
<td>77</td>
<td>3541</td>
<td>1.48</td>
<td>257</td>
</tr>
<tr>
<td>CH$_3$N$_2$H$_3$</td>
<td>74</td>
<td>3647</td>
<td>1.42</td>
<td>283</td>
</tr>
<tr>
<td>DETA</td>
<td>75</td>
<td>3599</td>
<td>1.47</td>
<td>267</td>
</tr>
</tbody>
</table>

**Bromine Pentafluoride (BrF$_5$)**  
Heat of formation, liquid, $\Delta H_f^\circ = -132$ kcal/mole  
Density of liquid, 2.466 g/ml at 298°K  
Melting point, -52.5°C  
Boiling point, 40.3°C
### Maximum Specific Impulse (1000 + 14.7 psia)

<table>
<thead>
<tr>
<th>Fuel</th>
<th>Wt %</th>
<th>Oxidizer</th>
<th>( T_p , ^\circ \text{K} )</th>
<th>( \rho(\text{g/cc}) )</th>
<th>Isp(Sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH(_3)_N(_2)_H(_3)</td>
<td>78</td>
<td>3264</td>
<td>1.76</td>
<td>235</td>
<td></td>
</tr>
<tr>
<td>B(_2)_H(_6)</td>
<td>92</td>
<td>4380</td>
<td>1.99</td>
<td>246</td>
<td></td>
</tr>
<tr>
<td>N(_2)_H(_4)</td>
<td>77</td>
<td>3323</td>
<td>1.85</td>
<td>244</td>
<td></td>
</tr>
<tr>
<td>UDMH</td>
<td>78</td>
<td>3215</td>
<td>1.68</td>
<td>231</td>
<td></td>
</tr>
<tr>
<td>DETA</td>
<td>78</td>
<td>3042</td>
<td>1.84</td>
<td>220</td>
<td></td>
</tr>
</tbody>
</table>

### Hydrogen Peroxide (H\(_2\)O\(_2\)) 100%

Heat of formation, liquid, \( \Delta H_f^\circ = -44.84 \text{ kcal/mole} \)
Density of liquid, 1.443 g/ml at 298°K

### Maximum Specific Impulse (1000 + 14.7 psia)

<table>
<thead>
<tr>
<th>Fuel</th>
<th>Wt %</th>
<th>Oxidizer</th>
<th>( T_p , ^\circ \text{K} )</th>
<th>( \rho(\text{g/cc}) )</th>
<th>Isp(Sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>H(_2)</td>
<td>88</td>
<td>2404</td>
<td>0.435</td>
<td>322</td>
<td></td>
</tr>
<tr>
<td>B(_2)_H(_6)</td>
<td>73</td>
<td>3400</td>
<td>1.06</td>
<td>316</td>
<td></td>
</tr>
<tr>
<td>N(_2)_H(_4)</td>
<td>67</td>
<td>2923</td>
<td>1.26</td>
<td>286</td>
<td></td>
</tr>
<tr>
<td>UDMH</td>
<td>81</td>
<td>3008</td>
<td>1.25</td>
<td>283</td>
<td></td>
</tr>
<tr>
<td>Polyethylene</td>
<td>86.5</td>
<td>2999</td>
<td>1.30</td>
<td>277</td>
<td></td>
</tr>
</tbody>
</table>

### Fluorine (F\(_2\))

Heat of formation, liquid, -3.47 kcal/mole at 85.2°K
Density of liquid, 1.51 g/cc at 85.2°K
Melting point, -219.6°C
Boiling point, -188°C

### Maximum Specific Impulse (1000 + 14.7 psia)

<table>
<thead>
<tr>
<th>Fuel</th>
<th>Wt %</th>
<th>Oxidizer</th>
<th>( T_p , ^\circ \text{K} )</th>
<th>( \rho(\text{g/cc}) )</th>
<th>Isp(Sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>H(_2)</td>
<td>89</td>
<td>3964</td>
<td>0.468</td>
<td>410</td>
<td></td>
</tr>
<tr>
<td>N(_2)_H(_4)</td>
<td>69</td>
<td>4688</td>
<td>1.31</td>
<td>363</td>
<td></td>
</tr>
<tr>
<td>B(_2)_H(_6)</td>
<td>82</td>
<td>5101</td>
<td>1.20</td>
<td>360</td>
<td></td>
</tr>
<tr>
<td>UDMH</td>
<td>71</td>
<td>4342</td>
<td>1.19</td>
<td>344</td>
<td></td>
</tr>
<tr>
<td>Polyethylene</td>
<td>72.5</td>
<td>4382</td>
<td>1.28</td>
<td>325</td>
<td></td>
</tr>
<tr>
<td>NH(_3)</td>
<td>77</td>
<td>4543</td>
<td>1.15</td>
<td>357</td>
<td></td>
</tr>
</tbody>
</table>

### Red Fuming Nitric Acid (RFNA)

Red fuming nitric acid contains 2 weight percent H\(_2\)O, 84.6% HNO\(_3\) and 13.4% N\(_2\)O\(_4\)

Heat of formation, liquid, \( \Delta H_f^\circ = -41.0 \text{ kcal/mole} \)
Density of liquid, 1.57 g/cc at 298°K
Melting point, -49°C
Boiling point, 66°C

### Nitrogen Tetroxide (N\(_2\)O\(_4\))

Heat of formation, liquid, \( \Delta H_f^\circ = -5.4 \text{ kcal/mole} \)
Density of liquid, 1.43 g/cc at 298°K
Melting point, -11.2°C
Boiling point, 21°C

### Written by E. L. CAPENER

Refs: 1) J.D. Broatch, Fuel, 24-5, 106 (1950)
2) S.V. Gunn, J. AmerRocketSoc 22, 33 (1952)
& CA 47, 8341 (1953) 3) M.L.J. Bernard,  
FifthSymp(Intntl)Comb, 217, Reinhold, NY  
(1955) 4) L.R. Rapp & M.P. Strier, JetPropulsion  
27, No 4, 401 (1957) & CA 51, 8405  
(1957) 5) M. Kilpatrick & L. Baker, Jr, Fifth  
SympComb, Reinhold, NY (1955) 6) G. Morell,  
"Summary of NACA research on ignition lag of  
self-igniting fuel-nitric acid propellants," NACA  
RM E57G19 (Oct 3, 1957) 7) R.P. Rastogi  
et al., IndianJChem 2, No 8, 301 (1964) & CA  
61, 14453 (1964) 8) A.F. Duncan et al, "A  
study of hypergolic bipropellant mass explosion  
for attitude control of spacecraft," Final Rept  
No 5915 (Jan 15-Aug 31, 1962), Marquardt

Hyperol or Perhydrate. A tradename for the compound of hydrogen peroxide and urea, OCNH₂₂, H₂O₂, solid, decomp above 82° (Ref 2). It is claimed to be an advantageous replacement for hydrogen peroxide in chemical analyses (Ref 1)

Refs: 1) J.R. Boer, Chem & Indus 44, 1137 (1925) & CA 20, 158 (1926) 2) E. Janecke, RecTravChim 51, 579 (1932) & CA 26, 4747 (1932)

Hypervelocity Gun. A patent by Clark & Boltz (1959) claims a device consisting of a compression chamber & piston energy absorber that produces shear projectiles at velocities of 10,000 to 15,000 ft/sec which is claimed to be 5 times greater than velocities that are produced by impact. Applications for this device include simulation of possible effects of meteor particles on missiles in outer space and general studies of high-velocity particles colliding with various materials

Hypervelocity firings into mixtures of hydrogen with air or with oxygen were studied by H. Behrens et al and reported at 10th Symp Combust (1964), pp 245-52

Refs: 1) A.B.J. Clark & P.T. Boltz, USP 2882796 (1959)
2) Anon, Ordnance 44, 468 (1959)

Hypervelocity Impact: Dependence of Crater Dimensions on Impact Velocity. Craters in copper and lead, produced by hypervelocity impact, were measured and the dimensions correlated with impact vel. The results indicate that craters scale with ca the 1.7 power of vel, in agreement with computer physics results based upon hydrodynamic calculations
HYPOCHLOROUS ACID AND HYPOCHLORITES

HyPOCHLORIC Acid (Unterchlorige Säure in Ger), HClO or HOCl; known only in aq solutions. Although chlorine water was prepdk as early as 1774 by C. W. Scheele (Ref 1; p 243), it was not known that chlorine reacted with water, Cl₂ + H₂O = HClO + HCl, forming a new acid, HClO. A. J. Balard first identified it; he prepdk not only an aq solution of the acid, but also isolated the anhydride, chlorine monoxide.

In the preparation of hypochlorous acid from chlorine and water, the yield is very low, on account of the small solubility of chlorine in water. The yield of HClO can be increased either by adding an insoluble oxide or carbonate which forms a sparingly soluble chloride. For instance, on adding mercuric oxide, the reaction proceeds; 2Cl₂ + H₂O + HgO = HgCl₂ + 2HClO, and with silver carbonate; 2Cl₂ + H₂O + Ag₂CO₃ = 2AgCl + 2HClO + CO₂.

Another way to increase the yield is to pass additional chlorine into soda or lime solutions of chlorine; Cl₂ + Na₂CO₃ + H₂O = NaClO + NaCl + H₂CO₃, followed by acidification with H₂SO₄ and distillation of the liberated HClO. The acid can also be made by the action of water on chlorine monoxide, Cl₂O. This reaction is accompanied by the evolution of 4 kcal of heat per mol.

Hypochlorous acid is a weak acid with a dissociation constant equal to 3 to 4 x 10⁻⁸, which is about 1/600th the strength of acetic acid. When it is stored it slowly decomposes according to the following equations:

2HClO = 2HCl + O₂ or 3HClO = 2HCl + HClO₃. Both reactions are greatly accelerated by light. The first of these reactions makes hypochlorous acid a strong oxidizing agent.

Hypochlorous acid reacts with inorganic bases with the formation of salts, called hypochlorites. In reactions with organic compounds, the acid may act not only as an oxidizer (see above) but also as a chlorinating agent (in the case of saturated compounds) or the addition of HOCl to the double linkage C=C to form a chlorhydrin; R₁C = CR₂ + HOCl = R₁C(OH),C(Cl)R₂.

Solutions carrying up to 1% hypochlorous acid, free from metals and protected from strong light, may be stored for a day without important loss. Solutions up to 20% may be kept for a short time at room temp, and 30% solutions may be kept for a long time at -20° (Ref 3, p 694).

For more information on the properties of hypochlorous acid, see Refs 1 & 2.

HyPOCHLORITES. The combination of the hypochlorite ion with a metallic ion or organic radical is called a hypochlorite. The inorganic compounds are called salts, while organic hypochlorites are esters.

The first hypochlorites (those of sodium and potassium) were prepdk in solution, in 1788, by C. L. Bertholet by treating solutions of the corresponding hydroxides with chlorine; 2NaOH + Cl₂ = NaCl + NaClO + H₂O.

In 1834, A. J. Balard prepdk solutions of various hypochlorites by mixing aq hypochlorous acid with alkalies, magnesium, copper oxide, zinc oxide, etc., avoiding an excess of acid.

J. L. Gay-Lussac dissolved two mols of chlorine monoxide, Cl₂O, in a solution containing 1 mol of K₂O and then evaporated the resulting solution in vacuo. This gave solid potassium hypochlorite.

The following inorganic hypochlorites are described in literature: aluminum (very unstable), ammonium (very unstable), barium, calcium, cupric, lithium, magnesium, potassium, silver (unstable), sodium, strontium and zinc. None of these salts is used in explosives or is explosive per se, but some of them are used in industry.

However, when small amounts of damp S and calcium hypochlorite are mixed, a brilliant crimson flash is observed accompanied by the scattering of molten burning S (Ref 9).

Also, according to J. Welchherz, Chem Ztg, 52, 729-30 (1928), a mixture of Ca hypochlorite with combustibles explodes on heating (See also Ref 2).

Calcium hypochlorite has been used to neutralize toxic explosion gas in mining (Ref 8): Explosion of 1 kg detonite 10A & ammonite No 62, yields 42-56, and 58-81 l poisonous gases, respectively. Use of ampules containing pyrolusite, NaHCO₃, or Ca hypochlorite (15-25% of the explosive wt) as neutralizing substances neutralized 50% of the poisonous gases &
decreased by 30% the required ventilation time of the mine galleries. Hydrogen peroxide (3%), Ca hypochlorite, or water were introduced in polyethylene ampules (vol 250 m³) into the bore hole, whose opening was hermetically sealed by a “hydroseal,” and water was introduced into the bore hole under pressure.

**Analysis of Hypochlorites.**

Methods of determination of hypochlorous acid and hypochlorites are given in Ref 2, pp 292-296.

According to Scott (Ref 5, p 262), hypochlorites decolitze indigo, but do not decolorize KMnO₄ solutions. If arsenous acid is present, indigo is not decolorized until all the arsenous acid has been oxidized to the arsenic form (distinction from chlorous acid and chlorites). A quantitative method of determination of hypochlorites in the presence of chlorine by using the KI method, is given in Scott (Ref 5, p 274).

Feigl (Ref 6) gives a spot test for hypohalogenates using benzidine reagent and Welcher (Ref 7) gives several qualitative and quantitative tests.


**Organic Hypochlorites** (Esters of Hypochlorous Acid). Methods of preparing alkyl esters of hypochlorous acid are given under individual compounds, but a general method, recently proposed, is via the reaction of HClO with the appropriate alcohol in solution (preferably in freon but pet ether, carbon disulfide or benzene can also be used) (Ref 8). The lower members are unstable, mobile, yellow oils and most of them explode when brought into contact with a flame or when exposed to bright light. In the absence of a flame or light, all of them, except the tertiary compounds, decompose spontaneously on standing (Ref 8). According to Comastri (Ref 6), they are much more stable if any HCl, formed as a result of the reaction, is immediately removed by neutralization with NaHCO₃ (See also under Ethyl Hypochlorite).

The decomposition reactions of the primary, secondary and tertiary hypochlorites are as follows:

1) RCH₂OCl  RCHO + HCl
2) RR₃CHOCI  RR₃CO + HCl
3) RR₃R₂COCl  RR₃CO + R₂Cl

In reaction (1), an aldehyde is formed, in (2), a ketone and in (3) a ketone and an alkyl chloride. Reaction (3) only proceeds on heating.

In general, the esters are strong oxidizing and chlorinating agents and they explode on contact with copper powder.

Some aliphatic hypochlorites are described below:

**Amyl Hypochlorites.** C₅H₁₁ClO; yellow oils; may be prepared by bubbling chlorine through an aqueous emulsion of the corresponding alcohol, or by shaking a CCl₄ solution of an alcohol with an aqueous solution of HClO (Refs 2, 4 & 5).

All of these esters, especially the isoamyl are unstable; the ester of the tertiary alcohol, (CH₃)₂C(Cl)O)C₂H₅, is more stable than the others; it boils at 76° at 752mm and decomposes in sunlight, but keeps fairly well in the dark.

Ref: Beil 1, 423 & [433]

**Butyl Hypochlorites.** C₄H₉ClO; yellow oils; may be prepared by methods similar to those for the amyl hypochlorites. These esters, especially those of isobutyl alcohol, are unstable.

The esters of n-butyl- and sec-butyl-alcohols explode when exposed to light. The tertiary
butyl hypochlorite, \((\text{CH}_3)_2\text{C}(\text{OCl})\), is more stable than the others. Its bp is 79.6° at 750mm and d is 0.9583 at 18° (Refs 1, 4 & 5) (Beil 1, [396]— the primary hypochlorite, \(\text{CH}_3\text{C}(\text{HOCl})\), [402]; sec-butylyl hypochlorite, \(\text{C}_4\text{H}_9\text{Cl}\) (\(\text{CH}_3\)OCl) and [415] — tertiary compound; [411]

Ethyl Hypochlorite. \(\text{CH}_3\text{CH}_2\text{Cl}; \text{mw 80.52}, \text{OB to CO}_2 \text{ and Cl}_2 -109.3\%, \text{d=1.013 at 6°}, \text{bp 36° at 758mm. Yellow, volatile, oily liquid; was prep'd in 1885 by Sandmeyer (Ref 3) by bubbling Cl}_2 \text{through cold ethanol or by mixing a cold aq solution of HClO with cold ethanol.}

Other methods of preparation are given in Refs 1 to 5

Comastri (Ref 6) gives the following method of preparation; Place in a separatory funnel a solution containing 23g of NaOH in 23g of ethanol and 200ml H_2O. Cool in an ice-salt bath and pass in chlorine through a tube reaching to the bottom of the container. The resulting ethyl hypochlorite collects in the form of an oil layer on the surface of the liquid and may be separated easily. It is washed with a cold saturated solution of NaCl, followed by a cold satd NaHCO_3 solution, and immediately dried (while cold) over CaCl_2. The yield is about 72% of theory. The following reaction takes place:

\[ \text{C}_2\text{H}_5\text{OH} + 2\text{HClO} = \text{C}_2\text{H}_5\text{ClO} + \text{HCl} + \text{H}_2\text{O} \]

Comastri claims that, contrary to the literature, the action of light on \(\text{C}_2\text{H}_5\text{ClO}\) is inappreciable and the material can be kept for several hours at room temp, provided that all HCl (formed as result of the above reaction) is promptly neutralized with NaHCO_3 solution.

If the HCl is not removed, the hypochlorite decomposes (or even explodes) spontaneously in a few minutes, on account of the following reaction:

\[ \text{HCl} + \text{C}_2\text{H}_5\text{ClO} \rightarrow \text{C}_2\text{H}_5\text{Cl} + \text{C}_2\text{H}_5\text{OH} \]

Ethylhypochlorite is insol in w; sol in bz, chf and eth. It explodes violently on heating or by the action of direct sunlight (Beil 1, 324 (164) and [325])

Methyl Hypochlorite. \(\text{CH}_3\text{ClO}; \text{mw 66.49}, \text{bp 12° at 726mm. Gaseous substance; was prep'd by Sandmeyer (Refs 2 & 3) by bubbling chlorine through a cold solution of NaOH in methanol. EXPLODES on contact with flame (Beil 1, 282 and [271])

Pentyl Hypochlorites. Same as Amyl Hypochlorites

Propyl Hypochlorites. \(\text{C}_3\text{H}_7\text{ClO}\); yellow oily substances. May be prep'd by methods similar to the ones mentioned under Amyl Hypochlorites (see above)

All the propyl hypochlorites, especially the isopropyl, are unstable. n-Propyl-, as well as the isopropyl hypochlorites, explode when exposed to light (Refs 2, 4 & 5) (Beil 1, [368] n-propylester; [381] iso-propylester)


 Hypophosphites are salts of hypophosphorous acid, H(\text{H}_2\text{PO}_3)_\text{H}

\[ \text{H:P:O:H} \]

They are generally prepared by the action of phosphorous on the corresponding salt in alkaline solution. The free acid is prep'd by treating Ba-hypophosphate with sulfuric acid. It is monobasic and is a strong reducing agent (Ref 6)

Hypophosphites have been claimed as ingredients of priming compositions: Lead hypophosphite/MF/Dinitrophenylazide/K_3\text{Ba(NO}_3)_4 & \text{Sb}_2\text{S}_3 (Ref 2); alkali, alkaline earth, \text{NH}_4 & \text{Mn hypophosphites mixed with oxidizers such as Pb or Ca nitrate (Ref 4); complex salt hypophosphites suitable for use in detonators are claimed to be formed from equimolar amounts of normal or basic Pb staphnate and Pb hypophosphite (Ref 5)

An early French patent claims the manufacture of explosives based on hypophosphites mixed with oxidizers (Ref 1)
Burrows has claimed that the double salt of Pb hypophosphite and Pb nitrate is a suitable ignition composition around the bridge wire of a blasting cap. He also claimed such mixture as Ca hypophosphite and K chlorate (Ref 3)


Hyporka. A glass fiber insulation used in rockets

“Hyros” Explosives. Prepd by nitrating a mixt of resin, and cereal or flour, adding KClO₃ and sufficient water to make a homogeneous paste. A typical compn is resin 53, wheat flour 27 & nitric acid of 20%B₆ 20%; the nitrated product is mixed with KClO₃ 3 parts & water to make a paste which is molded into blocks
Ref: Leckzinski, FrP 395635 (1908) & CA 4, 2733 (1910)
IASP. Initial Average Sustained Pressure (in rockets) is the average pressure during the first part of the run, after full pressure is reached. IASP is illustrated in the following Fig.

![Image of IASP illustration]

Definition of Values Measured on Film Strips

Ref: F. Bellinger et al., IEC 38, 166 & 167 (1946)

ICC. Interstate Commerce Commission (qv)

Ice Blasting by Means of Explosives. In order to prevent ice jams in rivers and keep them flowing during the thawing period, it is often necessary to help nature by using explosives. Considerable damage can be done to bridges, boats, piers, etc. if the ice jams are not removed at the right time. Ice jams also produce flooding.

An interesting description of breaking up the ice jams in the Raritan river in New Jersey is given by Berliner.

Ref: Berliner, Expls (1953), pp 24-28

ICI. Imperial Chemical Industries Ltd of London, England, has extensive equity holdings in the commercial explosives industry throughout the world. When ICI was formed in 1926 one party to the merger was Nobel Industries Ltd which involved major British explosives interests and particularly the factory set up by Alfred Nobel in 1871 at Ardeer, Stevenston in Scotland.

Nobel’s Explosives Company Ltd (NEC) with its headquarters at Stevenston, is a wholly owned subsidiary of ICI and has been given responsibility for the explosives interests of that company. NEC operates a complete industrial explosives industry in Great Britain and operates plants and services in other countries as well as supplying explosives and accessories to many customers throughout the world.

ICI has a majority shareholding in other companies with major explosives interests of which the most important are Canadian Industries Limited (CIL), ICI Australia Ltd, Indian Explosives Ltd and the Philippines Explosives Corporation. ICI also has a substantial holding in African Explosives and Chemical Industries Ltd, which operates the largest commercial explosives factory in the world at Mudderfontein (See Nobel’s Explosives Company in a future Vol).

ICT. Abbr for Igniter Composition for Tracers (qv)

Ideal and Nonideal Detonations. See Detonations, Ideal and Nonideal in Vol 4, pp D389-R-D390-R

IDENTIFICATION OF AMMUNITION AND EXPLOSIVE DEVICES

The identification of ammunition is a very complex art. It may be accomplished in any one of a number of methods such as examination of physical size, shape, color markings, stencilled markings, metal stampings, and data written on the packaging material.

Historically speaking, early types of ammunition were identified solely by size and shape. As the art of waging war became more complex and sophisticated, logistics support to the field also became more complicated. It was discovered that some means other than size and shape would be necessary for rapid identification of ammunition on the battlefield. According to available information the first distinctive identification of ammunition was developed by the French. This method was the application of paint to cannon projectiles to designate the various types used on board ship. This practice was abandoned prior to 1803 because of the lack of a suitable colored lacquer or paint which would hold its color and remain on the projectiles during shipboard storage at sea.

By 1803, the British had adopted an identification system consisting of various colored letters and numbers applied to the ends of powder barrels to designate the type of propellant, the size of grain, special ingredients used in its manufacture, and method of processing.
By the 1850’s, the British had a standardized color code for identifying smooth bore artillery projectiles. This code consisted of a black body color with white, red and/or yellow bands to designate various loadings. An all white body denoted steel shot and a black body with one white band designated chilled iron shot. About the time of the American Civil War the US Navy adopted a similar code for the rapid identification of projectiles aboard ship. Black, white, red, yellow and blue were used to permit rapid identification of different projectile loadings or functional features. In this same time period, in both the American and British service, metal stampings were applied to the exterior surface of projectile bodies as proof of acceptance by the inspectors and to identify the manufacturer. With the introduction of studded projectiles in the British Land Service, metal stampings were employed to indicate various arsenal modifications to the projectiles. The painting system employed for smooth bore projectiles was retained and modified to allow for various new types. One example was a black body with one blue band to designate a ring projectile.

With the advent of breech loading artillery, the British retained the same basic black projectile body color with colored bands until new special types of projectiles were introduced. An example of this was a yellow body with a red band on the nose to denote high explosive, “lydite” (Picric Acid) filled projectiles. When plaster kits were assembled to these projectiles a wide brown band was painted on the projectile nose and one narrow red band painted underneath it. In addition to the color markings, metal stamp and stencil markings were also employed for identification purposes.

The US Army did not adopt a standardized color code for artillery projectiles until the turn of the 20th century. At this time, there was one color code system for mobile army ammunition, one for seacoast ammunition and one for navy ammunition. Each color code was distinctively different, because there was no attempt to standardize identification among the three services.

By this time, breech loading weapons had been improved by development of the brass cartridge case, the base of which provided a good surface for placing information. At first, only the weapon bore size and/or the manufacturer’s name was applied to the cartridge case base. Later, as it became necessary to better identify ammunition components because of mishaps or malfunctions in the field, both metal stampings and/or ink stencil marks were applied to every component that had a surface large enough to accept such a mark without detrimental distortion occurring.

By the outbreak of World War I practically every major power in the world had one or more color code systems to identify their projectiles.

By this time the art of pyrotechny was so well advanced that many different types of signals were being employed. Because of the vast number of different signaling devices developed, various identification systems were developed by the major powers, which consisted of colors, raised dots and milled case edges and stencilling. In addition, when special purpose small arms cartridges were developed, some means of identification had to be developed. Colored bullet tips, partially or completely blackened cartridge cases and metal stamp base markings were employed for small arms ammunition identification.

Aircraft bombs were introduced in World War I, and soon, various special purpose types were developed. Therefore, a color code was developed. Again each service (land and sea) of the major powers developed their own identification system consisting of colored bodies, colored bands, metal stampings and ink stencil markings.

In the same time period, specialized fuzes for both artillery projectiles and bombs were developed. To designate differences in delay or weapon to be used with, artillery fuzes were color coded for rapid identification in the field.

In the early 1960’s, a concerted effort was undertaken by the NATO countries for the establishment of a standardized color code for the identification of ammunition of all types except small arms ammunition below 20mm in caliber. The following color code was adopted and is being used by most NATO countries. The colors adopted and their significance are as follows:

<table>
<thead>
<tr>
<th>Color</th>
<th>Significance</th>
</tr>
</thead>
</table>
| Yellow| 1) HE ammunition  
2) Presence of explosive  
a. sufficient for HE function  
b. particularly hazardous to user |
Color | Significance
--- | ---
Brown | 1) Rocket motors  
2) Presence of explosives  
a. sufficient for low-explosive function  
b. particularly hazardous to user
Gray | Ammunition containing toxic or irritant agents. Presence of high or low explosive components is indicated by yellow or brown bands
Gray with red band or bands | Ammunition containing irritant agents
Gray with dark green band or bands | Ammunition containing toxic agents
Black | Armor defeating ammunition
Silver or Aluminum | Countermeasure ammunition such as radar echo leaflets
Light Green | Smoke or marker ammunition. Presence of high or low explosive indicated by yellow or brown bands
Light Red | Incendiary ammunition or presence of highly flammable material designed to produce damage by fire
White | Illuminating ammunition or colored lights
Light Blue | Practice ammunition. Presence of high or low explosive components indicated by yellow or brown band
Orange | Ammunition used for tracking and recovery in tests or in training operations
Bronze | Dummy and training ammunition

OP 4 | Ammunition Instructions for the Naval Service May 1943
TM 9-1900 | Ammunition General, 1941  
Ammunition General, 1942  
Ammunition General, 1945  
Ammunition General, 1956
V 650 | Ammunition register för Armén  
Beställes hos KATF/TFB, Stockholm 80, 1960
TM 9-1901 | Artillery Ammunition, 1944  
Artillery Ammunition, 1950  
Artillery Ammunition, 1967
TM 9-1980 | Bombs for Aircraft, 1942  
Bombs for Aircraft, 1944  
Bombs for Aircraft, 1950
TM 9-1985-1 | British Explosive Ordnance, 1953
Supplement to Intelligence Bulletin No 85 | British Navy Projectiles and Fuzes, June 1945
Bomb & Hand Grenades by Bertram Smith, E.P. Dutton & Co NY 1918
Catalogue of Enemy Ordnance Materiel Vols 1 & 2, US Army Ordnance Depot 1 March 1945
Catalog of Cartridge Manufacturers and Base Markings by B.D. Munhall & H.P. White 1944
TM 5-280 | Foreign Mine Warfare Equipment, April 1965
TM 5-280 | Foreign Mine Warfare Equipment, July 1971
French Grenades and Grenade Fuzes Ministry of War, 15 Dec 1930
TM 9-1985-2 | German Explosive Ordnance, 1953
TM 9-1985-3 | German Explosive Ordnance, 1953
PATR 2510 | German Explosives, 1958
PATR 2145 | Russian Explosives, 1955
H.Dv. 454/6b | German Intelligence Data Sheets on World War I Chemical Ammunition British, French, Italian, Russian, 1918
List of a Few of the Publications Which Contain Additional Information on the Identification of Ammunition and Explosive Devices
OP2216 | Aircraft Bombs, Fuzes & Associated Components
26/Manuals/1473 Ammunition Handbook for the Royal Army Service Corps 1936
Geschosszünder Beschreibungen und Zeichnungen, July 3, 1938–Berlin, 1938
OPNAV 30-3M Handbook of Japanese Explosive Ordnance, 15 August 1945

Kemp D&PS Handbook of German Aircraft Ammunition by Bross & D&PS Aberdeen, Md, 1956

N 3353 Istruzione Provvisoria Sulle Bombe A Mano S.R.C.M. Breda—O.T.O. od 35, Roma 1938

No 647 Istruzione Sulle Munizioni, Parte I, Roma 1936

TM 9-1985-6 Italian & French Explosive Ordnance, 1953

TM 9-1985-4 Japanese Explosive Ordnance, 1953

TM 9-1985-5 Japanese Explosive Ordnance, 1953
L’Artificiere D’Artiglieria, 3rd Edition, 1941

Les Fusées Paris, 1918
Manuel Des Munitions—Imprimerie Nationale, 1932

AOP-2 Method of Application of the NATO Code of Colours for the Identification of Ammunition (except Ammunition of a Calibre Below 20mm). Prepared by the Department of National Defence, Canada Ottawa 1966

TM 9-1981 Military Pyrotechnics, 1943

TM 9-1981 Military Pyrotechnics, 1951

MIL-STD-709 Military Standard Ammunition Color Coding, 27 June 1960

MIL-STD-709B Military Standard Ammunition Color Coding, 13 October 1972

R. Moh C.3 Normas para el pintado y marcado de la munición de combate, ejercicio e instrucción, Buenos Aires 1959

Bulletins
216-R2 & 287 Notes on Ammunition by Maj Wm. C. Foote, C.A. Coast Artillery School, Fort Monroe, Va, Oct 31, 1918

SS 420 Notes on German Shells (second edition) General Staff (Intelligence) General Headquarters, 1 May 1918

D 460/9+ Ringbuck für Zünder, Band II, 5 June 1941

D 460/8+ Ringbuck für Zünder, Band I, 5 June 1941

Ordnance Magazine, Various Editions, American Ordnance Association, Washington, DC

TM 9-1950 Rockets, 1958


The Bombardier & Pocket Gunner by Ralph Willett Adye London 1803


Various German drawings of projectiles and fuzes 1939–1945

C.W.S. Intelligence Summary No 12

Markings of Japanese Chemical Warfare Munitions

Written by John F.W. Pflueger

Iditol. A synthetic resin obtained by the condensation of phenol & formaldehyde. It is used as a binder in many Russian pyrotechnics
Ref: Gorst (1957), 162

Iverler (or Jewler) Explosives. Various Sprengel-type explosive mixtures proposed in 1897-1900 by iverler of Russia, such as Prometéus. Other explosives of this class consisted of liquids such as nitrobenzene, methanol etc absorbed on potassium chlorate mixed with Portland cement and/or copper oxide etc
Ref: Daniel (1902), 328

Igdonit. A Russian explosive similar to ANFO. It consists of a mixture of AN and S to 8% fuel oil. It is usually mixed just prior to use. Max deton velocity is only 2000 m/sec for mixes with S to 6% oil (Ref)
See IGNADITY in Vol 5, pp D1744-R to D1746-L

Igelit. Trade name for a polyvinyl plastic used as a binder in some Ger pyro compns, such as listed in PATR 2510 (1958), p Ger 154-R

Igloo Magazines. This type of magazine is now standard for storing ammunition of the US Armed Forces because it permits considerable saving in space over other types of magazines. An Igloo is a concrete building of semi-cylindrical shape with the axis on the ground and with earth covering the arched roof and piled up against the sides. This earth acts as a barricade and also serves to conceal the Igloo as much as possible from view.

The usual sizes of igloos are: 26'6" x 60'8" and 13' to the highest part of the roof, or 36' x 62' and 11'6" to the highest part of the roof.

Usually igloos are built in groups of seven, so arranged that one is in the center and six others around it as on the corners of a hexagon (Ref 1). Each igloo holds 143,000 lbs of explosives, making a total for the group of one million pounds. With this arrangement, a 7-igloo group occupies a circle of 1000 ft diameter with about 1,920 ft between the nearest buildings of adjacent igloo groups, and occupies only about 33% of the space required by older types of magazines for the same amount of explosives.

According to Hall (Ref 2), the usual 500 feet, which has been prescribed as intermagazine distance for unbarricaded surface magazines containing 250,000 lbs of explosives, may be reduced to as low as 185 feet in the case of igloos or underground magazines.


Ignis Volatilis. (Flying Fire) (Feu volant in French). A kind of Greek fire used militarily, somewhat like an incendiary missile. Marco Greco in his 9th to 10th century book entitled: “Liber Ignium ad Comburendos Hostes” gives the composition as: sulfur 1 part, charcoal (of linden, or willow trees) 2 parts, saltpeter (crude) 6 parts.

Ref: Daniel (1902), 382

Ignitacord. Thermalite Ignitacord is the Dupont Inc name for a device for igniting safety fuse. It is cordlike in appearance and burns progressively along its length with an external flame at the zone of burning. The flame is short and quite hot, and offers a means of lighting a series of safety fuses in the desired rotation with the lighting time at the face being no greater than that necessary to light one fuse.

The product is available in two burning speeds: namely, Type A Cord burning at the approximate rate of 8 to 10 seconds per foot, and Type B Cord burning at approximately 16 to 20 seconds per foot. Type A Cord is intended for use in staking rounds where the blasts may contain a large number of holes or where the holes are normally collared from 2 to 4 feet apart. Type B Cord is better suited for use in heading or development rounds where the holes may be collared much closer together.

Both types of cord are clearly marked at one foot intervals so that the fuses may be easily spaced to insure positive rotation. Metal connectors are available for connecting safety fuse to the cord. These connectors are crimped to the safety fuse at the same time that the caps are crimped, and are so constructed that they protect the end of the safety fuse from moisture. They contain a small pressed charge of an ignition compound which lights the fuse when the cord burns through the connector.

Ignitacord is subject to ignition by open flame, sparks, friction or a sharp blow. The connectors can be ignited by heavy impact, as for example, from falling rocks. Ignitacord must not be used as a substitute for safety fuse, or in any place where an open flame is prohibited.

Ref: Blasters’ Handbook (1958), pp 100 & 225-27

IGNITERS

Igniter. Any device serving to ignite either explosives, propellants or pyrotechnic compo-
Igniter Bag or Pad. Separate-loaded ammunition used in cannon of larger caliber, consists of a propellant charge loaded in cotton or rayon (formerly silk) bags, which are loaded directly behind the projectile in the breach of the gun. At the base of the bags, a flat pad in the shape of a disc (igniter pad or bag) made of closely-woven cloth and filled with Black Powder, is sewn or laced. The charge of this bag is ignited by means of a primer located in the breach behind the powder and then the flame is transmitted to the propellant charge (smokeless powder).

Sometimes, in addition to the igniter bag, a charge of Black Powder running through the propellant in the form of a "core" is also used. 


Igniter Bag (Used in Rockets). A flat bag filled with Black Powder and placed near the propellant about the middle of the rocket motor. This serves as an auxiliary igniter to the main one (see Rocket Igniter), which is placed in the throat of the nozzle. For more information, see Ohart, (1946), p 334 or books on rockets

Igniter Composition. See Ignition Composition

Igniter Cord. Called Pyrocore and developed by the duPont Co, it consists of a small continuous metal tubing contg a detonating (HE) ignition core. The core is flame sensitive and promotes ignition at the speed of detonation. Upon functioning, the "Pyrocore" produces high-velocity hot metal particles capable of igniting a variety of igniter and propellant compositions. One use of this device is in combustible cartridge ammunition (see Vol 2, p C79, Ref 25), where instantaneous ignition is required

Ref: duPont Co's advertisement in Ordn 50, No 276, 583 (1966)

Igniter Cord. Extruded incendiary cords of thermoplastic consistency are prep'd by combining powd oxidizing and reducing agents in a plastic binder. Thus, 33 parts plasticized nitrocellulose (I) (consisting of high-N, low-viscosity nitrocellulose 50, di-Bu phthalate 48, Ph2NH 2, and citric acid 2 parts) was heated in a Werner-Pfeiderer mixer with 66 parts Si (specific surface 38,000 sq cm/cc) and 1 part N2N2-diethylcarbanilide (II). Another mixt contg 1 14, Pb3O4 60.6, KClO4 24.7, and 1 0.7 parts was prep'd separately. The 2 mixts were rolled into sheets, diced to 3/16-in cubes, and fed to a screw extruder in a 1:3 ratio. After extrusion at 100° to a diam of 0.065 in, the threads were coated with a 0.004-in polyethylene film, giving a product with a burning speed of 36 sec/yard. I can be replaced by a 60:40 mixt of polystyrene (mol wt 80,000) with di-Bu phthalate or a 40:60 mixt of poly(methyl methacrylate) with di-Bu phthalate

Ref: K. J. Brimley, BritP 787411 (1956) & CA 82, 7704 (1958)

Igniter Core. See Igniter Bag

Igniters, Delay, Electric. Devices used for igniting Black Powder charges. They consist of copper, other metal or plastic tubes provided with an ignition mixture (which can also be Black Powder) in which is embedded a piece of resistance wire (bridge), with two leads attached, similar to electric blasting caps. Pieces of safety fuse of various lengths, to provide different delays, may be attached to the closed end of tube, and the loose ends of the fuses are then dipped in wax for waterproofing.

When ready for firing, a small section of the fuse is cut away at the waxed end (to expose a clean surface) and imbedded in the main Black Powder charge to be used in the blast. When an electric current is passed through the bridge wire, the wire is heated to incandescence and ignites the flash powder in the igniter. The ignited powder burns with sufficient intensity to rupture the Cu tube, igniting the fuse (attached to it), which communicates the fire to the main charge of Black Powder. See also Sect 1, Part A in Vol 4, pp D733-34
Igniters, Electric. Devices intended for firing explosives of the Black Powder type, primarily in commercial blasting. Their construction is similar to that of electric blasting caps except that the capsules (tubes, shells) can be made of wood, paper, cardboard, plastic, etc instead of metal, such as Al, Cu, gilding metal or German silver. As a filler, these caps usually contained an igniting compound, such as mealed powder mixed with gumcotton, or other flash mixtures, packed around the bridge wire. See also description of Squibs in this section
Ref: Marshall 2, (1917), 545


Igniters for Fuses. Devices for igniting a safety fuse, which then explodes a blasting cap which in turn explodes the main charge, may be divided into the ones intended for use in blasting operations where an open flame or a spark is of no danger (such as quarries, non-gaseous mines, etc) and those for use in gassy or dusty-coal mines, where no open flame or spark should be allowed. The last named igniters may be called safety igniters (allumeurs de sûreté, in French)
Open Flame Igniters

Amadou or tinder igniter is the oldest of these devices, in which the heat produced by a glowing tinder was utilized to ignite the core of a fuse. The tinder itself was ignited by means of sparks produced on striking flint with steel

Wick igniter utilized the heat produced by a slowly burning special wick, which was ignited by flint-steel, as above

Ordinary matches are most frequently used at the present time, but they are not satisfactory in wind or rain. Neither is it convenient to use them for lighting a number of fuses trimmed to fire in rotation because these must be lighted quickly, surely and in a specified order (Ref 4)

In order to get better results when using ordinary matches, the free end of the fuse is slit open (about 1/4", lengthwise) and the head of a match is inserted in this slit so that it touches the core of the fuse. After this, the head is lighted by another ordinary match; by this means the fuse is ignited more readily than by a single match

Safety fuse match lighter consists of a paper tube 1¼" long and of the same diameter as the fuse. One end of the tube is closed and coated with the same composition as Swedish (safety) matches. The open end of the tube is slipped over the freshly cut end of a fuse so that the match composition is against the end of the fuse. By striking the match end of the tube against the coated side of a safety-match box, the match is ignited and this in turn ignites the fuse. The match end of the tube can also be ignited with an ordinary match (Ref 4)

The lead spitter, pull-wire & hot wire fuse lighters are described in Sect 1, Part A of Vol 4, p D733

Closed Flame or Safety Igniters

A device invented by Lagot (1881), consisting of a tube filled with pieces of charcoal impregnated with a compound (such as nitrate) which permits the charcoal to burn in absence of air and without producing flame. The tube has an opening just large enough for the insertion of the end of a fuse to be ignited (Ref 1)

Master fuse lighter consists of a waterproof fiber shell covered with flexible rubber sheeting. A charge of ignition composition is placed in the base of the shell and a freshly cut “pilot fuse” and six fuses leading to boreholes are inserted through the rubber covering. The pilot fuse is ignited from the outside and it ignites the ignition composition inside the shell which then simultaneously ignites the other fuses

The use of this lighter insures better results in rotation firing and allows more time for the blaster to retire before the detonation of the first charge, since he has only the pilot fuse to light. The use of this type of fuse has the disadvantage of requiring large amounts of fuse (Ref 4)

Heath and Frost proposed a modification of the miner’s safety lamp so that it was possible to use it to heat a small piece of iron to incandescence which could then be touched to the open end of a fuse. This operation was done in such a manner that no flame or spark was produced outside the enclosure of the lamp (Ref 1)

Bickford’s Collier Safety Lighter, invented in 1889, consists of a tube (tinned-iron or steel) closed at one end and containing a mixture of KClO₃ and sugar, pressed into a pellet not
exceeding 100mg in weight. This is placed in close contact with a hermetically sealed glass capsule containing concd HzSO4. A length of specially prepd and waterproofed safety fuse is fitted into the open end of the tube, the joint being cemented with tape to form a closed chamber. By squeezing the tube with pliers, the glass ampule is broken. This brings the HzSO4 into contact with KClO3 + sugar, ignites the mixture which in turn ignites the fuse (Refs 1 & 2)

This device may be considered as a modification and improvement of the devices proposed by Roth and by Zschoöke. In the latter, a glass capsule containing HzSO4 was wrapped in a piece of cloth impregnated with a concentrated solution of KClO3 + sugar (Ref 1)

Mortier used a small piece of metallic sodium coming in contact with a drop of water to ignite fuses (Ref 1)

Muller, about 1880, invented a device which was essentially as follows: the cut end of a fuse was inserted into a tightly-fitting priming cap, which in turn was placed inside a special, hermetically sealed pistol. The cap was fired when struck by the pistol firing pin and this ignited the fuse. The flame produced by the priming cap remained inside the pistol (Ref 1)

A similar device was proposed by Hohendahl in 1896

The Miner's safety match was patented by Pope and manufactured during the latter part of the 19th century by William Bennett Sons & Co of Camborne, England. It consisted of a slightly tapered metallic tube, the narrow end of which was closed. A small quantity of priming composition was placed inside the tube and then a tightly-fitting fuse was inserted so that the open end of the fuse touched the priming composition. By rubbing the closed end of the tube against a surface covered with a friction compound, sufficient heat was produced to ignite the priming composition inside the tube and that, in turn, ignited the fuse (Ref 1)

A device invented by Meinhard consisted of a metallic tube, 20cm in length, closed at one end. Inside the tube was placed a small quantity of priming explosive and on top of it the fuse was inserted so that it fitted tightly

On striking the end containing the priming mixture with a hammer, or other object, ignition of the priming mixture as well as of the fuse was achieved (Ref 1)

Hargreaves patented a device in which the fuse was ignited by friction (Ref 3)

Most of the devices described above are no longer used and are only of historical interest

Today, in gassy mines in the US, the explosive cartridges are detonated directly with electric blasting caps (Refs 4 & 5), without the use of safety fuse


Igniter Pad. See Igniter Bag

Igniters for Propellants. Devices acting as intermediates between priming charges and propellant charges and serving to produce a flame larger than the one produced by a priming cap

In small arms cartridges, where the propellant grains are of small dimensions and the charge itself is small, proper ignition may be achieved by a primer cap without introducing any auxiliary device, such as an igniter

On the other hand, in guns, where the propellant charge and the grain size of the powder are both large, it is impossible to achieve proper ignition unless either the size of the primer is increased considerably or an intermediate charge of easily ignitable material is introduced. Of these two methods, the former is undesirable because the use of large primers is not only dangerous but is also much more expensive. Thus it is the latter method that is commonly employed, provided a proper substance is used in the igniters. The usual propellant igniter is Black Powder because it is easier to ignite than smokeless powder and because it produces a large flame of uniform intensity independent of the pressure developed in the barrel of the gun

Igniters used for separate loaded ammunition of larger caliber cannons are called "Igniter Bags" and were described above. Igniters used in fixed ammunition of cannons of medium and small caliber consist of a long, narrow, per-
forated tube filled with grains of Black Powder, closed at one end and attached to the percussion primer at the other end. The tube penetrates the propellant charge to as much as 3/4 the length of the cartridge. In this arrangement, the percussion primer is struck by a blow of a firing pin. This sets the priming charge afire, which then ignites the Black Powder in the perforated tube. The flame of burning Black Powder shoots through the perforations and ignites large areas of the propellant powder, insuring proper propulsive action. When the igniter and percussion primer are assembled as one unit, the ensemble is called a combination primer, or primer-igniter. The function & location of primer-igniters in artillery ammunition is described in Sec 3, Part D in Vol 4, pp D777-803.

Igniter, Pyrotechnics. Any device used for igniting pyrotechnic compositions, such as illuminating, smoke, tracer, flare, photoflash mixtures, etc may be called “pyrotechnic igniter” and the compositions used in such igniters are called “igniting mixtures” or “igniting charges”

Igniter, Rocket. There are generally two igniters in a rocket motor, the main and auxiliary. The main igniter consists of a primer and a charge of Black Powder, or other igniting mixture, contained in a plastic body (generally ethyl cellulose) and placed in the throat of the nozzle of the motor. The primer is fired by the heat generated in a wire by means of electric current, thus igniting the Black Powder of the igniter, which in turn ignites the propelling charge, usually consisting of grains of double-base powder (Ref 1, p 333-34).

In addition to the main igniter, there is an auxiliary one, described under Igniter Bag for Rockets.

Igniter Squibs. The squibs are essentially electric detonators except that they produce a flash & not an explosion. The commercial type squib, which is also used for military purposes, consists of an aluminum shell 7/8 to 1-1/2 long with the flash charge in the bottom of the closed end. It contains an electrical firing element in the form of a bridge wire and two lead wires sealed in the other end with waterproofing compound or a deformable plug that is crimped into the open end of the shell. About 0.5 ampere is required to fire the squib; when the current is passed through the bridge wire, it is heated to incandescence and ignites the flash powder. The ignited powder burns with sufficient intensity to rupture the Al shell and the flash thus produced ignites the next element in the train, which can be either a Black-Powder charge, a slow-burning fuse or a flash-initiated detonator (Refs 1 & 2).

Electric squibs will ignite pelleted Black Powder which is too wet to be fired with a safety fuse. In blasting operations they permit tight tamping of the bore hole, the ignition of the powder charge at any desired point, the firing of a number of shots at the same time and definite control of the time of firing the charge. Electric squibs are considered the safest and most effective means of firing black blasting or pellet powder (Ref 3).

Electric squibs may also be of the delay type, varying in length from 3-1/8 to 5-1/8, depending on the delay time (Ref 1).


Igniter Tests. Assembled electric igniters intended for military use are usually subjected to the following tests, after selecting 10 samples for each test:

1) Material and Workmanship. All materials shall be of high grade and all parts free from chips, dirt, grease and other foreign matter. The seal between the two wires and the cup, tube, etc shall be continuous. If the seal is questionable, it is subjected to a pneumatic test by introducing an internal air pressure of 4 psi, held for 3 seconds.

2) Continuity of Circuit and Resistance. is to be determined by means of a Wheatstone bridge.

3) Waterproofness (Water Submersion Test). Ten igniters are kept under water for 18 to 48 hours (depending on specification requirements) at 70-75°F and then fired immediately.

4) Current and Functioning. The terminals of each of 10 igniters shall be connected, one at a time, in series with an ammeter, a rheostat and a source of direct current (new dry cell). Move the rheostat until the igniter is fired and note the current (amperes) required for it. Note also if the igniters function properly on firing; the metallic (or plastic) shell should be ruptured.
but the sealing plug should not be blown out:

**Note:** Sometimes, requirements call for conditioning the igniters at high temperatures, say 150°, for 1 hour before firing

5) **Ignition Delay.** If the igniter is of the delay type, it is necessary to determine the time elapsed from the start of current flow to the rupture of the igniter case. For instance, the M12 electric igniter has an ignition delay not exceeding 100 milliseconds (Ref 2)

6) **Detailed Examination of Parts.** Sometimes it is necessary to take the igniter apart and submit the shell (metal or plastic), the wires, sealing compound and igniter compound to a physical and chemical examination. The disassembly operation should be carried out behind a barricade, preferably by remote control. The tests are usually different for each type of igniter and are described in the corresponding specifications


**Igniters for Tracers and their Compositions**

Igniter compositions for Tracers are designed to be nearly gasless and to have low ignition temps. Some of these compositions are described in Vol 4, Sect 3, Part C, p D744. They are usually pressed into a cavity at the rear portion of a projectile after first loading most of this cavity with a tracer composition. See Fig 30 in Vol 4, p D744.

A tracer composition difficult to ignite consists of Sr(NO₃)₂ 65, Mg (powder) 30 and binder 5.

This composition has to be ignited by a compound which develops an extremely high temperature, such as BaO₂ 78, manganese powder 20 and calcium resinate 2%. Unfortunately, this composition develops a very bright light which might blind the gunners during night firing. In order to avoid this possibility, an additional charge, consisting of BaO₂ 78, manganese powder 20 and calcium resinate 2, is placed on top of the above composition. This mixture is easily ignited, but it does not burn with as hot or bright a flame as the magnesium mixture. In this kind of loading, the manganese mixture is called the "igniter mixture," while the magnesium mixture is called the "Subigniter mixture" or "Subigniter.

During WWI, in the so-called "perforated luminous ball cartridges," fired from airplane machine guns, the Germans used two charges, besides the propelling charge: an illuminating charge consisting of Mg 64.7, Sr(NO₃)₂ 19.3, Ca(OH)₂ 11.7 and resinous matter 4.3%; and above it, an igniter charge consisting of K₂MnO₄ 54.9 and iron powder 45.1% (Ref 3, v 1, p 192).

During WWII, the Germans developed two types of igniter mixtures:

a) Barium peroxide + metallic magnesium + binder;

b) Barium nitrate + metallic magnesium + binder (some compositions also contained Styplonic Acid)

The barium peroxide mixtures were similar to the American "Igniter K"

Examples of type a):

1) BaO₂ 78.4, Mg 19.1, binder 2.5%. A quantity equal to 0.67g was used in each 20mm HE self-destroying projectile

2) BaO₂ 75.0, Mg 22.0, binder 3.0%. A quantity equal to 0.30g was used for each 37mm APHV shell

3) BaO₂ 69.7, BaCO₃ 13.5, Mg 14.2, binder 2.6%. A quantity equal to 3.0g was used for each 88mm AP shell

Examples of type b):

1) Ba(NO₃)₂ 32.7, Mg 36.7, Na Picrate 11.9, binder 18.7%. A quantity equal to 0.30g was used for each 20mm APHV shell

2) Ba(NO₃)₂ 41.7, Mg 30.0, Sr Picrate 12.8, binder 15.5%. A quantity equal to 0.20g was used for each 50mm APHV short-case shell

Heiskell claims a tracer unit for a projectile, suitable for improving daylight visibility & having a non-flash slow-burning compn which delays ignition of the tracer compn for a predetermined time (Ref 5).

One of the Italian igniter compositions contained BaO₂ 75.0, Mg 23.0, binder 2.0% and 0.9g was used for each 47mm AP Round Nose Shell (Ref 2).

One of the Russian igniter compounds contained BaO₂ 60.0, Mg 13.6, binder (resin) 26.4%; 0.65g was used for each 76mm APHE shell (Ref 2).

*Refs:* 1) Faber (1919), vols 1 & 2 2) D. Hart, PATR 1335, 6 (Sept 1943) 3) "Data on American & Foreign Explosives," PB 11544 (1944) 4) Ohart (1946), pp 61-63 & 77 5) R.H. Heiskell, USP 2899291 & CA 53, 19391 (1958)
IGNITION

Ignition is the act of kindling or setting on fire any combustible substance. High explosives as well as Black Powder, Nitrocellulose, smokeless powder and pyrotechnic compositions can undergo ignition (burn) without detonating.

The consequences of ignition, ie burning or combustion, are described under "Burning & Combustion" in Vol 2, pp B343-B357, under "Combustion" in Vol 3, pp C425-C433 and "Detonation-Development from Burning or Deflagration" in Vol 4, pp D245-D252. For further discussion of the ignition of explosives, the reader is referred to High Explosives, Hot Spots, Pyrotechnic Propellants & Initiation Mechanisms in this Vol and to Thermal Explosions in Vol 9. Some discussion of Ignition, Igniters etc is also presented under Detonators, Igniters & Primers in Vol 4, pp D757-D807.

For convenience, we will divide the subject of ignition into the following sections which treat different aspects of the general problem: 1) Elementary Theory of Ignition 2) Ignition Cartridge 3) Ignition Compositions for use with Pyrotechnics 4) Ignition of Firedamp & Coal Dust 5) Ignition—Spontaneous 6) Ignition, Test for 8) Ignition Theory of Explosives

1) Elementary Theory of Ignition

Introduction

A highly simplified diagrammatic representation of ignition is shown in the following sketch:

Since heat loss varies essentially linearly with temperature difference, it predominates at small temp differences. Above some temp $T_1$, heat generation which varies exponentially with temp difference becomes larger than heat loss. In this sketch $T_1$ can be considered to be the ignition temperature of the system, although, more accurately, $T_1 = \text{ign temp} - \text{initial temp}$.

This simple model represents most of the situations of interest for explosives & propellants. For dust and gas explosions, however, heat generation, ie reaction rate, may be controlled by the availability of oxygen at the reaction site. Such reactions are said to be diffusion-controlled.

Combustible systems can be ignited by sparks, hot wires, flames, hot particles, heated surfaces and many other sources. Ignition by sparks & hot wires are of particular interest in connection with explosives and we will therefore consider them in more detail below

Ignition by Sparks. (See also Electricity, Extraneous and Hazards Associated with it in Vol 5, pp E36-E46). In spark ignition the duration of stimulus energy is very short. It is possible to pass small electric sparks through an explosive gas or explosive dust cloud without producing ignition. When the spark energy is increased, a threshold energy is eventually obtained at which the spark becomes incendiary, in the sense that a combustion wave propagates from the spark through the volume of gas. This minimum ignition energy is a function of experimental variables such as the parameters of the explosive gas and the configuration of the spark gap.

A versatile apparatus for determining minimum energies for electric-spark ignition of gases and dust clouds, developed by Blanc et al (Ref 1), is shown below:

Many data have been obtained with this & similar apparatus (see Ref 7).

Lewis & Von Elbe (Ref 7), in discussing the theoretical aspects of spark ignition, start with the supposition that a spark instantly establishes a small volume of gas within which the temperature is very high. The temperature within the spark volume decreases rapidly due to the flow of heat to the ambient unburned gas. In the adjacent layer of ambient gas the temperature rises and induces chemical reaction, so that a combustion wave is formed which propagates...
outward with approximately spherical symmetry. Whether the wave develops to the steady state depends on the size to which the inflated volume has grown at the time when the temperature at the origin has decreased to the order of the normal flame temperature. In order to continue to propagate, the flame should at that time have grown to at least such size that the temperature gradient between the burned gas in the core and the outer unburned gas has approximately the same slope as the temperature gradient in the steady-state wave. If the size is too small, viz, if the gradient is too steep, the rate of heat liberation within the inner more or less spherical zone of chemical reaction is insufficient to compensate for the rate of heat loss to the outer zone of preheated unburned gas. In that case the loss of heat to the unburned gas continuously exceeds the gain of heat by chemical reaction, so that the temperature decreases throughout the reacting volume, the reaction gradually ceases, and the combustion wave becomes extinct after only a small amount of gas around the original spark has burned.

Quantitative development of the above model of the ignition process is overwhelmingly complicated. Lewis & Von Elbe therefore chose to attempt correlation of experimental minimum ignition energies with some energy functions computed from minimal flame parameters.

One such energy value, denoted by $H'$, is obtained by computing the sensible heat in a sphere of diameter $d$, assuming that the content of the sphere is completely burned so that its temperature is $T_b$ and its density is $\rho_b$. The heat contained in the sphere in excess of the heat at the initial temperature $T_u$ is obtained from the equation

$$H' = \frac{\pi}{6} d^3 c_p \rho_b (T_b - T_u)$$

Another energy value which they use is denoted by $H''$ and is given by the equation

$$H'' = \pi d^2 \frac{\rho_b}{\rho_u} (T_b - T_u)$$

Further development of their model is summarized in their diagram of a minimal flame shown below.

Note that the temperature profile differs from the total heat profile because the heat per unit volume depends not only on the local temperature but also on the local density of the flame gas. In the preheat zone the profile of conducted heat coincides with the profile of total heat, whereas in the reaction zone the conducted heat gradually drops to zero. The difference between total heat and conducted heat represents the heat gained by chemical reaction. The volume integral of the total heat is approximately $H'$, and the volume integral of the conducted heat is approximately $H''$.

This model is claimed to agree with experiment in that experimentally determined minimum energies are smaller than either $H'$ or $H''$, and generally, but not always, $H' > H''$. For details, the reader is referred to Ref 7.

Ignition of gases resulting in detonation is treated in Vol 4, pp D732-32 under "Detonative Combustion" & "Detonative Ignition of Gases" and also under "Detonation (and Deflagration) in Gases" and "Detonation of Gases" etc; "Development from Burning" etc, pp D360-63.

More recent developments in the study of DDT (Deflagration-to-Detonation Transition) will be described under Transition to Detonation.

Ignition by Hot Wires. Now we turn to a discussion of ignition by hot wires, but we purposely exclude studies on EED's (See Vol 5, pp E63-E68) or initiation of detonation by hot wires (See Initiation, Hot Wire in this Vol). We quote Lewis & Von Elbe:
"Electric sparks are very hot and fast-acting ignition sources. Because the discharge time of an electric spark is very short (of the order of $10^{-9}$ to $10^{-7}$ second), the energy that is imparted to the gas at the end of the discharge period is highly concentrated, so that a very steep temperature profile with a very high temperature at the center is established. In this initial stage of flame development the chemical heat liberation is insufficient to maintain such steep temperature profile, so that the profile broadens and the temperature at the center decreases. Within a period of time which depends on the physical and chemical properties of the gas, and provided that the discharge energy is sufficient, the profile develops to the profile of the minimal flame and thence continues to propagate as a steady-state wave, while the temperature in the center settles down to approximately the value of the flame temperature.

"If the same amount of source energy were delivered by, say, an electric current over a time larger than the time of development of a minimal flame, the temperature at the core would drop below the flame temperature, the heat liberation in the reaction zone would not attain a balance with the outflow of heat into the preheat zone, and the flame would become extinct. On the other hand, if the current flow were continued for a longer period, the temperature profile ultimately would become sufficiently broad, and the temperature in the core sufficiently high, so that heat liberation within the reaction zone overbalances the outflow of heat and ignition occurs.

"We single out three interdependent quantities which characterize the ignition threshold of a slow source such as described. One is the total heating time (the time during which the current flows) which we shall call the critical heating period. Another is the total energy delivered during this time, which we shall call the critical source energy, and which defines the current strength. A third is the temperature $T_c$ in the core at the end of the heating period, which we shall call the critical source temperature. The general form of relation between these three quantities is shown schematically in Fig 168. Thus, the minimum ignition energy, which is denoted of the symbol $E_0$, corresponds to a very short ("zero") value of the critical heating period. The corresponding critical source temperature has a value smaller than the flame temperature $T_f$. As the critical heating period is increased (and the current strength correspondingly decreased) the critical source temperature decreases and the critical source energy increases. In terms of unit increase of critical heating period, these changes are initially large, as indicated in the figure, and gradually become smaller. The temperature curve finally becomes quite flat; this corresponds to the rule, which is inherent in Arrhenius' law, that at low temperatures a few degrees of temperature change produce large changes in the rate of the chemical reaction. In the present system this means that at low values of the critical source temperature a few degrees of temperature change in the reacting core of the heated volume change the rate of heat liberation very greatly. Hence, relatively large changes of critical source energy, viz, critical heating period, which correspond to large changes of the reaction rate at the ignition threshold, produce only small changes of critical source temperature."

Figure 168 is an idealization because it does not take into account the energy required to heat the source, i.e., the wire, to some critical temperature. This critical temperature should be close to that of the flat portion of the lower curve of Fig 168. If one assumes that the critical wire temperature remains constant in any given ignition system, and allows for heat losses by the wire:

$$E = A + Bt$$  \hspace{1cm} (1)

where $A$ is the minimum critical energy for ignition and $B$ is the combined rate of heat loss from the wire to its soldered ends & to the medium in which the wire is imbedded.

![Fig 168. Source energies and critical source temperatures versus critical heating period. Schematic illustration](image)
Figure 1 shows Jones' results (Ref 3) of his study of critical firing energies for electric fuseheads (also known as matchheads). The data in Fig 1 are for an 80/20 LMNR/KClO₃ ignition mixture (LMNR is Lead Mononitroresorcinolate).

**Fig 1. Relation between ignition energy, mean firing current, and critical heating period.**

The energy plot certainly agrees with the requirements of Eq (1). Now it would be expected that both A & B of Eq (1) would increase with wire length. That it is so is shown below (this Fig is also taken from Ref 3)

![Relation between cross-sectional area of wire and G=AJ/l (l=wire length) (Ref 3)](image)

We may therefore write

\[ A = G l \]  

(2)

where \( l \) is the wire length and \( G \) the minimum critical energy per unit length. Further,

\[ B = H + J l \]  

(3)

where \( H \) is the rate of heat loss to the wire ends and \( J \) the rate of heat flow from the wire to the medium per unit length of wire. The corresponding amount of heat absorbed by the medium per unit wire length is \( J t \). This does not represent the total heat absorbed by the medium, but rather the excess over the heat absorbed when the critical energy for ignition is a minimum, i.e. when \( t = 0 \).

It should thus be possible to divide the term \( G \) into two parts—one representing the heat per unit wire length which is imparted to the explosive medium when the critical energy is a minimum, and the other representing the heat content of the wire per unit length. The second part should be equal to the product \((T_c - T_u)c\), where \( T_c \) and \( T_u \) are the critical and the ambient temperature, \( c \) is the heat capacity per unit volume of wire material, and \( a \) is the cross-sectional area of the wire; it should therefore vanish when the wire diameter is made vanishingly small. The first part should not vanish.

**Fig 2. Relation between cross-sectional area of wire and G=AJ/l (l=wire length) (Ref 3)**

with wire diameter. The term \( G \) should therefore be of the form

\[ G = K + L a \]  

(4)

where \( K \) is that part of the minimum critical energy per unit length of wire which is absorbed by the medium, and \( L = (T_c - T_u)c \) is the heat content per unit volume of the wire at the critical temperature. Figure 2 shows values of \( G \) obtained by Jones for wires of various diameters. Each point is the minimum critical energy per unit length of wire for zero critical heating period and for a given wire diameter. It is seen that the plot of the experimental points versus the cross-sectional area yields a curve which is virtually a straight line. Thus, it would seem that \( K \) is determined by the point of intersection of the line with the \( G \) axis, and \( L \) by the slope. This suggestion should be treated with some caution because \( K \), though not vanishing with wire diameter, nevertheless is certainly a function of the diameter. However, the error introduced by considering \( K \) as constant is not large.

The results of Jones (Ref 3) are in accord with the expression

\[ E = (K + L a) + (H + J l) t \]  

(5)

which, at a constant wire length \( l \), is of the same form as Eq (1). In Eq (5) \( K \) & \( J \) should be functions only of the ignition composition and should not change when the wire material is changed. Conversely \( L \) and \( H \) (at least for
similar solder joints) should change with wire material but not with igniter composition. These expectations are in accord with the observations of Jones (Ref 3). His experimental findings are summarized in the tabulation below:

<table>
<thead>
<tr>
<th>Composition</th>
<th>LMNR/chlorate</th>
<th>Copper acetylide</th>
</tr>
</thead>
<tbody>
<tr>
<td>Critical temp, °C</td>
<td>380</td>
<td>440</td>
</tr>
<tr>
<td>K, cal/cm</td>
<td>0.5 x 10⁻³</td>
<td>0.6 x 10⁻³</td>
</tr>
</tbody>
</table>

**Propagation of Ignition.** As mentioned at the beginning of this article, burning, i.e. the propagation of ignition throughout the combustible medium, has been described in various articles of this Encyclopedia. However, since an important burning mechanism, proposed by Eyring et al (Ref 2), has not been described, we present this mechanism below (direct quote from Ref 2)

**Rate Laws of Surface-Burning Reactions**

It is of some interest to find the dependence of extent of reaction on time, for topochemical reactions of the surface-burning type, in which the linear rate of burning is constant.

**Example 1: Sphere uniformly ignited over its surface**

Here the boundary between burned and unburned material is given by

\[
\frac{R}{R_0} = 1 - \frac{t}{\tau}
\]

and the fraction of material reacted is obviously

\[
N = 1 - \left(\frac{R}{R_0}\right)^3
\]

Therefore the rate law in its integrated form is

\[
N = 1 - \left(1 - \frac{t}{\tau}\right)^3
\]

**Example 2: Sphere ignited at its center**

Here the boundary between burned and unburned material is given by

\[
\frac{R}{R_0} = \frac{t}{\tau}
\]

and the fraction of material reacted is obviously

\[
N = \left(\frac{R}{R_0}\right)^3
\]

Therefore, the rate law is

\[
N = \left(\frac{t}{\tau}\right)^3
\]

**Example 3: Sphere ignited at one point on its surface**

Here the boundary between burned and unburned material is given by

\[
\frac{R}{R_0} = 2 \cdot \frac{t}{\tau}
\]

The fraction of material reacted is found by a simple integration to be

\[
N = \frac{1}{2} \left(\frac{R}{R_0}\right)^3 - \frac{3}{16} \left(\frac{R}{R_0}\right)^4
\]

Therefore, the rate law is

\[
N = \left(\frac{t}{\tau}\right)^3 \cdot \left[4 - 3 \cdot \frac{t}{\tau}\right]
\]

The dependence of N on t has been plotted in the following figure for these three examples.

![Burning rate for various models](image)
General Observations on Ignition of Explosives.
In the preceding paragraphs we considered the general subject of ignition and not necessarily the ignition of explosives. Since the primary purpose of this Encyclopedia is the presentation of material relating to explosives, we present below a brief recapitulation of observations on the ignition of explosives.

Primary explosives can be ignited and will burn without detonating if they are essentially unconfined (Refs 4 & 6). Lead Azide appears to be an exception because its ignition results in detonation or at most a very short-lived combustion which almost instantaneously goes over into detonation (Ref 6). However, dextrinated Lead Azide can be made to flash without detonating (Ref 5).

In order to secure proper functioning of an initiating compound, it is important that it can be ignited with comparative ease. Some initiating substances, however, such as Lead Azide are difficult to ignite, and for this reason are either mixed with easily ignitable substances, such as Lead Styphnate etc, or are primed by small quantities (0.2g) of some easily ignitable substances as Lead Styphnate or Lead Mononitrosorosinate.

Secondary explosives will burn without detonating if unconfined or lightly confined. Some secondary explosives, such as PETN, are difficult to ignite and propagate burning only at slightly elevated ambient pressures.

Propellants are designed to burn stably. However, ignition of unconsolidated propellants (high porosity) or large masses of propellant can result in detonation.

Black Powder does not detonate, but under confinement it deflagrates (burns) so rapidly that its external effects are almost those of a detonation (Ref 6).

Written by J. ROTH


2) Ignition Cartridge. Some mortar ammunition, eg, US Army 60 and 81mm mortars, is propelled by the so-called ignition cartridge (which resembles sporting cartridges in form). It is placed in the fin stabilizer tube and additional propellant increments are fastened to the fins by means of special wire holders. Charges in ignition cartridges (14.5 grains for 60mm and 24.0 grains for 81mm mortar) usually consist of small grains of double-base propellant, NC 57.75 ± 1.50% with N = 13.25 ± 0.05%, NG 40.00 ± 1.50, K2SO4 1.50 ± 0.50 & DPhA 0.75 ± 0.10%

Grain dimensions are: diameter 0.035" and thickness 0.00325" for the 60mm mortar, and 0.059" (diam and 0.0100" thick) for the 81mm mortar. The grains are ignited by means of a Black Powder pellet which in turn is ignited by a percussion primer. The flame from the ignition cartridge ignites the propellant increments

Refs: 1) Ohart (1946), pp 55 & 192 2) US Specification MIL-C-20480A (July 1954) (M3 Ignition Cartridge for 81mm Mortar Ammunition)

3) Ignition or Igniting Compositions for Use With Explosives. Any substance or composition intended to ignite (but not detonate or explode) an explosive belongs to this class. This should include not only all the primary ignition compositions, but also any ignition leads which then transmit the flame to other portions of the explosive device. It should be noted that, as there is no strict distinction between priming (which can explode or detonate) and igniting compositions, some of the mixtures given below might belong to either type.

Numerous mixtures have been proposed for ignition of explosives. Of these, the most commonly used are mealed gunpowder, finely grained Black Powder, nitrocellulose mixed with other substances & finely divided smokeless powders mixed with other substances.

For instance, for igniting propellant charges of smokeless powder, grained Black Powder is used, which in turn is ignited by means of a primary ignition composition.

A primary igniting mixture of numerous US detonators consists of lead sulfocyanate 45±1
and potassium chloride 55±1%

The primary igniting mixture in the M19 electric igniter (used for simulating gunfire) contains dinitrodiazophen 40±3, potassium chlorate 58±3 and nitrostarch 2±0.5%. This mixture is ignited by heat produced by incandescent resistance wire and in turn ignites the charge of Black Powder placed next to it.

The following examples of igniting mixtures are taken from Chemical Abstracts and arranged chronologically. Some of these compositions may be also classified as priming compositions.

L. Fisher, 27, 1758 (1933); Igniter charge for blasting caps consisting of K₃Fe(CN)₆ 20-40 with an oxidizer such as KClO₃ 10 to 30 and a nitrated carbohydrate such as NC 70-30; USP 1890112 (1932)

H. A. Lewis, 27, 4931 (1933); Ignition composition used as a top (primary ignition) charge in blasting caps, comprising Pb thiocyanate, an oxidizer such as KClO₃ and a solid low-ignition-point fuel, such as pyro smokeless powder. This mixture ignited a primary charge of LA; USP 1918920 (1933)

H. A. Lewis, 28, 4235 and 5242 (1934); Ignition composition for blasting caps containing: K(SCN)₂ 30-70, KClO₃ 30-60 and sulfur (as an accelerator) 1 to 20%; USP 1964825 (1934) & CanPat 340569 (1934)

C. P. Spaeth, 29, 1252 (1935); Ignition charges for electric blasting caps containing Tetramethylene diperoxide either alone or with various admixtures; USP 1984846 (1934)

C. P. Spaeth, 29, 5660 (1935); Ignition composition for the top charge in blasting caps, comprised of NC impregnated with a liquid explosive nitric ester, such as NG and KClO₃; USP 2007223 (1935)

O. A. Pickett, 29, 6062 (1935); A thermally fired igniter charge for detonator caps contained zirconium mixed with KClO₃ and gum arabic or nitrostarch; USP 2008366 (1935)

C. E. Sosson, 29, 7078 (1935); An ignition composition for use in low-tension electric fuse heads, comprised of a mixture of finely divided metallic zirconium 35 to 39% and an easily ignitable salt of nitrophenol or its derivatives, such as Pb-2-nitroresorcinol, 65 to 61. The mixture may be used either in loose form around the electric bridge or may be agglomerated with NC gel; BritP 428872 (1935)

E. Jones, 29, 8336 (1935); A safety igniter suitable for use with burning (nondetonating) explosives, comprised of an electric fuse head (Zr + Pb-2-nitroresorcinol) and a combustible material, such as a mixture of finely divided Pb-Si + Pb₂O₄, or Ca silicide + PbCrO₄, or Zn + KMnO₄ or Fe + KMnO₄. Both fuse heads as well as the combustible mixtures are of the kind producing very little or no gas. They are sealed inside a metal container which does not burst when the mixture is ignited, but is heated only on a portion of the surface to a temperature sufficient to ignite the blasting charge and not sufficient to ignite the firedamp; BritP 430750 (1935)

C. E. Sosson, 30, 1564 (1936); Ignition composition suitable for use in delay detonators: zirconium 50 to 95 used with basic Lead-2-nitroresorcinolate, O₂N₃N₃C₆H₃(OH)₂·COO(PbOH) (or other lead salts of nitrophenols) 50 to 5; USP 2027208 (1936)

E. Jones, 30, 4324 (1936); Safety igniter for explosives, consisting of a Cu tube 33 mm long, 6 mm in diameter and 0.2 mm thick containing a charge of 0.75g of finely divided Zn and 40% KMnO₄, loaded at a pressure of 60 psi. A fuse head of metallic Zr and Lead Stypnate (LSI) is placed close to the composition; CanP 357600 (1936)

J. Taylor & W. Young, 30, 4324 (1936); A complicated device suitable for igniting safety blasting explosives without igniting firedamp; CanP 357532 (1935)

A. Stettbacher, 30, 2387 (1936) and Nitrocellulose, 6, 202 (1935); Chemical and mechanical delay igniters. A review, mainly of chemical delay incendiary bombs

L. A. Burrows, 31, 6467 (1937); An ignition composition in an electric blasting initiator, comprising at least one of the compounds selected from the group consisting of the silver and mercury derivatives of chlorinated azodicarbonamide; USP 2086533 (1937)

L. Burrows, W. Filbert & E. E. Reid, 32, 2357 (1938); Ignition charge suitable for electric blasting caps and containing bis-Nethyl Lead Stypnate, C₂H₅N₂O₃ [OPb(C₂H₅)₂]; USP 2105635 (1938)

B. Zielinski, 32, 3964 (1938); Ignition mixtures suitable for percussion caps, primers, etc and consisting of LSt 20-50, Ba or Pb nitrate 30-50,
Sb₂S₃ 5 to 30, a friction producing agent such as glass up to 20% and a sensitizer such as 2,4,6-Trinitro-1,3,5-triazido-benzene 0.25 to 10%; USP 2111719 (1938)

W. F. Filbert & W. E. Lawson, 32, 5630 (1938); An ignition composition suitable for use in electric blasting initiators, contains the nitrated product of Pb-diphenylolpropane mixed with Zr, KCIO₃ and nitrostarch; USP 2118501 (1938)

E. Jones, 32, 8147 (1938); A safety igniter suitable for blasting charges resembling the one described in CA 29, 8336 (1935); USP 2127603 (1938)

C. G. Storm, 34, 7605 (1940) and Army Ordnance, 21, 20-30 (1940); Methods of improving ignition of propellant charges are discussed

L. Burrows & C. Van Winter, 35, 2722 (1941); A slow ignition charge, comprised either of smokeless powder and an oxidizer, such as KCIO₃ or KNO₃ with or without a metal, such as Al, Te, Mg, Se or Zr, or a mixture of smokeless powders. It is suitable for electric squibs. The mixture is heated by means of a bridge wire of Ni-Cr alloy with a dia. of at least 0.00225", whereby certainty of firing is assured under conditions prevailing in series firing; USP 2228339 (1941)

E. I. duPont de Nemours & Co, 36, 274 (1942); Ignition composition for use in electric blasting caps, consists of or includes a double salt of Pb(NO₃)₂ with a Pb salt of dinitrophenol; BritP 519749 (1940)

L. A. Burrows & G. A. Noddin, 36, 2725 (1942); Ignition composition suitable for blasting explosives comprises of finely divided colloidal smokeless powder mixed with 15-25% of the Pb salt of a dinitrophenol such as 4,6-dinitrocresol-[CH₃-C₆H₄(NO₂)₂]₂ Pb; USP 2268372 (1941)

J. Gillies, 41, 6274 (1947); A primary ignition composition for igniting purposes, yielding little or no volatile matter among the products of its combustion, consists of a mixture of CsI₂ and MnO₂ with or without CuO; BritP 575506 (1946)

E. Jones, 42, 7532 (1948); Claims ignition device for use in military land mines, consisting of a priming compn consisting of a mixt of MgFulminate, Sb₂S₃ & KCIO₃. The container for the priming compn & other parts of the device are described & illustrated; BritP 574053 (1945)

Gévelot & Gauplait, 47, 8374 (1953); Claim an ignition compn for electrical primers, consisting of a mixture of dissolved Guncotton with PbO₂ (60%), Pb₃O₄ (65%) or KMnO₄ (50%). Addition of Sb₂S₃, C, KCIO₃ or KCIO₄ modifies ignition props of above mixture; FrP 881262 (1943)

F. B. Clay & R. A. Sahlin, 49, 14326 (1955); Igniter compns for tracer projectiles which include a Ca silicate fuel as a partial or complete replacement for Mg burn in such a manner as to give a dim or invisible trace to a min distance of 25 yds & a bright & properly colored trace at greater distances. A typical igniter comp is SrO₂ 78, BaO₂ 4, PbO₂ 4, Ca silicate 7 and Ca resinate 7%; USP 2709129 (1955)

R. H. Heiskell, 50, 5293 (1956); Claims a pyrotechnic compn as follows: A dark-burning, nonflash igniter mixt suitable to ignite the tracer compn within a projectile is composed of CuO 50, Mn 49, graphite or stearic acid 1% to act as binder. Other possible O-bearing compds include BaO₂, Sb₂O₅, Pb₃O₄ & PbCrO₄. Another igniter mixt is composed of 80% BaO₂ & 20% sulfide from the group CaS, Cr₃S₄ & SeS, & a small ant of graphite or stearic acid; USP 2726943 (1955) & USP 2726944 (1956)

R. H. Heiskell, 50, 8208 (1956); Claims highly stable, dark-burning igniter compns containing B₄O₃ 85-85, Mn 15-55, stearic acid binder up to 10, and graphite lubricant up to 10%; USP 2716599 (1955) & USP 2726944 (1956)

R. H. Heiskell, 50, 8208 (1956); Claims a nonluminous igniter for initiating ignition of the burster charge of a projectile, consisting of BaO₂ 60-7, Sb₂S₃ 21.3-35, asphaltum 1-4 & graphite 1-4%; USP 2714061 (1955)

T. J. Mulqueeney, 50, 9742 (1956); Claims: Ignition compns of the Pb-Se type, where the PbO is produced in situ, were prep'd & evaluated for use in blasting caps & detonators. This type of mixt afforded less erratic firing times than when the PbO was mechanically incorporated. The Pb is oxidized by heating in an internally baffled rotating drum in an oxidizing atm; USP 2740703 (1956)

J. H. McLain & T. A. Ruble, 51, 12494 (1957); An improved fuse for use in firing gas
grenades, smoke candles, etc, is provided by placing a standard
(II), a delay charge (III), & an ignition mixt
(IV) in a fuse body. II, which is relatively insen-
sitive to frictional impact, stable, substantially
glassless, easily ignited, & has a high burning
temp so as to act as a relay between I & III,
is composed of PbO_2 + 2, Si (V) 33.7 ± 2,
Mn 11.2 ± 1, celluloid (VI) 1.8 ± 0.2%. III,
which has substantially the same characteristics
as II except that on burning it produces a plug-
forming residue to prevent backfiring, is com-
posed of PbO 76.9, V 19.3, Fuller's earth 1.5,
VI 1.8 & graphite 0.5%. IV which has substi-
tually the same characteristics as II except that
it gives a large flash & produces a large vol of
gas, consists of PbCrO_4 59 ± 5, V 19.6 ± 3,
Mg 19.6 ± 3 & VI 1.8 ± 0.2%. In practice a
small pellet of II at about 0.188 inch diam &
0.2 inch height & weighing 0.3-0.4g & a pellet of
III of about 0.188 inch diam & 0.3 inch
height & weighing 0.6g are pressed in beneath
the primer under a load pressure of 400-600
lb, another pellet of III is similarly incorporated.
The cavity is filled with loose IV, & the whole
is sealed with a crimped-in closure plug; USP
2792294 (1957)

Th. Goldschmidt, 52, 3346 (1958); Safety
igniter for miner's lamps prep'd from Al-bearing
Ce alloys; CerP 926654 (1955)

W. J. H. Schneider, 52, 5825 (1958); Claims
an igniter composition consisting of Tribasic
Pb picrate, Pb(C_6 H_5 O_7)_2, 3PbO, 2.5H_2 O,
having an explosion point of 160-80⁰, & an
ignition delay of 0.6, 1.5, and 4.7 sec at 260,
250, and 240⁰, resp, is prep'd by reaction while
hot of a mol Pb salt and a picric acid (I) soln
contg 7.3-7.5 inoles NaOH per mole of I; FrP
1026869 (1953)

Mulqueeney & F. R. Seavey, 52, 7704
(1958); An explosive train for elec detonators
and blasting caps is described having a cyclonite
base charge, a Pb azide initiating charge, and
an ignition charge; the initiating charge is
placed between the ignition and base charges.
The ignition compn consists of 30-50% loose,
dry Hg fulminate (40-90% through 200-mesh
sieve) mixed with 50-70% ground, dense prop-
ellant (100% through 60-mesh sieve). The
propellant compn is diphenylamine 0.3-1.2,
graphite 5.0 max, ash 3.0 max, moisture &
volatile compds 2.0 max, ether extractable
compds 2.75% max, and Nitrocellulose (12.0
to 12.7%) balance. The av firing times in
millisecc for caps charged with the various
mixes are: 50/50 Hg fulminate/propellant
ignition 3.6, total 10.9; 40/60 mix, 3.9 and
13.2; 30/70 mix, 5.3 and 18.0; USP 2825639
(1958)

D. T. Zebree, 53, 17514 (1959); Claims an
improved delay fuse for igniting primary explo-
sive compositions in detonators & squibs.
The preferred delay mixture consists of 60% BaO_2,
Se (75/25) & 40% Pb-Sn alloy (85/15). Ingre-
dients should be in the 1-70μ particle size range;
USP 2889269

H. H. Williams & W. A. Gey, 53, 22957
(1959); Claim an igniter material for ignition
of solid propellants in a stable, reproducible
manner, which builds up pressure gradually so
as not to cause the propellant grain to chip or
crack, consists of a mixt of polytetrafluoroethylene
(I), B, & NH_4 ClO_4. The I, mol wt 500,000-
1,000,000, serves as a binder for the other 2
materials & as an oxidizer for the metal. The
particle size of the NH_4 ClO_4 is 1-5μ. The compn
is 5.5-7.5% B & the remainder NH_4 ClO_4 & I in
substantially equal amts; USP 2900242 (1959)

D. T. Zebree, 54, 2745 (1960); Claims an
initiating device having a delay fuse element &
a loose ignition mixt (Pb 72.4/Se 27.6% by
wt) in contact. Incorporation of 1-5% Si, which
increases the heat of combustion of this mixt,
compensates for deteriorations which occur in
ignition mixts, esp useful for blasting caps;
USP 2908559 (1959)

W. E. Schulz, 55, 3062 (1961); Claims igni-
ting comp, a mixt of Mg, Te & TeO_2 loosely
packed for use in elec actuated squibs. The
ignition temp of this mixt is higher than that
of other squib compns; a 3-5 amp, or larger,
firing current is typically desirable. Ignition
time, functioning at low & high temps & at
low pressures, storage stability etc are good.
Typical compns are: 5/88/7; 16/78/6; 27/67/6;
& 36/59/5% by wt Mg, Te & TeO_2 resp in
each case; USP 2953447 (1960)

E. J. Walden, 55, 25258 (1961); Claims
ignition delays at -65⁰F of a rocket propellant
from NH_4 NO_3 & a butadiene-methylnitrylpyridine
, copolymer (90:10) are reduced 50% by the
inclusion of pellets composed of KClO_4 26.5,
Ba(NO₃)₂ 16.6, Zr-Ni alloy (50:50) 53.9, & Et cellulose 3.0% embedded in the ignition-sustaining material; USP 2990683 (1961)

S. V. Peyton & E. Williams, 56, 14523 (1962); Claim a compn for ignition by stab-pin or flash, containing Pb 2,4-dinitroresorcinate 50, Ba(NO₃)₂ 45, & tetrazene 5% by wt. It has very good thermal stability, & its life in hot & moist storage is very much longer than that of fulminate mixts. It is highly compatible with Al, Cu, brass & Shaped Cu & also with Pb(N₃)₂; BritP 892741 (1962)

L. G. Herring, 56, 15720 (1962); Claims a pelleted igniter made by mixing a mixt of powd Zr & Ni with a powdered inorg oxidant & a small amt of Et cellulose; this mixt is pelleted & dried to remove excess solvent; USP 3017300 (1962)

J. D. Clark, 60, 7864 (1964); Claims an igniter for a rocket propellant consisting of a HNO₃ soln of an amine nitrate, such as disopropylamine nitrate, is provided by a dough-like mixt of about 85% of a metal hydride, such as LiH or LiAlH₄, mixed with about 15% of a dry rubber cement. The igniter presents a large contact surface & is supported within the rocket chamber in such a manner that, when the monopropellant is allowed to impinge on it, as through a spray nozzle, the heat of reaction increases the monopropellant to the ignition temp & heat of combustion causes decom of the igniter; USP 3115005 (1963)

R. L. Shimpagh, 62, 12969 (1965); Claims an igniton compn consisting of Ba chromate, boron & lead dioxide for use in electric detonators; USP 3123667 (1965)

W. A. Proell, 64, 12454 (1966); Claims a mixt of Black Powders 30-50 & smokeless powder 50-70%, compressed into a coherent cake to be used for igniting propellants; USP 3234059 (1966)

(No author cited), 67, 10158 (1967); Claims an igniton compn having excellent low pressure ignition properties, consisting of a monovalent or polyvalent metal powder, eg Al, Cu or Mg; a metal oxide, eg PbO₂ or BaO₂; & 15% B as an activator; GerP 1243067 (1967)

G. B. Young & S. J. Lubinski, 67, 55840 (1967); Claim a new ignition charge resistant to electrostatic discharges may be prepd from amorphous B & Pb oxides in the following mixt ranges: B 1.5-2.5, PbO 97.5-98.5; B 8-30, PbO 70-92; & B 15-30, PbO₂ 70-85%. Thus, 10g 90-7% pure amorphous B & 490g 5µ 99% pure PbO are ball-milled with 0.7kg H₂O for 16 hrs, dried at 212-25°F to <0.1% H₂O, & passed through a 20-mesh screen. The screened particles are electrostatically grained by tumbling in a glass jar at 65 rpm for 2.25 hrs. When incorporated into a cap, these compns exhibit an a c breakdown voltage of 1.4-2.2kv. They also show a consistent bridgewire sensitivity & burn in a gassy & non-violent manner; USP 3317360 (1967)

D. H. Lee & D. D. Evans, 66, 67510 (1967); Claim a simple, efficient means of ignition of the monopropellant N₂H₄, provided by use of a fluid bed consisting of carrier pellets of Al₂O₃ impregnated conventionally with a N₂H₄-decomp catalyst such as a mixt of the oxides of Fe, Ni & Co, a layer of the pellets ~ 0.25 in thick at the N₂H₄ feed end of the bed being coated with a powerful, H₂O-soluble, solid oxidizer such as KMnO₄, KClO₃, or, preferably, I₂O₅, which causes ignition of the initial N₂H₄. The heat thus developed raising the temp of the bed sufficiently to self-sustain N₂H₄ decomp; USP 3298182 (1967)

4) Ignition Compositions for Use With Pyrotechnics. This subject was treated in considerable detail in Vol 4 under Detonator, Igniters and Primers, Sect 3, Part B & Ignition Train Used in Pyrotechnics, pp D759-768, with the following additional reference:

Gordon & Campbell studied preignition & ignition reactions of the pyrotechnic system Zn—C₆Cl₆—KClO₄. They found the following reactions to be responsible for preignition, ignition, and combustion phases of this system: (I) 3Zn + C₆Cl₆ — 3ZnCl₂ + C; (II) 4C + KClO₄ — KCl + 4CO(+CO₂); (III) 4Zn + KClO₄ → KCl + 4ZnO. Reaction I is exothermic and together with reaction II raises the temp of the system to >500°C. At temps above 421°C the preignition reaction III becomes highly exothermic and propagation ensues in the neighborhood of 520°C.

5) Ignition of Firedamp and Coal Dust. Many disastrous coal mine explosions have been caused by the ignition of firedamp (methane/sir mixtures) and/or coal dust. The subject of firedamp-coal dust explosions has been studied extensively. Some of this extensive literature was reviewed in Vol 3 under Coal Mine Explosions, pp C360-378, & under Coal Mining Explosives, Non-Permissible, pp C437-444, & Coal Mining Explosives, Permissible, pp 444-459. Below we give recent references on the ignition of firedamp and present a cursory discussion of the mechanism of firedamp oxidation (ignition & combustion). This complex mechanism tends to explain the great variability in experimental results on firedamp ignition & to furnish some excuse for the unfortunate fact that coal mine explosions still occur even after almost a century of intensive international effort to eliminate them.

The oxidation of methane (as well as other hydrocarbons) proceeds via a chain branching mechanism. Each of the stages of the chain branching mechanism: initiation, propagation & termination, may be affected to a different degree by relatively slight changes in ambient conditions under which oxidation takes place. The cumulative effect of these "slight" changes on the over-all reaction may be quite large and lead to ignition or non-ignition under what appears to be very similar circumstances. Similarly, once ignition has been effected, its propagation (burning and/or explosion) is also subject to the variability described above.

The following reaction scheme is taken from Lewis & Von Elbe (Ref 1) to whom the reader is referred for details:

\[
\begin{align*}
\text{Initiation:} \\
\text{CH}_4 + \text{O}_2 & = \text{CH}_3\text{OOH} \\
\text{CH}_3\text{OOH} & = \text{H}_2\text{CO} + \text{H}_2\text{O}
\end{align*}
\]

\[
\begin{align*}
\text{Propagation & Chain Branching} \\
a) \text{H}_2\text{CO} + \text{O}_2 & = \text{free radicals} + \text{OH} \\
\text{OH} + \text{CH}_4 & = \text{H}_2\text{O} + \text{CH}_3 \\
\text{CH}_3 + \text{O}_2 & = \text{H}_2\text{CO} + \text{OH} \\
b) \text{OH} + \text{H}_2\text{CO} & = \text{H}_2\text{O} + \text{CHO} \\
c) \text{CHO} + \text{O}_2 + \text{M} & = \text{CHO}_3 + \text{M} \\
d) \text{CHO}_3 + \text{H}_2\text{CO} & = 2\text{CO} + \text{H}_2\text{O} + \text{OH} \\
(M \text{ is a third body})
\end{align*}
\]

\[
\begin{align*}
\text{Termination:} \\
e) \text{OH} + \text{wall} & = \text{destruction} \\
f) \text{CHO} + \text{O}_2 & = \text{CO} + \text{HO}_2 \\
\text{HO}_2 + \text{wall} & = \text{destruction} \\
\text{H}_2\text{CO} + \text{O}_2 + \text{wall} & = \text{destruction of H}_2\text{CO}
\end{align*}
\]

Without going into details, it is clear that the overall oxidation reaction will be affected by the fate of OH radicals (as well as other intermediates). For example, if the rate of step a) & d) exceeds that of steps b) & d) the oxidation of methane will proceed & conversely. Further complication is introduced because step d) depends on the relative rates of steps c) & f). Thus it is not surprising that minor changes in reaction conditions, which can affect different elementary reactions in different ways, can lead to major changes in overall reaction rate, and indeed be the difference between ignition & non-ignition.

The ignition of firedamp (coal-dust explosion are generally considered to be triggered by firedamp explosions, therefore we will make few further comments about coal dust explo-
sions) by explosives presents its own set of problems. Among these are the high temperatures of detonation products, the possible presence of radicals in these products which can promote ignition in the methane (as discussed above), and the adiabatic compression and heating of firedamp pockets by the blast. Presumably all these possible ignition mechanisms are minimized by proper stemming of the bore-hole (Ref 2), the presumption being that the products and blast now emerging from the bore-hole have been attenuated in temperature, radical concentration & blast pressure. Experience shows that, although good stemming may be helpful in reducing the incidence of firedamp ignition, it does not completely eliminate ignitions. To some extent this may be due to fissures in the coal seam, containing firedamp, crossing the bore-hole. Also the amount of stemming required to obtain the necessary degree of attenuation may make its use uneconomical.

Two approaches are now used in attempts to prevent firedamp ignition by explosives. The first attempts to reduce the temperature of detonation products of the explosive, and the second introduces flame-suppressing materials at and near the face where blasting
operations are carried out. The first scheme involves “sheathing” the explosive or incorporating into it materials that act as “heat sinks.” This has been discussed in Vol 3, pp C437-444 & pp 444-459. The most widely used practical application of scheme two is “rock-dusting,” ie dusting the inside of the mine with fine limestone. Rock-dusting appears to be quite effective in preventing coal-dust explosion but is not so effective in preventing methane-air explosions. Other dusts, such as lithium halides, appear to be more effective than limestone, but their use is not economical.

Modern coal-mining practice tends towards the use of mechanical devices, such as Cardox, Hydrox & Mechanical Miners rather than explosives. This will not necessarily reduce the number of coal mine explosions because these may be, and indeed are known to be, initiated by other means such as faulty electrical equipment, breakage of electrical cables or other means of producing a hot spark or arc

Written by J. ROTH


References, & brief abstracts when available, to studies of firedamp and/or coal dust ignitions & explosions, not listed in previous Encyclopaedia Vols are given below:


3) N. E. Hanna et al, US Bur Mines RI No 5463 (1959) & CA 53, 14517 (1959). The effect of 0, 3.6, 6.6 & 9.5% NaCl on the incendiency of permissible explosives was studied by the Bruceston up-and-down technique in gallery tests. The study showed that the addn of relatively small quantities of NaCl tends to reduce but does not eliminate igni-


“Relative efficacy of stemming materials in reducing incendiency of permissible explosives.” Stemming, used to confine explosives in shot holes in underground coal mines, minimizes the chance of ignition of firedamp or coal dust by the detonation products. Tests by the Bureau of Mines indicate that NaCl, pure H2O, gelled water (3% carboxymethyl cellulose), and a concd salt-H2O soln are more effective than an equal wt of wet or dry fireclay. The H2O, gelled H2O, and salt-H2O soln are equally effective. The approved stemming device is distinctly inferior to 1 lb of dry fireclay

6) N. E. Hanna et al, US Bur Mines RI 5867 (1961) & CA 55, 27886 (1961). “Factors affecting the incendiency of permissible explosives. Ammonium Nitrate and carbonaceous material.” Analysis of 15 specially prepd explosives, with 3 particle-size ranges of NH4NO3 and 5 types of carbonaceous material, showed that the particle size of NH4NO3 in permissible-type formulations had a highly significant effect on the incendiency of the explosive to firedamp, with coarse NH4NO3 producing less incendiency explosives than the fine. The type of carbonaceous material had no significant effect on the incendiency

7) W. J. Montgomery, CanMiningMetBull 55, 765 (1962) & CA 61, 11840 (1964). Review article on the explosive capacity of various dusts (coal & ore dust, esp sulfides). Measures for the prevention of explosions are described

8) J. Nagy & D. W. Mitchels, US Bur Mines RI No 6344 (1963) & CA 60, 6667 (1964). Data are given to assist in interpreting observations made after an explosion in an operating coal mine when evidence is collected to establish the cause & factors affecting ignition & propagation of the explosion


“Ignitability of gallery atmospheres.” Experiments show that a variation in the natural gas
content in the gallery can have a great effect on the ignitibility of the gallery atm; max ignitibility of the atm by permissible explosives occurs with \( \approx 8.1\% \) natural gas. For similar explosives, incendivity increases with detonation velocity or bulk strength

10) D. V. Stoyanov, Vuglishta (Sofia) 19 (9), 25 (1964) & CA 62, 10257 (1965). The factors affecting the explosive properties of coal dust are the chem compn of the coal, the dispersion compn & conc of the coal dust, the compn of the mine atm, the source of the flame, the natural ash content, the moisture content & the volatile content of the coal. A nomogram is given by means of which the min amt of inert matter to be added to prevent an explosion can be determined from the natural ash content.

11) J. Nagy et al, US Bur Mines Rept No 6597 (2) (1965) & CA 62, 11591 (1965). "Explosibility of carbonaceous dusts." Activated carbons & charcoals were tested. In general only dusts having \( > 8\% \) volatile matter do not present an explosion hazard. Carbon black presented a fire rather than explosion hazard. All bituminous coal dusts are explosion hazards. In the 119 samples tested none with \( < 13\% \) volatile content ignited by spark. Dusts having a 30% volatile content had 50% probability of ignition by spark. All dusts with \( > 40\% \) volatile content ignited by spark.

12) G. Yoshikawa et al, Kogyo Kayako Kyokai-shi 27 (6), 360 (1966) (Eng/Japan) & CA 67, 75024 (1967). Based on high-speed motion pictures of firedamp ignition by typical coal-mine dynamite shots, hydrodynamic-thermodynamic analysis is made on the expanding flow of gaseous detonation products with salts dispersing in them. Detonation products attained highest velocity at a distance from the charge at twice its diam, set up a shock wave in the surrounding firedamp atm, & decreased in velocity as expansion proceeded.

13) D. Rae, Safety in Mines Res Estab, Res Rept 253 (1967) & CA 69, 1140 (1968). "The main characteristics of slow coal-dust explosions and their relation to the testing of barriers." The main characteristics of slow coal-dust explosions in the new gallery at Buxton are a const initial flame front acceleration from practically zero velocity up to a speed of about 120 m/sec, followed by an increasing acceleration of the flame front. The initial acceleration is detd wholly by the effect of the igniter on the dust deposit and is made evident by plotting the square of the flame arrival time against distance, but the subsequent increasing acceleration is due to the flame of the explosion. This behavior is found also in other expl galleries where the ignition zone is short and where the main fuel in the ignition zone is the same as in the deposit in the rest of the gallery. The violence of an explosion can be characterized by the initial acceleration, which can be used to compare explosions in different galleries. The min acceleration that will sustain an explosion without oscillation of the flame along the gallery is inversely proportional to the length of the gallery, explosions in long underground roadways can accelerate more slowly without suffering oscillations than those in short expl galleries, and this may be important in the siting of explosion barriers.

14) R. W. VanDolah et al, US Bur Mines Rept No 7195 (1968) & CA 70, 13126 (1969). "Development of Slurry Explosives for Use in Potentially Flammable Gas Atmospheres." In this rept expls with 23AN-based slurry expls showed that it is possible to have them cap-sensitive and relatively nonincendive. All slurries that used flaked Al were sensitive to No 6 EBC (electric blasting cap) except one contg AN 47.2, NaCl 10, water 30, Al (particle size 12µ) 8, sugar 3.5, guar gum 1.0 & buffer 0.3%. Buffer (consisting of NaOH & KH₂PO₄ in 1:29 ratio) was added to hold the pH at 4.5-5.5.

15) V. P. Gorkovenko et al, VzryvnoyeDelo 1970 (68/25), pp 116-22 & CA 73, 89710 (1970). Evaluation of the permisibility of AN-based expls was conducted by comparing oscillograms of light emitted during explns. The amplitude and time of light emissions, dependence betw their values, and the probability of ignition of methane-air mixts were detd.

done by detg the "selectivity index," $S = (V_1 - V_2)/V_2$, where $V$'s are volumes of gaseous expln products ($V_1$, obt'd on expln in a steel tube and a sand wad in the Dolgov Bomb; $V_2$, obt'd on expln in a free space in the Dolgov Bomb; and $V_3$, calculated.

17) L. V. Dubnov & A. I. Romanov, Ibid 1970 (68/25), 127-31 & CA 73, 89721 (1970). Techniques for evaluating the combustion tendency of permissible expls, is discussed. The value $\psi = (S_2 - S_1)/S_1$, where $S_1$ is av (50%) transmission distance in air (gap) of initiation of deton and $S_2$ is av (50%) transmission distance of combstn, is taken as a basis for evaluation of the combstn tendency. The transmission distance was detd by expln of a composite cartridge consisting of active and passive segments with air gaps between them.

18) Anon (US Dept Instr) Fed Regist 28 Aug 1971, 36, 17336 & CA 75, 14265 (1971). Standards for preventing explosions from explosive gases other than methane & procedures for testing accumulations of such gases: the following gases shall not be allowed to accumulate in underground coal mines in excess of the concs (vol%) listed: CO 2.5; H 0.8; H_2S 0.8; C_4H_2 0.4; C_3H_8 0.4; & MAPP (methylacetylene-propylene-propadiene) 0.3.

Recent review articles are:
1) H. Kaffanke, Maîtreie Degagement Grisouteux, Amelior Climat Mines, Journees Inform. (Luxembourg), 1971, 217 & CA 76, 144547 (1972); prediction of firedamp emissions
2) M. Boutonnat et al, Ibid, 355 & CA 76, 144545 (1972); equipment for measurement of firedamp & ventilation control
3) H. Eicker, Ibid, 373 & CA 76, 144544 (1972); maintenance of firedamp meters & transmission of measured values

6) Ignition, Spontaneous, of Explosives.

This section presents some historical examples of spontaneous ignition. A more technical discussion of spontaneous ignition will be given under Thermal Explosion, Catalytic Effects in a future Vol.

Explosives containing improperly stabilized ingredients such as NC or NG, decompose in storage, especially at tropical temperatures, with evolution of nitrogen oxides, formation of nitric acid (and nitrous acid) and a rise in temperature. As these nitrogen compounds act auto-catalytically, the temperature and rate of decomposition increase progressively until a point is reached at which the powder ignites spontaneously, ie without any external stimulus.

For underoxidized compositions, stored in closed containers, the combustion process following spontaneous ignition may be incomplete. However, gases generated by this process will generally burst the container and after-burning may occur and contribute to the general conflagration which in turn can lead to explosion of any remaining unreacted material some of which ignited spontaneously or of other nearby explosive charges.

Propellants were especially susceptible to spontaneous ignition before it was learned how to stabilize the NC. This was done by including substances called stabilizers in the propellant formulations. These substances, such as diphenylamine, centralite etc absorb the nitrogen oxides as they are formed, thus removing them from the sphere of reaction. This prevents further decomposition and the rise in temperature which decomposition produces. For similar reasons, it is advisable to keep the storage magazines cool and, in fact, since the catastrophe aboard the French battleship Jéna, occurring in 1907, nearly all countries introduced refrigerating systems for cooling the magazines on ships.

Many disastrous explosions occurred at the end of the 19th and beginning of the 20th century, which were caused by spontaneous ignition of propellants either containing no stabilizers or containing those which were ineffective, such as amyl alcohol used in some French powders.

Among the disasters caused by spontaneous ignition of propellants may be cited:

Japanese battleship, Mikasa, destroyed in 1905 with 599 men lost

Brazilian warship, Aquidaban, destroyed in 1906 with 213 lives lost

French battleship, Jéna, exploded in 1907 with 114 lives lost

Japanese ship, Matsushima, destroyed in 1908 with 114 men killed

Land magazine of Batuco, in Chile, in 1908.
French battleship, Liberté, destroyed in 1911 with 204 men killed.

Italian warship, Benedetto Brin, in 1915.

The explosion of the Russian dreadnought of the Black Sea Navy, Empress Marie, which occurred during WWI, was probably due to the same reason, although there is a chance that it might have been due to sabotage.

Even as insensitive a material as Ammonium Nitrate, AN, may be subject to spontaneous ignition. The disastrous explosions of two ships at Texas City in 1947 could have been a consequence of the spontaneous ignition of paper-bagged fertilizer grade AN. The actual ignition may have occurred during rail shipment and later developed into a full-scale deflagration and subsequent detonation in the hold of SS Grandcamp. Spontaneous ignition may have resulted from the AN being bagged while it was too hot and/or improper removal of acid from the AN. Both these conditions could lead to autocatalytic decomposition which can result in spontaneous ignition. For alternate explanations of possible causes of the fire that resulted in the Texas City disaster see Vol 1, A 358-362.


7) Ignition Tests are historically divided into two categories:

A. Those testing the response of a substance to an open flame; i.e determination of the flame temperature at which a substance ignites and remains burning. Usually this temp is a few degrees higher than the so-called flash point of the substance

B. Those testing the behavior of materials exposed to high temperatures in the absence of an open flame. These tests are used to determine so-called ignition or explosion temperatures. Ignition or explosion temperature is an indefinite quantity since its evaluation depends strongly on the conditions of measurement. As an example of this variability and dependence on test conditions, the "explosion temperatures" for as standard an explosive as TNT are quoted from 290 to 370°C! (Ref 16)

C. Measurement of times to ignition of explosive exposed to hot flames.

In view of the above the following ignition tests & ignition temperature tests are primarily of historical interest. They also serve to outline the difficulties encountered in trying to characterize quantitatively the response of explosives to heat. Quantitative treatment of "explosion temperatures" and delay to explosion (induction time) and the parameters of the explosive that affect these quantities will be presented in a future Vol under "Thermal Explosions".

A. Flame Tests

The International Committee on Explosives and the German Railway Commission prescribed the following ignition tests, applied mostly to safety explosives. It is advisable to conduct these tests behind a protecting barricade

a) Fuse Test. Place 3g of powdered explosive in a short glass or paper tube, about 2cm in diameter, and tap it gently to give the explosive an even surface. Introduce a slow burning fuse (rate 1cm/sec) and ignite it. If the explosive ignites, it is classified as a deflagrating or readily inflammable explosive. Also observe whether the explosive burns partly or completely and note the time elapsed between lighting the fuse and ignition of the test sample.

b) Red Hot Iron Basin Test. If the explosive passed test a) without igniting, it is submitted in the following test to a much higher temperature in order to be sure that it is not liable to explode when exposed to fire.

By means of a burner, heat to red heat a hemispherical iron basin 12cm in diameter and 1mm thick and introduce a small quantity, not more than 0.5g, of the explosive to be tested. If no explosion takes place, introduce more of the explosive, gradually increasing the quantity until a weight of 5g is reached and repeat the test with 5g three times. Observe the manner in which the explosive burns and the time that elapses between adding the explosive to the basin to the extinction of the flame.

c) Red Hot Iron Test. This test is designed to ascertain the inability of safety explosives to explode on heating or to burn continuously.

Heat the end of an iron bar, 15mm in diameter, for a length of 10cm to a cherry red heat (about 900°C) and plunge it into a small quantity of explosive placed on a sheet of asbestos. If the explosive does not ignite or is difficult to ignite, and burns slowly without explosion,
and the flame dies the moment the hot bar is withdrawn, continue testing fresh portions of it, gradually increasing the amount of sample up to 100g. If the explosive still behaves as described above, it is considered to have passed the test.

This test is not mentioned among German Railway Commission tests, instead, the following two tests are prescribed:

d) Iron Basin with Wood's Metal Test. Fill an iron basin, 14cm diam and 7cm high, to within 2cm of the top with molten Wood's metal and insert an armored thermometer or a thermocouple in the middle of the molten mass. Heat the basin and, as soon as the temperature reaches 100°, place 3 test tubes (15 mm id and 120mm long) containing 0.2-0.5g of powdered explosive about 50mm away from the thermometer, spaced about 100m apart and immersed 20mm into the bath. Raise the temperature of the bath at the rate of 20° per minute until ignition occurs or until the temperature reaches 320°.

Note: The test may also be conducted by raising the temperature 5°/min (Ref 2)

e) Iron Box Test. Place 0.5 to 1kg of the explosive to be tested in a sheet iron, riveted box 1mm thick and 85 x 85 x 85mm in size. Fasten the lid by means of iron wire, bound crosswise around the box and place the box in a brisk wood fire (by means of a mechanical device operated by remote control). The explosive is considered to have passed this test if it does not ignite or explode within 10 minutes.

Notes:

a. This test has to be carried out in the open and not less than 100 meters from inhabited buildings.

b. If it is known that the explosive is not sensitive, the box may be laid by hand provided it has been wrapped up well beforehand in several layers of paper so that the explosive will not be ignited too soon.

Fleischer & Burtle (Ref 3) give results for a flame test for different types of Lead Azide. In this test a std safety fuse provides the flame and percent firing of the Lead Azides is recorded as a function of distance from the fuse as shown in the tabulation below:

<table>
<thead>
<tr>
<th>Lead Azide</th>
<th>Distances from Fuse (inches)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1/4</td>
</tr>
<tr>
<td>Dextrinated</td>
<td>100%</td>
</tr>
<tr>
<td>Pure</td>
<td>100</td>
</tr>
<tr>
<td>PVA</td>
<td>100</td>
</tr>
</tbody>
</table>


B. Ignition Temperature of Explosives, Tests. (Verpuffungstemperatur or Entzündungspunkte, in German and Inflammation, in French). When an explosive is heated by means other than an open flame, a temperature is reached at which the sample ignites and burns or deflagrates producing a flame or fumes off without any loud report, or finally explodes or detonates with a loud report. This test should not be confused with either the “flash point” or “ignition point” test (see Sect A) in which a substance, preheated to a certain temperature, is ignited by means of an open flame, whereas in the “ignition temperature” test no outside flame is used.

Numerous tests exist for determining ignition temperature, but few of them give the same result. Results may vary as much as 100° depending on the method used, on the rate of heating, on the size of the sample used, on the state of division etc. See Thermal Explosions for a discussion of why measurement variables affect ignition temp results.

One of the most reproducible of these tests is described under “Explosion Temperature Test,” an official test at Picatinny Arsenal (Ref 15). This test is applicable to the determination of ignition point, decomposition point, deflagration point etc for those explosives which do not really explode under conditions of the test.

Some other “ignition temperature” tests are shown below:

a) German Railway Commission Test (Refs 1, 2 & 3). A small sample of an explosive (0.1g for all explosives; except Black Powder, NH₄NO₃, and chlorates, for which 0.50g is used) is placed in a test tube, 125mm long, 15mm id and 0.5mm thick, which is tightly corked and placed in a paraffin or Wood’s metal bath provided with a stirrer and thermometer and preheated to 100°. Heating is continued with stirring so that the temperature
increases 5° per minute and the point is noted at which the sample ignites, fumes off etc.

A. Pérez Ara (Ref 3, p 109) gives the following temperatures for explosives tested by this method:

Black Powder 225 to 300, NC (13%N) 185, smokeless powder 185, NG 160, Nitrostarch 170, NH₄NO₃ 225, Tetryl 190, Mercury Azidé 200, Silver Azide 200, Lead Azide 340 to 350, Cuprous Azide 210 to 350, Cupric Azide 245, Mercuric Fulminate 160 to 200, Silver Fulminate 200, Chédîte 200, blasting gelatin 207 to 211 and Picric Acid 225 to 350°

b) A Test Described by A. Pérez Ara (Ref 3) and Datta & Chatterjee (Ref 6). The above apparatus is used but the empty test tubes are preheated in the bath to different temperatures depending on the explosive to be tested. At this point, a small (0.10 to 0.50g) sample is added to one of the empty tubes and if no ignition takes place the temperature is raised about 5° and a new sample is thrown into the tube. The test is continued until the sample either ignites, fumes off or explodes. If the sample ignites at the first trial, the temperature is lowered 5-10° and the test repeated and continued at progressively lower temperatures until one is reached at which the sample does not ignite. After this, the temperature is raised again and the point is noted at which the sample ignites.

Usually, this test gives higher results than the method a) (Ref 3)

Note: Datta & Chatterjee used a KHSO₄ bath which they heated up to 500°. Kast and Haid tested a number of initiating explosives by both methods (Ref 11) and found that, although most explosives give higher results by method b), there are many cases in which both methods give the same results, especially if heating is conducted at a rate of 20° per minute in method a)

c) Methods of Direct Ignition of Kast (Ref 3). A small quantity of explosive (up to 0.5g) is placed on a small iron plate and an attempt is made to ignite it by the flame of a splint of wood. Another small sample of explosive is touched with a thin iron rod, preheated to red heat. Another small sample is thrown into an iron dish of about 12cm diameter, preheated to red heat.

d) Spanish Method (Ref 1, v 2, pp 435-6). A long copper strip, 15mm thick and 80mm wide, is heated at one end with two gas burners. The other end of the copper strip has a circular holder with five holes containing copper test tubes. The central test tube is filled with mercury and contains a thermometer while the other four, placed around the first, contain 0.1g samples of explosives. Heating is continued until one of the samples in one of the test tubes nearest the heated end ignites or explodes and the temperature is noted. The remaining samples ignite a bit later. The average temperature of ignition of all the samples is taken as the "ignition temperature"

e) Ignition Temperature by Fisher-Johns Apparatus, by Koffer Micro Hot Stage, by Dennis Melting Point Apparatus or by the Maquenne Block. The first apparatus is described in Fisher's Catalogue No 80, p 619, No 12-142.

The second is shown in Arthur H. Thomas Catalogue (1950 ed.), pp 877-880.


Essentially, all of these devices consist of a metallic block or bar heated either by gas (as in the Maquenne Block) or by an electric heater. The test may be conducted by placing a few small grains of explosive on the preheated block and noting the temperature at which the sample ignites, degradates or explodes by means of a thermometer installed inside the block.

Tests conducted at Picatinny Arsenal with these devices showed that fairly consistent results may be obtained for a given explosive if all determinations are made under the same conditions (same rate of heating, size of grains,
size of sample etc)

i) Method of Snelling and Storm. In this method (Ref 5), a 2-3 g sample of NG was heated in a test tube immersed in a paraffin bath provided with a thermometer. The temperature of the NG inside the tube was measured by means of a Cu-constantan thermocouple. The experiment is conducted behind a barricade. The following results are for a period of heating equal to 5 minutes.

<table>
<thead>
<tr>
<th>Temperature of Bath</th>
<th>Sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>At the start of heating the NG</td>
<td>80</td>
</tr>
<tr>
<td>NG appears to boil owing to rapid evolution of water and oxides of nitrogen</td>
<td>174</td>
</tr>
<tr>
<td>Vigorous boiling (bubbling) of NG</td>
<td>186</td>
</tr>
<tr>
<td>NG becomes thick &amp; viscous</td>
<td>193</td>
</tr>
<tr>
<td>Temperature of sample rises faster than that of the bath</td>
<td>198</td>
</tr>
<tr>
<td>Violent detonation</td>
<td>201.5° 215-218°</td>
</tr>
</tbody>
</table>

If the rise in temperature is slow, the sample decomposes and partly chars at temperatures above 180° without producing any explosion.

g) Method of Micewicz and Majkowski. This method (Ref 14) may be considered to be a modification of the Snelling and Storm's method described above.

M & M examined several high explosives and found that in many cases the temperature of the explosives rises considerably above that of the bath during heating (on account of the exothermic decomposition that takes place) but sometimes it falls again before ignition takes place.

h) Koehler and Marqueyrol (Ref 8) determined the ignition temperature of NC in vacuo, in air, and in an atmosphere of CO₂. They came to the conclusion that the results are practically the same and are as follows:

- Guncotton (CP₁) with N=13% 180 to 190°
- Collodium cotton (CP₂) with N=12% 186 to 189°
- Poudre B 174 to 187°

i) Stettbacher (Ref 9) comments on the great variance of results for the ignition temperature of NG, ranging from 180° by Nobel to 220-255° by Staubinger, and criticizes the various methods of determination.

j) Taylor and Rinkenbach (Ref 12) determined ignition points of various explosives by dropping small portions onto a block of heated Wood's alloy:

<table>
<thead>
<tr>
<th>Explosive</th>
<th>Ign Temp</th>
</tr>
</thead>
<tbody>
<tr>
<td>Merccuric Fulminate</td>
<td>260°</td>
</tr>
<tr>
<td>MF + 10% KClO₃</td>
<td>240°</td>
</tr>
<tr>
<td>MF + 20% KClO₃</td>
<td>235°</td>
</tr>
<tr>
<td>Silver Fulminate</td>
<td>245°</td>
</tr>
<tr>
<td>Lead Azide</td>
<td>383°</td>
</tr>
<tr>
<td>Silver Azide</td>
<td>273°</td>
</tr>
<tr>
<td>Mercurous Azide</td>
<td>298°</td>
</tr>
<tr>
<td>Lead Stryphane</td>
<td>293°</td>
</tr>
<tr>
<td>Silver Acetylide</td>
<td>220-225°</td>
</tr>
<tr>
<td>Lead Picrate</td>
<td>281°</td>
</tr>
</tbody>
</table>

The following table gives values for TNT, PA and Tetryl:

<table>
<thead>
<tr>
<th>Explosive</th>
<th>Time* of Heating (min)</th>
<th>Temp of Bath at Ignition or Explosion</th>
<th>Max Temp of Explosive reached during Heating</th>
<th>Temp of sample at ignition</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>TNT</td>
<td>11.55</td>
<td>294°</td>
<td>299°</td>
<td>248°</td>
<td>Ignition</td>
</tr>
<tr>
<td>TNT</td>
<td>19.87</td>
<td>288°</td>
<td>296°</td>
<td>276°</td>
<td>Ignition</td>
</tr>
<tr>
<td>TNT</td>
<td>39.58</td>
<td>283°</td>
<td>294°</td>
<td>284°</td>
<td>Ignition</td>
</tr>
<tr>
<td>PA</td>
<td>13.25</td>
<td>289°</td>
<td>305°</td>
<td>243°</td>
<td>Ignition</td>
</tr>
<tr>
<td>PA</td>
<td>17.17</td>
<td>289°</td>
<td>301°</td>
<td>281°</td>
<td>Ignition</td>
</tr>
<tr>
<td>PA</td>
<td>19.25</td>
<td>293°</td>
<td>296°</td>
<td>247°</td>
<td>Ignition</td>
</tr>
<tr>
<td>PA</td>
<td>36.47</td>
<td>283°</td>
<td>298°</td>
<td>262°</td>
<td>Ignition</td>
</tr>
<tr>
<td>Tetryl</td>
<td>18.43</td>
<td>189°</td>
<td>202°</td>
<td>202°</td>
<td>Explosion</td>
</tr>
<tr>
<td>Tetryl</td>
<td>5.87</td>
<td>182°</td>
<td>232°</td>
<td>232°</td>
<td>Explosion</td>
</tr>
<tr>
<td>Tetryl</td>
<td>5.27</td>
<td>189°</td>
<td>214°</td>
<td>214°</td>
<td>Explosion</td>
</tr>
</tbody>
</table>

*Measured from the moment the bath reached 100°
Silver Picrate 335
Barium Picrate 403
Copper Picrate 373
Hexamethyleneetriperoxidediamine 200
Mannitol Hexanitrate 232
Dulcitol Hexanitrate 205
Cyanuric Triazide 252

k) Tammann & Kröger (Ref 13) examined numerous liquid and solid explosives and came to the following conclusions:

1) The explosion (ignition) temperature decreases with increasing size of sample and increases with rate of heating.

2) With volatile explosives which melt before exploding, the required explosive sample weights are appreciably higher (20 to 100 mg) than with non-volatile explosives (0.4 to 10 mg).

3) With volatile explosives giving gaseous products of explosion, the substance evaporates or decomposes before the explosive limits are reached.

4) The dependence of explosion temperature upon rate of heating is almost always linear, with the exception of potassium picrate.

5) TNT begins to decompose at 150° and on long heating at constant temperature, the explosion temperature is raised by the formation of decomposition products.

l) Method employed at Picatinny Arsenal (Ref 15) is given under "Explosion Temperature".

m) A Manometric-bomb Method for determination of the ignitability of an explosive is described by Rogozhnikov et al (Ref 17). By this method, the min. amount of igniter required to initiate steady combustion (at increasing pressure) of an explosive is determined; simultaneously, the pressure increase with time is measured. 20 explosion comps. were tested.

The method is useful for investigation of the effects of various additives to the explosives on their ignitability.


n) Ignition Temperature Test (Method of M. Kostevitch). In the methods described above as well as in the method practiced at Picatinny Arsenal (see Explosion Test in Vol 1, pp. XVI & XVII), in which an explosive is placed in a blasting cap shell which is then put into Wood’s metal preheated to a temperature close to the melting point of the explosive, the explosive itself cannot be seen during testing and any melting, decomposition or other phenomenon that takes place cannot be noted. In order to observe all stages of behavior of an explosive, Col Kostevitch proposed to use an all glass apparatus, originally developed by the Imperial Russian Ordnance Dept in 1908 for the determination of melting points of high explosives, such as TNT, PA etc. This apparatus is essentially the same as that described in US Specification MIL-T-248A and used for the determination of setting points of TNT and other high explosives.

It consists of a heavy-walled glass test tube, provided with a stopper and a thermometer. This tube is inserted by means of a rubber stopper into a 2nd test tube of larger diameter, serving as an air jacket. The ensemble is inserted, by means of another stopper, into a beaker filled with a nonflammable, transparent, high boiling point liquid (e.g. dibutylphthalate or silicones) and provided with a thermometer and a mechanical or hand stirrer with a ring at the lower end.

In the Kostevitch method a sample of explosive (about 0.1 g) is introduced into a small glass bulb (volume ca. 3.5 cc), resembling in appearance that used for weighing fuming acids, but made from a larger diameter tube in order
to facilitate the introduction of the explosive. The tube of the bulb is inserted in a hole drilled in the stopper of the outer test tube. This stopper also contains a thermometer. The bulb of the tube and that of the thermometer are so arranged that they are on the same level. After the liquid is preheated to about 100°, the bulb with the sample is introduced and heating is continued in such a manner that the rise of temperature is uniform, say 5° to 20° per minute. The behavior of the sample is observed during the entire period of heating and the temperatures are noted at which the sample begins to fume off, when the flash appears and finally when the tube fills with black smoke.

Refs: 1) M.M. Kostevitch, SS, 23, 156 (1928) 

C. Ignition Times for Flame Exposures.

In order to measure the time required to ignite explosives exposed to flames, several devices have been proposed. Among these the pendulum of Prof Cronquist of Sweden appears to be the simplest (Ref 2). It consists of a metallic pendulum, 2.5-3m long, making a swing of 75cm at a rate of 1cm in 1/50 second. A small sample of explosive is attached to the lower end of the pendulum and, as the pendulum swings, it passes through a row of 5 small flat gas burners arranged in the form of an arch. Each burner has a flame 15cm long. If, for instance, the explosive ignites after passing two burners, the time is 15 x 2 x 1/50 = 0.6 seconds.

Cronquist gives the following results: Small grain Black Powder 3.20 to 4.50 sec; single-base smokeless powder for small arms (nongraphited) 0.44 to 0.90 sec; same, graphited 1.96 to 3.28; Cordite (old type) 0.16 to 0.30 and Ballistite 0.16 to 0.40 sec

N. C. Hansen of Denmark (Ref 3) proposed an apparatus consisting of a pendulum 2.35 meters long swinging through a single gas burner. The duration of contact with flame is regulated by starting the swing from different positions—the greater the lateral displacement at the start of the swing, the shorter the duration of contact with the flame. Test results are expressed in centimeters from the start of the swing to the center of the flame, i.e. the maximum distance at which no ignition takes place in ten trials. These values can be easily converted to the duration of contact with flame.

Refs: 1) Marshall 2 (1917), p 437 2) A.W. Cronquist, SS, 1, 106 (1906) 3) N.C. Hansen, SS, 8, 165 (1913)

8) Ignition Theory of Explosives.

Some of this material was presented in Sect 1 above. Other material will be described under "Thermal Explosions" in a future Vol. Below we list and abstract several references specifically directed to the theory and/or mechanism of ignition of explosives.

On the basis of this study of the preignition, ignition & combustion reactions of Black Powder by thermoanalytical techniques, Campbell and Weingarten (Ref 1) proposed that the preignition reaction (1) is between S & KNO3, which is sufficiently exothermal to activate the propagative combustion reaction between charcoal & KNO3 (2): S + 2KNO3 + K2SO4 + 2NO (1) & 3C + 2KNO3 + K2CO3 + N2 + CO2 + CO (2). In (1) the S is in a molten state & the KNO3 is in its thermally active trigonal form. In (2) the charcoal is in the solid state & the KNO3 is in a molten condition.

Librovich (Ref 2) gives: A math treatment of the ignition of propellants and explosives. It is assumed that ignition is caused by a stream of hot gases and that evapn of the condensed phase results entirely from the heat liberated at the burning surface. Ethylene dinitrate is used as an example. With a surface temp of 473°K, 4.62 cal/cm² is required for ignition.

Khlevnov (Ref 3) analyzed the process of ignition of a nonmelting explosive by a hot metal plate for model conditions (specified). Equations were derived describing the max temp on the surface of the explosive, the temp distribution in the explosive at the time of ignition, the criterion of ignition, etc.

Illuminating Devices and Compositions. Any pyrotechnic device or composition producing intense light and serving to illuminate various objects such as enemy territory at night, either for observation or photography, or for locating targets for bombing etc., or illumination of airfields for landing at night, in rain or fog.

Among illuminating devices may be cited flares (ground or airplane), photoflash bombs, illuminating projectiles (star shells), position lights etc.

Illuminating devices generally produce either white or yellow-tinted light (although some colored lights are also used) of great intensity, ranging from several tens to several hundreds of thousands candle power per square inch. Among the older illuminating compositions used during WWI are:

White light mixture for airplane flares: $\text{Ba(NO}_3\text{)}_2$ 76, Al (powder) 9.75, Al (flake) 8.25, S (flour) 4.0, castor oil or vaseline 2.0%; Intensity about 350,000 candle power (Ref 1, v 2, p 111)

White light for wing-tip flares: $\text{Ba(NO}_3\text{)}_2$ 81.0, Al (flake) 13.7, sulfur 5.3% (Ibid p 225)

White light for position signals: $\text{Ba(NO}_3\text{)}_2$ 66.9, Al (flake) 15.5, sulfur 16.7, $\text{Sb}_2\text{S}_3$ 0.6, stearin 0.3% (Ibid p 206)

French white fire: $\text{KNO}_3$ 57.1, Sb (metallic powder) 19.1, $\text{Pb}_3\text{O}_4$ (red lead) 17.5, sulfur 6.3% (Ref 1, v 1, p 189)

German white illuminating compound: fired from special pistols, contained $\text{Ba(NO}_3\text{)}_2$ 61.5, Al (powder) 20 and sulfur 18.9% (Ibid p 191)

After WWI, most of the research and development in pyrotechnic compositions for the US Army was carried out at Picatinny Arsenal, where the development of white light compositions proceeded along the following lines: 1) Mg metal was introduced in 1931 to replace part or all of the Al in order to obtain higher candlepower 2) Study of white light compositions in 1933 showed that a mixture of Ba and Sr nitrates was more efficient than the Ba salt alone or other oxidizing agents, such as $\text{KNO}_3$, $\text{KClO}_4$ and $\text{NH}_4\text{ClO}_4$. Furthermore, the addition of some Sr to the Ba salt overcame the greenish tinge obtained with the use of the Ba salt alone 3) Further work on white light compositions up to 1945 was concerned mainly with compositions for special tactical purposes such as ground signals, mortar illuminating shells, flares with reduced smoke output, special airport flares and developing substitutes for critical material. The use of Mg in illuminating mixtures is also mentioned by Reilly (Ref 2)

Among the white pyrotechnic compositions used during WWII, the following may be cited:

White Flare Composition: $\text{Ba(NO}_3\text{)}_2$ 66, Al 26, sulfur 6, castor oil 2%. 500g of this powder loaded at 6,000 psi burns for 65 seconds, developing 60,000 candlepower

During WWII, the Germans developed several illuminating mixtures containing 14-28% of polyvinylchloride (either additionally chlorinated or not) together with 17-35% of magnesium powder, 50-61% of oxidizer and 1 to 5% of vaseline or synthetic wax (Ref 3)

Research conducted at Picatinny Arsenal has shown that for certain purposes yellow-tinted lights are preferable to white ones. Therefore, compositions containing sodium salts, such as sodium oxalate, were introduced, as for instance, in the following:

Yellow-tinted flare: $\text{Ba(NO}_3\text{)}_2$ 34, Mg (pre-coated with 6% linseed oil) 36, Al 8, Na oxalate 20, linseed oil 1, castor oil 1%. 300g of this composition pressed at 3,600 psi burned for 280 seconds and developed 556,000 candlepower

Faber (Ref 1, v 1, p 192) gives the following yellow light illuminating composition used by the French in WWI: Mg 67.9, Al (powder) 2.0, Na oxalate 17.4, $\text{KClO}_4$ 2.6, sawdust 9.6 & shellac 0.5%

The use of sodium oxalate for yellow illuminating compositions was also recommended by Reilly (Ref 2)

Brock (Ref 6) claims that the luminosity of flares is increased by the addition of 1-5% MgO. For example a mixture consisting of Mg 50.0, $\text{NaNO}_3$ 42.7, polyvinylacetate binder 4.5 & MgO 2.8% had a luminosity of 7.6 x 10^5 candlepower and a burning time of 2.24 secs

A detailed discussion of the theoretical aspects of illumination and the evaluation of several illuminating compositions is given by Tavernier (Ref 5)

Colored illuminating mixtures may also be used, but they do not produce light of such intensity as do white lights. Several new formulations were developed at Picatinny Arsenal before and after WWII
Some illuminating mixtures are listed in Chemical Abstracts, for example, L. Bohner, CA 32, 2754 (1938): German illuminating composition used in compressed form as a charge for "Very pistols" contained small amounts of cuprene in addition to the usual components, such as magnesium powder, barium nitrate and a binding agent such as shellac.

G. J. Schlatt, CA 33, 4426 (1939): Composition consisting of Ba(NO3)2 36-40, Sr(NO3)2 6-8, Mg (powder) 50-54%, coated with a mixture of linseed and castor oil.

Another colored composition which is wickless and burns with a colored flame consists of: 100 part metaldehyde, at least 5 part AN, and at least 4 parts of a mixture of chlorates and nitrates of flame-coloring metals (Ref 5).

High luminous intensity & color purity produced by flames resulting from the combustion of Group II perchlorates mixed with alcohols or other org solvents was noted in a previous study made to adapt these mixtures to rockets & flares for aerial use. Tried to improve Marine MK2-0 Marker but concluded that no improvement is possible unless MK2-0 is redesigned (Ref 7).


Illuminating Projectiles or Light Rockets (Star Shells). The star shells or light rockets, as used during WWI to light up the landscape and reveal the position of the enemy at night, consisted of a single star attached to a parachute. The illuminating composition consisted of 1 part powdered Al, Mg or Mg-Al alloy, mixed with 2 parts of oxidizing material such as Ba(NO3)2, NaN3, KNO3 and KClO3. Shellac or linseed oil was used as a binding material: oil reduced the rate of burning. Sulfur was added sometimes to make the mixture easier to ignite. These mixtures were ignited by means of a charge of Black Powder (Ref 1).

It should be noted that Ba nitrate is preferable to K nitrate because the latter compound has a tendency to lower the temperature of combustion of the mixture.

Some flares also had compositions similar to that of star shells.

More recently, an illuminating compn for projectiles was prepd by compressing a mixture of pulverized Al/Mn alloy with basic Pb nitrate into a rigid coherent pellet. The pellet was inserted in the nose of a projectile (Ref 2).


Imatrix. A Swedish explosive consisting of K chlorate of constant porosity which is impregnated with a combustible oil, such as petroleum, Diesel oil or Brennöl (d ca 0.8; bp 150-330°). Imatrix can be detonated by a blasting cap. Its temp of explosion is ca 3600°; QC 1163 kcal/kg; specific vol 450-490 l/kg; deton velocity 3000-4000 m/sec, exphn temp 370° ("Temperatur-empfindlichkeit")

Ref: Dr. Langhans, Explosivst, 1962, 86

1. M. Filling. Jellied gasoline filling developed during WWII by Standard Oil Co and used for filling incendiary bombs: isobutyl methacrylate polymer 5.0, fatty acids, such as stearic acid, 2.5, naphthenic acid 2.5, aqueous solution (40%) of caustic soda 3.0, and gasoline 87.


Imidazole or Glyoxaline and Derivatives. See 1,3-Diazole and Derivatives in Vol 5 of Encycl, pp D1165-R & D1166-L.
2-Imidazolidone or Diazacyclopentanone-1.
See Ethyleneurea or Ethylene carbamid in Vol
6, p E291-R

Imidazolines, Imidazolidines, Imidazolones,
Imidazolidones; Their Derivatives and Nitrated
Products. (Imidazolidine is also known as
1,3-Diazacyclopentane or Cyclotrimethylene-
1,3-diamine)

These compounds, cyclic derivatives of urea
or guanidine, are as follows:

2-Imidazoline, \( \text{H}_2\text{C} - \text{N} \searrow \text{NH}_2 \);
\( \text{H}_2\text{C} - \text{N} \nearrow \text{CNH}_2 \);
\( \text{H}_2\text{C} - \text{N} \swarrow \text{CNNH}_2 \);
\( \text{H}_2\text{C} - \text{N} \downarrow \text{CNNH}_2 \);

2-Imidazolidine, \( \text{H}_2\text{C} - \text{N} \searrow \text{C(NH}_2)_2 \);
\( \text{H}_2\text{C} - \text{N} \nearrow \text{CNNH}_2 \);
\( \text{H}_2\text{C} - \text{N} \swarrow \text{CNNH}_2 \);
\( \text{H}_2\text{C} - \text{N} \downarrow \text{CNNH}_2 \);

2-Imidazolone (or glyoxalon), \( \text{HC} - \text{N} \searrow \text{C=O} \);
\( \text{HC} - \text{N} \nearrow \text{CNNH}_2 \);
\( \text{HC} - \text{N} \swarrow \text{CNNH}_2 \);
\( \text{HC} - \text{N} \downarrow \text{CNNH}_2 \);

2-Imidazolidone (or imidazolidinone or ethylene
urea), \( \text{H}_2\text{C} - \text{N} \searrow \text{C=O} \);
\( \text{H}_2\text{C} - \text{N} \nearrow \text{CNNH}_2 \);
\( \text{H}_2\text{C} - \text{N} \swarrow \text{CNNH}_2 \);
\( \text{H}_2\text{C} - \text{N} \downarrow \text{CNNH}_2 \);

These compounds per se are non-explosive, but
they can be nitrated to give explosive and/or
thermally unstable nitrates.

The following substances were prepared &
studied by McKay & Wright (Ref 1) starting with
the reaction of nitroguanidine with ethylene-
diamine to give 2-nitramino-\( \Delta^2 \)-imidazoline (\( \Delta^2 \)
is used to indicate that a double bond is attached
at position 2):

\[
\begin{align*}
\text{CH}_2\text{NH}_2 + \text{HN} & \rightarrow \text{CH}_2\text{N} - \text{CNH}_2 + \text{NH}_3 \\
\text{CH}_2\text{NH}_2 + \text{HN} & \rightarrow \text{CH}_2\text{N} - \text{CNH}_2 + \text{NH}_3
\end{align*}
\]

Nitration of this product with mixed nitric-
sulfuric acid at \(-10\) gave 1-nitro-2-nitramino-
\( \Delta^2 \)-imidazoline,
\[
\text{CH}_2\text{N} - \text{CNH}_2 - \text{NO}_2 + 2\text{NH}_3
\]

If the nitration is carried out with one equivalent
of the nitric acid in acetic anhydride, the same
product is obtained, but if nitric acid is used
in excess, the so-called 1,3-dinitroimidazolidine,
\[
\text{CH}_2\text{N} - \text{NO}_2 + \text{CO}
\]

\( \text{CH}_2\text{N} - \text{NO}_2 \), is obtained. It seems that the
last reaction proceeds as follows:

\[
\begin{align*}
\text{CH}_2\text{N} - \text{NO}_2 \rightarrow \text{CH}_2\text{NH} & \rightarrow \text{CH}_2\text{N} - \text{N} - \text{NO}_2 \rightarrow \\
\text{HN}_3 & \rightarrow \text{CH}_2\text{N} - \text{NO}_2 \rightarrow \text{N} - \text{N} - \text{NO}_2 \\
\text{CH}_2\text{N} - \text{NO}_2 & \rightarrow \text{CH}_2\text{N} - \text{NO}_2
\end{align*}
\]

2-Imidazolidine, \( \text{H}_2\text{C} - \text{N} \searrow \text{C(NH}_2)_2 \);
\( \text{H}_2\text{C} - \text{N} \nearrow \text{CNNH}_2 \);
\( \text{H}_2\text{C} - \text{N} \swarrow \text{CNNH}_2 \);
\( \text{H}_2\text{C} - \text{N} \downarrow \text{CNNH}_2 \);

\( \text{H}_2\text{C} - \text{N} \searrow \text{C(NH}_2)_2 \);
\( \text{H}_2\text{C} - \text{N} \nearrow \text{CNNH}_2 \);
\( \text{H}_2\text{C} - \text{N} \swarrow \text{CNNH}_2 \);
\( \text{H}_2\text{C} - \text{N} \downarrow \text{CNNH}_2 \);

The 1-nitro-2-nitramino-\( \Delta^2 \)-imidazoline is an explo-
sive comparable in brisance and power to RDX,
but it is much more sensitive to impact. Its
thermal stability is low, since 97% is destroyed
by 5 minutes of boiling in water. When 1-nitro-
2-nitroaminimidazoline is dissolved in cold aq
ammonia and then acidified a stable product
responding to the formula
\[
\text{CH}_2\text{NH} - \text{NH}_2
\]

\[
\text{CH}_2\text{N} - \text{NO}_2 \text{NH}_2
\]

is obtained. It is
called 1-nitro-2-amino-2-nitroaminimidazolidine
(Ref 1). Some 2-imidazoline compounds were
reported in Ref 2, and among them
1-Nitro-2-amino-2-imidazole nitrate,
\[
\begin{align*}
\text{CH}_2\text{N} - \text{NO}_2 + \text{C(NH}_2)_2 & \rightarrow \text{NO}_3^{-} \text{N}=36.2\%; \text{mp 161 °}, \\
\text{CH}_2\text{NH} & \text{with decomp}
\end{align*}
\]

The 1-substituted-2-nitramino-2-imidazoline
derivatives were prepd and investigated by McKay
et al (Ref 3). Nitrulation of these products
yielded 1-substituted-2-nitrimino-3-nitroimidazoli-
dines, which proved to be stable at room
temperature. As examples of these compounds may
be cited:
1-\( \beta \)-Nitroxyethyl-2-nitrimino-3-nitroimidazolidine,
\[
\text{CH}_2\text{N} - \text{NO}_2
\]

\( \text{CH}_2\text{N} - \text{CH}_2\text{O} - \text{NO}_2 \); mw 264.6, N 31.82%;
crystals; mp 115–16° with decomp
1-(N-Nitroguanyl)-N-nitro-\( \beta \)-aminoethyl-2-nitrimino-
3-nitroimidazolidine,
\[
\text{CH}_2\text{N} - \text{NO}_2
\]

\( \text{CH}_2\text{N} - \text{CH}_2\text{N} - \text{NO}_2 ) \text{C(NH)NH} - \text{NO}_2 \); mw 350.22,
N 40.00%; mp 161–62° with decomp. McKay et al (Ref 3) do not mention that any of these materials are explosive

Other reactions of the Imidazolidines are described in Refs 4 & 5 and below
1-Nitroso-2-iminoimidazolidine or 1-Nitroso-2-
Imidazolidimine, mp 95–100° (with decomp)
(Ref 6)
1,3-Dinitro-imidazolidine or 1,3-Dinitro-1,3-
Diazacyclopentane,
\[
\begin{align*}
H_2C-N &-CH_2-NO_2 \\
H_2C-N &-CH_N=CH_2 \\
\end{align*}
\]
; mw 162.11, N34.56%, OB–49.4%; decomposes without igni-
tion at 205°, ignites immediately when dropped on a 350° surface. It is explosive & is claimed to be about as impact sensitive as Tetryl (Figure of Insensitivity, FI, of 0.78). Its power is 136% PA. It is prepd by nitrination of diendymethylen-
tetraazacyclicdecamonomohydrochloride,
HCl
\[
\begin{align*}
H_2C-N &-CH_2-CH_2-CH_2-NCH_2 \\
\end{align*}
\]
with 98% nitric acid in acetic anhydride (Ref 8)
1,2-Dinitro-iminoimidazolidine, Silver Salt. (No formula given). When 1,2-Dinitranoimidazolidine,
\[
\begin{align*}
H_2C-N &-CH_2 \\
H_2C-N &-C=NO_2 \\
\end{align*}
\]
, is treated with aq alc AgNO₃, its silver salt is precipitated. This salt exploded on a spatula over an open flame (Ref 7)

Wright, JACS 73, 2213–16 (1951) & CA 46, 1899 (1952) (Reaction of 1-nitro-2-nitroamino-
2-propoyrimidazolidine with acetyl chloride)
6) M.W. Kirkwood, JACS 76, 1936 (1954) & CA 49, 6927 (1955) 7) A.F. McKay et al,
JACS 76, 6374 (1954) & CA 49, 15861 (1955)
8) R.J.J. Simkins & G.F. Wright, JACS 77, 3157 (1955) & CA 50, 3496 (1956)

Imide. See Amides, Imides and Derivatives in Vol 1, pp A168–171 with the following additional entry: Two instances of explosions of ammoniacal Ag solns are described. Both are attributed to the formation and subsequent explosion of Silver Imide, Ag₂NH. In the first instance, a marking ink consisting of a mixt of solns of AgNO₃, NH₄OH, Na₂CO₃ & gum arabic exploded on warming. In the second occurrence, Ag₂O was precipitated by NaOH from an AgNO₃ sp’m; The precipitate was washed, dissolved in NH₄OH, and a few drops of AgNO₃ soln were added until a permanent precipitate reappeared. This soln was then stored in tightly-stoppered dark bottles. Two such bottles exploded, one of them violently, after 10–14 days storage

Explosion is ascribed to the formation of Ag₂NH via:
Ag(NH₃)₂OH = Ag₂NH + 3NH₃ + 2H₂O

The silver imide, Ag₂NH, in its amorphous form is claimed to be very sensitive to heat & shock even when wet
Refs: 1) H. Vasbinder, PharmWeekblad 87, 861 (1952) & CA 47, 4083 (1953) 2) No other refs to explosion or decomp of Imides or Imdid Acid & Derivs were found in CA 1927–
71

Iminobisaceticacid or Iminodiaceticacid or
Iminodiacetic Acid Diazide. See Diglycol-
amidic Diazide in Vol 5, p D1262

Iminodiethanol-dinitrate. See Diil(2-nitroxy-
ethyl)-nitramine or DINA in Vol 5, pp D1240–
1242

Iminohydropurines. See Aminopurines and Derivatives in Vol 1, p A154-L

3-Imino-5-phenylimino-1,2,4-triazoline (N⁹-
Phenylidihydroguanazol), C₆H₅N=C–NH₂;
N≡N:C–NH₂,
mw 173.18, N 40.44%; brownish-red powder, mp—explodes mildly at 138°. May be prep'd from 3-imino-5-phenylimino-1,2,4-triazolidine
Ref: Beil 26, [120]

Iminosuccinylazidic Acid (Iminosuccin-Azidsäure
in Ger.),

\[
\begin{align*}
&\text{HN} \\
&\text{CHCO}_2\text{H}
\end{align*}
\]

mw 156.10, N 35.89%, OH—51.3%; yellow ppt (from aq HCl). Prep'd by dropwise addition of coned aq NaNO₂ to iminosuccinylazidic acid dissolved in cold HCl. Explodes violently on heating
Refs: 1) Bell, not found 2) Th. Curtius & W. Dörr, JPrChem 125, 439 (1930) & CA 24, 3214 (1930)

2-Imino-1,3,4-thiadiazoline and Derivatives.
See Aminothiazole and Derivatives in Vol 1, p A262—263

2-Iminothiazoline and Derivatives. See Aminothiazole and Derivatives in Vol 1, pp A263—265 and the additional entry below:

Nitrosiminothiazoline. HC—S—C=NNO

\[
\begin{align*}
&\text{HC—} \\
&\text{NH}
\end{align*}
\]

129.14, N 32.54%; orange-red amorphous powder; explodes ca 140° & decomp on storage in moist atm; insol in water; sol in alc or eth.
Prep'd by reacting aminothiazole with NaNO₂ in HNO₃
Ref: Beil 27, 155

5-Imino-1,2,3,4-thiotriazoline. See 5-Amino-
1,2,3,4-thiotriazole in Vol 1, p A164-R

5-Imino-2-thion-1,3,4-thiodiazolide. See 5-
Amino, 2-mercaptopo-1,3,4-thiodiazole in Vol 1, p A224-L

IMP. Initial Maximum Pressure (in rockets) is
the value of the pressure when it reaches a
steady state. See Fig under IASP in this Vol
Ref: F. Bellinger et al, IEC 38, 166-67 (1946)

IMPACT, INITIATION OF EXPLOSION BY
Introduction

It is common knowledge that a blow (impact) can initiate explosion in certain substances usually referred to as explosives. Aside from superficial statements, such as the one above, the subject of impact initiation (also called impact sensitivity) of explosives has been shrouded in myth, confusion & misinformation. Until recently, more time & money has been invested, with less return on the investment, in studying impact sensitivity than any other aspects of explosion sensitivity. This was most aptly stated in the following quotation (Ref 4)

"It is now becoming more widely recognized that most of the standard impact and friction methods of measuring the 'sensitivity' of an explosion have little physical significance. This is fair enough: hitting a solid with a hammer or rubbing it with a piece of sandpaper is perhaps an experiment more proper to a carpenter than to a physicist"

F. P. Bowden, F.R.S.

Even the concept of impact sensitivity as a specific property of an explosive is questioned in a recent publication (Ref 17), at least for the response of confined explosive samples to impact

Nevertheless, examination of the response of explosives to more or less controlled impacts did provide qualitative information about the safety of handling these explosives. There is no question that Lead Azide is much more impact sensitive than TNT, and that TNT is much safer to handle than Lead Azide. It is the quantitative meaning of the "much more" that creates confusion and indeed it may have no quantitative meaning at all. Presumably some of the immense effort devoted in past years to impact testing is excusable because it was based on the fond hope (unfortunately unrealized) of finding the ideal explosive—one that is powerful but "insensitive"

We have thus far avoided distinguishing between impact sensitivity, projectile (or bullet) sensitivity and shock sensitivity which also involve impact loading of the test explosive. Shock sensitivity, as the name implies, is the response of an explosive to an externally generated shock. Measurements of shock sensitivity are very reproducible, although many existing measure-
ment techniques only measure the shock sensitivity of a material in the particular environment of the test, i.e., the measured "sensitivity" is not an absolute quantity. The characteristic time scale of a shock sensitivity test is of the order of 1 μsec and peak stresses in the test sample are of the order a few kilobars to several tens of kilobars. Characteristic response times in a falling weight impact test are at least two orders of magnitude greater than in the shock sensitivity test, while peak stresses, in condensed media, are at least two orders of magnitude less. Typical test results are anything but reproducible. Bullet sensitivity tests have times & stresses that lie between shock & impact test values. Generally, reproducibility is rather poor so that in this respect bullet sensitivity resembles impact sensitivity more closely than it does shock sensitivity. For a discussion of bullet sensitivity, see "Bullet Tests" in Vol 2, pp B332-340.

In what follows we will describe some impact machines and impact tests, i.e., the apparatus and methods for measuring impact sensitivity. We will then present and discuss impact sensitivity data for common explosives obtained with these machines by different laboratories. Then we will briefly consider how impact sensitivity tests have contributed to the development of the theory of initiation of explosives. Finally, we will examine impact testing from a theoretical point of view.

**Impact Machines and Impact Tests.** One frequently hears the remark that there are as many types of impact machines as there are explosives test laboratories. This is somewhat misleading. In reality, all laboratories use essentially similar equipment but test procedures, and test data analyses differ. Basically an impact machine is an apparatus that drops a steel weight from a pre-determined height onto a plunger or striker resting on top of the test sample which is placed on a steel anvil.

Procedural differences among laboratories consist mainly in different methods of confining the explosive sample subjected to impact. In some laboratories the explosive is simply spread as a thin layer between striker & anvil. In other laboratories fine sandpaper is placed under the explosive layer. Sometimes the explosive is confined in a small steel cup and the bottom end of the striker is shaped to fit snugly into the open end of the cup. In some tests a brass sleeve is placed under the striker & over the test sample contained in a cup.

Different laboratories use different criteria for determining whether a drop-test resulted in an explosion ("go") or failure ("no-go"). In many test facilities any visual observation of smoke, flash or flame, or any crackle, pop or bang detected by the operator is taken to signify a "go" result. Sometimes sound-meters are used to detect the audible signals of a drop test. In still more elaborate tests, the volume of gas, produced by a "go" result, is measured. In general, for "sensitive" explosives like PETN, distinguishing between "go" & "no-go" is quite easy. For "insensitive" explosives like TNT, this distinction becomes much more difficult, and it is for these "insensitive" materials that operator judgment becomes unreliable and instrumental judgment devices become a necessity.

Most test procedures are designed to obtain the so-called 50% height, i.e., the height of drop for which 1/2 the trials are "go" and 1/2 the trials are "no-go." The so-called "Bruceton" or "up and down" or "stair-case" test is an efficient method of obtaining 50% heights. In this method, if the first trial is a "go" the next trial (using a fresh explosive sample) is made at a pre-determined drop height that is lower than the height of the first trial. Drop heights are lowered in pre-determined steps (using a fresh sample for each trial) until a "no-go" is observed. Now the drop height is increased to that of the last trial before the "no-go." If this trial is a "go" the drop is again decreased in steps until a "no-go" is observed. If it is "no-go" then the drop height is increased in steps until a "go" is observed. This scheme, of increasing the drop height after a "no-go" & decreasing it after a "go," is continued until 20 or 25 drops are made. Statistical methods of choosing the drop height steps and analyzing the test data have been carefully worked out by Dixon & Mood (Ref 1). A concise description of test procedure and analysis of results is found in Ref 5.

The Bruceton test (as expected) gives a good value of the mean drop height, i.e, 50%
height, but \( \sigma \) the std deviation of this mean, derived by statistical methods, often disagrees with observations, ie observed “go’s” and “no-go’s” frequently lie outside a 2\( \sigma \) limit around the 50% height. Martin & Saunders used computer simulation of the Brueton test to assess its statistical validity (Ref 8). Their computer runs could produce the equivalent of 200,000 drop tests and thus provided a good check of the large sample theory on which the theory of the Brueton test is based (although in practice Brueton sample size is far from “large”). Martin & Saunders conclude that a Brueton test of 25 samples yields a good estimate of the 50% height but a poor estimate of its std deviation. To get a good estimate of the latter the minimum test sample size should be 100 samples. They also conclude that test height increments, ie the increase or decrease in drop height after a “no-go” & after a “go” should be equal to about 2\( \sigma \). In practice this means that one needs some estimate of \( \sigma \) before making the test in order to select the best height increments.

The Brueton method is also frequently used in other destructive tests, eg in functioning tests of detonators, gap tests for shock sensitivity, or in tests where the test alters a sample so that it is no longer in its original state.

Brief mention of impact machines is made under “Physical Tests” in Vol 1, p XVII. Because there are many literature references to the following impact machines, and not because they are inherently better than any others, we will now describe the following: Explosives Research Laboratory (ERL), Rotter, Bureau of Mines (BOM) & Picatinny Arsenal (PicArsn).

The ERL machine is used at LASL, LLL & NOL. A detailed view of the striker-anvil region of an ERL machine is shown in Fig 1 & an overall view in Fig 2.
It is claimed that the sandpaper under the explosive sample tends to minimize the effects due to variation in surface roughness of striker & anvil (Ref 9). However, it was also claimed (Ref 10) that sandpaper tends to cushion impact & give misleading results for very "sensitive" explosives like the azides. The Buceton method, usually involving 25 drops, is used at LASL, LLL & NOL. Samples are 30-40 mg and are usually unconfined, though LASL places them in a dimple in the sandpaper (Ref 11). LASL & NOL use a 2.5kg falling weight, but LLL appears to be using a 5kg weight (Ref 18). Microphones and/or pressure transducers are used to aid the operator in his judgment between "go" and "no-go" results for "insensitive" explosives.

The Rotter apparatus as used at AWRE is shown in Fig 3.

A 40mg sample is placed on a brass cap which is inverted over the anvil. The falling weight is normally 5kg. If the volume of gas evolved by a 0.03mg sample is more than 1ml, then the test is considered a "go." The Buceton method is used and results are reported as Figure of Insensitiveness or FI which is obtained by dividing the measured 50% height by the 50% height for a standard RDX sample and multiplying this ratio by 80 (Ref 12). It has been claimed that the volume of gas evolved can be related to the ease of propagation of explosion thru the test explosive. Hornby (Ref 13) presents data that show that increase in drop height (above the 50% height) has little effect on the amount of gas evolved by RDX or by 60/40/1 RDX/TNT/wax. For pure RDX gas evolution is always (with considerable variation) large (16ml). For the RDX mixture gas evolution is always small (1-2ml). This is interpreted to mean that explosion, once initiated, propagates readily in pure RDX but not in RDX/TNT/wax. For Picric Acid gas evolution increases with increasing drop height. According to Wilby, this shows that explosion in PA propagates more readily as impact energy input is increased.

Figure 4 shows a sketch of BOM machine with samples placed in confining cups. This
Fig 5-1. Picatinny Arsenal Impact Test Apparatus
machine is also used with samples spread between sandblasted anvil & striker surfaces. Drop weights vary from 5kg to 1kg but 2kg appears to be the weight most frequently used. The writer’s experience with BOM machines is that they are no better or no worse than the other types—at least for "sensitive" and "moderately sensitive" explosives.

A schematic diagram of the Picatinny Arsenal Machine is shown in Fig 5. The test sample is poured into a hardened steel cup and cup & sample are weighed. This weighing, together with a previous weighing of empty cup, gives the test sample weight. A brass cover is then pressed over the cup and the covered cup is positioned in a recess in the anvil. A vented plug is placed on top & in the center of the brass cover. The falling weight (2kg, 1kg & 1 lb weights are used) impacts the vented plug. The test is begun with a 12-inch drop. If an explosion is observed, drop heights are lowered in 2-inch steps until failure occurs. New sample assemblies are used for each drop. If the initial drop results in a failure, drop heights are increased in 4-inch steps until an explosion is observed. These preliminary tests are used to "zero-in" on the minimum drop height at which at least one "go" is observed in 10 drops (with fresh samples for every drop). One inch below this height no explosions should be observed in 10 drops. It is claimed that this machine is particularly suitable for impact-testing of "insensitive" military explosives such as TNT & Ammonium Picrate (Ref 16).

In addition to the impact machines just described, the search for a "perfect" impact machine has resulted in the development of many designs, most of which are not very different from each other, but are claimed to be "improvements in the state of the art" by their inventors. Below we list, in chronological order, references to and very brief descriptions of some of these machines.

C. E. Bichel "Fallapparat zur Bestimmung der Empfindlichkeit gegen Schlag und Stoss," SS 3, 407 (1908)

W. J. Williams, "Impact Tests with the Drophammer Method of Kast," JFranklinInst 169, 143 (1910)

Anon, "BOM Small Impact Machine," IEC 25, 665 (1933)

C. Hahn "An Improved Falling Weight Apparatus," ArchPharmChemi (Copenhagen) 55, 259 (1948) & CA 42, 6113 (1948)


J. C. Kopiuk describes an apparatus very similar to the PicArcn machine, ASTM Bulletin No 250, 61 (1960)


N. A. Kholevo, "Description of Type I, 2 & 3 Machines," in Theory of Explosives (transl from Russ Teoriya Vzryvnykh Khs Veshestev)


Note: Development of these machines was actually begun in 1946.
Impact Sensitivity Data. In view of the preceding section & the discussion to be presented in the next section, preparing tables of impact sensitivity data for various explosives (even for the common ones) is an exercise in futility. Reported results from different test facilities whether they are for 50 heights or no explosion in 10 trials, etc. and whether the impact stimulus is reported as fall height, fall impulse or fall energy, just do not agree. When internal “standards” are used and results are reported as Figures of Insensitivity or FI chances of qualitative agreement among test facilities are improved but quantitative agreement is rare. About the best one can hope for is a qualitative ordering of impact sensitivity, ie statements like: Lead Azide is more sensitive than RDX which is more sensitive than TNT

The following Tables of Impact Sensitivity data serve to justify the above statements:

Table 1 shows quantitative accord between NOL & LASL (they use the same type of equipment & test procedure), and some agreement between NOL, LASL & Picn Arsn for sensitive & moderately sensitive explosives such as PETN, RDX & HMX. The BOM 50% heights for explosives more sensitive than TNT are much larger than those of the other facilities. The Rotter data (Table 2) gives 50% heights that are even larger than the BOM data. Thus Tables 1 & 2 show that quantitative agreement of impact data among different laboratories is rare.

### Table 2 (Ref 12)

<table>
<thead>
<tr>
<th>Explosive</th>
<th>Median Ht (cms)</th>
<th>Figure of Insensitivity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sulphurless Gunpowder</td>
<td>253</td>
<td>194</td>
</tr>
<tr>
<td>TNT</td>
<td>197</td>
<td>152</td>
</tr>
<tr>
<td>Plastic explosive</td>
<td>197</td>
<td>152</td>
</tr>
<tr>
<td>RDX/TNT/Al/BWX, 41.5/40/8/0.5</td>
<td>190</td>
<td>146</td>
</tr>
<tr>
<td>RDX/TNT/Wax, 60/40/1</td>
<td>152</td>
<td>117</td>
</tr>
<tr>
<td>RDX/TNT, 60/40</td>
<td>137</td>
<td>105</td>
</tr>
<tr>
<td>RDX/Wax, 91/9</td>
<td>127</td>
<td>98</td>
</tr>
<tr>
<td>Octol</td>
<td></td>
<td>94</td>
</tr>
<tr>
<td>Gunpowder</td>
<td>117</td>
<td>90</td>
</tr>
<tr>
<td>Tetryl (CE)</td>
<td>112</td>
<td>86</td>
</tr>
<tr>
<td>RDX, Standard</td>
<td>104</td>
<td>80</td>
</tr>
<tr>
<td>RDX, Military Grade</td>
<td>98</td>
<td>75</td>
</tr>
<tr>
<td>HMX</td>
<td>73</td>
<td>56</td>
</tr>
<tr>
<td>PETN</td>
<td>66</td>
<td>51</td>
</tr>
<tr>
<td>Lead Azide*</td>
<td>113</td>
<td>20</td>
</tr>
<tr>
<td>Lead Stypnate*</td>
<td>69</td>
<td>12</td>
</tr>
<tr>
<td>Lead 2:4 dinitro-resorcinat*</td>
<td>61</td>
<td>11</td>
</tr>
</tbody>
</table>

* These explosives were tested with a 2kg weight, and Lead 2.4 dinitro-resorcinat was used as the standard for comparison with an FI of 11

* σ is the std deviation of the 50% height
** This is really the height at which at least one explosion was observed in 10 trials. One inch below this height there were no explosions in 10 trials
TABLE 3
Comparison of Impact Sensitivity Tests on the Basis of Figures of Insensitivity, FI

<table>
<thead>
<tr>
<th>Explosive</th>
<th>ARDE (Rotter Test)</th>
<th>NOL (ERL Test)</th>
<th>LASL (ERL Test)</th>
<th>Picn. Arsn (PA Test)</th>
<th>LLL (ERL Test)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TNT</td>
<td>152</td>
<td>522</td>
<td>560</td>
<td>150</td>
<td>228</td>
</tr>
<tr>
<td>Comp A-3</td>
<td>98</td>
<td>195</td>
<td>~ 250</td>
<td>140</td>
<td>128</td>
</tr>
<tr>
<td>Comp B</td>
<td>117</td>
<td>200</td>
<td>214</td>
<td>170</td>
<td>116</td>
</tr>
<tr>
<td>75/25 Octol</td>
<td>94</td>
<td>~</td>
<td>138</td>
<td>170</td>
<td>95</td>
</tr>
<tr>
<td>75/25 Cydotol</td>
<td>~</td>
<td>~</td>
<td>132</td>
<td>~ 140</td>
<td>95</td>
</tr>
<tr>
<td>Tetryl</td>
<td>86</td>
<td>126</td>
<td>153</td>
<td>80</td>
<td>80</td>
</tr>
<tr>
<td>HMX</td>
<td>56</td>
<td>87</td>
<td>95</td>
<td>90</td>
<td>94</td>
</tr>
<tr>
<td>RDX (used as Std)</td>
<td>80</td>
<td>80</td>
<td>80</td>
<td>80</td>
<td>80</td>
</tr>
<tr>
<td>PETN</td>
<td>51</td>
<td>40</td>
<td>44</td>
<td>50</td>
<td>31</td>
</tr>
<tr>
<td>Lead Azide</td>
<td>20</td>
<td>5</td>
<td>~</td>
<td>50</td>
<td>~</td>
</tr>
</tbody>
</table>

*ARDE values from Table 2; NOL, LASL & Picn. Arsn values computed by us from the data of Table 1; LLL values computed by us from the data in Ref 18

One might suspect that use of an internal standard might bring better agreement. Consequently, in Table 3, we have used ARDE's definition of Figure of Insensitiveness, FI, to arrange the data of Tables 1 & 2 on a common basis. As seen in Table 3, this procedure does not bring all the results into agreement. Differences in FI appear to be greatest for the least sensitive & most sensitive explosives.

Now even if we abandon all attempts at getting quantitative accord, and merely attempt a qualitative ranking of explosives, i.e. Expl A is more sensitive than Expl B etc, we still find disagreements. As shown in Table 4 there are reversals in sensitivity rankings for RDX, HMX & Tetryl, and Comp B, Comp A-3 & TNT.

To some degree the variation in impact sensitivity data among different test facilities can be attributed to differences in: drop weights (some laboratories use 5kg weights, others 2kg weights etc), in test sample weights, in sample confinement, impact surface roughness, methods of analyzing the data, etc. However when the effects of these variables are examined in any one laboratory, it is generally found that these effects are relatively small, and certainly not large enough to explain the large differences in 50% heights or even FI's obtained by different laboratories for the same series of explosives.

TABLE 4
Qualitative Ranking of the Impact Sensitivity of Common Explosives
(Based on FI data of Table 3)

<table>
<thead>
<tr>
<th>Explosive</th>
<th>ARDE</th>
<th>Picn. Arsn</th>
<th>LLL</th>
<th>LASL</th>
<th>NOL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lead Azide</td>
<td>1</td>
<td>1 (b)</td>
<td>~</td>
<td>~</td>
<td>1</td>
</tr>
<tr>
<td>PETN</td>
<td>2</td>
<td>2 (c)</td>
<td>2 (c)</td>
<td>2 (c)</td>
<td>2</td>
</tr>
<tr>
<td>RDX</td>
<td>4</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>HMX</td>
<td>3</td>
<td>5</td>
<td>4</td>
<td>5</td>
<td>4</td>
</tr>
<tr>
<td>Tetryl</td>
<td>5</td>
<td>3 (d)</td>
<td>3 (d)</td>
<td>6</td>
<td>5</td>
</tr>
<tr>
<td>Octol</td>
<td>6</td>
<td>9 (f)</td>
<td>6</td>
<td>5</td>
<td>~</td>
</tr>
<tr>
<td>Comp B</td>
<td>8</td>
<td>6</td>
<td>7</td>
<td>7</td>
<td>7</td>
</tr>
<tr>
<td>Comp A-3</td>
<td>7</td>
<td>8</td>
<td>~</td>
<td>8</td>
<td>6</td>
</tr>
<tr>
<td>TNT</td>
<td>9</td>
<td>7</td>
<td>9 (e)</td>
<td>9</td>
<td>8</td>
</tr>
<tr>
<td>Explosive D</td>
<td>~</td>
<td>9</td>
<td>~</td>
<td>10</td>
<td>9</td>
</tr>
</tbody>
</table>

(a) 1 is the most "sensitive" and 9 is the least "sensitive" explosive
(b) FI values make it appear much less impact-sensitive relative to PETN than FI values from other facilities
(c) Assuming that LA would have had lower FI value had it been tested
(d) Same FI as for RDX
(e) Assuming that Comp A-3 is more sensitive than TNT
(f) Same FI as Expl D


It might appear then that impact testing is a complete exercise in futility and should be abandoned. This is definitely the wrong conclusion. Impact “sensitivity” properly interpreted has its uses. This is aptly stated by Hornig (Ref 14):

“Sensitiveness measurements should be considered as dimensionless, although we often do give units to them. The numbers gain their significance only by comparison. Fortunately, we have today a considerable storehouse of experience in the handling and use of a large number of explosives and explosive compositions. We also have laboratory measurements of their sensitiveness. When a new compound or composition is tested, its standing can be compared with the known explosives and the new one can be categorized as being similar to one of the old ones. This gives a reasonable knowledge of the degree of hazard faced.”

**Theoretical Considerations.** In the preceding sections we have been rather negative about impact testing of explosives. We have stressed the great variability of impact data and the lack of a quantitative characterization of impact sensitivity. Yet impact testing has triggered important developments in the theory of initiation of explosives. The classic work of Bowden & his school on this subject is summarized in two excellent monographs (Refs 2 & 3) and in a more recent publication (Ref 15). However the hot spot theory, so successfully championed by Bowden, cannot explain all the strange and apparently contradictory effects observed in impact testing of explosives. Some of these weird phenomena led Kholevo (Ref 6) and later Bobolev et al (Ref 17) to develop a rather different theory of impact initiation than that of Bowden. We will now examine both of these theories in some detail.

Much of Bowden’s theory has already been described under Hot Spots in this Volume (we shall henceforth refer to this as Ref 19). Briefly what Bowden proposed was that most impact initiations are produced by very localized regions of high temperature ie hot spots. He showed that a falling weight impacting an explosive sample can generate hot spots in the following ways: a) by adiabatically compressing air (or vapor of the test explosive if impacts are made at sub-atmospheric pressure or at elevated temperatures) bubbles trapped in or purposely introduced into the explosive sample; b) by intercrystalline friction (for high-melting explosives); c) by friction of the impacting surfaces; d) by plastic deformation of a sharply-pointed impacting surface; e) by viscous heating of the impacted material as it flows past the periphery of the impacting surfaces—appreciable heating in this case is claimed to occur only for quite energetic impacts.

The many mechanisms of generating hot spots, coupled with complex nature of the development of a hot spot into an explosion (see Ref 19) or the decay of a hot spot without explosion, offer a convincing explanation for the observed variability of impact test results. Thus, in test method a), hot spot formation via adiabatic compression may be dominant. In test method b), viscous heating may be the main mechanism for heat generation. In test method c), conditions may favor rapid quenching of hot spots, and so forth. Even with a given test method several hot spot generating mechanisms may be operative and these can then develop into explosion by different paths. Consequently even slight changes in procedure or conditions may produce quite drastic changes in impact test results.

Some of the quantitative consequences of hot spot theory were presented in Ref 19. A qualitative discussion of heat flow in a compressed gas bubble hot spot was also presented in Ref 19. The necessity of having enough, but not too much, liquid or solid spray or foam within a compressed gas bubble (in order to have sufficient heat flow from the bubble to the surrounding condensed explosive) provides another hard-to-control variable in impact testing and thus increases the variability of test results.

Until now we have studiously avoided defining what constitutes an explosion or “go” result in an impact test, except mentioning that “go’s” are judged on the basis of sound, flash, gas volume etc. What physical events produce these observed manifestations? Based primarily on the studies of Bowden (Ref 2) it is clear that in an impact test “explosion” frequently starts as a relatively mild deflagration which can propagate as slowly as 10m/sec, which turns into a much more violent deflagration, (100-1000m/sec), in the confined (by striker & anvil) explosive. For many “insensitive” explo-
sives this is all that occurs. For the more "sensitive" explosives (PETN, RDX) the fast deflagration stage is followed by a true detonation. The only known exceptions to the sequence of deflagration-detonation are the heavy-metal azides. For Lead & Silver Azides impact creates either a full-blown detonation (no deflagration stage) or a failure (Ref 2).

During impact the explosive sample is at an elevated pressure, and the initial deflagration of the sample consequently occurs at elevated pressures. As pointed out by Andreev (Ref 7), the critical thickness for the deflagration of secondary explosives is much greater than the usual sample thickness in an impact test. Since the critical thickness decreases as ambient pressure increases, impact not only generates the heat required to initiate deflagration but also provides the necessary pressure for the deflagration. Highly localized "explosion" upon impact can be the result of too rapid a pressure drop in the impact system. Thus, deflagration may die out if the pressure in the system becomes too low (approaches atmospheric). These considerations re-emphasize the difficulty or even impossibility of separating the effects of initiation and propagation in an impact-generated explosion.

A rather different view of impact initiation to that of Bowden et al, is presented by the Russian school. The originator of this school appears to be Kholevo (Ref 6), supported by Andreev (Ref 7), with a recent and detailed presentation of this point of view provided by Afanas'ev & Bobolev (Ref 17) (from now on we will refer to these writers as A & B without listing the Ref). Briefly stated, A & B consider that the prime mechanism of hot spot generation in solid explosives is by inelastic deformation of the entire impacted explosive sample (Kholevo (Ref 6) calls this "flow").

To understand what follows we must briefly digress and describe three impact machines constantly alluded to in the Russian literature—the so-called Kholevo No 1, No 2 & No 3. Kholevo No 1 is quite similar to Kast's impact machine (see section on Impact Machines). The explosive sample is placed between a striker & an anvil which are contained within a sleeve. Since striker & anvil fit rather loosely into the sleeve there is a small air gap into which the explosive may be extruded by impact. Kholevo No 2 has no sleeve and the sample can "flow" or deform unimpeded upon impact. Kholevo No 3 is essentially the same as No 1 except that there is a very tight fit between striker/anvil & sleeve. If the sample is spread uniformly over the entire striker-anvil contact area, sample deformation is totally impeded in Kholevo No 3.

Unfortunately, the translation of A & B's book is rather uneven and it is difficult to capture all of the finer details of their presentation. Moreover their presentation is somewhat diffuse and lacks a good summary. Below we will summarize the main thrust of A & B's ideas and then select (from various portions of A & B's book) supporting evidence for these ideas.

**Summary:**
1) Impact Initiation is a thermal process that occurs heterogeneously at localized regions of high temperature—hot spots

2) As usual, the impact initiation process is divided into two stages; in the first stage part of the explosive is heated to a critical temperature and in the second stage these heated regions self-ignite.

The actual impact (at fall hammer velocities of 1-5 m/sec) can be considered as a pseudo-static process, i.e. the rate of loading is much slower than the sound velocity in either the apparatus or the compacted explosive sample.

3) The maximum hot spot temperature is limited by the melting point of the impacted medium, but this is the melting point at the elevated pressure of the impact and not the ordinary melting point at one atmosphere.

4) A necessary but not sufficient condition for initiation by impact is that impact pressure (stress) be sufficiently high so that the melting point of the explosive is raised above some critical temperature, $T_{CR}$. For $T > T_{CR}$ the explosive, in the hot spot, will decompose adiabatically in times of the order 10μsec, which has been observed experimentally. The relation between $T_{CR}$ & the critical stress is then expressed by:

$$ P_{cr} = (T_{cr} - T_M)/\alpha $$

where $P_{CR}$ is the max critical stress generated in the explosive by the impact and $T_M$ is the melting point of the explosive at one atmosphere. This relation is based on the assumption that
\[ T_m(P) = T_m + \alpha P \]  

(1a)

where \( \alpha \) is a constant which, according to Ref. 4a, has the approximate value of 0.02°C/atmosphere for all explosives. For most of the common high explosives \( 400 \leq T_{cr} \leq 600 \) °C and the corresponding \( P_{cr} \) is then \( \approx 10 \) kbar. Furthermore, the hot spot radius for these explosives is of the order of 10^{-4} cm. It must be emphasized that Eq. 1 says nothing about how the hot spot is created. All it says is that impact must generate a stress in the explosive at least equal to \( P_{cr} \) or \( P_{impact} \geq P_{cr} \). This criticality condition could be satisfied by slow hydrostatic compression of the sample under circumstances where explosion is known not to occur.

5) If, as stated above, \( P_{cr} \approx 10 \) kbar, then the question arises of how stresses of this magnitude can be generated by impact in explosive compacts whose compressive strengths are about 100 fold less than \( P_{cr} \). For unconfined explosive compacts (in Kholevo No 2) this can only occur with “thin” explosive compacts, i.e., with explosive layers whose \( h/D < < 1 \) where \( h \) and \( D \) are explosive thickness & diameter. The relation between, \( \sigma_u \), the ultimate compressive strength of the explosive compact & the average stress, \( \bar{P} \), at which the compact fails via brittle fracture (in compression) is:

\[ \bar{P} = \sigma_u(1 + D/3\sqrt{3}h) \]  

(2)

The stress varies over the sample (compact) surface such that the peak stress is (2 to 2.5)\( \bar{P} \) at the center and \( \sigma_u/3 \) at the periphery (where \( \bar{P} \) is the average stress on the sample). Since \( P_{cr} \ll \sigma_u \) for most explosives, and the criticality condition is \( P > P_{cr} \), effective hot spots cannot form near the sample periphery. Effective hot spots are also not generated at the center of the sample because inelastic deformation of the sample is a minimum near its center. As will be shown, A & B claim that non-uniform inelastic deformation of the entire sample generates hot spots.

Because of the variation in stress over the sample, A & B relate all stress effects to \( \bar{P} \) the average stress. Now if we assume that there exists an average critical stress \( \bar{P}_{cr} \), above which explosions occur, and that it lies on the \( \bar{P}_u \) vs \( h \) (or \( h/D \) if \( D \) varies) plot of Eq 2, then

\[ \bar{P}_{cr} = \sigma_u(1 + D/3\sqrt{3}h_{cr}) \]  

(2a)

defines a critical thickness \( h_{cr} \). This critical thickness is a specific property of the explosive if the impact is made on unconfined explosive samples (ie in Kholevo No 2 type impact machines) with a striker of fixed diameter.

According to Eqs. 2 & 2a, one should present experimental data in the form of plots of \( P \) vs \( h/D \) or \( h \) (if \( D \) is a fixed sample diameter). This will be done in a following section.

6) We will now present a summary of A & B's description of the processes that generate hot spots in impacted layers of explosive. The processes are complex but fortunately rather drastic simplifications lead to conclusions that are in accord with experimental results. In considering non-isothermal shear on plastic bodies, A & B conclude that this shear will result in a discontinuity in the straining rate and consequently in heat generation by the transformation of elastic energy stored in the sample into deformation work at the shear plane. These processes are always localized and continue spontaneously until all the stored elastic energy (and the kinetic energy of relative motion, along the shear plane, of the upper & lower portions of the sample) of the sample is used up. If there is sufficient energy for portions of the sample (along the shear plane) to melt the process continues as deformation work of the viscous layer. When the rapidly-deforming fluid phase is created, heat transfer in it occurs via convection and is more rapid than heat transfer in the solid which is by conduction. Thus the liquid phase tends to stay at temperatures that are near the melting point (at that pressure) because of rapid heat losses particularly to any solid material dispersed within the liquid. Some of this solid material will melt and absorb energy thereby (heat of fusion). This model is obviously in accord with Eq 1 which states that the critical temperature is the temperature of melting at \( P_{cr} \).

A & B also present arguments that indicate that inhomogeneities in the solid, unless they are as large as 0.1mm, do not affect the heating mechanism. They also argue that Kholevo's original viscoplastic model (Ref 6) is unrealistic and that their model of a brittle body is closer to reality.

7) Thus far we have presented A & B's ideas
on the initiation of solid explosives by impact. In the real world initiation and propagation are very difficult to separate. Consequently any experimental measurement of "impact sensitivity" will depend on the processes that control initiation and propagation under the particular conditions of the measurement. A & B point out that the deformation and fracture of the solid explosive, which creates the hot spots required for initiation, may also aid transition to detonation because the deflagration started by the hot spots propagates into regions of greater surface area (fractured regions) where it will accelerate. Unless some quenching mechanisms are also present, the accelerating deflagration will eventually become a stable detonation.

Based on an analysis of the penetration of a thick, rigid plastic body by a rigid cylinder (whose radius is smaller than the radius of the body), A & B arrive at

\[ \frac{P_{cr}}{\sigma_u} \leq 3 \text{ to } 4.8 \]  

(3)

and empirically

\[ P_{cr} = 2F_{cr} \]  

(4)

Since the force \( F \) on a sample is \( \pi/4PD^2 \), the minimum force for initiation is:

\[ F_m = \pi/4P_{cr}D_m^2 = (3-4.8)\pi/4\sigma_uD_m^2 \]  

(5)

where the minimum sample diameter, \( D_m \), must satisfy the conditions required for propagating detonation in the particular thick explosive sample. The obvious condition is that

\[ D_m = d_{cr} \]  

(6)

where the \( d_{cr} \) is the critical detonation diameter, ie the diameter below which detonation will not propagate no matter how it is initiated. Moreover a further limitation is that \( D_m \) must exceed some limiting dimension \( a_1 \) which is required for the generation of hot spots capable of producing self-propagating chemical reaction. For secondary explosives, one expects \( d_{cr} \gg a_1 \), but this may not be true for primary explosives, eg \( d_{cr} \approx 10^{-2} \text{ mm for Lead Azide}; \) in which case

\[ D_m = a_1 \]  

(6a)

The above considerations were for thick samples, and are inapplicable to the usual conditions of an impact test, except for impacts on a single crystal of a primary explosive. For most secondary explosives the condition of Eq 3 does not agree with experiment, ie for these explosives the observed \( P_{cr} > (3-4.8)\sigma_u \). Thus another model must be developed which takes into account the appreciable increase in \( \sigma_u \) that is possible in thin samples.

We quote A & B:

"Suppose a rigid flat surface supports an explosive layer, which is subject to impact by a rigid cylindrical body (of diameter \( D \)) with flat base. Fig 6a illustrates impact on a layer of high density, and Fig 6b impact on a layer of low density, for example bulk density. In the latter case, the explosive should be strongly compacted in the impact zone before a significant pressure rise begins.* The deformation

\[ \text{Fig 6a. Layer density close to maximum} \]

\[ \text{Fig 6b. Layer of low density (pressing of the material takes place at the beginning of the impact)} \]

*Note: Typically crystal density/bulk density \( \sim 2 \) and a typical striker velocity, \( u_0 \), is \( \sim 5 \text{ m/sec}. \) The minimum time for a loosely packed explosive to be compressed on impact to near crystal density is then \( \sim h/2u_0 \). For \( h = h_{cr} = 0.2 \text{ mm} \) this amounts to \( 2 \times 10^{-5} \text{ sec or considerably longer than the initiation time of less than the } 10^{-2} \text{ sec claimed by A & B for compacted explosives. A & B do not comment on this. If, as A & B suggest, no appreciable heating occurs until the sample is compressed to near crystal density, the "long" compression time should not affect their subsequent conclusions, but in practice one should observe longer times from impact to ignition for loosely packed explosives than for densely packed explosives.}
conditions in the impact zone are similar to the conditions in instrument No 2 (the lateral pressure of the explosive mass surrounding the striker can be disregarded, since the explosive ultimate strength even in the cast state is very low). When the critical explosion initiation conditions are satisfied, ignition takes place in the impact zone. Spreading out from the ignition spot, the explosive transformation should become detonative in order to advance beyond the compression zone. It is known that detonation can only propagate through a charge of diameter not less than $d_{cr}$, the critical detonation diameter (for a charge of cylindrical form). In our case the charge through which detonation should pass is a layer of thickness $h$; detonation is only possible when the thickness $h$ is not less than some critical value $l_{cr}$, where $l_{cr}$ is expected to be proportional to $d_{cr}$, i.e., for detonation to propagate beyond the impact zone it is necessary that $h > l_{cr} = kd_{cr}$ ($k$ being a proportionality factor, apparently close to 1/2). We draw attention to the fact that $l_{cr}$ in the compression zone (a two-sided shell) is smaller than in the uncompressed layer. Thus, the propagation of detonation is limited by the uncompressed explosive, in the first place by the emergence from the compression zone into the uncompressed explosive.

Now from Eq 2, $F_u \gg F_{cr}$ & $h < l_{cr}$ we obtain for thin explosive layers

$$F_m = \frac{\pi}{4} D_m^2 \bar{P}_{cr} \tag{6}$$

and

$$D_m = 3\sqrt[3]{l_{cr}} (P_{cr} - \sigma_u)/\sigma_u \tag{7}$$

If the diameter $D$ of the striker is less than $D_m$ or the force is less than $F_m$ then detonation of the whole charge is impossible (provided, of course, that the velocity of the striker is much less than the velocity of sound) and explosion may occur only in the impact zone. Consider the variations which may be encountered in the case of impact on a thin layer:

1) When $h < l_{cr}$, detonation of the whole layer is impossible; explosion in the impact zone is possible if

$$F > \frac{\pi}{4} D^2 \bar{P}_{cr}$$

2) When $h \gg l_{cr}$, then for $D < D_m$ detonation of the whole layer is impossible; explosion may occur only in the impact zone after one or several attempts if

$$F > \frac{\pi}{4} D^2 \bar{P}_{cr}$$

After each failure the layer thickness under the striker becomes less than $l_{cr}$, with each failure material is ejected from the compression zone and some of the explosive surrounding the striker is scattered); for $D > D_m$, detonation of the whole explosive layer may be initiated if

$$F > \frac{\pi}{4} D^2 \bar{P}_{cr} \gg F_m$$

Thus, for impact on a thin layer detonation can be initiated over some range of the impact parameters $F$ and $D$, respectively equal to or greater than $F_{min}$ and $D_{min}$, if the layer thickness $h$ is equal to or greater than $l_{cr}$.

Analogous to the requirement of $d_{cr} \geq d_1$ for thick samples, we now have $d_{cr} \geq h_1$. For $h_1 > d_{cr}/2$ (encountered only for primary explosives, if at all) $l_{cr}$ in Eq 7 should be replaced by $h_1$. Also, to a good approximation, for $d_{cr}/2 > h_1$ (the usual case for secondary explosives) we replace $l_{cr}$ by $d_{cr}/2$ in Eq 7, which then becomes

$$D_m = 2.6 \{\bar{P}_{cr} - \sigma_u(T_o)\} d_{cr}(T_o, \mu, \rho_o)/\sigma_u(T_o) \tag{8}$$

where the terms in ( ) indicate functional dependence, e.g., $d_{cr}$ is a function of initial temperature, explosive particle size, and initial density of the explosive sample etc.

Let us now examine how these initial conditions affect $D_m$ and consequently the force required for initiation, remembering that $F_m$ increases as the square of $D_m$, thus a fairly small increase in $D_m$ can lead to a substantial increase in $F_m$ and consequently a decrease in impact sensitivity.

For pure primary & secondary explosives (except for borderline HE such as Ammonium Nitrate or Ammonium Perchlorate) $d_{cr}$ decreases as $\rho_o$ increases until $\rho_o$ approaches very close single crystal density when $d_{cr}$ may increase drastically. Thus if we limit ourselves to $\rho_o < 0.9 \rho_{cryst}$, increases in $\rho_o$ (according to Eq 8) should result in a greater sensitivity to impact. This is quite the opposite of what is found for shock initiation and will be examined more closely later on. At a fixed density, $d_{cr}$ increases as $\mu$ increases (see A & B, p 90). This increase is fairly pronounced at small $\mu$ ($< 0.2$ mm) but levels off & becomes almost asymptotic at large $\mu$ ($> 0.4$ mm, except for cast TNT or TNT with 1% paraffin oil). Thus an increase in $\mu$, as expected, leads to a decrease
in impact sensitivity. The effect of $T_o$ is more complicated. As expected, $c_{cr}$ decreases as $T_o$ increases, but so does $\sigma_U$. Since $d_{cr}$ & $\sigma_U$ occur in the numerator and denominator of Eq 8, and also as a negative term in the numerator, it is not possible to determine the effect of $T_o$ without specific information on the values of $d_{cr}$ & $\sigma_U$ at a particular $T_o$ for a particular explosive. Furthermore, and this is not considered by A & B, a $T_o$ appreciably above room temperature means that less energy needs to be expended for hot spots to reach $T_{cr}$ than for the same system initially at room temperature.

Because of all these compensating factors, the effect of $T_o$ on impact sensitivity is expected to be fairly small, but there may be regions of $T_o$ where the effect could be appreciable.

The chemical reactivity of the impacted explosive is manifested primarily thru $P_{cr}$ which controls $T_{cr}$ (see Eq 1): the more "reactive" the explosive the lower the $T_{cr}$. Of course $d_{cr}$ also depends to some degree on chemical "reactivity." Increased reactivity affects $P_{cr}$ & $d_{cr}$ in the same direction, i.e. towards increasing the impact sensitivity.

8) All the above considerations were primarily for well-compacted solid explosive wafers impacted in a laterally unconfined state (Kholevo No 2 machine). How do these considerations change if impacts are made in a Kholevo No 1 machine which provides some lateral confinement of the explosive sample? The degree of lateral confinement depends on $g$ the thickness of the air gap between striker/anvil and the surrounding sleeve. Obviously if $g$ is large Kholevo No 1 impact results should be indistinguishable from Kholevo No 2 results. According to A & B, it is very difficult to initiate explosives in the Kholevo No 3 machine in which $g \rightarrow 0$. The expression, analogous to Eq 2, proposed by A & B for Kholevo No 1 impacts is

$$P_U = \sigma_U (1 + h/g2\sqrt{3} + D/h3\sqrt{3})$$ (9)

According to A & B, the critical stresses for the No 1 & No 2 machines are essentially equivalent, therefore, comparing Eqs 2a & 9 (where the $h$'s are replaced by $h_{cr}$'s) the critical sample thickness is greater in Kholevo No 1 than the critical thickness in Kholevo No 2.

The Kholevo No 1 machine (with small $g$) has the advantage that any localized ignition (because of confinement) spontaneously leads to explosion. Thus characterizing the impact sensitivity of a laterally-confined explosive requires only the determination of conditions necessary for initiation. But this determination depends on both the properties of the explosive and the conditions of deformation (magnitude of $g$). Thus no specific explosive sensitivity can be defined for laterally (more or less) confined impacts. By way of contrast, in unconfined impacts (Kholevo No 2) the concept of explosive sensitivity has meaning, but now must determine now only the conditions for initiation but also those for propagation.

9) The wide scatter and probabilistic nature of impact measurements are attributed by A & B not to any statistically controlled initiation or propagation processes but primarily to fluctuations in $\sigma_U$ for the explosive samples.

10) We have attempted to present A & B's views on impact initiation of solid explosives. In spite of the concerted effort of these authors to develop a quantitative or at least semi-quantitative picture for impact sensitivity, A & B realize that they have not achieved that goal. We quote:

"Consider briefly the relative character of sensitivity evaluation, namely, that it is possible only to speak of a higher or lower sensitivity of one explosive as compared with another. It would be meaningless, for example, to say that the sensitivity of some explosive is so many times higher than that of another explosive. No single sensitivity has this property: in this sense it is analogous with such a property of explosives as, for example, detonation capacity. It is most convenient to compare explosives with respect to sensitivity, against those explosives about whose sensitivity fairly firm empirical ideas exist, eg the explosives of a reference series."

This writer heartily subscribes to these conclusions.

Experimental Evidence: We will now discuss the evidence presented by A & B in support of their views of impact initiation described above. After a brief description of their main experimental techniques we will attempt to present experimental data that support each...
of the ten topics of the previous section. Since these overlap to some extent the experimental data will also overlap occasionally.

The best evidence in support of A & B's views comes from experiments performed in the apparatus shown in Fig 7. This apparatus yields stress-time records which are analyzed to give the average stress-time, \( P \) vs \( t \), striker velocity-time, \( u \) vs \( t \), sample displacement time, \( \Delta h \) vs \( t \), and displacement vs average stress curves, \( \Delta h \) vs \( P \), shown in Fig 8. The material from the compression zone. Explosions are also characterized by sharp pressure drops. For primary explosives A & B present convincing evidence that explosion occurs before fracture and in times of the order of 10\( \mu \)sec. For secondary explosives the evidence is not as clear-cut. Incidentally, because of the "slow" (relative to sound velocity) rate of loading the impact process is essentially a static process, i.e. stresses in the striker & sample are equalized and there is no effect of impedance mismatch found in shock loading.

1) No new evidence is presented in support of the thermal nature of impact initiation. However A & B give a detailed analysis, based on results obtained with the apparatus just described, of the energy distribution in the Kholevo No 2 machine. Thus,

\[
W_o = W^{\text{pl}}_m + W^{\text{el}}_m + W_S + W_{\text{ex}}
\]

where \( W_o \) is the kinetic energy of impact, \( W^{\text{pl}}_m \) & \( W^{\text{el}}_m \) are the work of plastic & elastic deformation of the striker & anvil, \( W_S \) are the irreversible seismic energy losses, & \( W_{\text{ex}} \) is the elastic and plastic work in deforming the explosive charge. In a typical case (for which explosion is observed) if all of \( W_{\text{ex}} \) goes into uniform heating of the explosive the temperature rise in the explosive will amount to only 30°.

2) There is no direct experimental justification for the assumption that the initiation process is separable into a heat-up stage (to \( T_{\text{Cr}} \)) & a self-ignition stage (at \( T_{\text{Cr}} \)). Justification comes from the self-consistency of the overall picture of impact sensitivity developed by A & B.

3) & 4) Here again evidence is circumstantial. The existence of \( P_{\text{Cr}} \) (ideally no explosions below \( P_{\text{Cr}} \) & no failures above \( P_{\text{Cr}} \)) is required by the model which is summarized by Eqs 1 & 1a. Experimental verification of the existence of \( P_{\text{Cr}} \) will be presented below. It must be emphasized, however, that the model of Eqs 1 & 1a is not the only one capable of explaining the existence of \( P_{\text{Cr}} \).

5) We will now present data in support of the existence of \( P_{\text{Cr}} \) and Eq 2. Since most of these data are shown as functions of \( h \), the original thickness of the explosive samples, we must first examine the validity of using \( h \) rather than the thickness of an impacted sample.

Pre-pressed explosive wafers (at around 90%
of crystal density) are shown to behave almost elastically for deformations of less than 10% and then fracture. If one assumes that explosion occurs before this first fracture (at least for \( h_{cr} \geq h \); see below & item 7 of this section) then thickness of the impacted sample just before explosion, is very close to that of the original sample.

\[ P_{cr} \, \text{kg/cm}^2 \]

Fig 9. Average pressures upon fracture (\( P_{ult} \) — solid curves) and upon explosion (\( P_{ex} \) — dashed curves) vs charge thickness of HMX (●), RDX (▼), Tetryl (▲) and TNT (⊗).

\[ P_{ex} \, \text{kg/cm}^2 \]

Fig 10. Average pressures \( P_{ult} \) and \( P_{ex} \) vs charge thickness for PETN (●) and Trinitophenol (▲).

Points marked ▲ refer to Dina.

\[ P_{ex} \, \text{kg/cm}^2 \]

& for single crystals. Particle size affects the spread in \( P \) (but not the average) in the sense that spread is decreased as particle size decreases. Furthermore, A & B state that pre-pressing the explosives at 200atm or 20,000atm has no effect on \( P \). The writer estimates that PETN & RDX at 200atm have a bulk density, \( \rho_o \), of about 80% of crystal density. Thus, according to A & B, \( P \) does not depend on \( \rho_o \) in the range 80% ≤ \( \rho_o \) ≤ 100% of crystal density.

\[ P_{cr} \, \text{kg/cm}^2 \]

Fig 11. Spread of the measured pressures upon fracture (\( P_{ult} \) — circles) and upon explosion (\( P_{ex} \) — crosses) for RDX.

Figure 11 shows the typical variation in experimental data and indicates the accuracy with which \( P_{cr} \) can be determined. Incidentally Eq 2 fits the data of Figs 9, 10 & 11 to within experimental error. Two other observations are of interest: \( \overline{P} \) does not depend on hammer drop height (or drop velocity).

The empirical relation,

\[ \sigma_{u} = 4.2 \, T_{m} (\sigma_{u} \text{ in kg/cm}^2 \& T_{m} \text{ in } ^\circ \text{C}) \] (10)

appears to be well-obeyed by explosives for which data are available.

The critical stresses and critical thickness, obtained from Figs 9, 10 & 11 are summarized in Table 5.
<table>
<thead>
<tr>
<th>Explosive</th>
<th>$P_{cr}10^{-3}\text{kg/cm}^2$</th>
<th>$h_{cr}\text{mm}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>TNT</td>
<td>11</td>
<td>0.08</td>
</tr>
<tr>
<td>Trinitrophenol</td>
<td>9.5</td>
<td>0.11</td>
</tr>
<tr>
<td>Tetryl</td>
<td>6.4</td>
<td>0.12</td>
</tr>
<tr>
<td>RDX</td>
<td>7.0</td>
<td>0.25</td>
</tr>
<tr>
<td>HMX</td>
<td>6.4</td>
<td>0.43</td>
</tr>
<tr>
<td>PETN</td>
<td>4.8</td>
<td>0.27</td>
</tr>
</tbody>
</table>

6) To show that shear, due to a discontinuous rate of straining, is at a maximum at the upper and lower (contact) planes of an explosive sample, impacted in Kholevo No 2, and zero at its mid-plane, A & B performed the following experiment. Two sets of 1mm thick HMX samples were prepared: set 1 contained a thin layer of Lead Azide at its mid-plane, set 2 had a thin layer of Lead Azide on the upper, lower or both sample surfaces. Set 1 samples behaved just like pure HMX in that only fractures and no explosions were observed ($h > h_{cr}$, see Fig 13). All set 2 samples exploded.

They also used set 2-type PETN, RDX, Tetryl & TNT to show that $P_{cr}$ for Lead Azide is $2.6 \pm 0.2$ kbar and that $P_{cr}$ (at least for Lead Azide) is independent of $h$ or $h/D$.

7) The experimental results discussed below deal with certain aspects of initiation and propagation of impact explosions but some of discussion overlaps topics 5 & 6.

The usual charge thickness ("standard" charge) for HE in Kholevo No 2 is ~0.4mm (~50mg of HE). At this charge thickness anomalous behavior is encountered with some of the "less sensitive" explosives. Figure 12 shows explosion frequency as a function of impact energy for three explosives (HMX, RDX & Tetryl) all at "standard" (or nearly standard) charge-thickness. Note that HMX at $h = h_{cr}$ gives the expected "S" shaped frequency curve. For RDX $h > h_{cr}$ and the frequency curve is flattened for $f > 1$. For Tetryl $h > h_{cr}$, results are erratic, and $f$ is almost independent of impact energy. Explosion frequency, $f$, never exceeds ~0.2 for an almost 4-fold increase in impact energy.

On the basis of experimental evidence and theoretical considerations, A & B conclude that the frequency curve ($f$ vs impact energy) should be narrowest for $h = h_{cr}$ and still narrow for $h < h_{cr}$. For $h > h_{cr}$, the condition $F \gg P_{cr}$ is not satisfied and at first glance explosion should not occur. That explosions do occur for $h > h_{cr}$ is shown in Fig 12. The explanation offered for this contradiction is that fracture of the sample occurs before explosion for $h > h_{cr}$. If the impact energy is sufficiently high sequential fractures (see Fig 8) can produce a "new" sample thickness $h_n$ such that $h_n \leq h_{cr}$. Now if $P_n \geq P_{cr}$ explosion can occur. This sequential process tends to increase the variability of the dependence of fractures on $P$ and consequently on impact energy. Thus $f$ vs impact energy curves should show more spread than those for $h \leq h_{cr}$. Presumably at $f + 1$, ie large values of impact energy (and consequently $P$) there are more fractures and the frequency curve gets more spread out (flattened) than in the lower range of frequencies. The non-dependence of $f$ on impact energy, observed for Tetryl & TNT, is explained by A & B as follows: "Suppose $h_1$ is not much smaller than $h_{cr}$ ($h_1$ will be recalled is the limiting sample thickness for which hot-spots can lead to self-ignition), then the charge thickness, decreasing during fractures & ejections, may turn out to be below $h_1$ in which case no explosion should occur. However large the impact energy, the explosion frequency no longer depends on impact energy but is determined by the frequency with which the charge thickness, decreasing in jumps, enters the interval $h_{cr}$ to $h_1$ ."

An interesting consequence of the above considerations is the prediction that $f$-impact energy curves should be greatly "flattened" in impact machines having small diameter strikers. This prediction is based on the expectation that $P_{cr}$ does not depend on $h$ or
h/D. Thus the curves of Figs 9, 10 & 11 could just as well have h/D instead of h for the abscissa. Consequently one can just as easily speak of (h/D)_{cr} as h_{cr}. Now if D is reduced h_{cr} must also decrease to keep (h/D)_{cr} constant. With a reduced h_{cr} and “standard” thickness samples the anomalous frequency effect, and indeed failures at all impact energies, should be observed more frequently than at larger D (ie larger h_{cr}). This expectation is in accord with experience. All the anomalous effects are more pronounced in the so-called Weller impact machine (D = 1.5mm) than in Kholevo No 2 (D = 10mm). It should be noted, however, that primary explosives, for which h_{cr} is large, are initiated at rather low impact energies in the Weller machine. This is so because h_{cr} is still quite large, in the Weller machine, and its small D makes it possible to attain P_{cr} at low impact energies (P = 4F/\pi D^2).

Useful information is also obtained by plotting f vs h at different impact energies—Fig 13.

![Graph](image)

Fig 13. Explosion frequency vs charge thickness in the case of HMX for various impact energies

From these plots, one can obtain h_{cr}. Table 6 shows that h_{cr} based on P measurement are in excellent agreement with those based on frequency measurements.

<table>
<thead>
<tr>
<th>Explosive</th>
<th>From pressure measurements</th>
<th>From the frequency peak</th>
</tr>
</thead>
<tbody>
<tr>
<td>HMX</td>
<td>0.43</td>
<td>0.42–0.43</td>
</tr>
<tr>
<td>RDX</td>
<td>0.25</td>
<td>0.25</td>
</tr>
<tr>
<td>Tetryl</td>
<td>~0.12</td>
<td>~0.10–0.13</td>
</tr>
<tr>
<td>PETN</td>
<td>0.27</td>
<td>0.27</td>
</tr>
<tr>
<td>TNT</td>
<td>~0.08</td>
<td>~0.10</td>
</tr>
</tbody>
</table>


Andreev and Terebelina (Ref 7) performed a large number of impact experiments, in Kholevo No 2 type machine, in which they varied sample weight and sample location, and observed the explosion frequency at a fixed impact energy. Sample location, for a fixed sample weight, was either in the central region between striker and anvil (D=5 mm) or spread uniformly between anvil and striker (D=19 mm). Obviously the centrally located samples were thicker than the uniformly-spread samples. For “sensitive” explosives, primaries, PETN, RDX, HMX, etc, they found high explosion frequencies for centrally-located samples and almost no explosion for spread samples (except Mercury Fulminate or Lead Pichate for which sample location had no real effect). A & B’s interpretation of these results is that for the “sensitive” explosives h/D = (h/D)_{cr} for the central charges and (h/D) < (h/D)_{cr} for the spread charges. In the latter case either the impact energy is too low so that explosion frequency drops to zero at small h (see Fig 13), or else h < h_{1} in which case there would be no explosion at any impact energy. Conversely for “insensitive” explosives (h/D) > (h/D)_{cr} for central charges and (h/D) < (h/D)_{cr} for spread charges. The effect of variation in sample weight can be interpreted similarly. A & B give no quantitative comparison between Andreev & Terebelina’s data and (h/D)_{cr} obtained by A & B. We have done this under the assumption that all of the Andreev & Terebelina’s charges are at a density of 1 g/cc.

Qualitatively the expected trends do show up but quantitative agreement is frequently poor.

According to Eq 8 and the subsequent discussion (in topic 7 of the previous section) impact sensitivity should: a) increase with increasing sample density, b) increase with de-
creasing sample size, c) probably be independent of initial temperature. The experimental findings of A & B (see topic 5 of this section) are not in accord with suppositions a) & b) since impact sensitivity was found to be essentially independent of sample density & particle size. No direct evidence of the effect of initial temperature is presented. However, according to Eqs 2a & 9, \( h_{\text{cr}} \) should decrease as \( T_o \) increases (because \( \sigma_u \) decreases as \( T_o \) increases). A & B's data for HMX confirm this expectation.

We return briefly to an important consideration touched on at the beginning of this section and in topic 4 of the previous section, namely the time delay between impact and explosion. The idealized diagram (Fig 19 of A & B) shown below will be helpful in the following analysis.

In this diagram time coordinates have been changed by the transformation \( t' = \tau_{\text{ad}} - t \) where \( t \) is real time and \( t' = \tau_{\text{ad}} \) or \( t = 0 \) is the time at the beginning of impact. Thus impact occurs at \( t = 0 \) and with the above transformation explosion occurs at \( t' = 0 \); also at \( t' = 0 \), \( P = P_{\text{ex}} \). If, for simplicity, we assume that the impacted sample is at \( T_m \) at \( t = 0 \), then (in accordance with Eq 1a) its \( T-t \) history will be given by curve I in the diagram. For each \( T \) there is a corresponding adiabatic explosion time \( \tau_{\text{ad}} \). Three typical \( T-\tau_{\text{ad}} \) plots (curves II, III & IV) are shown in the diagram. Curve II does not intersect curve I, therefore there can be no explosion at \( t' = 0 \). Curve III intersects curve I twice, therefore explosion should have occurred before it actually did, which is contrary to the assumption of explosion at \( t' = 0 \). Curve IV, which has a point of tangency with curve I, is thus the only possible curve that satisfies the requirements of the problem under consideration. This condition of tangency is obtained as follows:

From thermal explosion theory (eg see article on Hot Spots in this Vol)

\[
\tau_{\text{ad}} = \frac{C}{Q_{\text{ZE}}} \frac{RT^2}{E/RT}
\]

(11)

and from curve I of the diagram

\[
\tau_{\text{ad}} = -(\bar{P}_{\text{ex}} - \bar{P}) \frac{dt'}{d\bar{P}}
\]

(12)

From Eqs (1a) & (12) the tangency condition becomes,

\[
\frac{\alpha_d \tau_{\text{ad}}}{dT} = \frac{dt'}{d\bar{P}}
\]

and via Eqs (11) & (12) leads to

\[
\frac{(\bar{P}_{\text{ex}} - \bar{P})}{\bar{P}} = RT^2/E(T - T_m)
\]

(13)

and

\[
\tau_{\text{ad}} = \tau_d RT^2/E(T - T_m)
\]

(14)

The only effect of altering initial conditions to \( T = T_o \) at \( t = 0 \) instead of \( T = T_m \) at \( t = 0 \) (as assumed above) is to change the equality sign in Eqs (13) & (14) to less-than-or-equal (\( \leq \)) signs; proof of this is given by A & B. Now for most secondary explosives for \( 400 < T < 700^\circ \)

C, the expected range of \( T_{cr} \), the term \( RT^2/E(T - T_m) \) \( \leq \) 0.1 and consequently

\[
P_{\text{ex}} - \bar{P} < 0.1 \bar{P}_{\text{ex}}
\]

and

\[
\tau_{\text{ad}} < 0.1 \tau_d
\]

Typically \( \tau_d \approx 10^{-4} \text{ sec} \) and consequently the time delay between impact and explosion \( \tau_{\text{ad}} \) is \( \approx 10^{-5} \text{ sec} \) or less.

3. We have already alluded to the applicability of A & B's impact model to the Weller impact machine. Below we examine the applicability of their ideas to other impact machines that also create rapid shear in the explosive samples. Let us consider the following tabulation (Table 11 of A & B) where all the \( P \)'s refer to compression stresses in the explosive.

Here \( \bar{P}_{cr} \) is the same as that of Table 1; \( \bar{P}_{tric} \) is from the data of Kozlov & Marnauv for experiments carried out under rapid shear; \( \bar{P}_{tricM} \) (data of Muratov) was obtained under conditions similar to \( \bar{P}_{tric} \) experiments except that the explosive was confined by a tin ring; \( \bar{F}_{tp} \) are results of experiments with Kholevo No 1 in which the striker was rotated \( 10^\circ \) to produce...
### TABLE 11. Critical stresses (in thousands of kg/cm²) obtained in different instruments

<table>
<thead>
<tr>
<th>Explosive</th>
<th>$P_{cr}$</th>
<th>$P_{fric}$ (50%)</th>
<th>$P_{H-p}$ (50%)</th>
<th>$P_{fric}$ (100%)</th>
<th>$P_{H-p}$ (100%)</th>
<th>$P_{fric}$ M (100%)</th>
<th>$P_{H-p}$ (30%)</th>
<th>$P_{H-p}$ (100%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TNT</td>
<td>~11</td>
<td>7.2</td>
<td>6.8</td>
<td>11</td>
<td>9.6</td>
<td>10.8</td>
<td>9.6</td>
<td>~</td>
</tr>
<tr>
<td>Trinitrophenol</td>
<td>~9.5</td>
<td>5.7</td>
<td>~7.4</td>
<td>6.1</td>
<td>~6.8</td>
<td>~</td>
<td>~</td>
<td>~</td>
</tr>
<tr>
<td>Tetryl</td>
<td>~8.4</td>
<td>4.7</td>
<td>5.2</td>
<td>7.2</td>
<td>6.8</td>
<td>~</td>
<td>~</td>
<td>~</td>
</tr>
<tr>
<td>RDX</td>
<td>7.0</td>
<td>4.2</td>
<td>5.2</td>
<td>6.8</td>
<td>6.5</td>
<td>9.6</td>
<td>~</td>
<td>~</td>
</tr>
<tr>
<td>PETN</td>
<td>4.8</td>
<td>3.1</td>
<td>2.8</td>
<td>5.4</td>
<td>5.3</td>
<td>5.3</td>
<td>8.7</td>
<td>~</td>
</tr>
<tr>
<td>Lead Azide</td>
<td>2.6</td>
<td>1.6</td>
<td>~</td>
<td>3.1</td>
<td>~</td>
<td>1.8</td>
<td>~</td>
<td>~</td>
</tr>
</tbody>
</table>

“shear” (data of Bochkov & Kupriyanov); $P_{H-p}$ is the same as $P_{H-p}$ but with Kholevo No 2; bracketed numbers are explosion frequencies.

The trends of the various $P$ values obtained with six different impact systems are in obvious qualitative accord. Closer examination of the data reveals that $P_{fric}$ & $P_{H-p}$ for $f = 50\%$ are always lower than $P_{cr}$. This is to be expected since $P_{cr}$ is essentially for $f = 100\%$ (see Eq 11). Consequently it is not surprising that $P_{cr}$ agrees well with $P_{fric}$, $P_{fric}$ & $P_{H-p}$ for $f = 100\%$. According to A & B, Mamaev ($P_{fric}$) showed that the lowest compression pressure (for PETN) was obtained when the sample was placed near the center (not the periphery) of his apparatus. This is in accord with A & B’s argument (derived from the study of plastic metals) that peak stress $P$ occurs at center of a sample and considerably exceeds the average stress $P$ (see topic 4 of the preceding section). For “confined system” of $P_{H-p}$, the stress is uniform and has the value of $P$ rather than $P$. It should be noted that $P_{H-p}$, which is a peak stress, is roughly twice $P_{cr}$ which is an average stress. This is in accord with Eq 4.

The expectation that $h_{cr}$ in Kholevo No 1 is greater than $h_{cr}$ in Kholevo No 2 (see Eq 8) is fully confirmed by experiment according to A & B.

9) No experimental evidence is given by A & B in support of their contention that $\sigma_{cr}$ varies appreciably for supposedly equivalent samples. However, this contention seems logical for the highly non-isotropic nature of an explosive sample consisting, as it does, of a variety of different sizes and shapes of explosive grains.

10) In topic 7 of the preceding section we summarized A & B’s development of the concepts of minimum impact force and minimum impact diameter (Eqs 6, 7 & 8), and in topic 10 we indicated that quantitative agreement between theory and available experimental data is poor. Now present this comparison. For some strange reason, probably due to the unavailability of the appropriate data, A & B choose to compute $D_{min}$ & $F_{cr}$ from data for $d_{cr}$ for $p_0 \sim 1$ g/cc charges of roughly 0.1mm particle size, whereas their best measurements in Kholevo No 2 are for $p_0 \sim 1.5$ g/cc samples. Table 6 presents their computations and also summarizes their measurements with the Kholevo No 2 machine with a sample diameter $D \leq 10$mm.

### TABLE 7. Impact Characteristics of Some Common Explosives

<table>
<thead>
<tr>
<th>Explosive</th>
<th>$C_{ult}$ kg/m²</th>
<th>$P_{cr}$ kg/cm²</th>
<th>(h/D)cr $\times$ 10²</th>
<th>d/mm cm</th>
<th>$D_{min}$ cm</th>
<th>$F_{min}$ kg</th>
</tr>
</thead>
<tbody>
<tr>
<td>TNT</td>
<td>340</td>
<td>~11</td>
<td>~0.8</td>
<td>12</td>
<td>~75</td>
<td>~5.3 $\times 10^7$</td>
</tr>
<tr>
<td>Trinitrophenol</td>
<td>520</td>
<td>9.5</td>
<td>~1.1</td>
<td>7</td>
<td>~32</td>
<td>8.9 $\times 10^4$</td>
</tr>
<tr>
<td>Tetryl</td>
<td>520</td>
<td>~8.4</td>
<td>~1.2</td>
<td>5</td>
<td>19</td>
<td>2.4 $\times 10^5$</td>
</tr>
<tr>
<td>RDX</td>
<td>820</td>
<td>7.0</td>
<td>2.5</td>
<td>5</td>
<td>5.0</td>
<td>1.3 $\times 10^5$</td>
</tr>
<tr>
<td>HMX</td>
<td>1250</td>
<td>6.4</td>
<td>4.3</td>
<td>4</td>
<td>4.6</td>
<td>1.1 $\times 10^5$</td>
</tr>
<tr>
<td>PETN</td>
<td>600</td>
<td>4.8</td>
<td>2.7</td>
<td>2</td>
<td>3.7</td>
<td>4.8 $\times 10^4$</td>
</tr>
<tr>
<td>Lead Azide</td>
<td>1550</td>
<td>2.6</td>
<td>28</td>
<td>~</td>
<td>~</td>
<td>~</td>
</tr>
</tbody>
</table>
It is obvious that \( D_m \gg D \) where \( D \leq 10 \text{mm} \) is the diameter at which explosions were observed for all the explosives shown in the Table. Now for TNT, at just below crystal density, A & B quote an experimentally found \( d_{cr} \sim 2 \text{mm} \). Using \( d_{cr} \sim 2 \text{mm} \) will reduce \( D_m \) six-fold and \( F_m \) 36-fold over those shown in the tabulation, but even this reduced \( D_m \) is still much larger than \( D \). If this 6-fold reduction of \( d_{cr} \) as \( \rho_e \) increases is also applicable to the "sensitive" explosives (RDX, HMX, PETN etc) then \( D_m \ll D \) for these explosives and, as required by A & B's theory, explosion is possible outside the impact zone (which is of dimension D). For the "insensitive explosives (TNT, etc) in Table 6, \( D_m > D \) and (according to A & B) explosion can occur only within the impact zone. Since the impact zone is a property of the impact machine, ie the impact zone varies with striker diameter, we have a contradiction to A & B's assertion that "impact sensitivity," determined with Kholo Nov 2, is solely the property of the explosive \& not a combined property of the explosive and conditions of impact. A & B make no attempt to resolve this contradiction. What they suggest, as already discussed, is that little quantitative significance be ascribed to \( D_m \) and/or \( F_m \), but that these quantities be used to develop qualitative series of impact sensitivities, eg Lead Azide is more sensitive than PETN which is more sensitive than RDX etc.

A very important facet of the general subject of impact sensitivity is the artificial lowering-desensitization of the impact sensitivity of certain explosives. If we accept Eq 8 as being at least qualitatively correct \& take \( D_m \) as a measure of impact sensitivity, then from Eqs (1), (8) & (10) and the usual situation of \( \bar{F}_{cr} \gg \sigma_u \)

\[ D_m \sim K(T_{cr}-T_m) d_{cr}/T_m \tag{15} \]

where \( K \) is a constant. To "desensitize" an explosive, ie increase \( T_m \), one can increase \( T_{cr} \) \& \( d_{cr} \) (ie decrease its reactivity and capacity to detonate) or decrease \( T_m \). For a given explosive little can be done about increasing \( T_{cr} \). Increase in particle size should increase \( d_{cr} \) and should result in decreased impact sensitivity (however see previous discussion in topic 7 about the observed lack of a particle size effect for PETN, RDX \& HMX at high compaction). Decreases in \( T_m \) can be achieved by forming eutectic mixtures of two or more explosive components. This is done in practice, eg in cyclotolts, octolts, tetratols etc. From Eq (8), it follows that a reduction in \( \sigma_u \) (which we eliminated in favor of \( T_m \) for the purpose of the above discussion) will result in a decreased impact sensitivity—at least an increase in \( D_m \). A reduction in \( \sigma_u \) can be achieved by using appropriate additives such as surface-active agents, low viscosity materials etc. It seems that more development effort should be directed towards this than has been expended heretofore.

Written by J. ROTH


Impact Velocity. The actual velocity attained by a bomb or other projectile on reaching the target. This must not be confused with the so-called terminal velocity, which is the theoretical
velocity on impact for a given size and shape of projectile. This theoretical velocity cannot be attained in practice because of air resistance.

**Impedance, Acoustic and Shock** is the product of density and sound velocity, namely \( pc \). Analogously shock impedance is \( \rho_0 U \) where \( U \) is the shock velocity in a medium whose density (ahead of the shock) is \( \rho_0 \). Both acoustic & shock impedances are used to estimate the interface stress, \( \sigma \), and interface particle velocity, \( u \), for planar shocks moving from one medium into another medium. A simple method of doing this, based on the so-called *acoustic approximation*, is illustrated below.

From conservation of mass & momentum (See Vol 4, pp D 604-5)

\[
\sigma = \rho_0 U u
\]

(1)

Now if we assume that \( U = \text{constant} = c \) (the acoustic approximation)

\[
\rho_0 U \equiv \rho_0 c
\]

(2)

and for a given medium \( \sigma \) is directly proportional to \( u \). This is shown in the following diagram (for actual \( \sigma \)-\( u \) curves see Fig 3, 4 & 5 of article on Hugoniot in this Vol) which represents

![Diagram of shock impedance and velocity](image)

a shock in medium I entering medium II. The heavy black line is a geometric reflection of line I thru the point \( \sigma_1, u_1 \). These quantities, \( \sigma_1 \) & \( u_1 \), are respectively the stress & particle velocity at the interface of medium I. It is obvious from the diagram that

\[
\tan \alpha = \sigma_1/u_1 = \sigma_2/(2u_1-u_2)
\]

(3)

By substituting for \( u_1 \) & from Eqs (2) & (3) and simplifying

\[
\sigma_2/\sigma_1 = 2/(1 + \rho_1 c_1/\rho_2 c_2)
\]

(4)

and analogously (by substituting for \( \sigma_1 \) & \( \sigma_2 \))

\[
u_2/u_1 = 2/(1 + \rho_2 c_2/\rho_1 c_1)
\]

(5)

The diagram or Eqs (4) & (5) show that for \( \rho_2 c_2 > \rho_1 c_1, u_2 < u_1 \) & \( \sigma_2 > \sigma_1 \) and conversely. Moreover by setting \( \rho_2 c_2 = 0 \) (medium II is a vacuum; for air \( \rho_2 c_2 \sim 0 \) in comparison to \( \rho_1 c_1 \) for continuous condensed media) the particle velocity is \( 2u_1 \). This is the so-called *free-surface* velocity, \( u_{fs} \), of medium I shocked to the state \( \sigma_1 u_1 \). Conversely, when \( u_2 = 0 \), \( \sigma = \sigma_f = 2\sigma_1 \) (see diagram & Eq (3)). This means that the interface stress on a *rigid wall* (consequently \( u_2 = 0 \)), in contact with a condensed medium shocked to the state \( \sigma_1, u_1 \), is \( 2\sigma_1 \). That \( u_{fs} = 2u_1 \) is also obtained from the exact solution (analytical or graphical) of the conservation equations for shocks in condensed media (Refs 1, 2 & 3). The exact solutions, however, show that \( \sigma_f > 2\sigma_1 \), but usually not much greater. In fact, for a rigid wall

\[
\sigma_f/\sigma_1 > 2.4
\]

As examples of the degree of accuracy of Eqs (3) & (4), let us compute the stress & particle velocity for detonating Comp B in contact with Mg, and Comp B in contact with steel. In each case we will replace \( \rho_1 c_1 \) by \( \rho_0 D \) for Comp B but still use the appropriate \( \rho_2 c_2 \) for the metals. For Mg: acoustic approx \( \sigma_2/\sigma_1 = 2(1 + 1.71 x 8/1.74 x 4.7) = 0.75 \);

\[
u_2/u_1 = 1.25 \quad \text{"exact"} \quad \sigma_2/\sigma_1 \quad \text{(Ref 2)}
\]

\[
= 0.99; \quad u_2/u_1 = 1.03
\]

For steel: acoustic approx \( \sigma_2/\sigma_1 = 2(1 + 1.71 x 8/7.8 x 6) = 1.55; \quad u_2/u_1 = 0.45 \quad \text{"exact"}
\]

\[
\sigma_2/\sigma_1 \quad \text{(Ref 2)} = 1.61; \quad u_2/u_1 = 0.50
\]

Note that for the metals \( u_2 > c_2 \) and consequently \( \rho_2 U_2 > \rho_2 c_2 \), therefore the acoustic approximation calculation should usually give a lower limit value of \( \sigma_2/\sigma_1 \) & an upper limit value for \( u_2/u_1 \).


Imperial Chemical Industries. See ICI in this Vol
Imperial Schultz Powder. It consists of Nitro-
lignin 80.1, Ba nitrate 10.2, vaseline 7.9 &
volatile matter 1.8%. Its props are reported as
follows:
<table>
<thead>
<tr>
<th>Property</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calories per g</td>
<td>742</td>
</tr>
<tr>
<td>Permanent gas, cc/g</td>
<td>763</td>
</tr>
<tr>
<td>Water vapor, cc/g</td>
<td>152</td>
</tr>
<tr>
<td>Total vol of gas at STP</td>
<td>915</td>
</tr>
<tr>
<td>Relative temp</td>
<td>106</td>
</tr>
</tbody>
</table>

This is a shot-gun powder used at one time in England

Imperial Explosives. It consists of NH₄NO₃
65-85% & WSi-Al alloy 15-35%
Ref: R. Imperial, USP 1054777 (1913) & CA
7, 1419 (1919)

Implosion. The reverse of explosion—when the
walls of a confining vessel collapse inwards, in-
stead of bursting outwards. For instance, when a
thin-walled glass container is partially filled
with boiling water, sealed and then cooled, the
vessel may collapse with a loud noise. This is
due to the fact that water vapor condenses on
cooling, creating a partial vacuum inside the
vessel. This causes the atmosphere, which is at
a higher pressure, to push the walls of the
container towards the inside until they are
broken

Ordinary electric light bulbs are imploded
(and not exploded) when the walls are
weakened by an accidental crack or from a
blow

Precisely controlled detonations are used to
create an implosion of fissionable material to
achieve critical mass in the explosion of an
atomic bomb (See Vol 1, pp A499-504)

Impulse. The impulse I, associated with a
pressure pulse of duration \( t \) and pressure \( P \) is
\[
I = \int P \, dt
\]
which is the area under the \( P-t \) curve
of the pulse. For triangular pressure pulses of
peak amplitude \( P_m \), \( I = P_m t/2 \), and for an
exponentially decaying pressure pulses
\( I = P_m / B(1-e^{-Bt}) \) where \( B \) is the decay constant.
In the cgs system \( I \) has the dimensions of
dyne sec/cm²

The impulse delivered to a structure deter-
imines whether the structure moves, how far
it will move, and under certain conditions
what damage it sustains (see article on impul-
sive loading of structures in this Vol). Impulses
can be generated by a detonating explosive, a
burning propellant, a flying projectile or
indeed anything that creates a pressure pulse

The impulse delivered by a detonating ex-
plusive at some distance (in air) from the
explosive charge is briefly discussed in Vol 2,
p 181-1. A more detailed discussion of
explosively-generated blast pressures & impulses,
between under water, will be given in
a future volume under "Scaling Laws for Blast."
Below we will discuss explosively-generated impulses
delivered to structures or media in contact
with the explosive charge or separated from
it by a relatively thin layer of attenuating
material

Theoretical Considerations

To gain some insight into these phenomena,
without getting too involved in mathematical
complexities, let us first examine some limiting
cases of impulse delivered to: a) a rigid wall &
b) air (for HE detonations there is very little
difference in the final results in the explosive
impulse delivered to air or to a vacuum)

a) The rigid wall problem has been treated
in detail by Baum et al (Ref 2), on the basis
of a polytropic equation of state for the
detonation products and a polytropic coefficient
\( \Gamma = 3 \). They find that
\[
I = \frac{8}{27} c_p \rho D
\]
where \( D \) is the detonation velocity and \( c_p \) is the
explosive weight per unit area of explosive. In a
following section we shall obtain a similar result
by quite a different method

b) Baum et al also consider this case but only
indirectly & in a somewhat complex treatment
(Ref 3). As an illustrative example, we will now
present our adaption of Baum's treatment:

A semi-infinite explosive slab of thickness \( h \),
bounded on both free faces by vacuum (or air),
is initiated instantaneously at one free face. We
wish to compute the impulse delivered at the
other free face. From the polytropic equation of
state assumption it follows that:
\[
c = c_j (\rho_0/\rho_f) \quad (2)
\]
and
\[
P = P_j (\rho_0/\rho_f) \Gamma = P_j (\rho/c_j) \Gamma \quad (3)
\]
where the subscripts \( j \) indicate CJ quantities, ie
quantities at detonation equilibrium. If we place
the origin of our coordinate system at the plane of initiation, i.e., \( x = 0 \) at that plane & \( x = h \) at the other explosive face, then the usual solutions of the Riemann invariants (Ref 1) coupled with the above assumption about the equation of state of the detonation products, give,

\[
x = (u + c)t
\]  
(4)

where \( u \) & \( c \) are local particle & sound velocities at time \( t \).

From symmetry considerations \( u = 0 \) at \( x = h/2 \). Then from Eq (4),

\[
c = h/2t
\]  
(5)

For polytropic detonation products with \( \Gamma = 3 \),

\[
c_j = 3/4D
\]  
(6)

Substituting the values of \( c \) & \( c_j \) from Eqs (5) & (6) into Eq (3) we obtain,

\[
P = \left(8/27\right)P_j (D/D_t)^3 t^{-3}
\]  
(7)

In our problem, \( I = \int_0^t P_j dt \), and from Eq (7), with \( t = h/D, \ P_j = \rho_o D^2 \), & \( c_a = \rho_o h \), we obtain,

\[
I = 1/27 \ c_a D
\]  
(8)

or 1/8 of the impulse delivered to a rigid wall (see Eq (1)). This is also the result obtained by Baum (Ref 3).

The results of Baum et al (Ref 4) can also be interpreted to mean that if the above HE slab were initiated instantaneously at its midplane,

\[
I = 2/27 \ c_a D
\]  
(9)

or twice the impulse of Eq (8). From symmetry, this case is equivalent to an HE slab of thickness \( h/2 \) (thus \( c_a/2 \)), confined by a rigid wall at its plane of initiation.

To determine how the numerical factors in Eqs (1), (8) & (9) vary with \( \Gamma \) (the other terms are unchanged by variation in \( \Gamma \) as long as the equation of state is still of the polytropic form) we have re-computed Eq (1) in its generalized form,

\[
I = \rho_o D(\Gamma + 1)^{-1/\Gamma} / \Gamma (\Gamma - 1)
\]  
(9a)

for several \( \Gamma \)'s.

The tabulation below

<table>
<thead>
<tr>
<th>( \Gamma )</th>
<th>3</th>
<th>2.5</th>
<th>2</th>
<th>3/2</th>
<th>1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Factor</td>
<td>0.296</td>
<td>0.445</td>
<td>0.750</td>
<td>1.720</td>
<td>( \infty )</td>
</tr>
</tbody>
</table>

shows that the numerical factors increase as \( \Gamma \) decreases.

For many purposes, rather than using the total impulse \( I \), it is more convenient to use an impulse per unit area per unit explosive weight, i.e., \( I_1 \equiv l/c_a \), or \( I_c \equiv \rho_o l/c_a = \rho_o I_1 \) (where \( \rho_o \) is the density of the unexploded HE) which we shall call the impulse constant. The term \( I_c \) has the dimensions of velocity and is closely related to specific impulse which is commonly used as a measure of effectiveness of propellant systems. \( I_o \) has the dimensions of dyne sec/cm\(^3\) in the cgs system. Thus \( I_1 \) & \( I_o \) are readily obtained from Eqs (1), (8) & (9) by dividing by \( c_a \) to get \( l_1 \) and then multiplying \( l_1 \) by \( \rho_o \) to get \( I_o \).

So far we have only considered limiting cases and “head-on” detonations. How is the impulse generated by a given explosive system modified if the detonation is tangential (“running” detonation) rather than head-on to the medium in contact with the explosive? Also what charges are to be expected if the medium is neither a rigid wall nor a vacuum? A qualitative answer to the second question is obvious from the foregoing discussion; for a “non-rigid” wall the impulse delivered will lie somewhere between that delivered to a rigid wall and to a vacuum. However, a generally-applicable quantitative answer to this question is not available. As will be shown later, theoretical calculations can be made for specific HE/medium systems but these cannot be generalized to all HE/medium systems.

An alternate method of computing \( I \), or better \( I_o \), is based on the Gurney formula (See Vol 6, p G195-L)

For a running detonation, i.e., detonation initiated at one end of a HE slab, the velocity \( v \) it imparts to a plate of mass \( m \), where the other face of the explosive slab is bounded by another plate of mass \( m' \), is given by:

\[
v = \sqrt{2E} \left[ \frac{3}{1 + 3m/c + 1 + 2m/c + (1 + 3m'/c)(1 + 2m'/c)^2} \right]^{1/2}
\]  
(10)

where \( \sqrt{2E} \) is the Gurney constant for a running detonation of the particular explosive used (Ref 5). Now the momentum imparted to plate \( m \), whose area is \( A\), is \( mv \), and \( mv/A = I \) or \( \rho_m l m' v = I \) where \( \rho_m \) & \( \rho_m' \) are plate density & thickness. Since \( l = \rho_o l/c \), Eq (10) can be modified to give,
\[ I_o = \rho_o \sqrt{2E'} \cdot \left[ \frac{1}{2} \right] \]  \tag{11}

where the bracketed term is the same as in Eq (10) and both \( m/c \) and \( m'/c \) are on a per unit area basis. For the conditions \( m/c \gg 1 \) \& \( m'/c > 0 \), Eq (11) becomes,

\[ I_o = \rho_o \sqrt{2E'} \cdot \frac{3}{2} \left( 1 + 2m'/c \right) \left( 1 + 3m'/c \right)^{1/2} \]  \tag{12}

This reduces to,

\[ I_o = \rho_o \sqrt{2E'} \cdot \frac{3}{2} \]  \tag{13}

if \( m'/c << 1 \), eg, for a system in which one face of the HE slab is in contact with air.

The writer has shown (Ref 9) that for \( m/c \gg 1 \) \& \( m'/c << 1 \),

\[ \sqrt{2E'} = \frac{0.578D}{\Gamma - 1} \]  \tag{14}

for a "head-on" detonation and,

\[ \sqrt{2E'} = \frac{0.95 \times 0.578D}{\Gamma - 1} \]  \tag{15}

for a tangential detonation. Thus Eq (13) becomes, for \( \Gamma = 3 \),

\[ I_o = 0.250\rho_o D \]  \text{ for "head-on" detonation}

and \[ I_o = 0.238\rho_o D \]  \text{ for tangential detonation.}

These are to be compared with the numerical factor of 0.296 = 8/27 obtained by Baum's method discussed above (Eq 1).

The ratios of numerical factors obtained by Baum's method divided by those obtained by the Gurney method are:

\begin{align*}
\Gamma & \quad 4 & \quad 3 & \quad 2.5 & \quad 2 & \quad 1.5 \\
\text{Ratio} & \quad 0.98 & \quad 1.20 & \quad 1.29 & \quad 1.50 & \quad 1.72
\end{align*}

Thus in the expected HE range of 2.5 \( \leq \Gamma \leq 3 \), for a given \( \rho_o \) \& D, the impulse constant, \( I_o \), computed by Baum's method will be 20-30\% greater than that computed by the Gurney method. As will be shown later, \( I_o \) by Gurney's method is in better accord with measurements than \( I_o \) by Baum's method.

Equation (15) combined with Eqs (12) or (13) shows that the impulse delivered by a running detonation is only slightly less than that delivered by a head-on detonation. Equation (12) also suggests why it is permissible to neglect the differences between air and vacuum. For vacuum \( m'/c = 0 \). Even if \( m'/c \) for air is as high as 0.1 (we certainly expect it to be less for HE in contact with air) the error is only about 5\%. A more sophisticated and convincing argument leading to the same conclusion is given in Ref 10.

Before comparing the above theoretical results with experiment, let us return briefly to the question of "non-rigidity" of the medium in contact with an exploding HE slab. Erdman (Ref 6) using a numerical procedure based on the method of characteristics obtained values of \( I_o \) for various media in contact with EL506D sheet explosive containing about 70\% PETN & 30\% binder whose \( \rho_o = 1.4g/cc \) \& \( D = 7100 \) m/sec.

Erdman's results (shown in Table 1 of the next section) indicate that \( I_o \) decreases as medium density decreases. Erdman did not carry out his calculations for the rigid wall case to sufficiently large values of \( t \) (ie, his arbitrary calculation cut-off does not account for all the impulse in the "tail" of the \( P-t \) curve). Consequently, his values of \( I_o = 2.8 \times 10^5 \) dyne sec/cm\(^3\) for the rigid wall is expected to be somewhat low.

From Eqs (13) \& (15), using \( \Gamma = 2.6 \) (appropriate for PETN), we compute \( I_o = 3.0 \times 10^5 \) dyne sec/cm\(^3\) for EL506. In view of the above, this is in excellent accord with Erdman's result.

If we assume that the factor of 0.95 between head-on and tangential detonations (Eqs (14) & (15)) is also applicable to Baum's method of calculating \( I_o \) for a running detonation, then from Eqs (1) \& (9a), with \( \Gamma = 2.6 \), \( I_o = 3.8 \times 10^5 \) dyne sec/cm\(^3\) which is some 25\% greater than the \( I_o \) computed by the Gurney method.

Comparison of Experiment and Theory

Despurneaux & Jacques (Ref 7) present data for 65/35 Cyclotol \& Octol that can be used to check the theoretical considerations of the previous section. They used flash radiography to measure deflections, \( \varphi \), of a steel plate in contact with a detonating slab of HE. Their measurements correspond to what we have called tangential or running detonation. They found that \( 1/\varphi \) is a linear function of \( m/c \) namely, \( 1/\varphi = \ldots \)
A + B m/c, where A & B are dimensionless

Plate velocity is given by \( v = 2D \sin \phi / 2 \), and for the small \( \phi \) of their measurements
\( v \approx D \phi \) so that \( I_1 = \frac{Dv}{c} = \frac{m}{c} D \phi \)

Combining this result with the linear dependence of \( 1/\phi \) on \( m/c \), it follows that at large \( m/c \),

\[
I_1 = \frac{D}{B} \quad \text{(16)}
\]

Defourneaux & Jacques used \( I_{\infty} \) to designate \( I_1 \) at large \( m/c \)

Combining Eq (16) with Eq (13), in which \( \sqrt{2E^c} \) was eliminated via Eq (15), we get the ratio \( R, R = (I_\phi)_{exp}/(I_\phi)_{theor} = \frac{0.475B}{(\Gamma-1)} \)

Below we tabulate \( R \) as a function of \( \Gamma \), using \( B = 3.95 \) for both cyclotol & octol (from Ref 7):

<table>
<thead>
<tr>
<th>( \Gamma )</th>
<th>( R )</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>0.94</td>
</tr>
<tr>
<td>2.9</td>
<td>0.99</td>
</tr>
<tr>
<td>2.8</td>
<td>1.04</td>
</tr>
<tr>
<td>2.7</td>
<td>1.10</td>
</tr>
<tr>
<td>2.6</td>
<td>1.18</td>
</tr>
<tr>
<td>2.5</td>
<td>1.25</td>
</tr>
</tbody>
</table>

Clearly, agreement between theory & experiment is quite satisfactory in the expected range of 2.7 \( \leq \Gamma \leq 3 \) (Ref 9) for these explosives

The impulse imparted to various metal, plastic and even foam plates by running detonations of EL506D sheet explosive were measured at SRI (Refs 5 & 8) using a variety of measuring

---

**TABLE 1: Comparison of Experimental & Theoretical Impulse Constants**

<table>
<thead>
<tr>
<th>Medium</th>
<th>Density of Medium (g/cc)</th>
<th>( (I_\phi)_{obs} \times 10^{-5} ) dyne sec/cm(^3)</th>
<th>( (I_\phi)_{theor} \times 10^{-5} ) dyne sec/cm(^3)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rigid wall</td>
<td>-</td>
<td>-</td>
<td>2.8 (a); 3.0 (b)</td>
</tr>
<tr>
<td>Steel</td>
<td>7.8</td>
<td>2.9</td>
<td>-</td>
</tr>
<tr>
<td>Aluminum</td>
<td>2.56</td>
<td>2.5</td>
<td>-</td>
</tr>
<tr>
<td>Magnesium</td>
<td>1.74</td>
<td>2.45</td>
<td>2.3</td>
</tr>
<tr>
<td>Micarta</td>
<td>1.36</td>
<td>-</td>
<td>2.0</td>
</tr>
<tr>
<td>Neoprene</td>
<td>1.25</td>
<td>-</td>
<td>2.0</td>
</tr>
<tr>
<td>Neoprene foam</td>
<td>0.63</td>
<td>-</td>
<td>1.8</td>
</tr>
<tr>
<td>&quot;&quot;</td>
<td>0.40</td>
<td>2.4 (?))</td>
<td>-</td>
</tr>
<tr>
<td>Polyurethane foam</td>
<td>0.025</td>
<td>2.1 (?))</td>
<td>-</td>
</tr>
<tr>
<td>&quot;vacuum&quot;</td>
<td>0</td>
<td>-</td>
<td>0.39 (c)</td>
</tr>
<tr>
<td>Steel</td>
<td>7.8 (d)</td>
<td>4.9</td>
<td>4.8 (e)</td>
</tr>
</tbody>
</table>

(a) Somewhat low—see discussion in previous section
(b) From Eqs (13) & (15) for \( \Gamma = 2.6 \)
(c) From Eq (8) for \( \Gamma = 2.6 \)
(d) Had an 8-mil-thick Pb foil on the "free" HE face
(e) From Eqs (12) & (15) with \( m^2/c = 1.3 \)
techniques. In all cases m/c >> 1 & in all, but one experiment, the HE face not in contact with the test plate was in contact with air. The impulse constants, $I_o$, based on these measurements are compared, in Table 1, with theoretical $I_o$’s, primarily those computed by Erkman (Ref 5).

It is seen that agreement between experiment and theory is quite satisfactory except for low density foams where experimental results may be uncertain.

The writer has measured detonation velocities and impulses delivered for four different detonating gas mixtures. His results (quoted in Ref 5) are used for the comparison of experimental and theoretical $I_o$’s shown in Table 2. Each of the “slabs” of detonating gas, of necessity, had a thin plastic sheet on its “free face.” Both slab thickness and plastic sheet thickness were varied to get varying values of m/c (see Eq (12)). The measured results in Table 2 show that $I_o$ increases as m/c increases. The Gurney treatment (Eqs (12) & (15)) predicts this increase although the calculated $I_o$’s are in general some 50% greater than the observed $I_o$’s. A possible explanation for the larger-than-observed computed $I_o$’s may be based on the expectation that the computed $\gamma$’s may be too small. At the high CJ temperatures & low CJ pressures the detonation products may be more dissociated and/or ionized than we computed. More dissociation or more free electrons (ionization) will make $\gamma$ approach 5/3, the $\gamma$ for monoatomic gases.

The $I_o$’s computed by Baum’s method (Eq (1) corrected for $\gamma$ (for gas detonations $\Gamma = \gamma = \gamma_0$) are in reasonable agreement with $I_o$’s observed for the larger m/c’s. This is fortuitous since in deriving Eq (1) it was assumed that

<table>
<thead>
<tr>
<th>Gas</th>
<th>m'/c</th>
<th>(1+2 m'/c)(1/2)</th>
<th>Theoretical $I_o \times 10^{-5}$ (dyne sec/cm²)</th>
<th>Obs $I_o \times 10^{-5}$ (dyne sec/cm²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C₂H₂ + O₂ (d)</td>
<td>0.396</td>
<td>1.21</td>
<td>635 940 355</td>
<td>530</td>
</tr>
<tr>
<td></td>
<td>0.47</td>
<td>1.25</td>
<td>660 940 355</td>
<td>450</td>
</tr>
<tr>
<td></td>
<td>0.72</td>
<td>1.405</td>
<td>740 940 355</td>
<td>520 (e)</td>
</tr>
<tr>
<td></td>
<td>0.96</td>
<td>1.495</td>
<td>785 940 355</td>
<td>762 (e)</td>
</tr>
<tr>
<td></td>
<td>2.38</td>
<td>2.01</td>
<td>1055 940 355</td>
<td>727 (e)</td>
</tr>
<tr>
<td></td>
<td>2.53</td>
<td>2.20</td>
<td>1160 940 355</td>
<td>910</td>
</tr>
<tr>
<td>2H₂ + O₂ + 5He (f)</td>
<td>0.99</td>
<td>1.50</td>
<td>180 210 128</td>
<td>149 (e)</td>
</tr>
<tr>
<td></td>
<td>1.65</td>
<td>1.76</td>
<td>210 210 128</td>
<td>170</td>
</tr>
<tr>
<td></td>
<td>3.30</td>
<td>2.30</td>
<td>275 210 128</td>
<td>175</td>
</tr>
<tr>
<td></td>
<td>6.60</td>
<td>3.10</td>
<td>370 210 128</td>
<td>205 (e)</td>
</tr>
<tr>
<td></td>
<td>9.64</td>
<td>3.70</td>
<td>445 210 128</td>
<td>365 (e)</td>
</tr>
<tr>
<td>2H₂ + O₂ + 10He (g)</td>
<td>1.17</td>
<td>1.57</td>
<td>131 140 103</td>
<td>115 (e)</td>
</tr>
<tr>
<td></td>
<td>1.97</td>
<td>1.88</td>
<td>157 140 103</td>
<td>120</td>
</tr>
<tr>
<td></td>
<td>3.94</td>
<td>2.48</td>
<td>207 140 103</td>
<td>131 (e)</td>
</tr>
<tr>
<td></td>
<td>7.87</td>
<td>3.38</td>
<td>281 140 103</td>
<td>155 (e)</td>
</tr>
<tr>
<td>2H₂ + O₂ + 15He (h)</td>
<td>2.19</td>
<td>1.96</td>
<td>145 115 &lt;100</td>
<td>110 (e)</td>
</tr>
</tbody>
</table>

(a) Eqs (12) & (15) with the appropriate $\Gamma$, $\rho_o$ & D
(b) Eq (1) with the appropriate $\Gamma$, $\rho_o$ & D using 0.95 to convert from a “head-on” to a “running detonation”; m/c assumed to be zero
(c) For air in contact with “free-face” of detonating slab of gas
(d) $\rho_o = 1.18$ g/l, D = 2900 m/sec & $\gamma = 1.31$
(e) Single measurement
(f) $\rho_o = 0.284$ g/l, D = 3440 m/sec & $\gamma = 1.39$
(g) $\rho_o = 0.238$ g/l, D = 3400 m/sec & $\gamma = 1.47$
(h) $\rho_o = 0.214$ g/l, D = 3450 m/sec & $\gamma = 1.53$
\( m/c = 0 \), therefore computations based on Eq (1) should agree with measured results for small \( m/c \) and not for large \( m/c \). Erkman’s computations are not directly comparable with experiment since he assumed that the “free face” was in contact with air where in reality it was in contact with a thin sheet of plastic. Erkman’s \( I_o \)’s presumably are lower limit values since the plastic sheet provides more “confinement” than air.

**Effect of Attenuators**

By interposing an attenuator material of the appropriate shock impedance (See this Vol p 156) between the HE and the metal plate to be propelled, one can appreciably reduce the peak pressure at the attenuator/plate interface over that obtainable at the HE/plate interface in the absence of the attenuator. This, however, does not necessarily mean that the impulse delivered to the plate is reduced when an attenuator is used. Usually, an attenuator lengthens pulse duration so that the reduction in peak amplitude is compensated by an increase in pulse duration (\( I = P_{m}t/2 \) for a triangular pressure, and in general \( I \) is proportional to \( P_{t} \)). Thus, for thin attenuators, a simple-minded approach using Eq (13) (ie, \( m/c >> 1 \) and assuming that \( m_{att} << m_{pl} \)) leads to the conclusion that the impulse delivered by a given explosive system is about the same with or without the thin attenuator. This means that the reduction in peak pressure (at least for triangular or rectangular pressure pulses) is almost exactly compensated by an increase in pulse duration.

In practice, as shown in the accompanying figure, attenuators do reduce the impulse (Ref 8) delivered. What the above simple-minded picture does not consider is that the attenuator “traps” some of the momentum delivered by the explosive. Stated differently, a thin attenuator does not transfer to the massive plate all the momentum, delivered to it by the explosive, even after making allowance for the mass of the attenuator. This does not mean that conservation of momentum is violated. It simply means that the attenuator retains (traps) more momentum than would be expected on the basis of its weight relative to the weight of the plate. Conversely, Altshuler et al (JETP 34, 614 (1958)) have shown that a thin metal plate, propelled across a void by an HE charge, delivers its momentum to a massive target more rapidly than it acquired it from the HE. This produces a considerably higher pressure in the target than would be obtained if the target were in direct contact with the HE.

**Pulse Duration**

For detonation products that obey a polytropic equation of state (probably also true for other equations of state), pulse duration is

\[ t = n h / D \]

where \( 1 \leq n \leq \infty \). Thus, for a triangular pressure pulse

\[ I_o = P_{m}t/2 = n \rho_o D h / 2 (\Gamma+1) \]

From Eqs (13) & (14), with \( m/c >> 1 \) & \( m/c << 1 \),

\[ I_o = \sqrt{3 \left( \frac{0.578}{2 \Gamma-1} \right)} \]

Solving for \( n \) we get

\[ n \approx (\Gamma+1)/((\Gamma-1) = 2 \text{ for } \Gamma=3 \]

Baum’s calculations (Ref 2) for a “head-on” detonation against a rigid wall give \( n \approx 1.5 \), ie, the impulse obtained from Baum’s Fig 142 \( \approx 1.5 P_{m}h/D \).

Essentially the same value of \( n \) is obtained by comparing the calculated impulse of Aziz et al (quoted in Ref 5) with a triangular pulse to give same area as the area under their P–t curve. Thus it appears that the \( n \) based on Gurney is a little higher than the \( n \)'s of more sophisticated theoretical estimates.

For a running EL506D detonation against a rigid wall, Erkman's calculations (Ref 5) lead to \( n \approx 2.5 \). Gurney-type calculations for this system give \( n \approx 0.95(3.6/1.6) = 2.1 \) in reasonable accord with Erkman. For gas detonations, agreement between Erkman & Gurney is also satisfactory as shown below (except for the oxy-acetylene mixture):

<table>
<thead>
<tr>
<th>gas</th>
<th>( \gamma )</th>
<th>( n ) (Gurney)</th>
<th>( n ) (Erkman)</th>
</tr>
</thead>
<tbody>
<tr>
<td>( CH_2 + O_2 )</td>
<td>1.31</td>
<td>7.0</td>
<td>4.6</td>
</tr>
<tr>
<td>( CH_2 + O_2 + 5He )</td>
<td>1.39</td>
<td>5.8</td>
<td>5.4</td>
</tr>
<tr>
<td>( CH_2 + O_2 + 10He )</td>
<td>1.47</td>
<td>5.0</td>
<td>4.9</td>
</tr>
</tbody>
</table>

The disagreement between Gurney & Erkman \( n \)'s may be another manifestation of too low a computed \( \gamma \) for the oxy-acetylene mixture (see above).

**Conclusions**

Let us re-examine the general formula (Eq 9a) for impulse, \( I_o \), delivered by a unit thickness of an explosive charge to a massive \( (m/c>>1) \) incompressible plate.
For an explosive slab, one of whose faces contacts a massive plate, and the other is in contact with air,

\[ I_0 = \rho_g D (\Gamma + 1) \Gamma^{-1} \]

This equation shows that \( I_0 \) will increase as \( \rho_g D \) increases, e.g., \( I_0 \) will be much larger for HE than for gas detonations because \( \rho_{HE} \gg \rho_{gas} \). The impulse constant will also increase as \( \Gamma \) decreases, this means that for two HE's of comparable \( \rho_g D \), the one with the lower \( \Gamma \) will deliver the larger impulse. Comparison of Eqs (8) & (9) shows that location of initiation can appreciably change the impulse delivered by a given explosive at a given \( m/c \). Thus a system initiated at its free face delivers only 1/2 the impulse of the same system initiated at its midplane. On the other hand, comparison of Eqs (14) & (15), confirmed by experimental evidence, shows that there is rather little difference in the impulse delivered by “head-on” or “running” detonations. Experimental data, given in Table 2, show that confining the “free face” of a gas detonation (to keep the gas in place), even by thin layers of plastic, increases \( I_0 \) over that expected from a free face in contact with air. Examination of Eq (12) reveals that the effect of \( m'/c \) becomes significant for \( m'/c \geq 0.2 \). Such values of \( m'/c \) are encountered in practice even with relatively thick “slabs” of detonating gas “confined” by thin layers of plastic on the “free face” of the slab.

Attenuators may be used to reduce both peak pressure & impulse. They act as momentum “traps”

Computational procedures are presented for making estimates of pulse duration in terms of the quantity \( h/D \). The factor, \( n \), by which \( h/D \) is multiplied to get pulse duration is approximately \( (\Gamma + 1)/(\Gamma - 1) \)

Written by J. ROTH


Impulse of a Primer. When a primer is fired, the force of detonation or length of “spit” is called impulse. It is measured by the maximum displacement of a mercury column resulting from the firing of the primer in an apparatus shown in Fig 15, p 50 of T.C. O'hat, “Elements of Ammunition,” J. Wiley, NY (1946)

Impulse, Specific. Specific impulse defines the pounds of thrust produced in a rocket thrust chamber, rocket engine, or similar unit at the expenditure of one pound of propellant per second; the ratio of thrust to the propellant mass flow. It is one of the most used performance parameters in rocket technology. Because performance and weight are of prime importance, it is highly desirable to design for a high specific impulse. This makes possible a reduction in size of propellant tanks and a saving in the total wt of fuel & oxidizer carried aloft in these tanks.

The most common equation for determining specific impulse is:

\[ I_{sp} = \frac{F}{\dot{W}} \]

where

- \( I_{sp} \) = specific impulse in lb/(lb/sec)
- \( F \) = thrust in pounds
- \( \dot{W} \) = wt flow rate of propLT in lb/sec

Specific impulse must vary with changes in altitudes because both thrust & weight change with altitude and gravitational forces.

Rocket power plants use the heat liberated by the reaction of chemical propellants, liquid or solid, as the source of energy. Specific impulse is more in liq propLT applications; it is difficult to accurately measure the propellant flow rate of solid-propellant-rocket thrust producing units. The average specific impulse is occasionally calculated for solid-propellant units on the basis of thrust, duration, and propLT wt, but there are other parameters that are more convenient. Modern rocket power plants are capable of obtaining specific impulse values between 240-250 lb/(lb/sec)

**IMR Propellant.** Improved Military Rifle (IMR) Propellant is a US single-base extruded propellant for small arms ammunition. Its compn is NC (13.15%N) 97.4, tin 2.0 & diphenylamine 0.6%. Tin is present to act as an antiflashing agent. Dinitrotoluene (6.75%) is added as a coating to moisture-proof the grains, also causes the first phase of the burning process to take place at a relatively slow rate, and has some antiflashing action. A glaze of graphite is added to facilitate the uniform action of automatic loading machines and to avoid the development of large static charges in blending & loading.

IMR Propellant and its many modifications in compn are manifold in the same manner as FNL (flashless, non-hygroscopic) & NH (non-hygroscopic) propellants, except for the application of a coating of DNT and glazing with graphite. The coating is applied by rotating the grains in a sweetie barrel heated above the mp of the coating agent. The glaze of graphite is applied to the coated grains by tumbling in a rotating drum (Ref 2).

Lindner (Ref 1) has reported the following characteristics of a typical IMR Propellant:

### Nominal Composition

- **Cellulose Nitrnte (13.15%N)**: 100.0
- **Dinitrotoluene (added)**: 8.0
- **Potassium sulfate (added)**: 1.0
- **Diphenylamine (added)**: 0.70
- **Volatiles (residual)**: 2.50

### Physical Characteristics

- **Density, g/cc**: 1.62

### Thermochemical Characteristics (calcd)

- **Flame Temperature, °K**: 2835
- **Isochoric**: 2285
- **Isobaric**: 331000
- **Force, ft lb/lb**: 59
- **Unoxidized Carbon, %**: 3
- **Combustibles, %**: 668
- **Heat of Explosion, cal/g**: 0.042
- **Gas Volume, mole/g**: 1.24
- **Ratio of Specific Heats of Reaction Products**: 28.9

**IMR Propellants are used in caliber .30 & caliber .50 rifle ammunition.**


**Incendiary “Blue Pencil.”** This ingenious sabotage device, resembling a blue pencil in appearance, was invented by the Germans during WWI and used to cause fires on ships of the Allies. It consisted of a celluloid tube containing a long glass bulb filled with concentrated sulfuric acid. The lower part of the bulb was connected, by means of a very fine capillary, to a reservoir provided with a siphon below which was placed a charge of potassium chlorate.

In operation, a saboteur breaks the upper tip of the bulb containing sulfuric acid and inserts the pencil into flammable material, such as cotton, sugar, explosives etc., preferably aboard a ship. After the tip is broken, air enters the bulb, allowing the acid to trickle down through the capillary into the reservoir, located below. As soon as a quantity of the acid accumulates, it discharges through a siphon arrangement into the receptacle containing the potassium chlorate and causes a very violent reaction accompanied by flash and flame.

Ref: A.B. Ray, IEC, 13, 721 (1921)

**Incendiary Bombs & Incendiary Agents.** See under Incendiary Warfare

**Incendiary Bomb Fires and Their Extinction.** As the modern incendiary bombs, containing magnesium or thermite, are hard to extinguish by water, several mixtures have been proposed for use in lieu of water. However, none of these mixtures has proved very effective.

The US Office of Civilian Defense (OCD) during WWII stated that, in fighting incendiary bomb fires, the spread of the fire is more of a hazard than the bomb itself. If the bomb falls on a spot where it cannot start fires, the best procedure is to allow it to burn out. If it falls where it can start a fire, the fire-fighter should approach it as soon as possible, regardless of risk, attempting to remain behind some protective cover, such as a wall, in order to avoid injury in case the bomb explodes. From behind cover, a strong stream of water should be directed at the surrounding area and not at the bomb, which should be allowed to burn out.

If no cover is available, the fire-fighter should operate from a crouching or prone position, to minimize the danger in case of an explosion.

In extinguishing fires caused by phosphorous,
it must be remembered that, as soon as the water dries out, phosphorous will start to burn again. In order to prevent reignition the extinguished phosphorous, while still wet, should be scraped off; placed in buckets and dumped where there are no flammable materials. As phosphorous is very poisonous and causes extremely painful burns, all bodily contact with it must be avoided.

The OCD recommended the use of sand or any of the “bomb-extinguishing” powders, because water can do more damage than good.

If water is available only in small quantities, it is better not to use it at all because insufficient amounts of water would cause bombs containing thermite or magnesium to burn more intensely by supplying additional oxygen and the released hydrogen could add explosive fuel to the fire.

Ref: Anon, JChemEducation, 20, 59-60 (1943)

Incendiary Compositions and Compounds. See under Incendiary Warfare

Incendiary Devices. See under Incendiary Warfare

Incendiary Grenades. See under Incendiary Warfare

Incendiary Liquids and Gels. See under Incendiary Warfare and under Flame Thrower Liquids

Incendiary Missiles. See under Incendiary Warfare

Incendiary Mixtures. See under Incendiary Warfare

Incendiary Projectiles. See under Incendiary Warfare

Incendiary Shells. See under Incendiary Warfare

INCENDIARY WARFARE

(Incendiary Agents, Incendiary Compounds and Mixtures, Incendiary Devices, Incendiary Missiles, etc)

Incendiary warfare accomplishes the purpose of causing fires in enemy territory and burning enemy personnel, animals, etc. In “modern” warfare, incendiaries are unfortunately also used for “strategical” purposes against civilians and cities which have no military installations. Incendiary warfare has assumed ever-increasing importance in recent wars.

Enormous numbers of incendiaries were dropped by American aviation during WWII and, as the war came to an end, incendiaries exceeded explosives in the bombing of Japan. The total amount of incendiaries dropped by US aviation was 123,000 tons on Japan and 120,000 tons on Europe. Records of the late conflict prove conclusively that incendiaries, per unit weight, achieved greater damage to industrial and other structures than did high explosives.

Stettbacher (Ref 7, p 129) gives the following figures on some of the heavy aerial bombardments of Germany and Japan during WWII

<table>
<thead>
<tr>
<th>Number of Bombs Dropped</th>
</tr>
</thead>
<tbody>
<tr>
<td>High Explosive</td>
</tr>
<tr>
<td>Köln (Cologne), 6/28/1943</td>
</tr>
<tr>
<td>Berlin, 3/8/1944</td>
</tr>
<tr>
<td>Tokyo, 3/15/1945</td>
</tr>
<tr>
<td>German Hq, 5/25/1945</td>
</tr>
</tbody>
</table>

According to Stettbacher, the loss of life in Germany was not as heavy as the loss in buildings: 500,000 dead, or one per 4.9 tons of bombs dropped, against 3.6 million buildings partially or completely destroyed. The heaviest loss of life in a single raid was in Dresden—100,000 people or 20% of the population.

It is the conviction of military authorities such as Major General A. H. Waite (Ref 9) that incendiary attacks broke the Japanese will to resist before the advent of the atomic bomb.

Historical

Incendiary warfare is one of the most ancient methods of conducting war and was used as early as biblical times.

The history of evolution on incendiary warfare may be divided into three periods:

1) before the invention of gunpowder and firearms
2) after the invention of firearms but before the introduction of aircraft for war purposes
3) after introduction of aircraft, particularly the airplane.
First Period, Before the Introduction of the Gun

In this period of incendiary warfare, incendiaries were thrown first by hand and later by special machines, called catapults and ballistae. Bows and arrows were also used for throwing incendiary missiles. The earliest account of incendiary warfare may be found in the Bible, where mention is made of burning oil and ignited fireballs, consisting of resin and straw, that were thrown by both defenders and attackers of fortified towns. The Bible also mentions (Judges 15, 3-5) that Samson used foxes with firebrands attached to their tails to burn cornfields of the Philistines.

Beside the Bible, the first reliable record of incendiary mixtures was given by the Greek tactician Aeneas, who, at about 350 BC, compiled the first European treatise on the art of war. He listed sulfur, pitch, pinewood, incense and tow as principal incendiary ingredients.

Seven centuries later (AD 350), Vegetius, a Roman military authority, added resin, bitumen and petroleum oils to this list which indicates a significant advance in incendiary technique.

All of these mixtures, however, were of low efficiency because they burned quickly and could easily be extinguished with water.

According to Fisher (Ref 9), one of the earliest incendiary munition was the "fire club." It was a wooden club with sharp iron prongs attached to each end, the center being wrapped with incendiary materials such as tared rope. The tar was lighted and the club hurled against a wooden structure, where the pointed prongs held the fire-bearing club in place against the wall or roof of the building until the fire spread. This device was described in 350 BC by Aeneas and was supposed to have been used before his time.

Incendiary hand grenades were clay or glass vessels with a narrow throat. They contained a flammable mixture, which was ignited just before the vessel was thrown against enemy installations. The vessel broke, bringing the flaming mixture in contact with objects to be ignited.

A device used as early as the 5th century BC consisted of a hollowed tree and a cauldron with burning coal, pitch and sulfur which were mounted on wheels and brought close to the enemy's wooden fortification. By means of a bellows, air was forced across the cauldron and this projected flames against the wooden structure. This device may be considered a prototype of modern flame throwers.

This crude machine was replaced by the "blowpipe," used by Roman armies. Incendiary balls, made of resin and sulfur, were forced through a tube and were ignited at the muzzle as they were ejected from the tube.

A similar device was known later (about the 13th century) as "flying fire" (ignis volatilis). It consisted of a mixture of saltpeter 50, resin 25 and sulfur 25% & was shot from a reed or pipe.

As soon as catapult machines were adapted to siege warfare, larger and more effective incendiary missiles were developed. The first incendiary missiles consisted of large stones coated with flammable materials, such as pitch, tar, sulfur, etc.

Romans used the so-called "firepot," which was a kind of iron fire bomb (as large as 2 ft in diameter) filled with pitch, sulfur, bitumen, etc and either perforated or latticed so as to permit emission of flames from burning charges. One such bomb may be seen on exhibition in the Tower of London.

"Fire Arrow" was one of the earliest incendiary weapons. It was a regular arrow with a charge of combustible material such as oakum, resin, sulfur, bitumen etc attached to it. This was ignited just before shooting. In order not to extinguish the flame during flight, the arrow had to be shot slowly and this limited its range.

A much more effective mixture was the one invented in 668 AD by Kallinikos and called "Greek Fire" (see below) or "Sea Fire." The exact formula of the original composition is not known because the Greeks kept it a secret. However, it seems that it contained, besides the combustible materials such as naphtha, pitch and sulfur, some oxidizer, which could have been saltpeter. This mixture was liquid that was discharged against the enemy either from pots, tubes or siphons, installed in the bows of ships. The moment the liquid came in contact with the water it was ignited. This device was so effective that it caused several defeats of the Arabic fleets in the 7th and 8th centuries, of the Russian fleet in 941 and 1043 and saved Constantinople several times from invaders. Greek
Fire may be considered to be another prototype of the modern flamethrower introduced by the Germans in WWI. Later, the name “Greek Fire” was erroneously given to combustibles, which were ignited and then thrown by ballistae, or other machines, and were used on land. These compositions were semi-solid masses consisting of combustibles such as sulfur, pitch, naphtha, etc. and oxidizers, such as saltpeter. They were also known as “Wildfire,” and were used extensively by the Moslems during the Crusades and by the Moors during the conquest of Spain, eg. in the siege of Niebla in 1257.

Incendiary compounds were known in China in ancient times but the ones containing saltpeter seem to have appeared about the 13th century, when they were used successfully against the Mongols.

According to the book of Marcus Graecus, entitled “Fire,” which appeared about 1300 AD, the following composition was known in the 13th century in Europe: saltpeter 6p (66.7%), charcoal 2p (22.2%) and sulfur 1p (11.1%). It was used in two munitions, a “thunder bomb” and an incendiary rocket, called the “flying tunica.” The latter consisted of a narrow cylinder filled with the above “Black Powder” mixture which served as both propellant and incendiary for this prototype rocket.

The idea of propelling incendiaries by means of a composition containing saltpeter, charcoal and sulfur is presumed to have been discovered by the Chinese. The introduction of substances, resembling the present Black Powder as rocket propellants and incendiaries, was a great improvement in incendiary warfare and for a while these weapons were of prime importance.

Second Period, After Introduction of the Gun

With the invention of cannon (14th century) that used Black Powder as a propellant it was possible to hurl missiles much greater distances than by ballistae or catapults, thus causing greater damage to the enemy without resorting to incendiary compositions. And so, with the introduction of cannon, incendiaries became of secondary importance. However, attempts were made as early as the 14th century to shoot incendiaries from cannons.

The first incendiary “shot” was a stone ball, preheated on a fire or by quicklime, and propelled from a cannon by a charge of gunpowder. In order not to ignite the propelling charge prematurely, a wad of wet turf or earth was placed on top of the Black Powder so that the ball could rest on it until the deflagration of the propellant drove it from the cannon.

With the introduction of iron balls (shot), the same procedure was followed. Hot iron shots were particularly effective against naval sailing vessels. Sometimes, even bullets from small arms were preheated and used as incendiaries.

A variation of the preheated iron shot was the hollow projectile charged with incendiary mixtures, intended to bring the shell to a red hot temperature after impact. This device was not as effective, however, as solid iron balls, or the previously described missiles hurled by catapults.

With the introduction (about the end of the 15th century) of hollow type projectiles charged with Black Powder, these “shells” also became, to a certain extent, incendiary missiles. This was due to the fact that, when such a shell burst, the flame produced by the explosion of the Black Powder could ignite any closely-located inflammable object.

As Black Powder burns comparatively quickly, this shell could not be a satisfactory incendiary missile, unless part of the Black Powder could be replaced by slow burning compounds such as pitch, rosin, sulfur, bitumen, etc. and this was tried.

However, much better results were obtained with improved “surface-burning” projectiles such as the following:

a) “Incendiary Ball”—prepared by dipping a small iron ball in molten sulfur, wrapping it in oakum, redipping in sulfur and finally rolling it in fine Black Powder. The ball was wrapped in wire and its diameter had to correspond to the caliber of the cannon.

b) Incendiary Elongated Projectile—prepared by kneading a warm incendiary mix over a crossed iron frame, in a manner similar to the above.

c) “Carcass”—invented in 1672 and used until nearly the end of the 19th century, consisted of a thick, elongated, narrow sack containing pitch, oakum and Black Powder and enclosed in an open framework of iron straps. The composition burned with-
out scattering and its flame ignited surrounding materials.

Other devices, such as "pitch rings" and "incendiary hoops" (Ref 9, p 113) also belong to the surface burning class of munitions.

Military incendiary rockets, used in Europe in the 13th and 14th centuries, were forgotten until the British Col W. Congreve introduced an improved model at the beginning of the 19th century. The idea to resurrect rockets as incendiary weapons came to Congreve when he saw how successfully Indian troops used rockets against British troops at the end of the 18th century.

Congreve's rocket, which was primarily an incendiary weapon, could compete successfully in accuracy with the cannon of that period and was used until the introduction of rifled barrels and breach loading, which improved the accuracy of artillery and increased its range to such an extent that the Congreve rockets became obsolete (about 1860).

Hale's rocket superseded that of Congreve. It was provided with a screw-like tail in an endeavor to give it a truer flight.

With the exception of the Congreve and Hale rockets, very little progress was made in the 19th century in incendiary warfare and it might even be said that it was again forgotten. Incendiaries were not used during the Crimean or the American Civil Wars, but were reintroduced by the Germans during the Franco-Prussian War of 1870-1871. The Germans, using incendiary projectiles, burned several French cities and forced their capitulation.

After that war, the Germans decided to improve the compositions and weapons used in incendiary warfare and especially to construct an incendiary projectile suitable for shooting from rifled, high-muzzle-velocity guns, invented in the 1860's. After many trials, they succeeded in developing several types of incendiary shells, some of which were used later in WWI. Other countries followed the German example and also developed incendiary shells.

The following additional incendiary mixtures and devices were invented in Europe in the 19th century:

Mixture of Niepce consisting of powdered coal impregnated with carbon disulfide, gasoline or petroleum, was loaded into a bomb together with a piece of metallic potassium or calcium phosphide, Ca₃P₂ and a small closed glass container with water. The impact of such a bomb broke the glass container and the water ignited either the K or Ca₃P₂ which in turn ignited the main incendiary mixture.

Mixture of Fleck—similar to the previous one, but using metallic sodium as the priming substance.

Fenian Fire (Liquid Fire)—consisted of a solution of yellow phosphorous in carbon disulfide. On evaporation of the CS₂, the phosphorous self-ignited in the air and ignited nearby flammable objects.

Darapski Projectiles were charged with petroleum, which was ignited by means of a mixture containing 3 parts of phosphorous dissolved in 1.3 parts of CS₂.

Nikles Mixture (Feu lorrain- Lorrain Fire) consisted of a mixture of chloro-sulfuric acid (Cl₂SO₄) together with phosphorous dissolved in CS₂. This mixture was ignited by introducing a small amount of ammonia.

Gugot Mixture—same as above, but replacing the chloro-sulfuric acid with the bromo-sulfuric.

Sophronius Mixture consisted of 3 parts of concentrated H₂SO₄ and 2 parts of KMnO₄.

In addition to these mixtures, it was proposed to use phosphorous trichloride, which catches fire on contact with either ammonia or ammonium hydrosulfide, NH₄HS.

Third Period, Incendiary Devices Used After the Introduction of Aircraft in Warfare

Although aircraft was introduced as an auxiliary warfare weapon by various countries as early as 1910, their actual combat use did not begin until WWI. By this time, incendiary materials were used in the following devices:

a) small arms ammunition; b) shells; c) trench mortar projectiles; d) grenades and other hand weapons; e) aircraft bombs; f) flame projectors (described separately); and g) rockets.

An excellent description of all of these weapons is given by Ray (Ref 4).

Only a brief resume is given below.

Small arms ammunition. Incendiary small arms ammunition, while carrying only a very small amount of material, has proved to be of the greatest value in effectively attacking highly flammable targets, such as aircraft, which might be filled with hydrogen or carry tanks.
of gasoline fuel. Most incendiary ammunition is of regular caliber and fits standard machine guns. There are two classes of bullets which may have incendiary effect—real incendiary bullets, which usually function only on impact and tracer incendiary bullets, which begin to function either at the instant of expulsion from the gun or shortly thereafter. Although the primary use of tracer bullets is to aid in directing aim, some of them may have considerable incendiary action, provided the target is struck before the bullet ceases to “trace.” Bullets used in WWI by the US Army carried the following incendiary materials:

a/ White phosphorous; b/ BaO₂ & Mg; c/ Ba(NO₃)₂, Mg, charred linseed oil, with an ignition composition consisting of red lead and Mg.

One of the German perforating tracer-incendiary bullets carried incendiary material consisting of Sr(NO₃)₂ 75, Mg 13, Al 3, Fe 6, resinous material 3%, in the rear section. On top of this was pressed an igniting composition containing KMnO₄ 55 to 58, iron filings 45 to 42% and this was ignited by the flame produced by the propellant.

Another German incendiary armor-piercing bullet contained igniter and incendiary compositions in the front section. The advantage of this type of bullet was that the burning composition in front of and around the armor-piercing shell of steel was more likely to ignite gasoline from a perforated tank than if the composition were in a separate container, which might break away from the bullet on impact. The priming or igniting composition which ignited on impact consisted of Mg, KClO₃ and Sb₂S₃, while the incendiary composition contained Mg, Al, Ba(ClO₃)₂, sulfur and a little NC.

During WWII, the following incendiary and tracer small arms ammunition were used by the US Armed Forces:

a) Incendiary Ammunition (I) was used primarily against aircraft as a means of setting fire to enemy planes. The straight incendiary bullet has an incendiary charge in the nose and in a steel sleeve ahead of the lead slug. The incendiary mixture consists of barium nitrate and magnesium-aluminum powders. The bullet functions on impact, the mixture being ignited by the heat and sparks of impact. The gilding metal jacket is torn open and the incendiary composition burns for 10 to 40 milliseconds. Incendiary fillers weigh from about 20 grains, for cal .30, to 75 grains for cal .50 bullets. One of the most effective bullets is the cal .50 M23, a 500 grain bullet which is fired at 3400 ft/sec.

b) Armor-piercing Incendiary Ammunition (API), has the lead point filler replaced by an incendiary composition. It is also used primarily against aircraft. The API bullet is similar to (I), except that the incendiary is ahead of the steel core. The cal .50 API bullet contains 15 grains of the incendiary mixture consisting of barium nitrate and aluminum-magnesium powder. This mixture ignites on impact.

It is estimated that in WWII 50% of the planes lost to anti-aircraft fire and approx 75% list to enemy aircraft were brought down by incendiary ammunition.

c) Armor-piercing Incendiary Tracer (API-T), similar to API but having a tracer mixture in the rear of the bullet. Caliber .30 ammunition is expected to trace for about 1200 yards and caliber .50 about 1800 yards. A longer trace has been developed for .50 ammunition (2500 yards). A dim trace (dim for the first 200 yards to prevent dazzling the gunner) has also been developed for .50 ammunition. The incendiary composition for calibers .30 and .50, which ignites on impact, consists of barium nitrate and aluminum-magnesium powder, the tracer composition consists of strontium peroxide and nitrate, magnesium and other ingredients. The tracer is ignited by means of a barium peroxide and magnesium powder mixture, which is pressed on top of the tracer mixture at 70,000 psi. The pressing is done by a step punch in order to create a large surface to present to the hot propellant gas which ignites the barium peroxide-Mg mixture.

For .50 caliber bullets with a trace of 2500 yards, a slower burning tracer composition containing Sr(NO₃)₂, KClO₃ and Mg is usually used.

In the so-called “night-tracer cartridges,” which neither blind the gunner nor reveal his position to the enemy, a dim igniter (essentially SrO₂, Ca carbonate and Mg) burns for the first 200 yards for the .30 and .50 caliber bullets. A tracer bullet for .45 caliber submachine gun traces for 160 yards.
d) Tracer Bullet (T), similar to ordinary ball bullets, but having the rear half of the lead core replaced by igniter and tracer compositions, which are essentially the same as in the API-T bullets. There are tracer bullets for .30 and .50 caliber rifles

A special .50 caliber bullet, called "M21 headlite tracer" was developed to satisfy the air force requirement for a bullet to simulate a ball of fire. The object was, for psychological effects, to make the enemy think that he is under fire from much larger caliber weapons than small arms. For this, the standard M1 tracer bullet is modified by replacing the usual igniter and tracer composition with strontium peroxide alone. Although in this case, the tracer distance is reduced from the usual 1800 yds to about 600 yds, the visibility is about three times that of the usual tracer bullet

Among foreign ammunition, the Japanese caliber .50 "explosive-incendiary" bullet is of interest. It was provided with a point-detonating fuze and a charge of high explosive, with the incendiary filler placed behind the explosive charge. After the projectile penetrated the target, the explosive filler was detonated, the jacket or shell shattered and the incendiary composition ignited. The composition of the incendiary mixture was Ba(NO₃)₂ 38, NaN₃ 20, Mg powder 36, Al powder 6% (Ref 9, p 58)

e) Incendiary Shells.

The first satisfactory incendiary shells suitable for firing from modern high-muzzle-velocity, breach-loading, rifled cannon were developed by the Germans before WWI

It should be noted, however, that in modern warfare, the incendiary-loaded artillery shell has become of secondary importance. This is because good targets for incendiaries are nowadays seldom found within range of artillery. If such targets are found, they usually can be destroyed more readily by incendiary aircraft bombs than by artillery. Thus, the effective use of the incendiary shell is narrowed to certain specialized targets and to situations in which the air force cannot be easily utilized

For these reasons, many types of incendiary shells that were standard during WWI have since been discarded. Nevertheless, it is interesting, from a historical point of view, to give a short description of shells used during WWI because at that time such shells played an important role

During WWI, shells were produced ranging from 37mm to 17.5cm. The smaller sizes were used for attacking aircraft from the ground while the larger calibers were used against all sorts of ground targets. There were tracer shells which had appreciable incendiary action over a considerable portion of their trajectory. There were also incendiary shells which did not function as incendiary devices until at the moment of explosion when the incendiary material was ignited and scattered

The consensus regarding incendiary shells was that the small caliber tracer-incendiary shell was quite effective against aircraft, but that the larger type, designed for use against ground targets, was not as successful

A common type of tracer-incendiary shell, carrying red lead and magnesium, was produced in the US in 75mm and 3" sizes. It was used principally for shooting down balloons. This shell was of very simple construction—just a cylinder with a threaded neck, in which a fuse was inserted, and six vents below the threaded part. The fuse charge ignited the primer which in turn ignited the incendiary mixture and the incandescent products of combustion were emitted through the six holes in the head. The smoke from this combustion left a definite trace of the projectile, and the hot gases evolved for 6 to 25 sec possessed efficient incendiary action

A more complicated tracer-incendiary shell was used by the British and Germans. In this shell (usually 75mm), the flash from the time fuse was transmitted through a tube (located in the center of the cylinder and extending from the top, near the fuse, almost to the bottom of the shell) to the primer. The primer ignited the incendiary composition, containing Mg, Ba(NO₃)₂, and binder with or without Sr(NO₃)₂, and blew out the base of the shell. This resulted in emission of flame from the base of the shell, lasting about 15 seconds

There also existed several models of shells designed solely for incendiary effects. Most of them contained thermite with inflammable organic materials. Others contained phosphorous, sodium, etc

The British designed a 77mm shell which carried thermite (Al + Fe₂O₃) as an incendiary
material and ophorite (Mg + KClO₄) as an igniting and expelling charge. This type of shell acted not only as an incendiary, but also as shrapnel producer, because it scattered hot metal and slag produced by the thermite reaction.

For larger sizes, e.g., 15.3cm, a shell was designed which contained units of thermite or other incendiary materials held in perforated cases. These units were ignited from a central tube and expelled from the front of the shell by the explosion of the powder charge located in the base.

The Germans developed a 15cm shell, for use against ground targets, which contained 12 or more celluloid cylinders surrounded by yellow phosphorous imbedded in paraffin. A central tube of black powder was used as a bursting charge.

The largest incendiary shell used during WW1 was the German 17.5cm “Minenwerfer,” described in detail in Ref 4, p 716.

A 7.7cm shell developed by the Germans for use against aircraft scattered steel slugs and flaming incendiary units on bursting, called “flaming onions” by allied aviators.

Towards the end of WWI, the US Army developed a base-opening shell, which on functioning, ignited and expelled a consolidated large mass of incendiary material.

During WWII, the US Army retained only the white phosphorous shell, which was not officially listed as incendiary ammunition. Yellow phosphorous also served as a smoke and anti-personnel agent with only incidental incendiary value—it easily ignites light combustible material. As an example of such shells, Fisher (Ref 9, p 53) describes the 81mm smooth-bore mortar shell. This shell weighs 11.5 lbs, has a range of slightly under 2000 yds and carries 4 lbs of solid yellow phosphorous. On impact, the nose fuse detonates the burster charge of Tetryl (placed in a thin-walled tubing, extending into a cavity in the phosphorous), which breaks the shell and simultaneously ignites and scatters the burning phosphorous.

A 4.2” chemical mortar shell was also used for incendiary purposes. It contained a burster charge of Tetryl, placed in thin-walled tubing extending into a cavity in the phosphorous. The Tetryl broke the shell and simultaneously ignited and scattered the burning phosphorous (Ref 9, p 53). These shells were mostly used in trench mortars.

The Japanese Army incendiary shells were used in 75mm guns and 90mm mortars. The incendiary material consisted of rubber pellets impregnated with a solution of 88% phosphorous and 12% carbon disulfide. A burster charge of picric acid scattered the incendiary pellets, which ignited on exposure to air and burned for about five minutes. These shells were only effective against readily ignitable materials (Ref 9, p 53).

**Aircraft Incendiary Shells.** The anti-aircraft incendiary shells used during WWII were very successful against hydrogen-filled airships but did little damage to airplanes. The problem of successfully attacking airplanes by incendiary shells from the ground has not been solved up to the present time, although the Germans and Japanese developed some ground anti-aircraft shells, for example, the following:

**German 20mm Incendiary Shell**—carried an incendiary composition containing phosphorous, thermite and other compounds. This charge was pressed into the lower part of the shell by means of a stepped ram. A charge of HE placed on top of the incendiary charge was detonated by means of a booster initiated by a fuse. This shell, as well as the 20mm Japanese shell, described below, proved to be more successful in air-to-air than ground-to-air firing.

**German 88mm and 128mm Aircraft Shell.** Used during WWII. The incendiary pellets were designated as FK44 and FA252. The FK pellet supposedly weighed about 27g and was approx 15mm in diam and 30mm in length. The pellet was said to consist of a detonator, a striker and a diaphragm. Three holes allowed gasoline to enter the pellet and to create a hydrostatic head as the pellet entered the fuel tank of the target. The hydrostatic head detonated an explosive filler of pressed PETN/wax/Al. The pellet was intended to blow a hole in the fuel tank large enough to prevent resealing and to ignite the gasoline. The 88mm carried 72 pellets and the 125mm, 124 (Ref 16).

**Other German Incendiary Shells.** German munitions also included incendiary core fragmentation shells. These incorporated a central flash tube which ignited the incendiary filling of each fragment prior to detonation. The shell...
functioned in a manner similar to the Italian shell described below. The materials used for the core are as follows:

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>B1</th>
<th>Dog-Small</th>
<th>Dog-Powdered</th>
<th>B19S</th>
<th>B19K</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ba(NO₃)₂</td>
<td>49.0</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>BaO₂</td>
<td>—</td>
<td>70.0</td>
<td>69.2</td>
<td>8.4</td>
<td>3.5</td>
</tr>
<tr>
<td>Fe₂O₃</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>62.5</td>
<td>65.8</td>
</tr>
<tr>
<td>Al, flaked</td>
<td>—</td>
<td>—</td>
<td>25.0</td>
<td>25.0</td>
<td>20.8</td>
</tr>
<tr>
<td>Al, powder</td>
<td>—</td>
<td>5.0</td>
<td>—</td>
<td>5.0</td>
<td>—</td>
</tr>
<tr>
<td>Mg/Al alloy</td>
<td>50.0</td>
<td>—</td>
<td>0.8</td>
<td>8.3</td>
<td>8.8</td>
</tr>
<tr>
<td>Necresil</td>
<td>1.0</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>NC*</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

*NC added as a granulating material in the form of collodion (Ref 16)

**Italian Incendiary Shell**. Italian antiaircraft incendiary pellet projectiles, in WWI, consisted of a 76 or 90-caliber shell loaded with a number of incendiary-filled cylindrical pellets. The pellets were ignited by a flash from a central flash tube. A delay mixture activated a secondary explosive in the base which broke the shell into flak. Composition of the incendiary filling is not given in Ref 16.

**Japanese 20mm Incendiary Shell**—carried a small incendiary charge of Ba(NO₃)₂ 50, Mg 40 and Al 10%.

**Japanese Incendiary Shells of Larger Calibers** were loaded with yellow phosphorous and produced large spectacular aerial bursts, but their value was probably more psychological than destructive to aircraft (Ref 9, p 55).

**Japanese 120mm Incendiary Projectile** contained 48 steel pellets embedded in a canister filled with white phosphorous. Each pellet had an annular cavity to increase the amount of phosphorous that could be retained by each pellet. The shell was exploded by means of a bursting charge of HE, fired by a point-detonating fuse.

**Trench Mortar Projectiles**. During WWI, the following trench mortar projectiles were developed: 3” and 4” Stokes bombs and 8” Liven’s drums. They were used mostly to destroy grass, shrubbery or camouflage which could act as a screen for enemy movements, to clean out woods, to demoralize the enemy during gas projector attacks and to indicate the range in night attacks.

The Stoke’s bombs were loaded for best results with about 7.5 lbs of thermite as the incendiary compound, and ophorite as the explosive igniter. The ignition was started by a time fuse.

The 8” Liven’s incendiary drum was a metallic container, 20” long and 7 5/8” in diameter, resembling an ellipsoid in shape. It was closed at one end and open at the other. It contained balls of cotton or jute impregnated with “solid oil” (thickened oil), prepared by treating a mixture of fuel oils (bp 170-225°) with sodium stearate. The lighter part of the oil mixture was readily ignited by the flash from the explosive charge, which was loaded in a central tube, and this ignited the heavier oils. One end of the drum was fitted with an impact fuse which caused the drums to explode on landing, and to scatter burning material over an area 50 yds in diameter.

Some Liven’s drums contained cotton waste impregnated with a mixture prepared as follows: TNT was dissolved in benzene and fuel oil and gas tar oil and a solution of yellow phosphorous in Cs₂ were added. The mixture was insensitive to shock, had a very low coefficient of expansion (0.0174 per °C between -10° and 55°) and a low vapor pressure.

These drums were projected from a mortar-like, smooth-bore barrel, called the Liven’s Projector. It had an 8” bore and was 37.5” long. A propellant charge ignited by an electric primer was used to obtain a maximum range of 1450 yards.

**Grenades and other small devices**. Small portable incendiary devices, such as hand grenades, are very valuable because they can easily set fire to flammable targets. These grenades can also be used for throwing into dug-outs and for destroying material abandoned in retreat, such as damaged planes.

The following grenades were used during WWI:

- **Phosphorous grenade** consisted of a sheet metal can filled with yellow phosphorous and provided with an explosive device. Although the primary aim of such grenades was smoke-production they were also incendiary weapons.

- **Thermite grenade** consisted of a cylindrical sheet iron container filled with thermite (about 600g) slightly moistened with sodium silicate. In the center of the thermite was a cavity in which
was inserted an igniter bag of Al + Ba₂O₂. Strands of a quick match connected the igniter bag to a piece of Bickford fuse.

**Thermite-gelled oil grenade** was developed with the idea of combining the advantages of thermite and gelled oil. The former develops temperatures high enough to melt metals, while the latter produces a long duration flame. The grenade consisted of a heavy cardboard container whose lower half had a celluloid container filled with gelled oil, while the upper half contained thermite with a celluloid binder (4% by weight of celluloid dissolved in a small amount of acetone). The grenade was fired by withdrawing the safety pin and releasing the “bouchon” firing handles. The spit of the fuse ignited the booster, which set off the igniter (Al + Ba₂O₂) and then the thermite. The resulting molten iron and slag penetrated and ignited the celluloid container with the gelled oil.

During WWI, the Germans developed several portable incendiary devices consisting of a metallic or pasteboard tube filled with mixtures of oxidizer and combustibles (e.g., KNO₃ 62.5, sulfur 27.0, carbon 10.5%) and fitted with a friction-igniting device. Some of these devices were of larger size, containing as much as 1670g of the above mixture. In practice this device served either as an incendiary or smoke producer, depending on circumstances.

The French developed an incendiary can, for throwing into trenches or dugouts, containing about 3 liters of petroleum oil and weighing about 7 lbs. It was provided with a friction igniting and exploding device.

Although uses for incendiary grenades are limited, they were still used during WWII. The present incendiary grenades are of two types: 1) standard types, regularly produced and supplied, and 2) improvised flammable grenades. Standard grenades, although normally thrown by hand to a distance of 75 to 100 feet, can also be fired from rifles or grenade launchers up to 750 ft. When projected from rifles, using a special cartridge, the incendiary grenade is a valuable weapon for attacking tanks and other vehicles.

**Standard grenades.** According to Fisher (Ref 9, p 55-6), there are two kinds of standard grenades used at the present time by the US Armed Forces: a/ thermite type and b/ phosphorous type.

While the **thermite grenade** consists of a steel canister 2 1/2” in diameter and 4 3/4” high provided with vents in the upper part and containing about 20 oz of thermite. In the top there is a “bouchon-type” fuse with a lever held in unarmed position by a safety pin. A plastic cup with igniter (starter) composition is attached to the lower part of the fuse. When the safety pin is removed, release of pressure against the igniter fuse lever permits the firing pin to strike a primer (after a delay of about 2 seconds), which results in ignition of the starter composition followed by that of the thermite. Combustion continues for about 30 seconds and develops sufficient heat to burn difficultly ignitable materials and to melt metals (even steel). Such grenades may be used for the destruction of guns, airplane equipment, etc about to be abandoned.

By attaching an exterior Primacord detonator, the grenade may be exploded during burning of the thermite, thus scattering particles of molten iron a distance of about 75 feet.

A US Army, WWII, incendiary grenade, AN-M-14, contained a thermite composition designated as “Therm-8” having the following ingredients: Thermite* 70-81%, Ba(NO₃)₂ 14.85-15.15, Al flake 1.01-0.99, Al grained 2.77-2.83, sulfur 0.89-0.91, castor oil 0.303-0.297 (Ref 15)

*Thermite has the following composition (parts by wt): Al (granular) 1.0, Iron oxide, magnetic (Fe₃O₄) 2.75

A **phosphorous grenade** is constructed in the same manner as the thermite grenade, but functions as a bursting rather than a burning munition. On exploding, solid pieces of phosphorous are scattered up to a distance of 75 ft and burn for about 30 seconds. This grenade may be used for setting fire to easily ignitable material, such as gasoline, etc.

The Japanese, during WWII, used two kinds of grenades, those filled with yellow phosphorous and those containing rubber pellets impregnated with phosphorous-carbon disulfide solution (Ref 9, p 56)

**Frangible grenades.** According to Fisher (Ref 9, p 56) and Ohart (Ref 10, p 359), the frangible grenade is an ordinary small glass bottle filled with gasoline (preferably thickened) or other combustible liquid and tightly closed.
Some type of impact igniter is attached by means of a tape to the upper part of the outside of the bottle. The bottle breaks when thrown against a tank or other hard-surfaced object and the igniter sets fire to the liquid. If well aimed against a tank, the liquid may penetrate inside the tank and burn the personnel.

The so-called Molotov cocktail, used for the first time during the last Spanish Civil War, also belongs to the flammable type of grenades.

**Aircraft Bombs.** It was recognized as early as 1910 that bombing from airplanes was of the greatest military importance. It was also recognized that incendiary bombs possessed great potential destructive power. The first successful incendiary bombs were developed by the Germans during WWI and dropped for the first time (1915) from airships over London. The Allies soon followed their example. During WWII, the US Air Force used such bombs very effectively, more so than the Germans, to burn out most of the large German and Japanese cities.

The bombs developed and used from 1915 up to WWII may be divided into the following types:

1/ Large bombs, known as *intensive type*, which practically burn up where they are dropped.
2/ Large bombs, known as *scatter type*, which on functioning, scatter a number of incendiary units over a wide area.
3/ Small unit bombs, called *darts* in the USA, which can literally be rained down upon a target.

Most of these bombs are described in detail by Ray (Ref 4, pp 718-720) and only a short summary is given below.

1/ *Intensive type bombs* comprise a large variety of models and carry practically all types of incendiary materials. The most satisfactory bombs developed during WWI consisted of thermite and large amounts of highly inflammable materials. The first bombs dropped in 1915 by German Zeppelins were of the intensive type. These bombs weighed 20 lbs and contained gasoline, thermite and an ignition mixture. They were wrapped with tow rope impregnated with tar and Ba(NO₃)₂. A percussion fuse served to cause the first ignition.

These bombs were dropped from airships only.

A later type of German bomb weighed 10 or 20 lbs and could be dropped either from airplanes or airships. It consisted of a sheet-metal torpedo-shaped container filled with an incendiary composition consisting of a pasty mixture of gasoline and paraffin with or without K perchlorate. The igniter was a mixture of Al and Fe filings with Ba(NO₃)₂, placed in a central tube and a charge of Black Powder was used to disrupt the casing and scatter the incendiary material. The bombs were wrapped on the outside with tow rope and then were shellacked.

The German example was followed by the French and British and later by the Americans.

A 20 lb bomb developed by the French was called “Chenard” and was considered to be successful. It contained a mixture of rosin and celluloid as the principal incendiary material. The bomb ignited while falling and reached the target in flames.

The British developed several intensive type bombs, such as the one filled with gasoline and ignited on impact by a Véyé cartridge but none was as successful as the small unit bombs developed by them later.

The US developed a 50 lb bomb which consisted of a sheet zinc body in the shape of a prolonged ellipsoid, filled partly with thermite (bound with Na silicate) and partly with gelled oil. The bomb was provided with an impact fuse connected to an igniter. When the bomb functioned on landing, the thermite was ignited and burned through the casing. The enormous amount of heat liberated liquified and ignited the gelled oil to spread the conflagration.

2/ *Scatter type bombs.* A 20 lb impact-functioning bomb developed by the French, contained as incendiary material cotton impregnated with a flammable liquid mixed with K chlorate and paraffin. As a combination igniting and explosive material, a sulfur-bound thermite (Thermaloy) was used. Thermaloy can act as a mild explosive as well as a great heat producer, when ignited under confinement. Thus, on ignition of the thermite the bomb exploded, scattering burning thermite and the impregnated cotton. However, this bomb was not considered a great success.

A 40 lb air-burst bomb developed by the British was filled with yellow phosphorous and
was used against observation balloons and ground targets.

A 50 lb “scatter-type” bomb developed by the US is identical with the intensive type as far as outside dimensions are concerned. Incendiary material to be scattered consists of either cotton-waste balls impregnated with inflammable materials, such as turpentine or CS₂, or gelled oil held in small celluloid containers. When the bomb lands, ignition and ejection of the units are effected by an explosion of Black Powder in the nose.

This type of bomb was never used because by the time it was fully developed the small unit bombs, or “darts,” were considered more effective.

3/ Small unit bombs. The British originated the idea of dropping a large number of small unit bombs instead of one large scatter type bomb and consequently developed the 6.5 oz bomb.

In its action the little bomb resembles that of a mortar and projectile. It consists of a cartridge very much like a shotgun shell which is functioned on impact by a strike point in the base of the body of the assembly. The flash from the cartridge ignites an incendiary charge of “flaming thermite” consisting of 11 pts of thermite and 6 pts of Ba(NO₃)₂. The little bombs were loaded into containers hung underneath an airplane in such a manner that containers could be released individually. Each container carried 144 or 272 bombs and the total number of bombs carried by a single plane was 16,000.

Following the British idea, the US Army developed two types of small bombs and called them “darts.”

The first type, primarily intended for use against crops and forests, burned with a large flame but did not have great penetrating power. It was in the shape of an elongated shotgun shell and contained an incendiary mixture consisting of Ba(ClO₃)₂ 54, rosin 16, Al 14 and asphaltum varnish 16%. The primer was a mixture of reduced iron and KMnO₄.

The second type, intended for use against buildings, contained a thermite-gelled oil mixture as an incendiary. This bomb had sufficient penetrating power to pierce a roof.

The unit bombs were not produced in quantity during WWI but, in modified form, they found wide use in WWII.

**Bombs used during WWII**

A most important contribution to incendiary warfare was made by the Germans, who conceived the idea of using the magnesium alloy, called “Electron” (see below) for the construction of bombs. Although this bomb was invented as early as 1917, it was not used in WWI, but was first employed in the military operations of WWII, beginning in 1939 (See Elektron Bomb in Vol 2, pp B234 to B237). It caused considerable damage to London and other British cities during the early part of WWII but later in the war when the Allies, especially the US, developed their own incendiary bombs of better quality and in larger quantities than those of Germany, it was the Germans and their Japanese allies who suffered most. The Italians were spared incendiary warfare.

The US Chemical Warfare Service produced four types of incendiary bombs during WWII. One type, the magnesium bomb, was copied after European designs, while the other three types were developed after Pearl Harbor, entirely by American scientists.

The M50A2 4 lb Incendiary Bomb was similar to British and German models. It was hexagonal in shape and had a body of cast-magnesium alloy (1.25 lbs), an iron nose plug and a sheet-metal tail. It was 21.3" long and 1.69" across the flats. It was filled with 265 grams of “Therm 64-C,” which has the following composition: Al (granular) 16.0, Al (flake) 9.0, Fe₃O₄ (iron scale) 44.0, Ba(NO₃)₂ 29. sulfur 2.0%. This filling is ignited by a “first fire” composition (igniting mix). It burns fiercely for one minute (without consuming outside oxygen) at 2985°C and melts and ignites the magnesium alloy of the casing. Some of the magnesium vaporizes and the Mg vapor mixed with air burns with a very hot flame (2000°C). The magnesium continues to burn for a maximum time of 10 minutes and due to the fact that it requires little outside oxygen (0.9p per 1p of Mg) it can burn even in closed buildings. Fires produced by such bombs are difficult but not impossible to extinguish. The disadvantage of such bombs lies in the fact that the fire is mostly confined to a very limited area. These bombs are generally released in clusters.
In order to spread fires over a larger area, an explosive element is sometimes added to
the bomb so that when it explodes, after being
partly burned, it scatters burning pieces over an
area having a radius of as much as 50 ft. This
greatly hampers fire-fighting because it is never
known just when the bomb will explode and
cause injuries to fire-fighting personnel. In the
US M50X2A3 bomb, 36 grams of tetryl is used
as the explosive charge and a delay fuse and
detonator is included in the ensemble (Ref 10,
p 235 & Ref 15, parts III & V)
The M-69 (6 lbs) incendiary bomb is a
hexagonal case of sheet steel 19.5” long and
2.9” across the flats. Its central portion con-
tains a charge of 2.2 lbs of “gelgas” or “Napalm”
held in a cheese cloth sack and an ejector-
igniter charge (placed near the fuse) of 0.4 oz
of Black Powder and magnesium. This was later
replaced by a Tetryl booster surrounded by
phosphorous, serving as igniter for the gelgas
or Napalm. Attached to the outside of the bomb
are four lengths of green-colored cotton gauze,
40” long and 3” wide. These break free as the
bomb starts to fall and act to stabilize the
fall and to provide drag, so that the descent
slows down to 225-250 ft per second. This
prevents the bomb from smashing to pieces
when it lands, yet the bomb has sufficient force
to pierce tile, slate, wood or galvanized iron
roofs. After the bomb lands, the impact activ-
ates a delay fuse which, after 3 to 5 seconds
delay, ignites the “igniter-ejector” charge. This
jects the sack with burning gelgas or Napalm
from the tail of the casing to a distance of 75
to 200 feet. The gel scatters around as “gobs”
of fire, which cling to surrounding objects and
continue to burn for the next 8-10 minutes.
The bomb may be fused either for delayed
action so as to permit penetration into struc-
tures before functioning, or for immediate action
so as to scatter flaming material over roofs or
other surfaces
In the 6 lb M69X bomb, 0.4g of the jelled
gasoline was replaced in the nose of the bomb
by a charge of Tetryl to act as a burster charge.
The bomb is so constructed that when the
main ejection-ignition charge forces the “gelgas”
from the bomb case, a delay fuse leading to
the Tetryl charge is also ignited. After a delay
of from 1 1/2 to 6 seconds, the explosion of
the Tetryl fractures the entire nose of the
bomb, producing more than 400 fragments.
This bomb acts not only as an incendiary, but
also is effective against personnel (Ref 15, parts
III & V)
In the M69-WP bomb, 0.4g of the gel con-
tained in the nose of a regular M-69 bomb
was replaced with a plastic cap containing white
phosphorous. When the “gelgas” is ejected from
the casing, the force of explosion breaks the
cap and scatters the WP about the area. The
smoke produced by WP obscures objects, makes
breathing difficult and hampers fire-fighting
All the types of bombs described above were
packed together in closed containers (containing
from 25 to 100 bombs) which fell like a single
bomb from a plane and broke open a few
thousand feet above the target, releasing the
individual bombs in a tight pattern. The hexa-
gonal shape of these bombs allowed packing
them in larger quantities per unit space (Ref
15, part III)

**Bomb Release Individual** weigh 70 lbs
and up, for example, the M47A2 and the M76
US bombs. These were designed for precision
bombing. A particular advantage of such bombs
is their ability to penetrate heavily-roofed
structures that would resist the impact of light
(cluster) bombs. When dropped from high alti-
itudes, these bombs have impact velocities of
about 1000 feet per min and may pass through
reinforced concrete up to 15” thick

The M-47 incendiary bomb weighed approx
70 lbs and consisted of a sheet-steel cylinder with
a rounded nose and tail. It was 45” long and
8-1/8” in diameter and was filled with jelled
gasoline. Originally, a Black Powder central
burster was used, but it was not as satisfactory
as the later model with a TNT-Tetryl burster,
which was surrounded by white phosphorous
used as the igniter for the gasoline gel. This
bomb was initiated by a nose fuse

The M76 bomb, also called “goop bomb” or
“Blockburner,” was the largest incendiary bomb
used in WWII. It weighed 475 lbs. It consisted
of a metallic cylinder with rounded nose and
tail and was loaded with 180 lbs of “pyrogel”
(see below). It was provided with standard nose
and tail fuses and a standard igniter and
burster. On reaching the target, the burning
pyrogel was scattered by the force of the
bursting charge to a distance of 150 to 500 ft and burned with an intensely hot flame. The pyrogel mixture combines the good features of both jellied gasoline which scatters and penetrates into hard-to-reach corners of structures, and of magnesium which concentrates a white-hot flame on the target (Ref 15, parts IV & V).

For other standard incendiary bombs (1946) see Ref 15, part V and table of contents of part I. For incendiary compositions, both experimental and standard, see part II.

**Jettisonable Airplane Fuel Tank Bombs.**
Another type of US incendiary bomb was the coverted auxiliary airplane fuel tank of 75 to 300 gallon capacity. They were filled with jellied gasoline and provided with a fuse and an igniter. Such bombs spread fire over a large area and were very effective against Japanese targets (Ref 10, p 239 & Ref 15, part IV).

In addition to the "electron" bomb previously described, the following foreign incendiary bombs used in WWII might be mentioned:

a/ **German Incendiary-Explosive Bomb** was a magnesium incendiary bomb which was lengthened in the nose to provide space for a high explosive charge. The total weight of the bomb was 5 lb and the total length, including the tail, was 21". On landing, the incendiary part of the bomb ignited but the explosive part remained intact and exploded up to 7 minutes later. In some bombs, the explosive part became detached on landing and rolled away from the incendiary section.

b/ **German 50kg Incendiary-Explosive Bomb** consisted of a casing, 30" long and 8" in diameter, which contained, besides the incendiary charge, a 12 lb charge of TNT lodged in the nose. On impact, the bomb ejected 60 small metal containers with thermite-type filling and 6 larger tumbler-shaped fire pots containing a magnesium-type filling. This was followed almost immediately by explosion of the TNT charge.

c/ **German 50kg Phosphorous Incendiary Bomb** consisted of a casing similar to the previous bomb, filled with a mixture of yellow phosphorous 86.5%, benzene 13.1 and polystyrene 0.4%. The bomb was split open on impact by the bursting charge and the incendiary was scattered in the form of a sticky self-igniting liquid.

d/ **German 50kg Petroleum Solvent Incendiary Bomb** was similar to (c) but contained petroleum solvent 87.7%, polystyrene 11.7 and phosphorous 0.5% as the incendiary mixture.

e/ **Japanese 1kg Incendiary Bomb** consisted of a casing, 10" long and 3½" in diameter, filled with red phosphorous and containing a burster (exploder) tube filled with Picric Acid. The bomb was exploded on landing by the PA. This ignited and scattered the phosphorous. Fragments of the bomb were thrown as far as 150 ft.

f/ **Japanese 50kg Rubber-Pellets Bomb** consisted of a casing, 40.5" long and 4.5" in diameter, filled with many rubber pellets or balls impregnated with yellow phosphorous in carbon disulfide and containing a burster tube with high explosive, such as PA. On landing, the bomb was exploded by the burster and pellets were scattered to a distance of up to 150 ft. As soon as the CS2 evaporated, the phosphorous ignited the rubber and this continued to burn for 5-7 minutes.

g/ **Japanese 60kg Thermite Bomb** consisted of a casing, 40" long and 8" in diameter, containing three "electron" inserts filled with thermite and a burster charge of high explosive. On landing, the burster charge scattered the burning inserts as separate thermite units, thus increasing the radius of damage.

h/ **Japanese 60kg Gelled-Oil Incendiary Bomb** consisted of a casing, 42" long and 9.5" in diameter, filled with gelled oil, such as kerosene and paraffin wax. Inside the filling there was a tube filled with thermite which had a quick match running through its center. On impact, the oil was ejected and was ignited by the thermite.

i/ **Japanese 560 lb Bomb** consisted of a cylindrical, metallic casing loaded with pieces of 1-inch iron tubing, each 2-3/4" long. It was filled with a special thermite mixture. On landing, the burster charge fractured the casing, ignited the thermite and scattered the pieces of tubing to a distance of about 150 ft.

j/ **Flame Thrower—described separately in Vol 6 of Encycl.**

k/ **Miscellaneous Incendiary Ammunition.** Among these may be mentioned "incendiary cans and boxes" used during WWI as a means of defense against gas attacks. The heat generated by these devices produced air currents.
which dispersed and deflected the gases

Various incendiary devices were designed for sabotage purposes, among them the ingenious device called "incipient blue pencil," previously described

Special devices were attached to gasoline tanks of airplanes by means of which the airplane could be quickly destroyed in case it was forced down in enemy territory

I. Rocket Incendiary Projectiles. During WWII several rocket incendiary projectiles were developed, among them may be cited:

German 32cm Rocket Projectile Incendiary
carried a charge of 13 gals of flammable oil.
It was used in anti-aircraft fire (Ref 9, p 55)

5-inch Incendiary Rocket, used at ranges of about 5,000 yds, was an effective weapon when fired from landing craft in amphibious operations. The rocket head was loaded with thermite or oil incendiary mixtures

German R100BS Air-to-Air Rocket. The rocket weighed 100 kg and was 210 mm in dia and 1,800 mm long. It was propelled by 25 kg of conventional solid propellant which gave the missile a specific impulse of 4,200 kg/sec with a burning time of 0.9 sec. The rocket attained a speed of 550-600 m/sec and had a range of about 2,000 m. The warhead weighed about 30 kg and contained 460 thermite-filled cylinders each weighing 55 g. The pellets were given an initial velocity of 500 m/sec in addition to the velocity of the rocket. The rockets were mounted under wings of a plane (Ref 14)

German Enzian Ground-to-Air Rocket. The rocket weighed 1,800 kg which included the weight of four assisted take-off units weighing 80 kg each. These units functioned for 5 sec and were then jettisoned. The missile had a length of 12 ft and a wing spread of 14 ft. The power supply had a duration of 73 sec which resulted in a vertical range of 16,000 m and a horizontal range of 25,000 m. The payload was either 300 or 500 kg. These warheads consisted of a metal shell, 1 1/2 mm thick containing cylindrical pellets of mild steel, 20 mm in dia by 30 mm long containing an incendiary core. This rocket appeared near the end of WWII and large-scale production was never attained before hostilities ended (Refs 14 & 15)

According to Fisher (Ref 9, p 55), the applicability of incendiary-filled rocket projec-
tiles to conditions of modern warfare is still somewhat uncertain despite the increasing importance of the rocket itself. At extreme ranges, the rocket cannot compete with the bomber in attacking incendiary targets because of the diffuse pattern of rocket impacts


a. TM3-215, Military Chemistry & Chemical Agents (1942), pp 149-150
b. TM9-1980, Bombs for Aircraft (1942)
c. TM3-325, Livens Projector MI (1942)
d. TM3-330, Incendiary Bombs (1942)
e. Flame Throwers (See below)


The following references on incendiary compositions, devices and uses are given in Chemical Abstracts:

G. Fox & G. Quayle, 13, 1766 (1919), Igniter compound for "thermite": Na$_2$O$_2$ 50, Mg powder 10, sulfur 5, KNO$_3$ 5, carboxic acid 30. It ignites on contact with H$_2$O

J. Buckingham, 13, 1931 (1919), Incendiary projectiles containing yellow phosphorous with or without powdered aluminum

G. Vautin, 13, 2284 (1919), Binding material for thermite, etc, consisting of Na and K silicates alone or mixed with borax or gelatinous Fe(OH)$_3$

G. Webb, 13, 2284 (1919), Incendiary mixtures suitable for projectiles: MgO 1.1p, Mg (wax-coated) 3.7p, Ba(NO$_3$)$_2$ 7, H$_2$O—sufficient to make the mixture set

G. Webb, 13, 2284 (1919). Incendiary composition, such as MgO 3.45p, Mg (wax- or resin-coated) 3.7, Ba(NO$_3$)$_2$ 7. This mixture was dry-pressed. The alkaline earth serves as a retarder of combustion and, by varying the quantity of this ingredient, the rapidity of combustion can be fixed within wide limits

A. Chanard, 14, 350 & 1045 (1920). An incendiary material prep'd by incorporating a small amount of celluloid or NC in combustible substances such as resin dissolved in gasoline, heavy oil, tar, etc

W. T. Scheele, 15, 3751 (1921). An incendiary mixture adopted for use in bombs and shells contains hexamethyleneetramine ((CH$_2$)$_6$N$_4$) 22.5 to 33.33 and Na$_2$O$_2$ 77.5 to 66.66

Another mixture contains hexamethyleneetramine (CH$_2$)$_6$N$_4$ 25, Na$_2$O$_2$ 50 and paraffin (or cylinder oil) 25

J. H. Hammond Jr, 17, 472 (1923). Incendiary shell containing a thermite charage ignited by a mixture of BaO$_2$ and Al, fired by a concussion fuse

W. L. Clay & A. H. Hallowell, 17, 473 (1924). Incendiary shell-structural features

A. Wolfson, 26, 4720 (1932). Incendiary compound, which may be ignited by a spark produced by friction, is prep'd by mixing solid fuels, such as hex (CH$_2$)$_6$N$_4$ or metaldehyde, with readily flammable substances, such as NC, P or nitrophenols. The composition may be pressed into tablets

A. Stetzbacher, 30, 1636 (1936) and 32, 8781 (1938). Review of chemical and mechanical incendiary devices

T. A. Craven, 31, 1618 (1937). Incendiary projectile containing a charge prep'd by coating yellow phosphorous with a solution of a vegetable resin and a second coating of KClO$_3$—S$_2$S$_3$ composition. The charge may be wrapped in cotton gauze impregnated with NaNO$_3$ with or without Mg composition to facilitate ignition

J. L. Nayler, 33, 1941 (1939). Description of various incendiary bombs, particularly those containing oils, Mg and P, and ways of combating the fires

E. Fisher, 37, 5241 (1943). Incendiary suitable for filling bombs and flares is prep'd by mixing thoroughly Mg dust with pulped paper in water and then filtering to produce a uniformly flat sheet or ribbon. After drying, the sheet is cut into pieces

W. R. Bluedorn & R. N. Nelson, 38, 257 (1944). Incendiary bullet of special construction containing finely divided Ti as combustible material

Chemical Warfare Service, 38, 1295 (1944). Brief history and description of modern incendiaries

C. W. van Hoogstraten, Chem Zentr 1941, II, 3272 & CA 38, 1880 (1944). Various types of incendiary bombs as well as means of combating the fires produced by them are described

W. C. Kabrich, 39, 2649 (1945). Research in the Technical Division of Chemical Warfare Service in incendiaries, etc

W. K. Griesinger, 39, 1284 and 4742 (1945). Canadian Pat 424914 (1945). Gelatinization of hydrocarbon distillates in order to prepare solidified fuels suitable as incendiaries, is achieved by mixing a hydrocarbon distillate with 0.5-10% by volume of an aqueous solution containing 10-40% by weight of petroleum sulfonate and
5-15% by weight of an alkali metal hydroxide. Petroleum sulfonate was derived from acid sludge separated from partly cracked gasoline-oil fractions which had been treated with \( \text{H}_2\text{SO}_4 \)

Ford Motor Co, 40, 210 (1946). Incendiary material is prep'ed by reducing MgO to Mg vapor and quenching the vapor in oil. This produces a thin slurry consisting of finely divided metallic Mg and oil. The slurry is allowed to settle, the oil is decanted and the thick slurry is filtered, leaving a thick, heavy mud

L. F. Fieser et al, 40, 5567 (1946) and Ind EngChem, 38, 768-73 (1946), "Napalm" (Detailed description of preparation and properties). Some other thickened fuels are described

A. Grobstein, 40, 5568 (1946). Incendiary with high penetrating power is prep'ed by placing a capsule with Li nitride (\( \text{Li}_3\text{N} \)) into an incendiary bomb filled with thermite or other composition. On burning, the thermite produces sufficient heat to fuse the nitride into a mass, which then becomes a penetrating, hot, corrosive agent. Nitrides of Ce, K, Ca or La are also suitable

J. Billing & J. W. Fisher, 40, 6818 (1946). Incendiary projectiles are charged with a highly inflammable thickened fuel, prep'ed by dissolving a quantity of metallic soap of a fatty acid containing 10 or more C atoms per mole, in a liquid hydrocarbon, preferably of the benzene series


H. R. Dittmar & D. E. Strain, USP 2443378 (1948) & CA 42, 6538 (1948). Incendiary gels containing a liquid hydrocarbon (such as gasoline, benzene, etc), 3% or less \( \text{H}_2\text{O} \), some alkali (such as \( \text{NH}_3 \), \( \text{NaOH} \), \( \text{Ca(OH)}_2 \), etc), 1 to 10% of a salt of a hydrocarbon-sol, salt-forming, acrylic resin (such as polymers of alkyl acrylates, or alkyl acrylates contg small amts of free COOH groups), with or without salts of fatty acids (such as alkali metal stearates, palmitates, oleates, etc

H. H. Cooke & E. J. Holzclaw, USP 2445311 and 2445312 (1948) & CA 42, 7985-6 (1948). Incendiary mixtures consisting of flammable naphtha thickened to form a highly viscous mass by means of isoolesfin polymers (such as isobutylene polymer, mol wt 50,000 to 100,000). Small amounts of metal soaps may replace part of the polymer. Na, K, P and thermite mixtures may be included. The bomb contains a nose fuse, a black powder ignition charge, a thermite nose charge and a tube of thermite extending through the incendiary charge

D. P. O'Brien, USP 2451864 (1948) & CA 43, 1190 (1949). An explosive charge for incendiary bombs consists of Mg 32p, \( \text{Ca}_3\text{P}_2 \) 2p and \( \text{KClO}_4 \) 1p

D. L. Woodberry et al, USP 2452091 (1948) & CA 43, 1190 (1949). Thermite mixtures can be improved by incorporating \( \text{Ba(NO}_3\text{)}_2 \), small amount of an oil and addnl Al beyond that required by the thermite; eg, Fe oxide scale 61.2, granular Al 19.2, grained Al 2.8, flaked Al 1.0, \( \text{Ba(NO}_3\text{)}_2 \) 14.6, S 0.9 and castor oil 0.3%. Particle sizes are of importance and are specified in detail

Wm. F. VanLoenen, USP 2530489 (1950) & CA 45, 2670 (1951). It is disclosed that a mixture of Mg, C and MgO obtained by the Hensgirg process (see below) protected by hydrocarbon oils and a metallic carbide, such as \( \text{CaC}_2 \) or a phosphide, form an incendiary composition which ignites in water or similar oxidizing liquid

Note: The USP 2530491, 2530492 and 2530493 describe various modifications of the above incendiary mixture

F. Hensgirg, USP 1884993 (1932) & CA 27, 942 (1933). A mixture of Mg, C and MgO prepared by a special process

L. Finkelstein, USP 2553668 (1951) & CA 45, 7354 (1951). Gelled gasoline for filling incendiary bombs is prep'ed by dissolving in 88.75p gasoline, 3p stearic acid, keeping the temp at 26°. Isobutylmetacrylate 5p was next stirred into the mixture, then CaO was added and finally water 1.25p

M. E. Barker, USP 2558726 (1951) & CA 45, 9864 (1951) describes an incendiary device consisting of a mass of NC in the form of a disc, or leaf, with a hollow center filled with white P and sealed with a material such as gelled Na silicate. The disc is stored under water, or in atmosphere of saturated steam. When exposed to air the coating of silicate loses its water of gelation thus making it porous. Air then penetrates through the pores and causes
the P and then the NC to ignite. The thickness of silicate coating is so adjusted that a time delay of one to 3 hours is obtained after start of air exposure.

J. A. Southern et al, USP 2570990 (1951) & CA 46, 1768 (1952). An easily combustible mixture suitable as fuel for incendiary bombs and grenades, flame throwers, etc. Consists of 7-14% volatile hydrocarbon fuel such as gasoline, and 93 to 86% of soap-type gelling agent, which is composed of Al oleate 50-75, Al stearate 25-50, to which is added about 1% of oxy-aromatic antioxidant compound.


L. F. Fieser, USP 2606107 (1952) & CA 47, 1392 (1953). Incendiary gels prepared by mixing light hydrocarbon fuels with thickening such as Al soap of a soap-forming carboxylic or unsatd fatty acid or the Al salt of such acids as the other component; eg, gasoline 90, Al coconut soap 5, Al napthenate 5%.

M. D. Banus & J. J. McSharry, USP 2688575 (1954) & CA 48, 14210 (1954) specify that Ti powder for use in first-fires mixtures for incendiary bombs or shells should have a density of 0.4 to 1.7 and contain <0.425% hydrogen. Method of attaining these specs are described.

L. E. Medlock, BritPat 742283 (1955). No CA listing found. These non-detonating, waterproof compositions consist of pulverulent oxidizing and reducing agents distributed uniformly through rubber or artificial rubber. Used for such purposes as fuses, igniter cords, and incendiary charges.

H. Z. Cier & H. G. Schutze, USP 2794003 (1957) & CA 52, 1602 (1958). Incendiary gels, solid fuels, etc are prep'd without external gellation agents when Na, K, Ca, Ba, Hg or Al derivs of oil-sol sulfonyl chlorides of satd paraffin or napthenic hydrocarbons contg 6-16 C atoms are formed in the hydrocarbon to be gelled. Most of the gels contain 0.1-5.0% water, alkali and 1-5% metal. The hydrocarbon, SO2, and Cl are exposed to light at 150°F for 15-360 min to give the sulfonyl chloride; addn of a basic Al soln or Al foil or powder with stirring forms the gel.

C. Blake & Wm. S. Graff, USP 2819163 (1958) & CA 52, 5826 (1958). An incendiary for military purposes is described. It contains a core of Zr, Ti, and Pb which sparks intensely when abraded and an enveloping Fe case. A compn contg 35% Zr, 30% Ti and 35% Pb (Zr and Ti <15 μ, Pb -200 mesh) is mixed with a xylene-4% Perbunan (synthetic rubber) soln in the presence of 0.875 cc/g of powd metal. The particle size of the electrolytic Fe powder for the case is important in order that the porosity be controlled.

C. B. Linn, USP 2881066 (1959) & CA 53, 14522 (1959) claims incendiary fuels consisting of gels formed by the condensation of 2 moles of an aromatic hydrocarbon with a ketone and gel-formation in benzene. The condensation of d-fructose with 2 moles of toluene, in an autoclave with HF catalyst, is given as an example.

Raymond E. Schaad, USP 2891852 (1959) & CA 53, 17513 (1959). Materials useful as rocket fuels, semisol or gel fuels for bursting and tail-ejection-type bombs, and incendiary fuels for flame throwers and hand grenades are described. They are made by mixing 0.1-25% by wt of satd, unsatd, or aromatic nitrohydrocarbons or their mixts, such as nitro- or dinitromethane, -ethane, -propane, or -butane with divinylated ketones or diaryl deoxyketitols prep'd by reaction of C5-H ketose sugars with C5-54 aromatic hydrocarbons. The latter include C6-H6, toluene, naphthalene, anthracene and their alkylated derivs.


P. Chereau, FrP 1318773 (1963) & CA 58, 13702 (1963), claims: rocket fuel or incendiary composed of a combustible metal, eg, Al, Mg, or Li or a liquid fuel such as kerosine, in fine grains or droplets encapsulated in situ by formation of a polymer skin. Thus, 0.18 of 2,4-tolyene diisocyanate is dissolved in 41.8g of paraffin oil. A portion (24.5g) of this mixt is added drop by drop to a stirred soln contg 2g of ethylene glycol in 250g water. Discrete spherical particles
are formed which are encapsulated by a skin of polyurethane. The particles are sepd by
centrifugation

A. D. Coates & E. O. Baicy, USP 3120459 (1964) & CA 60, 9094 (1964) claim the
following: A storage-stable powder useful as an incendiary component, a solid rocket-fuel additive, or a metal heating powder is obtained by coating a powd material which contains excess O or liberates O on heating, such as KClO₄ (preferred), NaClO₄, Ba(ClO₄)₂, KNO₃, NaN₃, Ba(NO₃)₂, or NaClO₄, under conditions which avoid decomn with 30-80% of final coated product) of an exothermic metal, such as Al (preferred), Mg-Ti alloys, or certain Mg-Al alloys, by treatng the O-contg material with metal vapor at 5 x 10⁻⁴ mm. Thus, 40-80 mesh KClO₄, treated as described with Al vapor produced by heating Al ribbon or wire with Ti wire filaments, yields a product which, when initiated by flame or an elec impulse, reacts in a self-sustaining manner

F. N. Vannucci, USP 3126259 (1964) & CA 60, 15676 (1964) claims a gelatious incendiary material is made by mixing 1 part of a powdered material consisting of SiO₂ & MgO with an equal amt of water, adding 1 part of a hydrocarbon such as kerosene or gasoline, and separating. The resultant incendiary can be used to ignite coal or other fuels or in incendiary shells, bombs & flame-throwing devices. The powdered material can be made from serpentine rock

R. J. Laran, USP 3157464 (1964) & CA 62, 2530 (1965) claims: HClO₄ & TiCl₄ were combined at -10°C in a mole ratio of 8:1. The product was 97-8% Ti(ClO₄)₂. This compound exists in 3 cryst form; it can be stored without decomposition if air & H₂O are excluded & temp is kept low, eg, with dry ice. Also, freshly sublimed BeCl₂ & anhyd HClO₄ were allowed to react for several hrs at -15°C. A white powder Be(ClO₄)₂ was obtained in 16% yield, excess acid having been removed in vacuo. Close temp control (-75 to 25°C) & the absence of H₂O are essential to these reactions. The products are powerful oxidants, insensitive to shock & thermal decomn, useful as igniters for propellants & incendiaries. Ti(ClO₄)₂ explodes on contact with Et₂O & ignites on contact with HCONMe₂ or HCONH₂

P. Huber, BelP 644290 (1964), claims poly-styrene and/or polyisobutylene (3-5%) can be used as a gelling agent for inflammable hydrocarbons & other org liquids, such as CS₂. Depending on the intended use, the compositions can be made shock-resistant, adhesives, and/or productive of thick black smoke

Review article: Chemical warfare: incendiaries. L. W. Greene, Kirk-Othmer Encycl Chem Technol, 2nd Ed, 4, 895 (1964) & CA 65, 3661 (1966); including a survey of incendiary requirements, agents & bombs; 8 refs

J. E. Claiborne, USP 3250652 (1966) & CA 65, 2057 (1966) claims: exothermic composns useful as propellants, incendiaries & nuclear blast simulators, which upon addn of small amts of H₂O produce flame & hot expanding gases, are provided by intimate mixts of 10-100 mesh powdered metal (preferably at least half Zn), 10-250 mesh NH₄NO₃, & 10-250 mesh NH₄Cl as a loose mass or as blocks, grains, or strands bound with an anhyd heat-softening gum or a thermost plastic resin or lacquer. Conventional diluents may be included to reduce reaction rate if desired. Thus, a mixture of 40-80 mesh Zn 30/powd NH₄Cl 20/powd NH₄NO₃ 47/ & polyvinylchloride molding powder 3% was heated to the softening point & pressed into blocks which, when exposed 6 hrs to 85% relative humidity, spontaneously ignited, or, when immersed in 10% of their wt of H₂O yielded copious amts of H, NH₃, Cl, N, & NO₂

H. W. Koch & F. Poppler, NASA Access No N66-31925, Rept No T-37/65 (1965) & CA 66, 117, 557, (1967), discuss the chem compsn & incendiary characteristics of napalm & Phosphor-Mg compounds. Test arrangements for obtaining temp rates in various environments are described. Burning time is measured as a function of amount (sic) & spatial extension of the flame

A. Lachs, USP 3314836 (1967) & CA 66, 117565 (1967) claims a military flame producing compsn containing colloidal yellow P in a flammable medium

M. Picone, USP 3396060 (1968) & CA 69, 78900 (1968) describes a long-burning, low d incendiary compsn containing Ti/Al-Mg alloy/oxidizer. The oxidizers used were Ba or Amm nitrates and K-perchlorate

J. R. E. Pleasant et al, USP 3414443 (1968) & CA 70, 39441 (1969) claim an incendiary
compn, which may be self-igniting for emergency kits & survival & distress fires. Metals or Metal hydrides are mixed with wax & a gelling agent

S. Schiff, USP 3416899 (1968) & CA 70, 39432 (1969). A napalm-type material is prep'd by hydrogenation of a hydrocarbon soln of a 3:1 butadiene-styrene block copolymer

P. G. Yur'ev, Zh. Vses Khim Obshchest 13 (6) 648 (1968) & CA 70, 69710 (1969), presents a general review of chemical incendiaries with particular emphasis on US Army weaponry; with 20 refs thru 1967

G. H. Custard, USP 3421439 (1969) & CA 70, 89345 (1969). An incendiary projectile is claimed which consists of a 20mm hollow body equipped with an impact-responsive primer, a booster charge & a main HE charge. The body of the projectile is of Ti or Zr which on ignition burns at temps of 2725-3325°C for up to 1 min

S. Samuel et al, USP 3441955 (1969) & CA 71, 23400 (1969). White P is mixed with hexamethylenetetramine to produce a smoke screen mix which is also an incendiary. Al or Mg may be added to the mix to raise flame temps or paper pulp, cellulose or high-melting hydrocarbons to produce longer burning times

J. Winkler, USP 3460922 (1969) & CA 71, 93351 (1969) claims a thixotropic-gelled flammable hydrocarbon compn, containing powdered metals and/or AN, for use in flame-throwers, rocket fuel and napalm. The hydrocarbon gel is prep'd by an in situ cross-linking of a polyurethane

V. P. Wystrach et al, USP 3464869 (1969) & CA 71, 103746 (1969) claim a match-ignitable, tacky compn for use in flares, signals & incendiaries. Example compn is made by mixing Mg, AN, Napalm B with an acrylonitril/e acrylic acid/Me-methacrylate copolymer. It burns with a bright yellow flame

V. A. Lehikoinen, USP 3498857 (1970) & CA 72, 113395 (1970) claims thermites and thermates (thermites + flake Al, S, hydrocarbon oil and/or oxidizers such as Ba(NO₃)₂, KMnO₄, etc) of improved ignitability due to inclusion of 2-50% of ferrocene-type compounds. Varying the proportion of ferrocene modifies the burning rate of the incendiary compn

Incendiary Flash Tests for Small Arms Ammunition. The purpose of these tests is to investigate the ability of incendiary bullets to develop, upon impact, an incandescent flash, sufficiently hot and sustained to ignite explosive vapors, or to initiate combustion in other readily combustible materials. These tests also establish the relative size, position and persistence of the flash as compared with a standard chart. Tests are made under controlled conditions against a target of fixed design

Detailed description of these tests is given in Ordnance Proof Manual 7-19 (1945) (11 pps)

Incident Wave in Blast. See Vol 2, p B182-L & Fig in p B183 under Blast Effects Due to Reflected Shock Waves

Increments. See under Base Charge of a Propellant in Vol 2, p B24

Increment Appoint (Fr) or Zusatzladung (Ger). An additional quantity of propellant used when the charge contains some flash reducers in order not to impair the ballistics of a weapon

Ref: Davis (1943), 326

IncT or IT. Incendiary Tracer

Indazolone, 5,7 Dinito.

\[
\text{O}_2\text{N} \quad \text{CO} \quad \text{NH} \quad \text{NH}
\]

mw 304.3, N 18.4%, OB -58.0%; yellow prism (from w), decomposes 185-215°C without melting; sl sol in water, benz & chl; moderately sol in boiling alc. Prep'd by reacting 3,5-dinitrobenzene with hydrazine hydrate. The Na-salt, Na₃C₆H₂N₄O₂, cryst, sol in w, alc or acet, explodes on heating. Reaction of the dinitro indazolone with HgCl₂ gives a crystalline needle-like product that is very unstable

Ref: Beil 24, 115 & (239)

Index of Damage. See under Damage Effects & Damage Potential of Air and Ground Blast Waves in Vol 3, pp D3-4

Index of Ignition. See Index of Inflammability under Physical Tests in Vol 1, p XVII
India Saltpeter. Same as Potassium Nitrate

Indicator Test Paper for Detecting Stability of Double-Base Propellants. Approx 60 commercially available dyes were investigated in experimental indicator test papers. Laboratory & surveillance tests indicated that 3 of these selected for further evaluation were less sensitive to double-base propellant degradation products than N/10 Methyl Violet paper. The order of sensitivity was as follows: N/10 Methyl Violet, 0.1% Ethyl Violet, 0.1% Trypan Red & 0.1% Benzoazurine. On exposure to four different types of partially degraded but still serviceable double-base propellants for a period of 1 yr, strips of each of these experimental indicator papers remained either unchanged or became only slightly faded. The N/10 Methyl Violet paper became totally bleached within 3 months.

However, strips of the experimental indicator papers, when exposed to oxides of N fumes generated in the lab and to double-base propellant fumes generated at an elevated temp, became either entirely bleached or totally faded. (Ref: 1) S. Helf, PATR 1782 (Oct 1950). 2) P. Rochlin, PATR 2242 (Sept 1952) (Concluded that the 3 experimental test papers appeared to be superior to N/10 Methyl Violet paper for detecting instability in stored propellants)

Indice Nitrométrique or Indice de l’Action Stabilisatrice is a value introduced by M. Gua & G. Guastalla and reported in AttiAccadScienze Torino 60, 73 (1925) & Chim & Ind 29, 63T (1933)

It was shown by the above authors that nitrogen content of nitric esters such as NG or NC cannot be determined by the nitrometer method, if stabilizers such as centralite are present, because centralite absorbs some nitrogen oxides thus giving a low result for N. The difference between the nitrometer N content of NG in absence of a stabilizer and N content of NG in presence of a stabilizer may be taken as an index of stabilizing action of the stabilizer; the bigger the difference, the better the stabilizer. This difference is called indice nitrométrique

Indigo, 5,5'-Dinitro.

Indigo, 6,6'-Dinitro. A dark-red powder, insol in w, s in alc, eth or benz, s in hot aniline. Prepd by heating 2,4-Dinitrophenyllic acid, methylketone with Na₂CO₃. Sublimes on glass surfaces, puffs off on platinum surfaces (Ref: Beil 24, 429 (382) & [245])

Indole-carboxylic acid, 5,7-Dinitro (called 5,7 Trinitroindol carbonsüre or Dinitrotrychol in Ger).

\[
\text{N} 16.7\%, \text{OB} -91.7\%; \text{pale yellow cryst (from alc), darkens 250°, mp ~284° (with decomp);}
\]

Insol in most solvs except hot alc. Prepd by heating 5,7-Dinitro-indol-dicarboxylic acid in water. Its Et ester-Trinitro derivative melts at 202-205° and is stable (Ref 2)


Indole-carboxylic acid, 5,7,x-Trinitro (called 5,7 Trinitroindol carbonsüre or Trinitrotrychol in Ger).

\[
\text{N} 18.9\%, \text{OB} -63.0\%; \text{leaflets (from water), mp 218-220° (with decomp); sol in hot water or}
\]

Insol in benz or eth. Prepd by nitrating 5,7-Dinitro-indole-dicarboxylic acid with fuming nitric acid. Puffs off on heating

(Ref: Beil 22, [49])

Induced Ground Waves. See Vol 3, p D4-L under Damage Potential of Air and Ground Blast Waves

Induction Period. See under Delay to Ignition in Vol 3, p D53-R; also under Hot Spots in this Vol

Indurite. The first successful American smokeless propellant invented by C. E. Munroe in about 1891 and patented in 1893 (USP 489684). It
was prep'd by washing ordinary Guncotton (GC) with methanol, in order to remove the lower NC's, and then gelatinizing (colloiding) it with NB (nitrobenzene) in the proportion, GC 40 and NB 60. It was made with or without oxidizing salts, such as KNO₃ etc. The resulting gelatinous mass was rolled into sheets of desired thickness and cut into squares or strips which were hardened ("indurated") by the action of hot water or steam. This treatment also distilled off most of the NB and left a hard and tough colloided substance.

When tested, it gave satisfactory results in guns ranging from the one-pounder to the 6" gun. The reason why it was not adopted by ordnance was that at a slightly later date Lt Bernadou introduced in USA pyro powder, which was the same as Mendeleev's "pyro-collodion" powder, but with slightly higher nitrogen content. Refs: 1) C. E. Monroe, JACS, 18, 817-46 (1896) 2) Davis (1943), 296

Industrial (or Commercial) Explosives, also called Civil (or Civilian) Explosives. See Agriculture and Forestry Use of Explosives in Vol 1, p A112-R; Blasting Explosives in Vol 2, p B202-L; Commercial or Industrial Explosives in Vol 3, p C434-R; and Dynamites in Vol 5, p D1584-L

Industrial Disaster Control is briefly discussed in the "Dangerous Properties of Hazardous Materials" by N. Irving Sax, Reinhold, NY (1957), pp 166-169


"WHAT IS FIRE?" Fire, combustion, or burning requires three things: (1) a fuel (any oxidizable material), (2) oxygen (usually air), and (3) a certain temperature (heat). Fire is the chemical union of oxygen with fuel, accompanied by evolution of thermal energy, indicated by incandescence or flame. If any one of these three constituents is not present in the proper proportions or degree, no fire will occur. If a fire exists and even one of them is sufficiently altered, the fire will go out. Therefore, in its simplest form, all fire control or extinguishment reduces to a manipulation of these three essential constituents.

Classification of Fires

Class A Fires. These are fires in ordinary combustible materials where the quenching and cooling effects of quantities of water or solutions containing large percentages of water are of first importance. Ordinary combustible materials tend to produce glowing embers after burning, and these must be quenched to prevent rekindling.

Class B Fires. These are fires in flammable liquids (oils, gasoline, solvents, etc.), where a blanketing or smothering effect is essential to put the fire out. This effect keeps oxygen away from the fuel, and can be obtained with carbon dioxide, dry chemical (essentially sodium bicarbonate), foam, or a vaporizing-liquid type of extinguishing agent. Water is most effective when used as a fine spray or mist.

Class C Fires. These are fires in electrical equipment, where the use of a nonconductive extinguishing agent is essential. Water spray, carbon dioxide, dry chemical, or vaporizing liquid is satisfactory.

Metal and Gas Fires. Such fires have not yet been classified. These require special agents and techniques. For a discussion of extinguishing agents and techniques for these classes of fire see p 206

Definition of Terms

The meanings of fire control terms as used in this book are as follows:

Flash Point (flash p). This is the lowest temperature at which a liquid will give off enough flammable vapor at or near its surface such that in intimate mixture with air and a spark or flame it ignites. The flash point of liquids is usually determined by the Standard Method of Test for Flash Point with the Tag Closed Cup Tester (ASTM D56-52, available from the American Society for Testing Materials, 1916 Race St, Philadelphia, Pa). This method is also the standard of the American Standards Association (ASA Z11.24-1952, available from the American Standards Association, 70 East 45th St, New York, NY). The Interstate Commerce Commission uses the Tag Open Cup (TOC) Tester giving results 5–10°F higher (less flammable). Other methods frequently used are
Cleveland Open Cup (COC) and Pensky-Martens (PM). The closed cup flash point value is usually several degrees lower (more flammable) than the open cup, as the test in the former case is made on a saturated vapor-air mixture, whereas in the latter case the vapor has free access to air and thus is slightly less concentrated. For this reason, open cup values more nearly simulate actual conditions (see below).

**Fire Point** (fire pt). This is the lowest temperature at which a mixture of air and vapor continue to burn in an open container when ignited. It is usually above the flash point. Where the flash point is available, only it is given; if it is not, the fire point may be given. It is at least as significant as the flash point as an indication of the fire hazard of a material.

**Autoignition Temperature** (autoign temp). This is the temperature at which a material (solid, liquid, or gas) will self-ignite and sustain combustion in the absence of a spark or flame (ASTM Designation D286-36). This value is influenced by the size, shape and material of the heated surface, the rate of heating (in the case of a solid), and other factors.

**Vapor Density** (vap d). This value expresses the ratio of the density of a vapor to the density of air. The vapors of most flammable liquids are heavier than air, thus they can readily flow into low areas, excavations and similar localities. Hence, ventilating outlets in a plant should be located near ground level. For combustible gases and vapors which are lighter than air, ventilating outlets should be near the ceiling.

**Melting Point** (mp). This is the temperature at which the solid and liquid forms of a substance exist in equilibrium. This value indicates at what temperature flammable materials that are solid at room temperature may become flammable liquids.

**Boiling Point** (bp). This is the temperature at which a continuous flow of vapor bubbles occurs in a liquid being heated in an open container. The boiling point may be taken as in indication of the volatility of a material. Thus, in the case of a flammable liquid, boiling point can be a direct measure of the hazard involved in its use.

**Formula.** In the event of a lack of information regarding a material, its formula can give a clue to its fire hazard. For instance, all materials composed solely of carbon and hydrogen are combustible and in some degree flammable. If they are liquids with a low boiling point they can be assumed to be fire hazards.

**Underwriter’s Laboratories Classification** (ULC). This is a standard classification for grading the relative fire hazard of flammable liquids against the following standards:

<table>
<thead>
<tr>
<th>Class</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ether class</td>
<td>100</td>
</tr>
<tr>
<td>Gasoline class</td>
<td>90–100</td>
</tr>
<tr>
<td>Ethyl alcohol class</td>
<td>60–70</td>
</tr>
<tr>
<td>Kerosene class</td>
<td>30–40</td>
</tr>
<tr>
<td>Paraffin oil class</td>
<td>10–20</td>
</tr>
</tbody>
</table>

Where this value is known it is an excellent measure of the relative hazard of a flammable liquid. Unfortunately, it is available in only a few instances.

**Susceptibility to Spontaneous Heating.** Many materials combine with atmospheric oxygen at ordinary temperatures and liberate heat. If the heat is evolved faster than it is dissipated due to poor housekeeping, a fire can start, particularly in the presence of easily ignited waste, etc. [“Factory Mutual Modified Mackey Method,” Industrial and Engineering Chemistry (March 1927)].

**Explosive Range or Flammability Limits.** These values expressed in percent by volume of fuel vapor in air are the ranges of concentration over which a particular vapor or gas mixture with air will burn when ignited. If a mixture within its explosive range of concentrations is ignited, flame propagation will occur. This range will be indicated by LEL for lower explosive limit or UEL for upper explosive limit. The values given, unless other wise indicated, are for normal conditions of temperature and pressure.

**FIRE PROTECTION**

The two main aspects of fire protection are **prevention** and **loss limitation**.

**Prevention.** Fire prevention is an inseparable requirement of fire safety. Since, in order for a fire to start, all three necessary constituents—fuel, oxygen, heat—must be represented, effective fire prevention simply boils down to manipulation of these constituents to the extent that a fire cannot start. For instance, where a flammable material such as acetone is used out in an open
work shop, two of the three needed constituents are immediately present, ie the fuel and a supply of oxygen. Now the only thing lacking to start a fire is heat. Thus by referring to acetone in Section 12 it is found that the flash point is 14°F, which means that at any temperature above 14°F, acetone can evolve enough vapor to form a flammable mixture with air which will catch fire if exposed to a spark, flame or other source of ignition. Thus, strictly from the standpoint of fire prevention in an installation using acetone, the following avenues are open:

1. The working (ambient) temperature must be kept below 14°F, or
2. The supply of atmospheric oxygen must be cut off, or
3. Sources of ignition, such as flames, glowing cigars and cigarettes, sparks, etc must be eliminated from the area, or
4. The area must be ventilated so that even though the acetone gives off enough vapor to form a flammable mixture with air, the vapor will be drawn out of the area by means of fume exhaust equipment as rapidly as it is evolved, thus preventing the build-up of dangerous concentrations of vapors.

Naturally, since conditions (1) and (2) above are relatively difficult to attain on an industrial scale, conditions (3) and (4) are the ones most likely to be used.

Furthermore, although total removal of any one of the necessary conditions for a fire will absolutely prevent its occurrence, such stringent restrictions on industrial operations are seldom economically feasible. Industrial materials are, however, studied with a view to ascertaining just how much leeway there is, so that a compromise between absolute fire prevention and economy of operation may be reached. It is for this reason that, while we know how to prevent fires, they still do start, and why loss limitation is such an important part of industrial fire protection.

Below is some discussion of the three essentials of fire.

**Oxygen.** Although under certain unusual circumstances it is possible to produce combustion-like chemical reactions with materials such as chlorine or sulfur, it is safe to say that nearly all combustion requires the presence of oxygen. Also the higher the concentration of oxygen in an atmosphere, the more rapidly will burning proceed. Industrially it is difficult to manipulate the oxygen concentration in a working area, particularly since a concentration of oxygen far enough below normal to keep fires from starting would also be too low to support human life.

When industry has found it necessary to work with materials so sensitive to oxygen that they would catch fire at ordinary temperatures merely upon being exposed to air, it has found it possible to isolate such materials from air, either in a vacuum chamber or in a chamber filled with an inert atmosphere, such as argon, helium, or nitrogen. In Section 12 the materials which require such isolation are so noted.

**Heat.** As a necessary component of fire, this is often manipulated to render an industrial set-up safe from fire. The most difficult aspect of controlling the heat component of a fire is the easily overlooked fact that to start a fire it is often necessary to heat to a sufficient degree only a very small quantity of fuel and oxygen mixture. Then, since fires are by definition exothermic, the very small fire started by a tiny heat source supplies to its surroundings more heat than it absorbs, thus enabling it to ignite more fuel and oxygen mixture, and so on, until very quickly there is more heat available than is needed to propagate a large fire. The heat may be provided by various sources of ignition, such as high environmental (ambient) temperatures, hot surfaces, mechanical friction, sparks, or open flame.

**Fuel.** The third aspect of fire prevention will be discussed in detail in the following section.

**Sources of Ignition**

The following are the chief sources of ignition and suggestions for reducing the hazard due to them:

1. **Open Flames.** At or near a flammable-liquid installation it is necessary to check for such sources as burners, matches, lamps, welding torches, lighting torches, lanterns, small furnaces, and the possibility of broken gas or oil lines becoming flaming torches. Ample isolation may often be obtained by means of partitions. In this respect the partition should be substantial enough to contain the fire while the sprinklers or other fire-fighting apparatus put it out. Fire-
resistant construction (brick or concrete walls) is generally recommended.

It is important to confine the flammable liquid while it is in use. Safety cans should be used for transporting small quantities of flammable liquids about a working area, as well as for storage at a bench. Wherever possible, closed systems should be used to prevent the spread of fumes, etc. In the event of a fire, it is imperative to prevent spreading of the fire. Hence all tanks should have trapped overflow drains leading to a safe place. Dikes must be used to contain the overflow of burning liquid; otherwise fires could easily spread over large areas, trapping personnel and causing great damage. The principle behind this form of protection is to contain the fire at all costs. Installations of flammable liquids in upper stories should be made only in such a fashion that burning liquid will be prevented from flowing down stairwells, pipe openings, cracks in walls, etc, by means of waterproof floors, dikes, overflow pipes, etc

(2) Electrical Sources (electric power supply and generating equipment, heating equipment, and lighting equipment). The following precautions for good maintenance are suggested. A complete listing cannot be given here. The provisions of the National Electrical Code are the recognized standard and these should be carefully observed in installing electrical equipment in hazardous locations.

(a) Use special wiring and conduit
(b) Use explosion-proof motors, particularly if located at ground level or in pits or low places
(c) Use only specially engineered heating units, keeping in mind the autoignition temperature of the material in use (hot water or steam heating units are much to be preferred)
(d) Controls for motors, thermal cut-outs, switches, relays, transformer contacts, etc which are liable to spark or heat up should not be installed in flammable liquid storage areas. Use only explosion-proof, push-button control switches within such an area
(e) In dangerous atmospheres and for storage, only vapor-tight globes with electric lamps may be used. In well-ventilated areas, ordinary lamps will do. Fixed lamp installations are to be preferred to extension cords. Also, approved safety flashlights are preferred to portable lamps
(f) Do not install fuses or circuit breakers in hazardous locations except in explosion-proof cases

(g) Motor frames, control boxes, conduits, etc should all be grounded in accordance with the general requirements for installation of electric power as outlined in the National Electrical Code.

(3) Overheating (excessive temperatures at points requiring heat). Such processes should be kept out of combustible buildings and closely supervised. The use of automatic temperature controls and high temperature limit switches is recommended, although supervision is still important

(4) Hot Surfaces. The incomplete immersion of hot metal in quenching baths, the contact of flammable vapors and hot combustion chambers, hot dryers, ovens, boilers, ducts and steam lines all are frequent causes of flammable vapor fires. Care should be taken that material whose auto-ignition point is lower than the temperature sometimes reached by operating equipment be kept at a safe distance from such equipment. This equipment should be carefully supervised and maintained to prevent accidental overheating, etc

(5) Spontaneous Ignition. Many fires are caused by spontaneous heating of materials, accelerated by external heat from processes such as dryers, ovens, ducts, impregnating or steam lines adjacent to piles of waste materials. Sometimes the accumulated heat in a closed, unventilated warehouse will be sufficient to accelerate oxidation to the point of an actual fire. Wherever flammable liquids are handled, particularly those which are known to be liable to spontaneous heating, it is important to pay particular attention to housekeeping and ventilation. Fires are almost sure to follow neglect of these matters. All equipment and buildings should be kept free of deposits and accumulations of wiping rags, waste materials, oil mops, etc

(6) Sparks, etc. Sparks from mechanical tools and equipment, hot ashes from smoking, unprotected extension lights, boilers and furnaces, backfire from gasoline engines, are all potential causes of fire. Smoking should be prohibited in areas where flammable liquids are stored or are used in the open. All equipment in such areas should be maintained in first class condition. Wherever possible, spark-proof or non-sparking tools and materials should be used
(7) Static Electricity. This is due to electrical impulses generated on the surface of a material by friction, such as calendering, printing, and the like. Many fires are caused in the rubber and paper industries by this means. Most of these occur during the months when humidity is relatively low, and artificial heat is used. Maintaining a relative humidity of from 40–50 percent in rooms where flammable liquids are used will greatly reduce the chance of static sparks. Electrical grounding, static discharge devices, etc. should be mandatory, and all flammable liquid tanks, piping and equipment should be so interconnected and grounded that the chances for static sparks are minimized. In all this type of equipment, belts should be eliminated and direct or chain drives used wherever possible. If belt drives must be used, the belt speed should be kept below 150 ft/min, or a special belt dressing should be used which will reduce the possibility of the formation of a static spark.

(8) Friction. Many fires are caused by mechanical friction, ie from fan impellers rubbing on casings, poorly lubricated fan bearings, grinding processes and machining, etc. Fans and other equipment should be frequently inspected and maintained in the best possible condition. Other processes known to generate a good deal of heat due to friction should be well separated from locations where flammable liquids are stored or used.

It is vitally necessary that a complete program for the handling, transfer and use of flammable liquids be set up and maintained. This program should start when the process is initially under construction. Where flammable liquids are called for in the original write-up of a process, the first question is to determine whether the flammable material can be replaced by a non-flammable one. If the question of cost arises it should be remembered that to the possibly low cost of the flammable liquid should be added the cost of special protection needed to use it safely as well as its effect upon the insurance rate. It may well be that in the final analysis, the cost of a flammable material is not as favorable as it seemed at first. However, there are many flammable materials in constant use for which there are no substitutes, but even they can be safely handled if proper precautions are taken.

(9) Fuel. Combustion takes place most readily between oxygen and a fuel in its vapor or other finely divided state. Solids are most easily ignited when reduced to powders or vaporized by the application of heat, but except in a few cases the temperatures required for the vaporization of solids are well above normal ambient temperatures.

Liquids present a different case. Some liquids will give off dangerous quantities of flammable vapors well below normal room temperature (the vapor pressure of a liquid is a measure of this effect); others do so at points only slightly above room temperature, and still others at much higher temperatures. It is apparent that the temperature at which a liquid evolves vapors which can form flammable mixtures with air is a measure of its hazard potential. This is indicated by the flash point.

So indicative of fire hazard is the flash point of a liquid that the Interstate Commerce Commission rates any liquid whose flash point is 80°F or below as a high fire hazard, the theory being that 80°F represents the upper limit of normal or “room” temperatures; any liquid which will flash at or below this point is dangerous. A flash point of from 80–350°F indicates a moderate fire hazard; above 350°F the fire hazard is considered slight. The National Fire Protection Association rates as a high hazard a liquid whose flash point is less than 20°F; moderate from 20–70°F, and slight from 70–200°F. Only liquids having a flash point less than 200°F are generally called flammable by the National Fire Protection Association.

In this book we use the ICC classification for liquids of flash point at 80°F or less as being dangerous fire hazards; we use the range from about 80–225°F to indicate moderate fire hazard; flash points in excess of about 225°F are rated as slightly hazardous. It should be understood, however, that practically all organic materials will burn if exposed to sufficiently high temperatures. The ratings given above are merely as indication of the risk involved in handling or storing them.

It is important to isolate a potential fire hazard. Thus it is necessary to use closed and vented tanks to hold flammable liquids. In this way the possibilities of igniting the tank of liquid are greatly reduced, as is the chance of materials at a distance from such a tank becoming involved in
fines. It is also important that flammable materials be housed in fire-resistant structures because burning liquids can generate great heat and often set fire to the buildings in which they are burning.

A vital point in flammable liquids safety is the prevention of the accumulation of explosive concentrations of vapors in closed off areas. Wherever either moderately or highly flammable liquids are used or stored, ventilation is a very important consideration.

The amount needed, whether natural or mechanical (fans and blowers) depends upon the materials and the conditions involved. No dependence should be placed upon the odor of the material as a warning, because some flammable vapors are heavy and tend to settle and because smell is deceptive. The safe procedure is continual testing with an explosion or flammable vapor indicator.

Besides flammable liquid fires, the results of which can be somewhat mitigated by effective loss limitation techniques, there are two more types of disaster, protection from which is nearly entirely dependent upon prevention.

10) Dust Explosions. Practically any combustible, when in the form of dust and mixed with air in the proper proportion, will burn so rapidly as to cause a severe explosion if ignited by heat, a spark or flame. Ignorance of this fact has led to many serious disasters. Grain, flour, coal dust, and metal powders all constitute hazards in this regard. Explosions have been known to occur in plants handling fertilizers, wood dust, powdered milk, soap powder, paper dust, cocoa, spices, cork, sulfur, rubber dust, and many other products. For the prevention of dust explosions good housekeeping is of the utmost importance. All equipment must be dust-tight and kept so. Explosion vents should lead outdoors to a safe location, and the vent ducts themselves should be strong enough to withstand the force of the explosion. Vacuum cleaning is superior to sweeping. The use of compressed air to blow dust off equipment and thus create dust clouds should be forbidden.

Ledges, exposed piping, beams, etc in the ceilings should be kept free from accumulation of dust. Where a dusty operation is to be installed in a location where there is piping and projections overhead, it is often erroneously con-

sidered satisfactory to install a smooth ceiling below the piping and other projections. This does not eliminate the hazard and may intensify it; unless the ceiling is extremely well designed and installed, dust will penetrate it and settle not only on the piping but on the upper side of the ceiling itself. Then a shock may be sufficient to fill the entire false space with a combustible dust cloud which a spark may set off. If piping cannot be relocated or eliminated, it would be better to leave it exposed and provide for a regular cleaning program.

It was for reasons such as the above that a starch dust plant recently constructed in the southwest, where weather conditions are moderate the year round, has been built entirely of open construction so that there is no confinement of the force of any explosion and the constant flow of air through the plant provides little opportunity for dust layers to build up.

As in the case of flammable liquid fires and explosions, the control of dust explosions is based upon prevention of ignition and secondarily limitation of damage in the event ignition does occur.

To prevent ignition, open flames, smoking, and cutting or welding are prohibited until the area is made dust free. Electrical wiring should be of the type suitable for a dusty atmosphere and static electricity, too, must be eliminated. Highly dangerous materials of this sort are handled most satisfactorily in enclosed systems in which suitable inert gases are introduced into the system to replace the air normally present. This precaution is particularly applicable to the field of powder metallurgy. The kind of inert gas used must be chosen on the basis of its suitability for the operation in question.

11) Salt-bath Explosions. The third type of disaster in which after-the-fact protection is much less important than prevention is the molten salt-bath explosion. There have been serious disasters involving such baths, because personnel involved on both the management and the operating level failed to appreciate the potential hazards of the situation. Due to mechanical failure or human failure, or a combination of both, molten salt baths have been allowed to explode. The hazards of molten salt baths may be summarized as follows:

(1) Violent generation of steam due to water
introduced as "carry-over" on a piece of work from a preliminary cleansing or quenching bath, condensation on overhead service piping, leaky roofs and operation of automatic sprinklers, also contact with liquid foods placed on ledges near the baths for "warming-up" by workmen.

(2) Sudden and explosive expansion of air occluded in blow-holes of castings and that trapped in tubes, closed piping, or hollow metal work when immersed in molten baths without pre-warming.

(3) Violent and uncontrollable chemical reactions between nitrate baths and carbonaceous materials such as oils, soot, graphite, and cyanide carry-over from adjacent carburizing baths.

(4) Vigorous and explosive reaction between overheated nitrate baths and aluminum alloys.

(5) Explosive reaction between normally heated nitrate baths and carelessly introduced magnesium alloys.

(6) Thermit-like reaction between aluminum alloy articles lost in bath and the iron oxide sludge blanketing and insulating the bottom of bath container.

(7) Structural failure of bath container while in operation under conditions tending to lower the normal durability; reaction between metal of bath container and nitrate due to localized overheating.

(8) Failure of temperature controls, with consequent overheating of nitrate bath.

(9) Storage and handling of bulk supply of sodium nitrate, and careless disposal and storage of waste nitrate without regard to active properties of the salt.

(10) Accidental or uninstructed setting of temperature control above safe operating limits.

The precautions for safe operation of molten salt baths are summarized as follows:

(a) Guard against the introduction of any extraneous matter.

(b) Protect completely from overheating by automatic control and temperature readings taken at regular intervals.

(c) Isolate the operation as far as practicable.

(d) Instruct all personnel thoroughly and completely in regular and emergency procedures.

From the foregoing it can be clearly seen that the handling of flammable liquids, flammable dust or molten salt baths are the three chief operations in which prevention is the most important phase of fire protection. In each of these cases whatever action is taken after the act and whatever physical protection is provided can only furnish some degree of mitigation of loss and it is often insufficient to prevent a large scale disaster.

**Loss Limitation**

The other aspect of a realistic fire protection program is limitation of loss due to a fire which includes a provision for the *prompt discovery* and equally *prompt extinguishment* of the fire. It is certain that everyone has at one time or another wondered why a particularly destructive fire was allowed to happen, when supposedly a great deal of effort is constantly being devoted to the prevention of such fires.

It may be that too often the prevention aspect of fire protection has been the total or nearly total effort at protection, with the result that when even a small fire starts it has a good chance to become a calamity. However, even the utmost vigilance would have been of no avail in guarding against some of the fires which are on record, and once the fire has started its cause is immaterial. The cause of the *loss* is much more important, and the facts which determine *that* are the physical conditions and those measures which have or have not been taken to limit the extension of the fire.

One of the means of preventing the extension of fire is to segregate hazardous processes and storage into separate buildings. But even where hazardous processes are not involved, the concentration of too much value in one fire area must be guarded against. This is best accomplished by the erection of separate buildings, adequately spaced, which in turn presents problems of maintenance and operation. Suppose a major plant has an operation involving flammable liquids. Such processes as spray painting or dipping use tremendous quantities of flammable solvent; extraction processes and the manufacture of products of which flammable solvents are major constituents are typical examples of processes which require subdivision. The flammable liquid operations and all its appurtenances must be physically separated from the rest of the plant for maximum safety. Care should be taken that the separate buildings are no larger than production efficiency demands; in
other words, only as much should be put under one roof as is necessary to be in one building. Where space or production requirements preclude separate buildings, the area in which flammable liquids are handled must be physically separated from the rest of the plant by approved fire walls or, where these are not practicable, by water-curtain type sprinklers.

Subdivision of one large risk into smaller fire areas may also be accomplished by means of fire walls which stop the spread of fire from one area to another. To accomplish this, the wall must be carried through the roof and either go through the side walls for a distance of at least 36 inches or turn back on both ends for a distance of several feet to provide a barrier around which the fire cannot travel.

Many otherwise sound fire walls have failed because holes were made in the walls to permit the passage of pipes, conduit, etc., and then never properly closed. Every hole in a fire wall must be sealed at the time such work is being done. The weakest point in a fire wall is the fire door provided to permit access from one section to another. At any given time a high percentage of such doors are found upon inspection to be useless as fire barriers. They must be test operated regularly. The chief deficiencies are missing fusible links, damage to the doors by materials handling equipment, which damage would prevent their operation, or blocking by material left in the doorway so that the door cannot close.

Proper maintenance includes regular inspection, physical guarding to prevent damage, the painting of "keep clear" lines on the floor, and a constant program of education. Wherever practicable, such doors should be closed at night to insure that they will be closed in the event of fire. Where the use or occupancy of the building has changed and fire doors are not needed any longer, the openings should be bricked up to the same thickness as the original walls. Even a pair of fire doors, one on each side of the opening, is less resistant to the passage of fire than the fire wall in which they are installed. In normal operation, because of the aisle space leading to the opening, the heat on the door should be less than elsewhere in the building. If the doors are closed and contents are piled against the doors, fire may be transmitted from one side of the wall to the other. In all cases, the doors provided should be of the type approved for the opening in the wall. In many cases, doors approved for use only on vertical enclosures such as stairways are installed in fire walls and will not serve their intended purpose in the event of fire. If openings are necessary in fire walls for the passage of conveyors and no type of door installation is practical, then the openings should be specially protected by hooded automatic sprinkler heads directly over the opening on each side of the wall. A fire wall should be thought of as a dam which any small leak can cause to fail.

The spread of fire from floor to floor in a building is prevented by the proper enclosure of vertical openings such as stairways, elevators, shafts, and process openings through the floor. The question of stairways deserves particular comment because in so many cases, self-closing stairway doors are found to be wedged open to permit easy passage from floor to floor. Such examples of poor management entirely negate the cost of closing the stairway off by providing the doors in the first place.

One of the primary reasons for enclosing stairways is to permit the passage without injury of personnel from upper floors to the street level past the floor which is on fire. If the stairway doors are wedged open this may be impossible. The stairways can immediately become choked with hot air, smoke and gases from the stairwell.

Provision of fusible link arrangements to close such doors is not very satisfactory because fumes and smoke will pass through without operating the link. In at least one laboratory the problem of stairway doors being wedged open versus the desirability of having them closed in a hurry has been solved by providing for electric latches. These latches hold all the doors open, but connected to the fire alarm system is a relay which causes all the electric locks to release when a fire alarm is sounded, thus closing the doors.

A further loss-limiting device which is useful where flammable dusts and vapors are used is the explosion vent. It is important to install explosion vents in areas where flammable liquids or dusts are used because of the possibility of great damage due to explosive ignition of such mixtures and air. Therefore, on a practical basis,
properly designed explosion vents are a suitable safeguard, as they reduce the chances of destruction indoors by allowing the force of the explosion to be transmitted outdoors.

In order to fully relieve the pressures produced by explosions in vapor and air mixtures, a vent area as large as 1 square foot for every 10 cubic feet of room volume would be necessary. However, it is unlikely that more than a fraction of the total volume of a room will at any time be within the explosive range. Therefore for a small room with a floor area of about 200 square feet, the venting area should be at least 1 square foot for each 30 cubic feet of room volume. For larger areas, this proportion may not be obtainable, but in no case should the vent area be less than 1 square foot for each 50 cubic feet of volume.

Approved explosion venting windows are available. Also, sky-lights, roof hatches, or light windows hinged at the top and carefully installed to swing outward under even slight pressure can be useful. Under some conditions, doors equipped with releasing latches may be utilized as vents.

Furthermore, where the conservation of heat in a plant is important, and the walls of the building are otherwise of strong construction, a section of exterior wall may be built of light wood, or hollow tile or some other material which is relatively weak compared to the rest of the building so that in case of an explosion, these sections will give first.

It is important that snow or ice be kept from collecting on explosion vents so that they can operate freely in case of an explosion.

In discussing the extinguishing of fires, Sax makes the following remarks about metal fires:

"Fires caused by burning metals are very difficult to extinguish and cannot be handled in the ordinary manner. For instance, to spray water upon burning metal might cause an explosion which would spatter flaming particles of metal to great distances. Usually the best way to attack burning metal is with specially formulated dry type fire-extinguishing agents. For instance, when one is planning to use a metal in a form in which it might readily become ignited, it is wise to discuss the situation with the manufacturer or supplier of this metal and obtain from him explicit instructions for the storage, handling, and fire extinguishment of the metal. This is particularly applicable to the use of sodium, potassium, lithium, zirconium, uranium, thorium, and magnesium. For instance, it has been found that ordinary sand, even when dry, is a very poor material for extinguishing metal fires; it may react with the hot metal and add more heat to an already intense fire. Often salt (sodium chloride), sodium bicarbonate, graphite, magnesium carbonate, magnesium oxide, or mixtures of all of these materials have been found effective; in every case the supplier of this material will know how it must be handled. Water should never be applied to burning metals."

**Industrial Hygiene (Industrial Health, Occupational Diseases, Occupational Hazards).** Industrial hygiene is a branch of medical science dealing with principles and rules for preservation and improvement of health of those working in industry.

Due to the fact that most explosives are toxic, particular care should be taken in handling them and the rules prescribed in books on Industrial Hygiene should be observed. The War departments of each country also have special rules applying to war plants, arsenals etc.

See also Health Hazards of Explosives and Propellants in this Vol.

**Refs:**
1. W. G. Thompson, Occupational Diseases, Appleton, NY (1914)
2. Great Britain Ministry of Munitions, Industrial Health and Efficiency, HMPO (London) (1918)
5. G. Lutz, Gewehrhygiene, Wissenschaftliche Verlags, Stuttgart (1933)
7. L. B. Chenoweth & W. Machle, Industrial Hygiene; A Handbook of Hygiene and Toxicology for Engineers and Plant Managers, Crofts & Co (1938)
8. J. B. Ficklen, Manual of Industrial Health Hazards, Service to Industry, West Hartford, Conn (1940)
9. L. I. Dublin & R. J. Vane, Occupational Hazards, US Govt Printing Office,

**Industrial Toxicology.** See under Toxicity & Toxicology

**Inert Gases’ Explosive Compounds.** Until quite recently, the noble gases (Xe, Kr, A, Ne, Ra & He) were considered chemically inert and incapable of reaction with other elements. While this concept remains generally unchanged, at least a few noble gas compounds can be prepared.

Xe fluorides and oxyfluorides constitute the principle noble gas compounds made to date. Scores of explns have been experienced during prep & use of Xe fluorides, of which perhaps the following were the most serious:

1) An expt was being conducted in which a soln contg 0.39g of XeF₄ dissolved in 1.5ml of w was being evapd under vac at room temp to yield a white solid thought to be Xe(OH)₄ or XeO₂₂H₂O. While the silica container (under vac) was being heated gently with warm (30-40°C) air from a blower, a deton took place which shattered the container & inflicted serious injuries (Ref 1)

2) In Oct 1963, an expl occurred during heating of a 3" OD, 1/8" wall, 6½" long cylndrical Ni reactor contg Xe and F gases. The incident was attributed to failure of a thermocouple-activated regulator which permitted pressure buildup beyond the rating of the vessel (Ref 5)

3) An expln during Nov 1963 was attributed to reactions of moisture which entered a liq N₂ cold trap following failure of a mechanical pump over a weekend. It was presumed that the moisture reacted with XeF₆ in the trap to form expl XeO₃ (Ref 5)

The case with which highly expl XeO₃ may be formed at room temp by reaction of Xe fluorides with moisture has been strongly emphasized (Refs 2, 3 & 4). Xenon trioxide, XeO₃, is a wh solid comparable in deton sensitivity to nitrogen trioxide and in expl force to TNT

Inert Simulants for High Explosives. In certain physical tests on weapons & explosives, eg the effects of humidity and temp cycling on the dimensional stability of the weapon, it is desirable to simulate the explosive filler by an inert filler. Many such inert simulants have been proposed and tested (Refs 1, 2 & 3). Simulants for Comp B & TNT have been patented (Ref 4). The simulant for Comp B consists of a mixt of 30% of 1,2 hydroxystellar acid, 5% wood rosin & 65% dead-burned gypsum. The simulant for TNT is a 40/60 mixt of hydroxystellar acid & dead-burned gypsum

A series of mock explosives to simulate PBX 9404, LX-04 & LX-10 has been developed in AEC laboratories

The composition, density, thermal properties & mechanical properties (elastic, viscoelastic & failure behavior) of these mock explosives are given in Ref 7.

Inertial, Terrestrial and Celestial Guidance Systems for Missiles. An inertial guidance system is one which is independent of information, other than gravitational effects, obtd from outside the missile

Missile guidance, wherein the predetermined path set into the control system of a missile can be followed by a device in the missile which reacts to some property of the earth such as magnetic or gravitational effects, is known as terrestrial reference guidance

A system wherein a missile, suitably instrumented and containing all necessary guidance equipment, may follow a predetermined course in space with ref primarily to the relative positions of the missile and certain preselected celestial bodies is called an inertial-celestial guidance system

Infallible Powder. One of the old duPont explosives
Ref: Marshall, I, p 330

Infallible Propellant. A double-base proplnt consisting of NC (13.25%N) 59.25, NG 40.00 & DPhA 0.75%, coated with graphite. Grains 0.055 inch in diam & 0.007 inch thick when subjected to 65.5° Surveillance Test and 120° & 134.5° Heat tests were of satisfactory stability
Ref: W.H. Rinkenbach, “The Stability of Double-Base Powders,” PATR 1359 (Oct 1943), p 2 & Table IV

Infantry Rocket Weapons. The title of a paper by Major Brill in Ordnance (Ref 1), where a brief description is given of the following weapons:

1) Rocket Launcher, 2.36 inch, known as Bazooka, developed in 1942. It was 54” long and weighed 12 lbs; could penetrate up to 4” armor at close range. Used successfully during WWII (Ref 1, p 629) (See also Bazooka in Vol 2, p B26)

2) Rocket Launcher, 3.5 inch, is the modern successor of original 2.36 inch launcher. Used successfully during Korean War

Note: Both launchers used non-guided missiles until 1959, when US Govt adopted French, antitank, wire-guided missiles SS10 & SS11. The SS10 is no longer in the US Army inventory and SS11 has been adopted to helicopter armament as the M22, but for ground use it was replaced by the Entrace Missile (Ref 1, p 630)

3) Entrace/Engin Téléguidé Anti-Char (Tele-guided Antitank Device). A French boxlike launcher using a solid-propelled, wire-guided missile. The target is optically tracked by the infantryman. The missile is designed for use against tanks, armored combat vehicles and some defensive installations. It is armed with a HE shaped charge warhead. Total weight of the
launcher and missile is 37.5 lbs (Ref 1)

4) Redeye. A 3-inch weapon developed in US during the Vietnamese war & designed to provide combat troops in the forward battle area the capability of destroying low-flying aircraft. The length of its tubular-type launcher is 4 ft and the total wt of missile and launcher is ca 28 lbs. In its nose the missile carries an infrared sensing device which enables the guidance system of the missile to "home" on the heat of the enemy's aircraft engine. A small charge ejects the missile from its launcher and, at a distance far enough to protect the soldier from the rocket blast, a fuse ignites the major rocket charge. Miniature computer circuitry within the missile directs a set of steering fins which enable the missile to change direction as necessary and chase the target at supersonic speeds until it intercepts it (Refs 1 & 2)

5) Light-Antitank Weapon (LAW), M72, also known as M72 Rocket Grenade, is the smallest antitank weapon because its total wt is only 4.5 lbs and its length is 25 inches. It can be carried and fired by one man using its disposable packing container as a launcher. A telescopic aluminum inner section is extended prior to firing; the outer section is made of fiberglass-plastic composition. A solid-fuel rocket motor furnishes propulsion and burns out before the rocket leaves the launcher. When the missile emerges, several narrow magnesium fins (folded against the motor case when packed in the tube) spring into position and stabilize the missile (Ref 1, p 630)

6) TOW (Tube-launched, Optically-tracked, Wire-guided) Weapon has been developed to provide greater firepower than possible with Entac. Although similar to other antitank missile systems, TOW uses somewhat different guidance principles. The optically tracked missile is automatically guided in flights by commands transmitted by means of a 2-wire link between the gunner and the missile (Ref 1, p 630)

7) MAW (Medium Assault Weapon). Two missiles have been under development during the Vietnamese War—one by industry and the other by the Research and Development Directorate of the Army Missile Command (Ref 1, p 630)


Infernai Machines (Höllenmaschinen in German; Engins Criminels in French; and Adskiyé Mzhiny in Russian). Various devices used by anarchists, revolutionaires and criminals against kings, presidents, politicians and rich persons; also for sabotage. These devices consist of bombs, grenades and mines, and can be exploded either by instantaneous or time fuzes


Inflammability or Flammability (latter is preferred in modern usage). The ease with which a material (gas, liquid or solid) will ignite, either spontaneously (pyrophoric), from exposure to a high-temperature environment (autoignition), or to a spark or open flame. It also involves the rate of spreading of a flame once it has started. The more readily ignition occurs, the more flammable the material; less easily ignited materials are said to be combustible, but the line of demarcation is often indefinite, and depends on the state of subdivision of the material as well as on its chemical nature

Closely related to inflammability is the general subject of inflammable or flammable (preferred) materials. These are defined as: any solid, liquid, vapor or gas that will ignite easily and burn rapidly. Flammable solids are of several types: (1) dusts or fine powders (metals or organic substances such as cellulose, flour, etc), (2) those that ignite spontaneously at low temperatures (white phosphorus), (3) those in which internal heat is built up by microbial or other degradation activity (fish meal, wet cellulosic materials); (4) films, fibers, and fabrics of low-ignition point materials
The National Fire Protection Association divides organic liquids into three classes: (1) those having a flash point (Tag closed cup) below 100°F (flammable, dangerous fire hazard); (2) those having a flash point at or above 100°F and below 140°F (flammable, moderate fire hazard); (3) those having a flash point at or above 140°F (combustible, slight fire hazard). Thus the critical flash point temperature for a flammable liquid is 140°F (Tag Closed Cup). However, shipping regulatory authorities (ICC, CG, IATA), as well as the Manufacturing Chemists Association, use 80°F (Tag Open Cup) as the critical flash point temperature.

Flammable gases are ignited very easily; the flame and heat propagation rate is so great as to resemble an explosion, especially if the gas is confined. The most common flammable gases are hydrogen, carbon monoxide, acetylene and other hydrocarbon gases. Oxygen, though essential for the occurrence of combustion, is not itself either flammable or combustible; neither are the halogen gases, sulfur dioxide or nitrogen. Flammable gases are extremely dangerous fire hazards, and require precisely regulated storage conditions.

Note: The terms “flammable,” “nonflammable,” and “combustible” are difficult to delimit. Since any material that will burn at any temperature is combustible by definition, it follows that this word covers all such materials, irrespective of their ease of ignition. Thus the term “flammable” actually applies to a special group of combustible materials that ignite easily and burn rapidly. Some materials (usually gases) classified in shipping and safety regulations as nonflammable are actually noncombustible. The distinction between these terms should not be overlooked. For example, sodium chloride, carbon tetrachloride and carbon dioxide are noncombustible; sugar, cellulose and ammonia are nonflammable.


Inflammability (Relative) of Dust Clouds. See also Dust Explosions in Vol 5, p D1578-R and Ignition Temperature Tests in this Vol. Relative Inflammability of a dust is defined as the percentage by weight of an inert dust, such as Fuller’s earth, required in a mixture with the flammable dust to prevent ignition and flame propagation when the mixture is dispersed into a dust cloud in the presence of a standard source of ignition. The more inflammable the dust, the higher will be the percentage of inert dust needed to prevent ignition. This classification was adopted in the Bureau of Mines study of coal-mine dusts. The equipment used in these tests was the Godbert-Greenwall apparatus and an open-spark inflammability apparatus.

The former consists of an electrically heated vertical cylindrical tube (furnace), the top of which is connected to a small brass chamber containing the dust sample to be tested. A pneumatic system is used for blowing the dust downward thru the heated tube. In the usual ignition test, the lowest tube temp at which a flame appears at the lower mouth of the tube is taken as the ignition temp of the dust cloud. In the relative inflammability test the temp of the furnace is held constant at 700°C (1292°F) and only the amounts of Fuller’s earth is decreased until no ignitions are observed.

In the open-spark apparatus, a high voltage continuous spark having an average power of 20 to 24 watts is passed through a spark gap between tungsten electrodes in a pyrex tube, and the cloud of powder or dust being tested is blown thru the tube. The relative inflammability is determined as in the furnace test by finding the minimum amount of Fuller’s earth needed in a mixture with the inflammable dust being tested to prevent ignition of the mixture by the spark.

Ref: E. Hartmann & J. Nagy, USB of M, RI 3751 (1944)

Inflammability of Explosives Tests (Flammability Tests). See also Index of Inflammability in Vol 1, p XVII. These tests are designed to ascertain the behavior of explosives towards open flame. The following tests are described by Reilly (Ref 1):

Test No 1: Same test as Fusel Test described under Ignition Sensitiveness

Test No 2: A small sample of explosive is placed on an asbestos board and a small flame of a Bunsen burner is directed against it for 10 seconds. It is considered nonflammable if it does not ignite in this period. Control samples of known explosives should be included for comparison.
Test No 3: If the explosive withstands the Fuse Test, it is subjected to the Red Hot Iron Test described under Ignition Sensitiveness Tests.

Test No 4: Wood Fire Test: In some cases, larger quantities, i.e. a pound or two, are burned in a wood fire to ascertain the degree of danger the explosive may offer in case of fire. The observations should be made under precautions at a safe distance. (Compare with “Iron Box Test,” described under Ignition Sensitiveness.)

Test No 5: Ignition Time Test: A small quantity of the explosive is fastened to the end of a pendulum which swings through a flame in such a manner that the time required to ignite the sample can be measured. (For details, see under Ignition Time of, Test.)

Andrew & Kostin (Ref 2) describe an inflammability (or ignition) test for explosives in which a “donor” explosive is used to ignite an “acceptor” explosive placed below the donor in a 1cm diam vertical glass tube. As the density of the donor is decreased its igniting ability also decreases.


Inflammability or Flammability (preferred)

Limits. Gaseous fuels mixed with gaseous oxidants (usually air or oxygen) will ignite only within certain composition ranges. The max fuel-rich composition capable of ignition is known as the upper flammability limit and the min fuel-poor composition is known as the lower flammability limit. These vary widely with fuels and oxidants and are discussed under the individual gaseous fuels or vapors, eg see acetylene gas in Vol 1, p A59-L, or acetone vapor in Vol 1, p A34-L. General references on this subject include:


Influence Tests. See Detonation of Influence or Sympathetic Detonation Tests in Vol 1 p X with additional information given below:

These tests are designed to measure the distance (usually in air) over which detonation may be conveyed from one explosive (donor) to another (acceptor).

In blasting operations it is important that an explosive be capable of transmitting detonation across an inert gap which interrupts the continuity of the explosive charge. The inert gap can be created in practice by air gaps, dirt or wads of paper getting between cartridge (sticks) of an explosive charge.

It is important that transmission shall not be too poor, because this involves the danger that, if for any reason different cartridges are separated from one another by too great a distance, the entire charge might not be detonated. Some explosives transmit so poorly when frozen that detonation is incomplete even when there is no break in the continuity of the charge. Total or partial failure to detonate is a frequent cause of accidents, as the explosive is apt to go off in subsequent handling of the material being blasted. Accidents also occur in subsequent drilling of bore holes near unexploded charges.

According to Marshall (Ref 1), a simple method of determining the distance of transmission, is to set out a row of small cartridges at increasing distances apart and to determine how many of them explode when the end cartridge is detonated.

The distance at which the explosion is transmitted from donor to acceptor cartridge depends not only on the nature of the explosive itself (its sensitivity, brisance, power, velocity of detonation etc) but also on the following factors (provided both cartridges are from the same material and are of equal size):

1) Density of the explosive
2) Kind of medium through which the explosion is transmitted

Transmission of detonation is different for air gaps and dirt gaps. Transmission distances are much greater in a wet soil than in dry soil. In the open, distances (across air gaps) are shorter than in bore holes. In the open, the medium on which the cartridges are laid also
influence transmission distance: shortest gaps on porous media (loose soil) & longest gaps on dense media (steel)

3) Gaps increase as the cartridge diameter increases

**Test Methods**

The **gap test** or **halved cartridge test** is described in Vol 1, pp XIV & XV. Examples of the gap sensitivity (inches of air across which detonation is transmitted from donor to acceptor) are given below for US dynamosics (Ref 6). Also see Vol 5, pp D1591 & D1737 (Table 5)

36 inches—for 60% ammonium gelatin containing NG 26.0, NC 0.4, NH₄NO₃ 32.5, NaN₃ 29.5, wood flour 2.0, starch 2.7, ivory-nut meal 3.9, sulfur 2.0, lime 1.0. Vel of Deton 4900 m/sec; wt strength 67%

26 inches—for ammonium gelatin containing NG 26.0, NC 0.4, NH₄NO₃ 57.2, red dog 4.0, clear flour 3.0, ivory-nut meal 4.4, starch 3.0, sulfur 2.0, lime 1.0. Vel of Deton 4800 m/sec; wt strength 61%

20 inches—for a dynamite containing NG 9.0, NH₄NO₃ 42.0, NaN₃ 26.8, ivory-nut meal 13.7, fine buckwheat hulls 5.0, sulfur 3.0, lime 0.5. Vel of Deton 2700 m/sec

15 inches—for 20% ammonia dynamite containing NG 8.5, NH₄NO₃ 16, NaN₃ 55.0, sulfur 6.5, fine buckwheat hulls 6.0, ivory-nut meal 7.0, lime 1.0. Vel of Deton 2500 m/sec; wt strength 20.2%

10 inches—for dynamite containing NG 9.0, NH₄NO₃ 75.0, NaN₃ 3.2, straw 12.3, lime 0.5. Vel of Deton 2700 m/sec; wt strength 67%

8 inches—for a permissible dynamite containing NG 10.0, NH₄NO₃ 9.0, salt 2.5, oat straw 12.0, lime 0.5. Vel of Deton 2600 m/sec

4 inches—for a dynamite containing NG 10.5, NH₄NO₃ 73.0, NaN₃ 4.0, wood pulp 9.0, balsawood 3.0, lime 0.5. Vel of Deton 2200 m/sec; wt strength 63%

2½ inches—for Pic Ars Low Velocity Dynamite (Ref 8)

1 inch—for Hercules Medium Velocity Dynamite (Ref 9)

From the above it is clear that gap sensitivity depends strongly on the NG content of the dynamite

Cook (Ref 7) suggests that \( S_0 \), the air gap over which detonation will be transmitted solely by shock, is given by:

\[ S_0^2 = k \text{M} \]

where \( k \) is a different constant for each explosive, and will depend on the density and physical state of the explosive; \( M \) is the weight of explosive

This equation will apply, on the other hand, only if one uses a primer of fixed \( L/d \) (length/diam), and \( k \) will be a maximum for any explosive of fixed composition, density, and granularity for \( L/d \sim 1.0 \). Also this relationship may be readily upset if there is a chance for the explosive to hurl solid fragments, because flying particles can set off an explosive over far greater distances via hot-spot mechanisms than the blast wave itself. In addition, \( k \) will depend upon how one expresses sensitivity results. For example, the maximum distance for consistent detonations (100 per cent D, 0 per cent F) is usually roughly about one-half the minimum distance for consistent failures (0 per cent D, 100 per cent F). It is usually best to adopt the 50 per cent F point as the reference point for defining \( k \), because this point can be most accurately established. That is, the per cent detonation versus gap distance curve is in general an S-shaped or probability curve


**Modified Influence (Propagation) Test.** Gawthrop (Ref 1) and others used a modified gap test to determine the relative ability of a shielded donor charge to transfer detonation over an air gap to an acceptor charge. Clark (Ref 2) also used this method for the determination of the gap in the transmission of detonation from a small charge (0.5 to 2g) of HE to a cartridge of 40% straight dynamite

For these tests, Clark placed a small charge of HE in a No 8 detonator shell and pressed it under a reinforcing capsule at 3400 psi. The detonator was placed centrally in a cylindrical oakenshield, with its long axis parallel to and
coinciding with the long axis of the shield and with its base flush with the end of the shield. The shield containing the detonator and the cartridge of acceptor explosive (40% dynamite), with the cut end facing the detonator, were then wrapped in three layers of heavy paper

The maximum gap through which detonation would be transferred with certainty was determined in four trials

Modified Gap Test results are given in the following tabulation (which shows that DDNP is an efficient donor):

<table>
<thead>
<tr>
<th>Detonator Base</th>
<th>Charge (g)</th>
<th>Wt of Base Charge (g)</th>
<th>Max Air Gap over which Detonation of 40% Straight Dynamite Occurs (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>DDNP</td>
<td>0.50</td>
<td></td>
<td>350</td>
</tr>
<tr>
<td>MF</td>
<td>0.50</td>
<td></td>
<td>150</td>
</tr>
<tr>
<td>MF</td>
<td>1.00</td>
<td></td>
<td>300</td>
</tr>
<tr>
<td>Tetryl primed with LA</td>
<td>0.66</td>
<td></td>
<td>400</td>
</tr>
<tr>
<td>PA primed with MF</td>
<td>1.25</td>
<td></td>
<td>300</td>
</tr>
<tr>
<td>MF-KClO₃ (80:20)</td>
<td>2.12</td>
<td></td>
<td>90</td>
</tr>
<tr>
<td>MF-KClO₃ (80:20)</td>
<td>1.05</td>
<td></td>
<td>100</td>
</tr>
<tr>
<td>Lead styphnate</td>
<td>1.74</td>
<td></td>
<td>50</td>
</tr>
<tr>
<td>Nitromannite</td>
<td>1.16</td>
<td></td>
<td>90</td>
</tr>
<tr>
<td>Mercury fulminate</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Results are strongly influenced by the tube in which tests are conducted. The following tabulation (Ref 3) shows that tests in iron tubes result in much longer gaps than tests in paper tubes:

<table>
<thead>
<tr>
<th>Explosive</th>
<th>Charge (g)</th>
<th>Weight of Paper Tube</th>
<th>Gap, in meters through Pipe</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pentryl*</td>
<td>0.50</td>
<td>4.00</td>
<td>16.5</td>
</tr>
<tr>
<td>DDNP</td>
<td>0.50</td>
<td>3.50</td>
<td>--</td>
</tr>
<tr>
<td>MF</td>
<td>0.50</td>
<td>1.50</td>
<td>--</td>
</tr>
<tr>
<td>MF</td>
<td>1.00</td>
<td>3.00</td>
<td>--</td>
</tr>
<tr>
<td>Tetryl primed with MF</td>
<td>1.34</td>
<td>2.50</td>
<td>12.5</td>
</tr>
<tr>
<td>Tetryl primed with LA</td>
<td>0.66</td>
<td>4.00</td>
<td>--</td>
</tr>
<tr>
<td>PA primed with MF</td>
<td>1.25</td>
<td>3.00</td>
<td>12.5</td>
</tr>
<tr>
<td>MF-KClO₃ (80:20)</td>
<td>2.12</td>
<td>0.90</td>
<td>3.5</td>
</tr>
<tr>
<td>MF-KClO₃ (80:20)</td>
<td>1.05</td>
<td>1.00</td>
<td>--</td>
</tr>
<tr>
<td>Lead Styphnate</td>
<td>1.74</td>
<td>0.50</td>
<td>3.5</td>
</tr>
<tr>
<td>Nitromannite</td>
<td></td>
<td>1.16</td>
<td>0.90</td>
</tr>
<tr>
<td>Mercury fulminate</td>
<td></td>
<td>1.06</td>
<td>12.5</td>
</tr>
</tbody>
</table>

*Primed with 0.20g of MF (reinforced)

Refs: 1) D.B. Gawthrop, JFrankInst, 214, No 6, p 647 (1932) 2) LeRoy V. Clark, IEC 25, 668 (1933) 3) ibid, p 1388

Infrared Guidance Systems. See German section under Guidance Systems for Missiles in Vol 6

Infrared Spectroscopy, Application to Explosives & Propellants. The infrared region is that portion of the electromagnetic spectrum which is located between visible light and the microwave radio region. It is conveniently divided into near infrared, which extends from the edge of visible light (0.0008mm, 0.8μm or 8000 angstroms) to about 0.025mm (25μm or 250,000 angstroms) and the far infrared, which extends from the near infrared to about 1mm (1000μm or 10 million angstroms). The region up to about 0.002mm (2μm or 20,000 Å) of the near infrared is called photographic or photoelectric because radiation in this region is readily detected photographically or photoelectrically.

Infrared (IR) radiation can be dispersed into a spectrum by means of coarse diffraction gratings or prisms of special material, such as rock salt, which is transparent to much longer waves than glass, after which the rays can be detected with a device which is sensitive to small heating effects such as bolometers, thermocouples and various other radiometers.

The infrared spectrum (especially the near-infrared) has assumed great importance in chemical and biological research because of the highly specific absorption of chemical compounds at these wavelengths. The infrared absorption of a given organic compound may be used to characterize that particular compound. The infrared spectrum of a mixture of several compounds among which there is no interaction, does not lie between the spectra of the individual compounds, but consists of a direct superposition of the spectra of the individual compounds.

A general reference on IR spectroscopy is Ref 7

Application of infrared (IR) spectroscopy to the study of explosives & propellants can be divided into three general areas:

1) Identification and quantitative estimation of unreacted explosives, propellants & inert ingredients in expl & proplnt compositions

2) Study of the composition of combustion or explosion products. With rapid scan spectro-
meters such studies can provide kinetic data.

3) Study of the intermediates or equilibrium composition in the preparation of explosives

Most of the published work belongs in area 1). Particularly pertinent are the studies of Pristera et al (Refs 2 & 12) on IR spectra of explosives & the identification of the ingredients of a complex propellant (Ref 2). In Ref 12, Pristera gives the IR spectra of: EDNA, MEDINA, α, β & γ HMX, Tetryl, BTN, NC, DEGN, Molten Hexanitrate, Metriol Trinitrate, NG, PETN, TEGN, 1,5-Dinitonaphthalene, 1,8-Dinitonaphthalene, 1,3,8-Trinitronaphthalene, 2,4-DNT, TNT, AN, Ammonium Picrate & 1,3,6,8-Tetranitrocenezole. Further information on other explosives, explosive mixtures and the inert ingredients in these mixtures is contained in Ref 1.

The following tabulation (from Leclercq in Ref 8) shows the symmetric & antisymmetric stretch frequencies of the N-O bond in various compounds:

<table>
<thead>
<tr>
<th>Compound</th>
<th>Sym freq (cm(^{-1}))</th>
<th>Antisym freq (cm(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>RNO(_2)</td>
<td>1500-1550</td>
<td>1320-1360</td>
</tr>
<tr>
<td>RNNO(_2)</td>
<td>1550-1600</td>
<td>1200-1260</td>
</tr>
<tr>
<td>RONO(_2)</td>
<td>1600-1660</td>
<td>1250-1300</td>
</tr>
<tr>
<td>Metal NO(_3)</td>
<td>one band only at 1350-1400 cm(^{-1})</td>
<td></td>
</tr>
</tbody>
</table>

IR spectra of nitroaliphatic compounds are reviewed by Slovakskii (Ref 13). This review contains 237 references thru 1969.

In the absence of conjugation N\(_3\) absorption bands of organic azides occur in the 2100-2110 cm\(^{-1}\) region (Ref 9). Conjugation or presence of electron acceptor groups shift the bands to 2135-2166 cm\(^{-1}\). It is claimed that for organic azide measurement of band intensity is a more sensitive indication of compound structure than measurement of band position. Electron donor groups in the molecule increase band intensity and electron acceptors lower it.

The spectra of inorgan azides are given in a review by Yoffe (Ref 10).

IR and Raman spectra of NF compounds are reviewed by Moskvitina & Kuzyakov (Ref 14). Their review contains 47 references.

Pristera has developed an IR method for determining α, β & γ TNT as well as 2,4-DNT in mixtures such as found in exudates (Ref 1).

Castelli et al (Ref 3) worked out a rapid IR method for determining NG, TA & 2-Nitrodiphenylamine in Casting Solvent.

Examples of studies belonging to area 2) are:

- Studies by Crawford & Rotenberg (Ref 4) who used a rapid-scan spectrometer in conjunction with a strand burning apparatus to examine NG-NC "low" temperature decomposition and flames. During the decomposition of commercial double-base propellants, the gas products were: NO, N\(_2\)O, CO\(_2\) & CO. When these propellants were burned under 100-150 psi nitrogen pressure at a linear velocity of ~100 cm/sec, CO\(_2\) & CO absorption bands appeared even at 2 cm away from the burning surface. Nitric oxide was barely detectable and N\(_2\)O was completely absent.

- Leclercq gives some IR spectra for residues of explosive that had been burned (Ref 8).

Zirkind reviews investigations of radiation emission of rocket exhausts since 1940 (Ref 11). Emission characteristics of the exhaust plume are strongly dependent on rocket engine parameters & the propellant system. If these & motor operating conditions are stipulated, machine computations can give exhaust composition & sometimes temperatures.

Examples of studies pertaining to area 3) are:

- Simecek's study of the preparation of the lower nitrates of PETN and measurement of their IR spectra (Ref 6).

- Leclercq's study of the nitration of amyl alcohol and cellulose by following the amount of OH remaining after nitration by IR absorption methods (Ref 5)


Influence Test. See Gap Test in Vol 1, p XIV

Infusorial Earth. See Kieselguhr in this Vol

Ingelite. See Antigel de Sûreté in Vol 1, p A466-R

Ingold, C.K. (1893-1970). English physico-organic chemist whose pioneering research helped to lay the basis of modern organic chemistry. His work on nitrination reactions did much to elucidate the mechanism of these reactions, particularly the nitrination of aromatic compounds. He became a fellow of the Royal Society in 1924 & was knighted in 1958. He was the recipient of many scientific honors & medals Refs: 1) “World Who’s Who in Science,” (A. G. Debus, Ed), Marquis, Chicago (1968) 2) C & EN 49 (3), 41 (1971) & CA 74, 106995 (1971)

Inhabited Building. In the American Table of Distances for Storage of Explosives, an “Inhabited Building” is defined as: “A building regularly occupied in whole or in part as a habitation for human beings, or any church, schoolhouse, railroad station, store or other structure where people are accustomed to assemble, except any building or structure occupied in connection with the manufacture, transportation, storage, or use of explosives” Ref: American Table of Distances for Storage of Explosives as Revised & Approved by the Institute of Makers of Explosives, Sept 30, 1955

Inhibiting of Solid Propellant Grains. To prevent surface burning and/or contact of the propellant grain with its container, various methods of deactivating and/or coating of the grain surface are used. This is called “Inhibiting” the grain.

Inhibitors or restrictors are applied to those surfaces of the grain where burning is to be prevented. They consist of plastic materials which are sometimes charged with an inert refractory filler (chalk powder) and burn very slowly if at all. Sheets of cellulose acetate or ethyl cellulose, 2 to 4mm thick, have been widely used for that purpose but cellulose acetate is not very satisfactory with double-base propellants because of the exudation of nitroglycerine during storage. With composite propellants the fuel, with an inert filler substituted for the oxidizer, is also often used. Restrictors can be applied by dipping the grain or by spraying the inhibiting material on. These two methods are not very satisfactory because each coat must be very thin to dry well and a prohibitively large number of applications are required to build up the desired thickness. Preferably inhibitor sheets are bonded to the grain, possibly under pressure at fairly high temperature, with a suitable glue like polyurethane, or a restricting tape is wrapped around the grain.

The mechanical and physical properties of the inhibiting material must be fairly similar to those of the propellant in order to minimize the differential expansion. In this respect the use of several layers with possibly different compositions is favorable. With long burning timers (more than 20 sec) the development of a reliable inhibitor poses a rather difficult problem, especially with end-burning grains. The situation is greatly improved in this respect when the inhibitor is not in contact with the hot gases. This is the case with case-bonded grains where the thin chamber liner acts as restrictor (Ref 2)

Inhibiting by Denitrating. This consists of the removal of the nitrate groups from the surface of the propellant grain by treating it with ammonium hydrogen sulfide and ammonium polysulfide. Previous to this treatment, however, the surface of the grain is swelled by a one minute dip in acetone (contg 5% by wt of AcOH), followed by one hour drying.

The denitration solution is prep'd by dissolving 8 parts by wt of ammonium hydrogen sulfide in 92 parts of alcohol (30% soln by volume). This aqueous solution is circulated over the grains for about 2 hours during which time the soln is kept up to strength by adding H₂S gas. The grains are then removed from the
denitrating bath and, after rinsing them quickly with water, placed in a dilute glycerin soln at 40° for two hours. Finally the grains are dried by wiping and then stored. This method gives grains with denitrated layers as thick as 0.018". These layers do not burn during firing but are left as a residue. Various kinds of grains were treated by this method, including Mk31 grains and triangular catapult grains. The powders prep'd by this method proved to be fairly non-hygrosopic and stable in storage.

Spiral Wrapping Process consists of wrapping around large cylindrical grains of powder (such as Mk31) a thin tape of inert, non-explosive material (such as cellulose acetate, ethyl cellulose) in order to create a protective, non-combustible layer. Wrapping is done on machines which resemble a shop lathe. Usually 6 layers of tape are applied to make a layer about 0.045 inches thick.

Inhibiting of Cast Double-Base Grains. A good inhibitor should bond to the grain with a strength equal to the grain strength itself. It should also have about the same coefficient of expansion as the grain material.

Alleghany Ballistic Laboratory found that casting grains in convolute-wound or extruded beakers is quite satisfactory. In the spiral wrapping process they had to use solvent-type adhesives, but would prefer to use a non-solvent type.

When using large grains with internal perforations, the problem of NG migration is not of great importance, because the outer inhibited cylindrical surface forms only a small percent of the total operating surface. After up to two years in storage at 120°F, booster grains were still in good condition.

Inhibiting of Double-Base Catapult Propellants with Ethyl Cellosolve Gel Lacquers. Propellants for slotted tube catapults are designed to produce a const press system in the catapult tube. Catapult propellants usually have non-circular cross-sections and are completely externally inhibited.

An ideal inhibitor is one which burns through at the same time that the powder web is exhausted so that no residue remains in the catapult. A dipping process was employed to coat the grain with inhibitor. Viscous gel lacquers were used in this process with viscosities of 8,000-10,000 centipoises at the dipping temperature. The solvents were adjusted so that there was a sharp increase in viscosity as the temp was lowered towards room temp. As the dipped grain is withdrawn from the warm lacquer the coating sets due to solvent evaporation and gelation caused by the temp drop. In this way running and sagging are minimized and a substantial coating can be applied with a single dip. If a very thick coat (such as 0.10") is required, 3 to 4 dips are necessary.

Most of the grains are 15 to 18" in length and the major diameters range from 0.8" for small experimental catapults to 3-3/4" for Field Installations. Inhibitor thicknesses are between 50 and 100 mils.

_Gel lacquer formulations_ used for dip coating. The early catapult propellants were inhibited by dipping in a lacquer consisting of toluene 80%, acetone 20% and a total solids content of 15-18%. The grains were dipped twice and had a cured inhibitor thickness of 20 to 30 mils. As the size of experimental slotted tube catapults increased, coatings of 100 mils were required. This necessitated an increase of the solid content to 25-27%. As this soln was too viscous some methylol was incorporated in solvent formulation. In order to improve the room temp gelling characteristics an aliphatic hydrocarbon was added.

With this system there is no serious curing problem for coatings under 50 mils thick, but for coatings of the order of 100 mils, about three weeks are required to remove the volatile solvents.

_Castable Cellulose Acetate Inhibitors for Rocket Propellants._ During the joint development of the propellant for the T34 Jato by PA & Hercules, it became desirable to investigate methods of restricting the ends of the powder grains. Since it was difficult to obtain uniformly good bonding and satisfactory inhibiting by cementing cellulose acetate sheets to the ends of large prop grains, the possibility of using some form of inert powder for this purpose was investigated. The inert casting powder adopted consisted of 90% cellulose acetate and 10% dimethyl phthalate in 0.30" x 0.030" granulation.
A layer of this inert powder was placed in an extruded cellulose acetate tube, partly filled with OGK casting powder (NC of 12% 88, 2-nitrodiphenylamine 2, dioctyl phthalate 5 and lead stearate 5%) and was then covered over with more OGK powder.

OGK casting solvent (72% NG, 27% triacetin and 1% nitrodiphenylamine) was then fed through the entire mass, following which the assembly was cured at 140°F. After curing the charge was sectioned and examined. The granules of inert powder had gelled and hardened, forming a well consolidated mass which was very well-bonded to, and almost undistinguishable from, both the OGK powder and the extruded tube.

A solid casting prep consisting of inert casting powder and OGK casting solvent. The chemical analysis of the cured charge gave NG 28.0, 2-NDPhA 0.4, cellulose acetate 55.1, DMePh 6.1, Triacetin 10.4%. The heat of combustion was 3790 cal/g. The heat of explosion was 260 cal/g.

Strands of this material would not burn in the inert atmosphere of the Crawford type bomb, but they burned in air at 70°F at the rate of 0.023" per second. This compared with a burning rate for H9 propellant of 0.097"/sec at 1 atm and 0.36"/sec at 1000 psi & 70°F.

This cast cellulose acetate material has been employed for end restricting 8¼" and 20" diameter perforated cast OGK grains and thermally cycled JATO motors at -65, +70, +165°F. No cracks or separations of the cast acetate restrictions were observed but some softening and deformation was noted after conditioning at +165°F. These rounds functioned satisfactorily when fired at either -65 or +165°F.

Inhibiting Double Base Rocket Propellants at Picatinny Arsenal. Most of the work at Picatinny Arsenal in connection with inhibiting of solventless extruded or cast double-base propellant has been with convolute wrappings of grains or tubes with cellulose acetate. There has been a trend toward use of extruded tubes made by the casting process.

When convolute wrapping of the long grain for the T200 Rocket (4.5" diam and 72" long) proved unsuccessful, a procedure was investigated for inhibiting with Thiocol composition developed at PA (see 6th JANAF Sol Prop Meet), but this method proved to be not very satisfactory on account of the appreciable migration of NG into the inhibitor, which resulted in its deterioration at elevated temp. Migration occurs also when using cellulose acetate inhibitors.

An attempt to prevent migration of NG by inserting a layer of cellophane after the first cellulose acetate wrap proved to be unsuccessful because of the poor bond between cellulose acetate and cellophane.


Inhibitors. Substances which slow down or stop a chemical reaction. For instance, some substances act as corrosion inhibitors, others are added to explosives to prevent their decomposition in storage (see below)

Inhibitors in Explosives. Substances added to explosives in order to take up the nitrogen oxides formed during storage or to neutralize residual acids or acids formed during storage. The presence of NO₂ and other nitrogen oxides and/or acids in explosives (especially in smokeless powders and dynamites) is very undesirable because they act as catalysts and promote further, more rapid decomposition. Most of the stabilizers used in smokeless powder, such as diphenylamine, centralites, urethanes etc are really inhibitors because they react with NO₂ and other nitrogen oxides to form nitros- and nitrocompounds. Acid neutralizers such as chalk are added to dynamites.

Inhibitor Strips for Smokeless Powders. In order to control the burning area of large grain smokeless powders and to prevent the grains from coming into contact with the rocket motor walls, inhibitor strips (consisting of ethylcellulose or cellulose acetate) 74, dibutyl phthalate 4, diethyl phthalate 16 and dimethyl phthalate 6%) are pre-glued to the surface of grains. This is done by means of adhesives consisting of solns of NC in ethyl acetate, Cellosolve, mesityl oxide, Bu acetate etc, which act as a mutual plasticizer for the
powder and the material of the inhibitor strip
Ref: A.M. Ball, USP 2643611 (1953) & CA 47, 9016 (1953)

**Initial Velocity.** Same as Muzzle Velocity—see under Ballistic Tests and Ballistics in Vol 1, p B5-R & B7-L & additional French refs:
1) R. Rousselet et al, MAF 22, 171-190 (1948), Measurement of initial velocities of projectiles at the battlefield
2) R. Rousselet, MAF 22, 649-655 (1948), Use of a fluxmeter by Grassot for measuring initial velocities of projectiles at the testing range
3) M.C. Thatcher, MAF 23, 305-308 (1949), Measurements of the initial velocities of projectiles (Translated from the English)
  Pepin Lehalleur, p 110, gives initial velocities for some French and foreign propellants

**Initiating Devices.** See Detonators, Igniters, Primers, and Other Initiating Devices used for Nonmilitary and Military Purposes in Vol 4, pp D733-D928

**Initiating Explosives.** (Initial Detonating Agents; Primary Explosives) (Zündsprengstoffe in Ger) Sometimes erroneously called Priming Compounds, are substances used in small quantities, which on being subjected to the action of flame (produced by primers or other means), heat, impact, friction or an electric spark, generate a detonation wave. The detonation of the primary explosive initiates detonation in a larger quantity of explosive (“main” charge or “base” charge) which usually possesses higher brisance, power and velocity of detonation than the initiating explosive

The main requirements for initiating explosives are: 1) sufficient sensitivity to heat, flame or impact etc that they can be readily detonated, but not so sensitive as to make them unsafe to handle and transport; 2) sufficient stability at elevated temperatures that they will not decompose while stored in ammunition, thus rendering it unusable. Brisance, power and velocity of detonation are of secondary importance in these explosives

There are many explosives which could serve as initiating agents if they were not too dangerous to handle and too unstable in storage. The most frequently used initiating substances are:
  Lead Azide, Mercuric Fulminate with or without Potassium Chlorate, Lead Styphnate, Cyanuric Triazide, Tetracene and Diazodinitrophenol
  The following substances are also good initiating agents but have not found much use:
  Silver Azide, Cadmium Azide, Cupric Azide, Triazidotrinobenzene, Chlorotrimercuraldehyde, Nitrogen Sulfide & Hexamethylenetriperoxidediamine

Until WWI, Mercuric Fulminate was the principal initiating agent used, but Lead Azide has now replaced it. Lead azide is not the most powerful azide, but is more stable and less dangerous to handle than some of the other ones. Cadmium Azide, for example, is more powerful than Lead Azide but is unsuitable as an initiating agent because it is difficult to prepare and is soluble in water.

Marshall (Ref 13, p 158) gives the effectiveness of various azides and fulminates as initiators. No 1 is the most effective and No 11 is the least effective:

1) Cd Azide. 2) Cd Fulminate 3) Ag Azide 4) Ag Fulminate 5) Lead Azide 6) Cuprous Azide 7) Cuprous Fulminate 8) Mercurious Azide 9) Mercuric Fulminate 10) Tellurium Azide 11) Tellurium Fulminate

For the preparation and properties of initiating agents, look under the individual compounds

(1939)  16) H. Muraour, MAF 18, 895
(1939) & CA 34, 4905 (1940)  17) Davis
(1943)  400–58 (under Primmers)  18) L.V.R.
Clark, USP 232608 & 2325742 (1943) & CA
38, 488–90 (1944)  19) R. Schmitt, SS 38,
133 (1943) & CA 38, 2822 (1944)  20) K.K.
Andreev, DRAcadSci(Russia) 44, 18 (1944)
& CA 39, 813 (1945)  21) Pérez Ara (1945),
557 & 647  22) C. Francais, FrPat 856366
(1946) & CA 42, 2774 (1948)  23) Stett-
bacher (1948), 95  24) Bowden & Yoffe
(1952)  25) Bowden & Yoffe (1958)
26) Baum, Stanyukovitch & Shekhter (1959)
27) W.H. Rinkenbach in Kirk & Othmer, Vol
8 (1965), 583–93 (Initial Detonating Agents)
28) B.A. Bydal, “Percussion Primer Mixes”,
Ordnance, Vol LV1, No 309, 230–33 (Nov–
Dec 1971) (A review of the history of initiating
explosives from their discovery early in the seven-
teenth century up to the development of today’s
safer noncorrosive, nonmercuric mixes)
29) O.E. Sheffield & W.R. Tomlinson Jr,
“Properties of Explosives of Military Interest”,
AMCP 706-177 (Jan 1971)  30) Anon,
“Principles of Explosive Behavior”, AMCP
706-180 (April 1972)

INITIATION

In its strictest sense Initiation is the generation
of initial conditions that lead to stable
detonation. However, it is common usage also
to refer to the first stages of deflagration and
explosion as initiation. Thus it is often difficult
to differentiate between “initiation” and “igni-
tion.” It is generally accepted that initiation is
a thermal process which requires an external
stimulus (except for “spontaneous” ignition) to
get it started. Regardless of the nature of this ex-
ternal stimulus, a necessary condition for initia-
tion is that heat generated in the explosive
(that is “initiated”) exceed all heat losses by
the explosive. Since heat generation (chemical
decomposition) increases exponentially with
rising temperature, but heat losses are roughly
proportional to temperature to the first power,
y any stimulus that generates a high enough
temperature in an explosive is apt to “initiate”
that explosive. Conversely, an explosive in
which exothermic reaction is readily started by
a variety of external stimuli, is said to
be a “sensitive” explosive. Thus the ease of
initiation is closely connected with explosive
sensitivity, but it must be emphasized that
“explosive sensitivity” depends not only on
ease of initiation but also on the ease of
growth and propagation of whatever process
was “initiated.” It is entirely possible that
highly localized regions of an explosive are
“initiated” but the processes in these regions
die out without propagating.

Because initiation is such a fundamental
part of explosion and detonation phenomena
various aspects of initiation have already been
described in previous volumes and in this
volume of the Encyclopedia. Particularly per-
tinent are the articles on Deflagration, Vol 3,
pp D38-40; Detonation (and Explosion), Initia-
tion and Propagation, Vol 4, pp D402-419;
Hot Spots & Ignition in this Vol.

In what follows we will give brief definitions or
descriptions (whenever necessary) of various
modes of initiation of condensed explosives,
ie, various external stimuli that produce initia-
tion, and provide references to Encyclopedia articles
where these modes of initiation are discussed in
detail.

Initiation by Booster. See below under Initiation
by Primers.

Initiation by Bullet Impact is a special case of
Initiation by Projectile Impact. See Bullet Tests
in Vol 2, pp B332-340 and Vol 4, p D153; also
under Projectile Impact below.

Initiation by Electric Fields. Silver Azide
may be initiated by DC fields of ca 70v (min field
strength for explosion ca 250v/cm) at extremely
low current flow (Bowden & Yoffe “Fast Re-
Maycock & Grabenstein [Science 152, 508
(1966) & CA 65, 562 (1966)] have postulated
that piezoelectric effects generated by compress-
ing explosive crystals may generate sufficiently
large electric fields to initiate these crystals.

Initiation by Electron Beams. According to
Bowden & Yoffe (“Fast Reactions in Solids”
Acad Press, 1958, p 114) the initiation of
lead & silver azides by an electron beam is
at least in part due to bulk heating of the
explosives by the beam. Pulsed electron beams have been used to initiate explosives and study their shock properties (J.H. Shea et al, ONR Symp Det (1970) p 351). See also *Electrons & Neutrons Action on Explosives* in Vol 5, p E77

*Initiation by Electrostatic Discharge.* This is described in detail under *Electricity; Exogenous Hazards Associated With It* in Vol 5, pp E36-54. See also *Initiation by Sparks*


*Initiation by Fission Fragments.* All attempts to initiate explosives by nuclear fission fragments such as α-particles, protons, Ar or Hg ions, γ-rays, X-rays, mesons and pions have resulted in failures [Bowden & Yaffe quoted above, & J. Cerny & J.V.R. Kaufman, JChemPhys 40 (6), 1736 (1964)].

*Initiation by Flame.* Various aspects of this process were described under *Burning and Combustion,* Vol 2, pp B343-346; *Dead-Pressed Explosives,* Vol 3, p D20; *Ignition* in this Vol; *Thermal Explosion* in a future Vol

*Initiation by Friction.* See *Friction Sensitivity Tests* in Vol 6, p F204-L and *Hot Spots & Impact* in this Vol

*Initiation (of Condensed Explosives) by Gas Detonations* does not involve the same phenomena as *Initiation by Primers or Boosters.* The pressures generated in gas detonations are too low to produce sufficiently intense shocks in condensed explosives for shock phenomena to play a major role in the initiation of the latter. Even the energetic C_2H_2 + O_2 detonation produces a detonation pressure of only ca 43 atm for an initial gas pressure of 1 atm. The max shock pressure produced by reflecting the oxy-acetylene detonation wave is only ca 0.2 to 0.3 kbars. Thus gas detonations initiate condensed explosives (if at all) by some direct heat transfer process. In oxy-acetylene detonations, the equilibrium temperature (CJ temp) is quite high ~4500^K.

The writer (unpublished results) was able to initiate Lead Azide pellets of ca 2.5 g/cc density with oxy-acetylene detonations. However ca 1.2 g/cc PETN pellet could not be initiated under these conditions. Gordon et al [Nauchn-Tekhn Probl Gorenya, Vzryva (1965) p 12 & CA 64 1894 (1966)] succeeded in initiating liquid mixtures of tetrynitromethane (TNM) and benzene with stoichiometric methane-oxygen detonations. For 1.5 vol parts of TNM & 1 vol p of benz the initial pressure, P_o, of the detonating gas mixture had to be greater than 2 atm to initiate the liquid. Initiation delays decreased as P_o increased; delays were 350, 10 & ~0 μsec for P_o of 2, 12 & 24 atm. For 4:1 by vol TNM/benz initiation of the liquid was observed for P_o > 0.7 atm. At P_o ~ 0.7 atm the initiation delay for this liquid mixture was ~70 μsec

*Initiation by Heat.* See *Thermal Explosion and Ignition & Hot Spots* in this Vol

*Initiation by Hot Fragments* is probably the primary initiation mechanism in the so-called *Halved Cartridge Gap Test* (see Vol 1, p XV) and in other systems in which initiation occurs by transmission across an air gap from a donor to an acceptor charge. The hot fragments are products and reacting material thrown off by the donor; see also *Detonation by Influence* in Vol 4, p 397-R

*Initiation by Hot Wire* is the usual method of initiation in *Electro-Explosive Devices* such as electric blasting caps, electric detonators and squibs. In general this method is only applicable to the initiation of primary explosives (unless very special conditions are used). Various aspects of *Hot-wire Initiation* are described under *Ignition* in this Vol; *Detonators, Igniters, Primers and Other Initiating Devices* in Vol 4, pp D737-742, D806-807, D846-850 & D854-856 and *Electro-Explosive Devices* in Vol 5, pp E63-68. Addit-

_Initiation by Heat._ See Thermal Explosion in a future volume and _Hot Spots & Ignition_ in this Vol

_Initiation by Impact_ is described in detail in this Vol under _Impact_. Tests in which initiation may occur as a result of dropping an explosive device are described under _Drop Tests_ in Vol 5, pp D1549-1554

_Initiation by Influence_ usually called _Sympathetic Initiation_, is the initiation of an explosive charge by the detonation of a nearby charge separated from the first charge by an inert medium; see _Detonation by Influence_ in Vol 4, pp 395-398 and also _Initiation by Hot Fragments & Initiation by Shock_.

_Initiation by Ionizing Radiation._ See _Initiation by Fission Fragments_.

_Initiation by LASER._ See _Initiation by Light_ in this Vol

_Initiation by Light._ See _Light, Initiation by in this Vol_.

_Initiation by Photochemical Effects._ See _Initiation by Light_.

_Initiation by Precursor_ is a phenomenon encountered in low velocity detonations, LVD, in liquid explosives. It depends primarily on cavitation of the liquid by the shock traveling in the container ahead of the shock in the liquid. For a description of this effect, see _Low Velocity Detonation_ in this Vol.

_Initiation by Primers (and Boosters)_ is the standard method of initiating secondary explosives. Thus _hot wires_ (or other means) are used to initiate the primer charge (Lead Azide, Mercury Fulminate etc) explosive whose detonation then initiates the _main charge_ of PETN, RDX etc. For “insensitive” explosives such as TNT, ANFO etc an intermediate _booster charge_ (PETN, RDX, Tetryl, NG Dynamites etc) may be necessary between the primer charge and the main charge. For details, see _Minimum Priming Charges_ in Vol 8; _Booster_ in Vol 2, pp 243-246; _Contact Detonation Sensitivity Test_ in Vol 4, pp D186-190; and the added reference of C. H. Johansson & T. Sjölin, ONR Symp Det (1965) p 435.

_Initiation by Projectile Impact_ is a complicated process which depends on a combination of shock initiation, impact initiation and hot fragment initiation effects. For fast projectile impact, shock initiation effects are predominant. With slow projectiles the initiation resembles _Impact Initiation_. Further complications are introduced if the impacting projectiles are hot; a special case where the explosive is part of the projectile is the so-called _Susan Test_—see _Barrier Tests & Their Comparison with Shooting Tests_ in Vol 4, pp D145-147 and _Detonation (and Explosion) Experimental Procedures_ in Vol 4, pp 333-335.

_Initiation by Radio Frequency (RF) Radiation_. RF radiation, ie, radio wave & radar transmitters can, under certain circumstances, initiate electro-explosive devices. This topic will be discussed under _Radio Frequency_ Radiation, _Effects on Explosives_. Also see articles on _Electromagnetic Compatibility_ & _Electromagnetic Field Hazard, Simulated_ in Vol 5, pp E70-71 and _Electric Blasting Caps_ & _RF Energy_ in Vol 5, p E25-L.

_Initiation by Shock_ will be described in detail under _Shock Sensitivity_. Some of the tests for determining the shock sensitivity of explosives have already been given under _Detonation (and Explosion) Experimental Procedures_ in Vol 4, pp D318-321, pp D322-331 & D344. Some preliminary discussion of shock initiation is also given in Vol 4, pp D520-522.

_Initiation by Sparks_. Primary explosives, Black Powder & possibly some dynamites can be initiated by sparks of a non-electrical nature such as those caused by metal-on-stone friction etc. Manufacturers of dynamite use non-sparking tools in their plants. This is probably...
a carry-over from the days when Black Powder was an important commercial explosive. Initiation by non-electric sparks is obviously a special case of initiation by heat.

**Initiation, Spontaneous.** This is really a misnomer. What it refers to is the self-heating of an explosive as a result of autocatalytic decomposition eventually leading to a thermal explosion. “Spontaneous” initiation may also occur during crystal growth e.g. in Lead Azide; see *Detonation (and Explosion), Spontaneous* in Vol 4, pp D561-563.

Written by J. ROTH

Innesco. Some Italian percussion fuses of WWII consisted of two parts—the “spolella” and the “innesco.” The innesco was a detonator holder without which the fuse could not operate.

Ref: Ordnance Sergeant, August 1943


**INOSITOL AND DERIVATIVES**

*Inositol or Hexahydrohexamethylenene.* 1,2,3,4,5,6-Cyclohexanehexanol, Mesoinositol (Inosite, Cyclohexanhexanol); C₆H₁₂(OH)₆, mw 180.16, d 1.524 at 15°/4° for the dicyclate, 1.752 anhydrous, mp (anh) 200-225°. White crystals, usually obtained from seeds of various plants, such as barley, peas and beans. Soluble in w, insol in abs alc & eth.

It is a cyclic hexahydric alcohol, which is optically inactive. Exists also in two optical modifications:

- **d-Inositol**: mp 247°. Colorless crystals; may be prepd by heating its monomethyl ether, pinene, C₆H₁₂(OH)₂(OCH₃) [described in Bellstein, 5, 1193 & (587) with hydriodic acid. Pinene occurs in resins and leaves of numerous plants, for example, senna leaves and the Madagascar rubber plant.

- **l-Inositol**: mp 238°. Colorless crystals; may be prepd by heating 75 parts of its monomethyl ether, *quebrachitol*, with 100 parts of hydriodic acid (d 1.70), placing the mixture in a long-neck flask provided with a tube for conducting gas away. After one hour of heating at 130-140° the temperature is raised to about 150° and heating is continued for an additional 3 hours. The mixture is then poured into a beaker and allowed to crystallize. The crystals are ground, washed with alcohol and ether, dissolved in water and filtered through activated charcoal and concentrated to a syrupy consistency. A mixture of alcohol 90% and ether 10% is then added in order to precipitate inositol. The resulting crystals are washed with alcohol and ether and dried. W. deC. Crater proposed nitrating the optically active inositols in order to prepare the explosive hexanitrate (see below).


**Inositol Hexanitrate (IH).** C₆H₁₂(ONO₂)₆, mw 450.16, N 18.67%, OB to CO₂ + 10.7%; d 1.411 pressed at 3,000 psi; mp 120-122° (dec), bp ignites at 195°. Colorless plates or prisms (from hot alcohol). May be prepd by nitrating water-free inositol with a mixture of 1 vol conc HNO₃ and 2 vols of concd H₂SO₄. The resulting crude hexanitrate is dissolved in boiling alcohol and crystallized on cooling.

The inositol trinitrate, which is formed as a by-product of nitration, remains in alcoholic solution and does not contaminate the hexanitrate. IH is insol in water; v sol in methanol and ethanol, acetone, ether and concd H₂SO₄.

It is an explosive, comparable in impact sensitivity to MF and more brisant than tetryl, as determined by the sand test (52g of sand crushed compared with 47.7g for tetryl or 109% of tetryl). Its thermal stability is low (Ref 4) and for this reason is not suitable for military purposes, but can be used as a base charge in commercial blasting caps (Ref 3).

When dropped on a heated plate, IH flashes but does not explode, although it may be detonated by a commercial blasting cap.

Crater (Ref 2) prepared hexanitrate of optically active d- and l-inositols by nitrating
20 p of these with 100 parts of mixed nitric-sulfuric acid at about 7° (45°F). The resulting products had nitrogen contents about 18.3%, as compared with 18.67% theoretical. They are explosives with properties resembling those of inactive inositol hexanitrate. (The dextro- product is a solid, while the levo- product is a liquid). They are better gelatinizers than NG or NC and have been recommended as ingredients of double-base smokeless powder. They can also be used in dynamites or other commercial explosive mixtures in combination with NG or aromatic nitrocompounds. Crater has also claimed the use of IH in non-headache dynamites (Ref 5)

The same inventor proposed to nitrate a mixture of 20 p inositol and 80 p of glycerin with the same mixed acid as used for nitrating glycerin. The resulting product is fairly stable, giving a 12-minute KI test at 82.2°; nitrogen content 18.4% (Ref 2)

An interesting property of IH is that it can be detonated by the flash of a match-head (eg Ba-nitrate/Mg powder/Pb-hyphosphite). Because of this Crater (Ref 3) patented its use as a base charge in electric blasting caps containing either a MF/KClO3 initiating charge or a matchhead. It can also be used in fuse-type blasting caps if a match-type ignition composition is placed over a prep'd charge of IH (Ref 3)

Ficheroule & Kovache (Ref 6) propose a method of preparing IH that is essentially similar to the one described above. They do, however, suggest the addition of 85 p of lactose to 15 p of wet IH before drying as they consider the drying of pure IH (without 85 of inert material) to be dangerous

They find IH to be more impact-sensitive than Hexamethylenetriperoxycyanamide and very sensitive to friction. IH explodes violently when placed on a sheet of paper when one of the corners of the paper is ignited and its flame reaches the IH


Inspection of Ammunition and of Explosives. The process of examination of explosives etc., usually of a visual, physical, chemical or ballistic nature are performed by specially assigned persons, called Inspectors. The idea of this inspection is to determine the quality of the product from the point of view of the specifications pertaining to it

Inspections may be divided into Initial, and Periodic or Subsequent Inspections

For more info regarding regulations & procedures connected with the inspection and testing of exps for the Government, see the following Refs


Instantaneous Fuse. This fuse, burning at the rate of 100 to 300 feet per second, was formerly much used in blasting operations where a number of charges had to be fired "simultaneously." It has now been replaced by detonating fuses and electric detonators

Instantaneous fuse was made by wrapping several strands of quick match in waterproof tape. The fuse was fired by means of a percussion cap in a special pistol

The term is sometimes erroneously applied to Cordeau Défomant or Primacord

Ref: Marshall 2 (1917) p 540

Instantaneous Photography. See High-Speed Photography in this Vol and Cameras, High-Speed Photography in Vol 2, pp C13-19

Instrumentation and Instrumental Methods of Analysis (see also next topic). The principle purposes of instrumentation in the chemical industry are to measure and control physical changes and chemical reactions. Instrumental methods of analysis concern the application of
various instruments to the solution of specific problems of analytical chemistry. Under the
title come spectroscopy, polarography, X-rays,
photometry, supersonics, various electronic
devices, such as oscillographs etc

Each of these subjects is mentioned under
separate headings. References on instrumentation
run into the thousands. In the opinion of Dr.
R. H. Müller, there should be a distinct subject
known as analytical instrumentation, which
should be concerned with a study of all known
physical phenomenon for their possible use in
analytical chemistry. General references on this
subject are given below
Refe: 1) M. F. Behar, Manual of Instrumentation,
Instrument Publishing Co, Pittsburgh (1932) 2)
R. H. Müller, IEC (Anal Ed), 11, (1939) (Review
on photoelectric methods in chemical analysis,
including 239 refs) 2a) R. H. Müller, Ibid, 13,
667-754 (1942) (A review of instrumental
methods of analysis) (313 refs) 3) L.A. McColl,
Fundamental Theory of Servomechanisms, Van
Nostrand, NY (1945) 4) D.P. Eckman, Principles
of Industrial Process Control, J Wiley, NY (1945)
5) H.W. Bode, Network Analysis and Feedback
Amplifier Design, Van Nostrand, NY (1945) 6)
Greenwood and Collaborators, “Electronic
Instruments,” Vol. 21, MIT Radiation Laboratory Series,
20, 389 (1948) (Instrumental methods of analysis)
8) R.H. Müller & J.J. Lingane, AnalChem 20,
795 (1948) 9) A.L. Chaplin, Instruments 21,
532-40 (1948) 10) H.M. Schmitt, Automatic
Chemical Process Control, Minneapolis-Honeywell
Regulator Co, Phila (1949) 11) R.H. Müller,
12 refs are included) 12) A. Camp & L. Slater,
ChemEng 57, No 9, 108-111 (1950) 13) D.M.
Considine & S.D. Ross, Chem Engg Prog 46,
518-22 (1950) 14) D.A. Smith & J. Procopi,
Instrumentation 4, 32-34 (1950) 15) J.A.
Parker, Instrumentation 5, 4-7 (1950) 16) R.
Rosenthal, Instruments 23, 664-69 (1950) 17)
E.F. Pollard et al, IEC 42, 748-52 (1950) 18)
M.F. Behar, Ed, The Handbook of Measurement
and Control, The Instrument Pub Co, Pittsburgh
(1951) 19) R.E. Kirk & D.F. Othmer, Eds,
Encyclopedia of Chemical Technology, Inter-
science, NY, v 7 (1951) pp 908-926; J-Proopi,
Instrumentation (11 refs) 20) R.H. Müller,
Analytical Chemistry, “Instrumentation”; Short
article, each month 21) H.H. Willard et al,
Instrumental Methods of Analysis, Van Nostrand,
NY (1951) 22) D.F. Boltz, Selected Topics
in Modern Instrumental Analysis, Prentice Hall,
NY (1952) 23) W.G. Holzbock, Instruments
for Measurements and Control, Reinhold, NY
(1955) 24) E.B. Pierson, Technology of
Instrumentation, Van Nostrand, Princeton, NJ
(1957) 25) P. Delahay, Instrumental Analysis,
Macmillan, NY (1957) 26) H.H. Willard et al,
Instrumental Methods of Analysis, Van Nostrand,
NY (1958)
of reinforcement and cancellation depending on how much the two light beams are out-of-phase (Ref 15).

*Interferometers* are instruments for measuring displacement (distance) by utilizing light interference phenomena. We will illustrate the operation of interferometers by describing a method of measuring the velocity of a specimen (with a reflecting surface), propelled by a detonating explosive charge (free-surface velocity), by using a *Laser Interferometer*.

A laser interferometer is shown schematically in Fig 1. The parameter measured is the free surface velocity of the specimen material. The principle of operation is as follows. Light from the single frequency gas laser is focused on the surface of the target by means of a lens L1. The reflected light is recollimated by L2, and then split by a beam splitter B1. Half the light traverses the delay leg and is recombined with the undelayed half at beam splitter B2. The photomultiplier then records a signal whose brightness depends on the relative phases of the two beams. Since the delay leg is fixed and the wavelength of the input light is a function of free surface velocity (Doppler shift), the number of fringes recorded at the photomultiplier is related to the free surface velocity. The relationship can be derived as follows (Ref 17). The Doppler shift is given by

\[ \Delta \lambda \text{(t)} = \frac{2 \lambda}{c} \mu \text{(t)} \]  

Fig 1. Laser Interferometer System (The velocity of the reflecting surface is related to the number of fringes observed by the photomultiplier)
where $\lambda$ is the laser wavelength, $c$ the speed of light, and $u(t)$ is the speed of the reflecting surface at time $t$. The delay lag length $\Delta \lambda$ is

$$N\lambda = c\tau$$  \hspace{1cm} (2)

where $\tau$ is the time for light to traverse the delay leg. Differentiating Eq (2) gives

$$\Delta N(t) = -\frac{c\tau}{\lambda^2} \Delta \lambda(t)$$  \hspace{1cm} (3)

and substituting for $\Delta \lambda$ in Eq (1) from Eq (3) gives

$$u(t) = \frac{\lambda}{2\tau} \Delta N(t)$$  \hspace{1cm} (4)

The number of fringes $\Delta N$ as a function of time are thus related to the reflecting surfaces velocity by a constant $\lambda/(2\tau)$.

The major advantages of the laser interferometer are high time resolution and the high surface velocity capability. The limits in rise time are limited by the capability of the photomultiplier or oscilloscope recording system. The bandwidth of each system is typically 600 MHz or a response time of $\approx 1$ nsec (Ref 13).

Many other interferometers have been described in the literature (see references). Among these are the Fresnel, Fabry & Pérot, Jamin, Rayleigh, Michelson and Mach-Zehnder interferometers. Some application of interferometers are given below.

E. Mach and his son L. Mach were the first to apply an interferometer to ballistics (Refs 3 & 4). They modified the Jamin Interferometer and used it to determine the densities of air near a projectile. A schematic arrangement of their apparatus is given in Ref 4, p 274, fig 105 and in Ref 10, p 1360. Zehnder (Ref 2) modified Mach’s apparatus and the so-called Mach-Zehnder apparatus is still used.

Bergdolt (Ref 12) used the Mach-Zehnder interferometer and a short duration light source with a rotating mirror camera to obtain interferograms of projectiles and the air flow patterns around them.

Ladenburg et al (Ref 10) described the application of the interferometer techniques to the study of faster than sound phenomena. Weimer et al (Ref 11) applied these techniques to the study of shock waves. Bennett et al (Ref 15) gave the theory of interferometric analysis and described the procedure used by them at Ballistic Research Laboratories, Aberdeen Proving Ground, to study air flow around projectiles in free flight.

Bundy et al (Ref 13) measured the velocity and pressure of gases in a rocket flame by determining the small shift in wavelengths of the flame luminosity with a Fabry-Perot interferometer.


Interior Ballistics. See Ballistics in Vol 2, p B7

Intermittent Detonator Device. An analytical & experimental investigation was conducted to determine the operating characteristics of an intermittent-combustion device employing gaseous detonative combustion. Gaseous propane & air
were used as propellants in the experimental investigation.

Results indicated that the intermittent-detonation device was capable of operating on at least two "resonant" spark-energization frequencies — approx 20 & 50 cps. Reaction thrust, noise output, and actual combustion frequency were noticeably greater at resonant conditions than at other operating points.

Reloading of the tube with fresh charge occupied most of the cycle time, which limited the operating frequency of the device. This intermittent-detonation device — which consisted of three assemblies: air manifold, fuel injector, and detonation tube — did not appear promising as a thrust producing mechanism because of its low operating frequencies.

It was recommended that the investigation of intermittent-detonative combustion be continued to determine the conditions under which resonant operation occurs. Also the effects of varying the physical characteristics of the detonation device should be determined experimentally.

Ref: L.I. Krzycy, "Performance Characteristics of an Intermittent-Detonation Device", NAVWEPS Rept 7655, US Ordnance Test Station, China Lake, Calif (June 1962)

Intermittent Illumination. See Cameras, High Speed in Vol 2, pp C13-19 & High-Speed Photography in this Vol


Internal Energy is the energy contained within a system. It is an extensive property of the system and the increase or decrease in internal energy between two states is independent of the way the change between states is brought about.

The First Law of Thermodynamics is usually written in the form $\Delta E = \dot{Q} - \dot{W}$ where $E$, the internal energy, $S$, the entropy, and $V$, the volume, are extensive properties of the system, and $P$, the pressure, and $T$, the temperature, are intensive properties of the system. Extensive properties depend on the mass of the system while intensive properties are independent of mass.

The internal energy, or rather the change in internal energy is an important variable in Hydrodynamics. It occurs in the Rankine-Hugoniot (RH) equation (Vol 4, p D531-R, Eq 8 or p D605-L, Eq 13) and as parameter in various equations of state for detonation products (eg see Vol 4, pp D285-287 or pp D294-295 or p D275).

International Committee Tests. The tests designed by the International Committee for blasting explosives are as follows:

1) Preliminary Test. Same as International 75° Test, as described under Physical Tests in Vol 1, p XVIII

2) Falling Weight (Impact) Test. Ten tests with 0.05 to 0.10g samples, using an apparatus having an anvil 1.27cm (0.5") in diameter, should show that the explosive is less sensitive than Picric Acid (PA)

3) Friction Test. A small sample of explosive rubbed in an unglazed porcelain mortar should not show greater sensitivity than PA at room temp

4) Sensitiveness to Ignition. The same tests are applied as given under Ignition Tests in this Vol. These include: Fuse Test, Red Hot Iron Basin Test, and Red Hot Iron Test

Ref: Marshall (1917), 2, p 437

International Heat Test. See Physical Tests in Vol 1, p XVIII

International Regulations. Following is a list of International Regulations and where they may be purchased:

"The International Carriage of Dangerous Goods by Road (ADR)"
"The International Regulations Concerning the Carriage of Dangerous Goods by Rail (RID)"
(approx $6 each)
From: Her Majesty’s Stationary Office, 49 High Holborn, London, WC 1, England

“International Maritime Dangerous Goods Code”, Volumes I, II, and III [approx $25 per set], Sales No IMCO 1972.9 (E)
From: Inter-Governmental Maritime Consulta-
Active Organization, 101-104 Piccadilly, London,
WIV OAE, England

"IATA Restricted Articles Regulations",
16th Edition, ($6 per copy)
From: International Air Transport Association,
PO Box 315, 1215 Geneva 15 Airport, Switzerland

"Transport of Dangerous Goods", Volumes I, II, III, and IV ($7.25 per set)
From: United Nations, Sales Section, New York,
NY 10017, OHM Newsletter, Vol III, No 4,
July 1973, Department of Transportation, Office of
the Secretary, Washington, DC 20590
Ref: G. Cohn, Expls & Pyrots 7 (1), 1974

Interrupter Burner Test. In studying the effects of
burning surface, grain consolidation, inhibitor
effectiveness, etc on the intermediate stages of
propellant combustion, it is frequently desirable to
interrupt the burning process. The Interrupter
Burner is a cylindrical combustion chamber with
a firing plug at one end and a rupture disc at
the other end. The rupture disc is designed to
rupture at some pre-selected pressure generated
by the burning propellant test sample. This
sudden release of pressure extinguishes
("interrupts") the burning of the test sample
Ref: C. deFranco, PicArsn Testing Manual No
2-3 (1951)

Interstate Commerce Commission (ICC) Regu-
lations. The transportation of explosives and other
dangerous articles in interstate commerce within
the limits and jurisdiction of the United States,
is regulated by Federal Law, Act of March 4,
1909, Chapter 321, Sect 232 and 234 (35S 1134)
as amended by the Act of March 4, 1921, Chapter
172 (41S 1444-1445)

Violations of this Act are punishable by severe
fines and imprisonment

Under authority of the Act quoted above, the
ICC has published regulations which can be ob-
tained from them by writing to Dept of Com-
merce, ICC, Washington, DC

Note: In addition to the regulations issued by
the ICC, the regulations of the Bureau of
Explosives (for rail shipments) as well as various
harbor, state, municipal regulations should also
be consulted. This complicates the situation, as
the various States, municipalities etc pass laws
that apply to the particular location, but there
are no universal laws governing the handling,
storage etc of explosives etc. For a summary of
labeling and shipping regulations for explosives,
see Sax "Dangerous Properties of Industrial

Intraplant Distances. All explosive operating
buildings and service magazines must be separated
in conformity with the so-called "intraplant
quantity-distance tables." If the hazards involved
require dividing an operating line into separate
buildings, such hazards are great enough to re-
quire the use of full intraplant distances between
buildings unless effective separate barricades
are provided, in which case the distances may be
halved

Each state has its own regulations regarding
distances. For instance, the table on p 810 of
TM 9-1904, gives the "Intraplant Quantity-
Distance Table" as prescribed by New Jersey
State laws

Intrinsic Pressure. See under Detonation (and
Explosion), Equations of State, Introduction
in Vol 4, p D269-L

Inulin (Alant Starch, Dahlin, Alantin). (C4H10O5)n
H2O, mw 990.86. White, hygroscopic powder or
hairy amorphous lumps; mp 178° (decomposes
beginning at 160°), d 1.35-1.4 at 20°/4°. Sl sol
in cold w, sol in hot w; sl sol in alc. May be ob-
tained from the bulbs of Dahlia variabilis or
other plants. On nitration, it gives an explosive
compound

Refs: 1) Bell, not found 2) Fieser & Fieser,

Inulin Trinitrate. [C4H9O2(NO3)3]n, mw
(297.14)14, N 14.14%, OB to CO2 -24.2%;
colorless, amorphous solid; mp softens at 90°
and melts at 102°, bp decomposes beginning
about 110°. When heated rapidly, it ignites
without exploding at 228-230°. First prep by
the nitration of inulin with mixed nitric-sulfuric
acid by W. deC. Crater of Hercules Powder
Co

When mixed acid contains 48.50% HNO3 and
51.20% H2SO4 is used the resulting nitrate contains
about 13.75%, while nitration with MA contains
20.8% HNO3 and 62.8% H2SO4 gives a 12.80%
product
Inulin nitrate is insol in w, eth, ethyl and methyl alc, toluene and carbon tetrachloride, but dissolves in acetone, ether & alc mixt, concd H₂SO₄, ethyl acetate etc. It does not crystallize from solutions but forms a film
Inulin nitrate is compatible with NC, nitroelast etc
It is a fairly stable explosive, having an impact sensitivity of about 30cm with a 2kg weight
Inulin nitrate was proposed as a base charge in blasting caps, as a sensitizer for ammonium nitrate in dynamites and as an ingredient of smokeless powders
W. deC. Crater gives the following examples of explosives using inulin nitrate primarily as a sensitizer:
Ammonium nitrate dynamite. NH₄NO₃ 65,
Inulin nitrate (13.75%N) 20, NaN₃ 11, Pulp (such as wood pulp) 4. This dynamite has a rate of detonation of 2310 m/sec, weight per 1¼ stick - 120g; cartridge cont 378 per 100 lb; sensitiveness (gap test) 4"; weight strength 53.5%. It flows and packs easily, is not dusty, does not freeze and does not cause headaches
Base charge. A standard blasting cap shell may be charged with 16g of nitroinulin (13.75%N) pressed under 7840 psi, and provided with a primer charge of 0.3g of 80/20 fulminate-K chlorate mixt. Such a cap was found to be satisfactory for the detonation of explosives
Ref: 1) W. deC. Crater, USP 1992123 (1933) & CA 27, 5190 (1933) 2) No further refs found in CA 1947-1971
Iodamide. Nl₃; see Nitrogen Iodide under Iodides in this Vol

IODATED AND NITRATED PARAFFINS
Some paraffin compounds containing both I and NO₂ groups are explosive, for example:
Iododinitromethane. CHI(NO₂)₂; mw 231.96, N 12.07%
Its potassium salt, KCl(NO₂)₂, mp (explodes violently at 154°); orange-yellow plates. It was first prepd by Villiers (Ref 2). Gotts & Hunter (Ref 4) dissolved 10g of K ditnitromethane in 300ml water containing 4g of KOH (one equivalent), cooled the solution and slowly added powdered iodine (18g) to it with shaking. The solution was evaporated to a small bulk (50ml), which caused the separation of about 163g of crystals. These were dissolved in hot water and then cooled—the pure salt crystallized out
The salt is nearly insol in cold water; sol in hot w. It is a violent explosive
Its silver salt, AgCl(NO₂)₂, mp (explodes at 109-110°). Pale yellow glistening leaflets; may be prepd by the action of AgNO₃ on an aqueous solution of potassium iododinitromethane. Insol in w (Ref 4)
Diiododinitromethane. I₂C(NO₂)₂; dark-colored, pungent smelling oil. Was obtained on acidifying an ice cold solution of potassium iododinitromethane. It could not be properly examined because it decomposed immediately with the evolution of gas and iodine (Ref 4)
Iodotrinitromethane (Sometimes erroneously called Iodopicrin). Cl(NO₂)₂, mw 276.95, N 15.17%; OB to CO₂ & I₂ +23.1%, mp 55-58°, bp 48° at 13mm. Bright yellow leaflets; may be prepd by the action of an ethereal solution of iodine on the silver salt of nitroform
It decomp on storage (Ref 1). Its hydrolysis const is 4 x 10⁻⁷ (Ref 5). Insol in w; sol in hot bz, ligroin and alc. No mention of any explosive properties (Refs 1, 3 & 5)
No further mention of explosive props of iodonitromethanes were found in the CA Formula Indices 1920-1971

Some interesting work on the decomposition kinetics of IC(NO₃)₃ has been reported. In the gas phase this compd decomposes with homogeneous first order kinetics over the temp range of 100-160° (Ref 6). The activation energy obtained (E = 34.4 kcal/mole, log Z = 15.25) suggests that the primary step is the rupture of the C–N bond followed by a radical (non-chain) reaction (Ref 7). Addn of a large excess of NO, one of the decomn products, increased reaction rate 20-30%, addn of a large excess of NO₂, another decomn product, lowered the reaction rate 10%. Addn of I₂, also a decomn product, had no effect on rate (Ref 7)
The strength of the C–I bond is claimed to be 34.8 kcal/mole (Ref 7)
Mass spectrometric studies fully confirmed the previous manometric studies quoted above (Ref 8). In the liq phase the pyrolysis of IC(NO₃)₃ at 80-140° has the same kinetic as in the gas phase. Replacing a NO₂ group with
F stabilizes the molecule. Thus the kinetic parameters for the decomp of FIC\(\text{NO}_2\)_2 are \(E = 39.7\) kcal/mole & logZ = 15.7 over the range 170-214°C (Refs 9 & 10).

Mass spectra of positive and negative ions of IC\(\text{NO}_2\)_3 have been obtained (Ref 11).


**Iodates and Periodates.** Many inorganic iodates & periodates are known. Although many iodates & periodates are thermally unstable only the ammonium salts appear to be explosive. However, iodates and periodates are powerful oxidizing agents and can react vigorously with many reducing agents (Ref 5).

Varhelhi & Kekedy studied the thermal decomposition of iodates & periodates by thermogravimetric methods (Ref 2). They found that there is no similarity in the decomposition of univalent iodates, but most divalent iodates decompose similarly since their decomp temps are similar. The thermal stability (ie their resistance to decomposition into iodates) increases as follows: of the alkali periodates Li<Na<K<Rb(Cs. The thermal stability of alkaline earth periodates increases in the order Ca<Sr<Ba.

**Ammonium Iodate.** NH₄IO₃; mw 192.96, N 7.25%, rhombic or monoclinic crystals, d 3.31; sol in w; decomposes on heating.

Solymosi et al (Ref 3) studied the thermal decomp & explosion of Ammonium Iodate. They found that the decomposition of ammonium iodate starts at 140°C. The reaction yielded a solid reaction product which according to analysis, consisted of iodine pentoxide. In the gas phase, iodine, oxygen, nitrogen and nitrogen oxide were found. The reaction occurred without time lag. The course of the curve and the kinetics were independent of whether the reaction was followed by the pressure of the gases formed or by the change in weight of the sample. The decomposition was described, by the unimolecular decay equation from the beginning of the reaction to 85% completion. The value of the activation energy was 66.4 kcal.

Explosion of ammonium iodate took place at a temperature of 185°C, i.e., very much higher than for the other halates (81°C for NH₄BrO₃ & 90° for NH₄ClO₃). The explosion was preceded by a readily measurable and gradually accelerating decomposition. Contrary to the two other halates, self-heating was not observed before the explosion. The explosion was accompanied by a violet colored flash and produced a temperature increase of only 10°C in the reaction zone. The activation energy of the process leading to explosion was 38.4 kcal.

From these measurements it appears that the stability of the compounds does not follow the expected order of chlorate < bromate < iodate. This behavior, however, cannot be regarded to be a characteristic property of ammonium halogenates only, because according to our experiments the same order of stability has been found in case of potassium halates. It is fairly difficult to propose a reliable reaction mechanism merely from kinetic data.

Solymosi et al suggest that the rate-controlling step in the thermal explosion of NH₄IO₃ is the rupture of an I—O bond.

They also make the statement (no data or refs) that ammonium halates (-ClO₃, -BrO₃ & -IO₃) are impact & friction sensitive. This is in line with the thermochemical calculations and conclusions therefrom made by Shidlovskii (Ref 4) for both NH₄IO₃ & NH₄IO₄.

**Ammonium Periodate.** NH₄IO₄; mw 208.96, N 6.7%; tetragonalless crystals, d 3.06, sol in w. Decomposes on heating.

Solymosi et al (Ref 2) imply that NH₄IO₄ is more stable than NH₄IO₃. However, a serious explosion occurred when NH₄IO₄ was scooped from one container to another (Ref 1).
Hydrazinium Iodate. \( \text{N}_2\text{H}_4\cdot\text{HIO}_3 \) has been postulated to exist only in cold solutions (Ref 4). In general, hydrazinium salts of oxidizing acids are much less stable than the corresponding ammonium salts


IODIDES

Most metals form iodies eg KI, PbI\(_2\), BI\(_3\), SnI\(_4\) etc. The iodies of the transition elements are usually in their lowest oxidation state eg FeI\(_2\), CoI\(_2\), MnI\(_2\), CrI\(_2\) etc. All of these metal iodies are stable and nonexplosive

Iodine forms numerous interhalogen compds, eg IBr, IC\(_2\), IF\(_4\) etc but these are not really iodies since the iodine in these compds (if they are not covalent) plays the part of a "metal" ion

The only explosive iode is Nitrogen Iodide which is described below

Nitrogen Iodide (also known as Nitrogen Triiodide). \( \text{NI}_3 \); mw 394.77, N 3.55%; black cryst. explodes on heating, sublimes in vac, insol in w, decom in hot water (Ref 15). Prepd by treating I\(_2\) with concd aq NH\(_3\), probably via

\[ 3\text{I}_2 + \text{NH}_3 = 3\text{HI} + \text{NI}_3 \]

By passing NH\(_3\) over I\(_2\) a series of ammoniated NI\(_3\) compds can be prepd. These lose NH\(_3\) under vac except that the last NH\(_3\) cannot be removed in this manner and the compd thus formed is NI\(_3\).NH\(_3\) (Refs 1, 2 & 3)

Suspensions of NI\(_3\) in NH\(_3\) at -78° give dark ppts with solns contg low concentrations of NH\(_2\)_\(_2^+\). The ppt w/ AgNH\(_2\) is Ag

\[ \text{Ag} \cdot \text{NH}_2 \cdot \text{NI}_3 \]

It is explosive when dry, but inert to excess Ag NH\(_2\) & stable in liq NH\(_3\) or in H\(_2\)O at 0°. At room temp the H\(_2\)O soln decomposes (accelerated by light) to Ag, AgI, NH\(_3\), & N (Ref 13)

Nitrogen Iodide is an exceedingly sensitive explosive when dry. It explodes upon the slightest touch and also upon very mild heating such as in a warm air stream. It must be stored wet, preferably under ether (Ref 15). Because of its phenomenal sensitivity NI\(_3\) is not a practical explosive, but by the same token it has been studied rather extensively to determine why it is so sensitive. Since these studies contribute to the understanding of explosion phenomena they are described below

Bowden & Yoffe (Ref 12, p 73) state:

"Nitrogen iodide (NI\(_3\), NH\(_3\)) is a very unstable material and will explode under the action of very weak shocks. Even at liquid air temperatures, initiation occurs at an energy of 0.60g/cm (Ref 4). The stability of nitrogen iodide under normal conditions depends to a great extent on the presence of ammonia which retards the decomposition of the pure NI\(_3\) (Ref 8).

Meekerämer (Ref 9) has found that the sensitivity to impact is also reduced in an ammonia atmosphere. Presumably the ammonia is rapidly adsorbed on the freshly exposed surface of the crystals of nitrogen iodide and prevents the decomposition from spreading"

Nitrogen Iodide & NI\(_3\).NH\(_3\) can also be decomposed and ignited by light (Refs 10 & 11). Light absorption is strong in the visible and weaker (but constant) in the IR (Ref 12, p 104)

Meekerämer (Ref 9) irradiated nitrogen iodide in low-pressure atmospheres of air or ammonia at various temperatures and determined the quantum yield of the decomposition. He finds that this value is not constant but increases to a limiting value as the irradiation is continued. This limiting value depends on the ambient temperature. The results suggest that the decomposition is mainly thermal due to the conversion of the light energy into heat. There is a small (5 to 10%) photochemical effect particularly when the nitrogen iodide is irradiated with blue or red light

When the intensity of the light from the flash-lamp is sufficiently high nitrogen iodide explodes. The amount of light energy required for ignition depends on the temperature (see Figure). Extrapolation of the curve back to zero light energy gives a value of 80°C which should be compared with 50° which is the 'dark' ignition temperature
Figure. Relation between the light energy required for ignition (joule) and temperature 
\(^\circ\text{C}\) for nitrogen iodide (Ref 9)

Again quoting Bowden & Yoffe (Ref 8, pp 105-106): "The time at which the explosion occurs relative to the duration and the intensity of the light flash has been determined by means of spark photography. Meerkämper (Ref 9) finds that explosion occurs just before or at the time of maximum intensity of the light flash. This means that only a fraction of the total light energy is used for ignition. This result is similar to that observed for Silver Azide. The light energy, \(E\), passing into the surface layer which is used for ignition is such that

\[
E_t = \text{const} = 1.35 \times 10^{-6}\text{J/sec/cm}^2
\]

where \(t\) is the time between the beginning of the light flash and the beginning of the explosion. For a light flash lasting 200 \(\mu\text{sec}\), the energy \(E = 0.023\ \text{J/cm}^2\), and the time \(t = 50\ \mu\text{sec}\). This time is close to that required for the lamp to reach its maximum intensity.

Nitrogen iodide also explodes when it is irradiated with light from a spark discharge in air. A glass plate placed between the spark and the nitrogen iodide sample hinders the passage of the shock wave from the spark. The duration of the spark (4 \(\mu\text{F}, 12\text{kV}\)) is ca 200 \(\mu\text{sec}\).

The light-time curve for such a spark rises very rapidly and explosion begins 6 \(\mu\text{sec}\) after the beginning of the light from the spark. When the discharge is from 0.4 \(\mu\text{F}\) at 38 kV the time from the beginning of the spark to explosion is 4 \(\mu\text{sec}\).

If a tube filled with Xenon is used instead of the spark in air the explosion begins 3.7 \(\mu\text{sec}\) after the first appearance of light. This value of 3.7 \(\mu\text{sec}\) means that ignition takes place ca 0.5 \(\mu\text{sec}\) after the flash has reached its maximum intensity. Again it is clear that only a small part of the incident light is used for igniting the nitrogen iodide.

The results described above, in particular the relation between ignition energy and temperature, suggest a thermal mechanism for the initiation. The light will not be absorbed in a uniform manner in the surface layers. However, a rough estimate of the thickness of the surface layer which is heated to the ignition temperature may be obtained by assuming that the whole of the surface layer is heated to a uniform temperature. If \(d\) is the thickness of this surface layer, then

\[
d = \frac{E}{A\Delta T_\text{B}}
\]

where \(E\) is the energy absorbed, \(A\) the surface area of the nitrogen iodide on which the light falls, \(\Delta T\) the temperature difference, \(g\) the density of the nitrogen iodide and \(C\) the specific heat. Experimentally it is found that \(E/A = 8 \times 10^{-2}\ \text{J/cm}^2\)

Since ignition occurs in less than 50 \(\mu\text{sec}\) for a flash of 200 \(\mu\text{sec}\) duration, it is assumed that only the energy \(E/3A\) is used for ignition

\[
\frac{E}{3A} = 2.7 \times 10^{-2}\ \text{J/cm}^2
\]

\[
= 6.4 \times 10^{-3}\ \text{cal/cm}^2
\]

For an ignition temperature of 80\(^\circ\text{C}\), \(\Delta T = 60\(^\circ\text{C}\), \(g = 3.5\ \text{g/cm}^3\), and \(C = 0.12\ \text{cal/g}^\circ\text{C}\),

\[
d = 2.6 \times 10^{-4}\ \text{cm}
\]

or if the ignition temperature is taken as 55\(^\circ\text{C}\),

\[
d = 4.4 \times 10^{-4}\ \text{cm}
\]

This value for the thickness of the hot layer is in agreement with the size of hot spot necessary for initiation by shock.

In view of the assumption made, however, this agreement should not be regarded as providing positive evidence for the thermal theory of initiation.

Nitrogen iodide cannot be ignited under concentrated ammonia solutions even with a very strong flash. Under pure water local explosions

---

**Note:** The text contains a series of equations and figures which are integral to understanding the context and the scientific explanation provided. The natural text is a simplified representation focusing on key points. For a full understanding, the original scientific context, figures, and equations should be reviewed.
do occur, but the explosion is not propagated to the surrounding material. These observations may be explained in terms of the thermal theory. Water prevents the propagation of explosion simply by abstraction of heat. In the case of ammonia solutions, the inhibition is probably due both to cooling and to reaction with hydrogen iodide. The removal of hydrogen iodide prevents the secondary reactions with nitrogen iodide from taking place and so reduces the heat liberated during reaction."

Poole has ignited Nitrogen Iodide by a single \( \alpha \)-particle from a radium source (Refs 5 & 14)

Bowden & Yoffe have suggested (Ref 12, pp 127-128 & Ref 14) that the abnormal sensitivity of \( \text{NI}_3 \) or \( \text{NI}_3 \cdot \text{NH}_3 \) is due to its being inherently unstable, and that the very slight external stimuli (slight touch, warm air, irradiation etc) primarily act to remove adsorbed \( \text{NH}_3 \) from the iodide surface. Apparently this adsorbed \( \text{NH}_3 \) stabilizes the iodide and its removal causes the iodide to decompose spontaneously.

Again we quote Bowden & Yoffe: "Nitrogen iodide will explode when heated in air at 50°C. The equation representing the decomposition is (Ref 4)

\[
8\text{NH}_3 \cdot \text{NI}_3 = 5\text{N}_2 + 6\text{NH}_4 \cdot \text{I} + 9\text{I}_2
\]

Garner & Latchem (Ref 6) found that when nitrogen iodide (\( \text{NI}_3 \cdot \text{NH}_3 \)) is dried over a layer of \( \text{P}_2\text{O}_5 \) at a pressure of \( 2 \times 10^{-3} \text{ cm} \) Hg, thermal decomposition could be studied at temperatures below 0°C. However, when the pressure fell below \( 2 \times 10^{-3} \text{ cm} \) Hg, explosion of the nitrogen iodide took place at temperatures as low as -11°C. At low pressures Meldrum (Refs 7 & 9) has shown that the equation representing the decomposition is different from that given above,

\[
2\text{NH}_3 \cdot \text{NI}_3 = 2\text{NH}_3 + \text{N}_2 + 3\text{I}_2
\]

Ammonia and water vapor retard the decomposition of the nitrogen iodide, a pressure of \( 10^{-3} \text{ cm} \) Hg of ammonia being sufficient to stop the reaction.

The main effect of gas pressure is to slow down the rate of diffusion of ammonia away from the decomposing surface and so increase the stationary concentration of ammonia at this surface

The activation energy for Reaction (2) is 18-19 kcal/mole. It has been assumed that this is the energy required to remove ammonia from the surface of the solid \( \text{NH}_3 \cdot \text{NI}_3 \). After removal of the ammonia the \( \text{NI}_3 \) that is left is unstable and decomposes with the liberation of heat. The explosion then grows from this hot region; the reactions proposed being either

\[
\text{NI}_3 + \text{NH}_3 \cdot \text{NI}_3 = \text{NH}_3 + \text{N}_2 + 2\text{I}_2 + 21 + 33.1 \text{ kcal}
\]

or

\[
\text{NI}_3 + \text{NH}_3 \cdot \text{NI}_3 = \text{NH}_3 + \text{N}_2 + 3\text{I}_2 + 68.1 \text{ kcal}
\]

If iodine atoms are produced as in Equation (3) these could react further with ammonia to give ammonium iodide according to the equation (see Equation (1))

\[
8\text{NH}_3 + 6\text{I} = 6\text{NH}_4 \cdot \text{I} + \text{N}_2
\]

It is clear from the results described above that the presence of ammonia can under certain conditions prevent the explosion of nitrogen iodide when the decomposition is brought about by heat, light, shock, or ionizing radiation.

The retarding action of ammonia has also been observed in the decomposition of a number of ammonium salts, e.g. ammonium perchlorate (\( \text{NH}_4 \text{ClO}_4 \)) and ammonium nitrate (\( \text{NH}_4 \text{NO}_3 \)) (Ref 10).

iodides in Smoke Compositions. DeMent claims mixts of elements or cmpds consisting of or contng halogenes, halide-forming elements or groups, an oxidant & a combustible material, such as iron, produce smokes of a large variety when ignited. More than 370 formulations are described. Suggested uses include military screening & signaling
Ref: J. deMent, USP 2995526 (1961) & CA 65, 25100 (1961)

iodine. Iodine added in quantities 1 to 5% to a liquid organic peroxide eliminates the danger of explosion and fires in its use, storage and shipment. Before use of the peroxide, the iodine is removed by an oxidizing agent or other suitable means
Ref: H.C. Stevens, USP 2415971 (1947) & CA 41, 3114 (1947)

iodine Azide. See under Azides, Inorganic in Vol 1, pp A542-543

iodine Pentfluoride. The controlled reaction of IF₅ with dimethyl sulfoxide, Me₂SO, was found to be quite violent. Although small scale tests in the cold (0°) proceeded without incident, delayed violent explosions occurred upon scale-up or upon allowing the temp to rise to 10-15°

iodoazide. See under Azides, Inorganic in Vol 1, pp A542-543

iodosooazidobenzene. See "Benzene and Derivatives" under Azido Halogen Derivatives of Benzene in Vol 2, p B44-R

iodoxyazidobenzene. Ibid

iodoazidothane. N₃CH₃CH₂I; no explosive props mentioned in Beil 1, (33)

iodoazoinide. N₃·2NH₃. See Iodides in this Vol. This name is used by Antelman (JChem-Educ 30, 134 (1953) & CA 47, 5741 (1953)) in describing some spectacular lecture demonstrations of incendiary & explosive reactions

iodobenzene and derivatives

iodobenzene or Phenyl Iodide. C₆H₅I; mw 204.02; pale-yel liq, fr p -31.4°, d 1.832 at 20°, refractive index 1.621 at 20°. It is almost insol in water, but sol in alc., eth & chlf. Iodobenzene is readily prepd by reaction of iodine with benzene in the presence of an oxidizing agent, or from benzenediazonium sulfate & potassium iodide
Refs: 1) Beil 5, 215, (18) [167] & {571} 2) Kirk & Othmer II (1966), 864

1-Azido-2-iodobenzene, C₆H₄N₃I; mw 245.03, N 17.15%; strong smelling oil, bp 90-91° at 0.9 mm Hg, d 1.893 at 25°, refractive index 1.6631 at 25°; prepd by diazotizing aniline with NaN₃. No expl props are reported. The 1-Azido-3-iodo- and 1-Azido-4-iodo-derivs are also known (Ref 2)
Refs: 1) Beil 5, (142) 2) Beil 5, 278 & (143) Mononitroiodobenzene, C₅H₄NO₂I; mw 249.02, N 5.63%. The following isomers are known:
1-Nitro-2-iodobenzene, citron-yel ndls, mp 54°, bp 288-89° (sublimes), d 1.9186 at 75° (Ref 1)
1-Nitro-3-iodobenzene, monoclinic crysts, mp 36-38°, bp 153° at 14 mm Hg, d 1.9513 at 75° (Ref 2)
1-Nitro-4-iodobenzene, dk-yel ndls (from alc), mp 172°, bp 287° at 726 mm Hg, explodes on heating in a tube at 605° (Ref 3)
Refs: 1) Beil 5, 252, (133), [190] & {621} 2) Beil 5, 253, (133), [91] & {622} 3) Beil 5, 253, (133), [91] & {623} Dinitroiodobenzene, (O₂N)₂C₆H₄I; mw 294.03, N 9.53%. Six isomers are known:
1,2-Dinitro-3-iodobenzene, lt-yel ndls (from alc), mp 138°, distils without decompn (Ref 1)
1,2-Dinitro-4-iodobenzene, yel pilits (from alc), mp 74.5° (Ref 2)
1,3-Dinitro-2-iodobenzene, orn-yel tablets (from alc), mp 113.7° (Ref 3)
1,3-Dinitro-4-iodobenzene, yel ndls (from alk or benzene), mp 88-90° (Ref 4)
1,3-Dinitro-5-iodobenzene, golden-yel plates (from 60% alc), mp 99° (Ref 5)
1,4-Dinitro-2-iodobenzene, lt-yel prisms (from alc & eth) or almost colorless ndls (from alc), mp 117.4° (Ref 6)
Other props & methods of prepn are given in the Refs

1,3,5-Trinitro-2-iodobenzene or Picryl Iodide, 
(O_2N)_3C_6H_4I; mw 339.02, N 12.39%, golden-
yel tetragonal ndls, mp 164-65º, d 2.285 at 22.5º; was first prep'd from 2-chloro-1,3,5-
trinitrobenzene & KI in the presence of alc. Other props & methods of prepn are given in the Refs. No expl props are reported
Ref: Beil 5, 275, {207} & {647} Iodochlorobenzene HCl salt or p-chloro
Phenyliodide hydrochloride, (C_6H_4Cl)_2HCl, yellow needles (from chlf); sol in chlf, glc 
HAc & benzene; sl sol in eth & petr eth; 
decomp slowly in stoppered dark vessel – rapidly in sunlight. Decomp suddenly 
between 110º & 136º. Prepd by reacting 
phenyl iodide (in chlf) with chlorine 
C_6H_5IClO_4 yel ppt which turns orange-red 
on drying explodes at 66-67º (no prep given)  
Ref: Beil 5, 218

1-Iodo-2-Ethoxy-3 Butene, CH_2ICOOC_2H_5CH_3; 
mw 225.96. In the course of prepn by Petrov's 
method (Ref 1), the reaction can violently explode without 
certain undefined conditions. Ref 2 relates 
that several runs on a 0.16M scale were completed 
without incident. However, during a 1M run, the 
reaction exploded while ethanol was being dis-
tilled under sl vac at 35º. Controlled attempts to 
determine the cause were unsuccessful
Ref: 1) A.A. Petrov, ZhurObshchelKhim 
(JGenChem) 19, 1046–62 (1949) & CA 44, 
1003 (1950) 2) J. Trent & P.G. Gassman, 
C&EN 44, No 43, 7 (1966)

IODOMETHANE AND DERIVATIVES

IODOMETHANE or Methyleneiodide, CH_2I; mw 267.87; colorl liq or leaflets at 0º; mp 5–6º, bp 
180º (decomp), d 3.325; sol in alc or eth; sl 
sol in w. Prepd by reacting methylene chloride 
with NaI. Not explosive by itself but if mixed 
with K or KNa alloys it explodes on shock 
Ref: 1) Beil 7, 71, (18), [37] & [38] 
2) CondChemDict (1971), p 573

IODOFORM or Triodomethane, CHI_3; mw 393.78; yellow leaflets, mp 115º, bp 260º (decomp), 
d 4.08; sol in w, alc, eth & chlf. Prepd by 
heating acetone (or methanol) with iodine in 
the presence of alkali. There is considerable 
fusion in the literature whether iodoform is an 
explosive. Beilstein (Ref 1), Sax (Ref 2), and CA 
1947–1971 do not mention any explosive prop-
erties. Ref 3 states that iodoform decomps 
violely at 400ºF. Friction makes a mixt of 
CHI_3+AgNO_3 decomp explosively (Ref 1) 
Ref: 1) Beil 7, 18, (19), [38], {102} & 
97> 2) Sax (1968), 840-R 3) CondChem-
Dict (1971), 473-L

IODONITROMETHANE, CH_2NO_2; mw 186.9, N 7.50%; 
prepd by reacting methylidide in eth with AgNO_2 
& a trace of iodine. Its Na salt, NaCH(NO_2)_2, 
white powder, explodes on heating 
Ref: Beil 1, 79 & {115}

IODONITROMETHANE, CH(NO_2)_2, mw 231.9, N 
12.1%. Its K-salt, prepd by slowly adding iodine 
to K-dinitromethane in eq KOH, orange-yellow 
platelets (from w); explodes violently at 154º. 
The Ag-salt, pale yellow leaflets, insol in w, expl-
odes at 104–106º
Ref: Beil 1, 79 & [45]

IODONITROMETHANE or Iodopicrin, CI(NO_2)_3; 
mw 277.0, N 15.2%; no explosive props mentioned 
for it or its salts
Ref: Beil 1, 79 & [47]

2-Iodo-2-nitro-1,3-indandione, 
C_6H_4CO_2CI; CH(NO_2)_2; mw 317.05, N 4.42%, 
OB –80.7º, mp 128º. Vanag & Lipman (Ref 
2) warn that an explosion might take place when 
iodonitroindandione is prepd by heating iodine 
with the silver salt of 2-nitro-1,3-indandione in 
a sealed tube at 130º. The silver salt of 2-nitro-
1,3-indandione decomposes explosively at 239º 
Ref: 1) Beil 7, 694, (375) & [632] (describes 
its parent compound 1,3-indandione) 2) G.Ya.
Iodonitrophenols

The 2-Iodo-4,6-dinitrophenol, C₆H₃N₂O₄I; mw 310.02, N 9.04%; mp 106–107⁰ has been prep (Ref 8). No explosive props are mentioned. Other iodo-dinitrophenol isomers are described in Bell; none is explosive:
4-Iodo-2,3-dinitrophenol (Ref 1)
5-Iodo-2,4-dinitrophenol (Ref 2)
6-Iodo-2,4-dinitrophenol (Ref 3)
4-Iodo-2,5-dinitrophenol (Ref 4)
3-Iodo-2,6-dinitrophenol (Ref 5)

The 2-Iodo-2,4,6-trinitrophenol or 2-Iodopicric Acid, C₆H₃N₂O₇I; mw 355.02, N 11.84%; is also known; mp 175⁰. It is prep by nitrating iodosobenzene with mixed acid. No explosive props are mentioned for it or its salts (Refs 6 & 7)

Refs: 1) Beil 6, 263 2) Beil 6, [252]
3) Beil 6, 263, (129) & [252] 4) Beil 6, 263 & [252]
5) Beil 6, 264 & [252]
8) D.B. Murphy et al, JACS 75, 4289 (1953) & CA 48, 12075 (1954)

IODOSO COMPOUNDS

These are substances of the type ArI0. Apparently only aryl iodoso compds are known since no alkyl iodoso compds were found in the literature. Vibration spectra of several ArI0 compds are given in Ref 2. Individual explosive iodoso compds are described below. No other refs to explosive iodoso compds were found in CA 1947–1971

Iodosobenzene, C₆H₄IO; mw 220.02; yell, amorphous pd; sl sol in hot water & alc; almost insol in eth, acet, benzene, petr eth & chl. It explodes on heating to 210⁰, and explodes in the presence of conc nitric acid. Iodosobenzene was prep by oxidation of iodosobenzene with ozone

Refs: 1) Beil 5, 217, (118), [166] & (575)
2) C. Furlani & G. Sartori, AnnChim (Rome) 47, 124 (1957) & CA 51, 8533 (1957)

Azidiodiosobenzene, N₃C₆H₄IO; mw 261.03, N 16.10%. Three isomers are known:

1-Azido-2-iodosobenzene, known in the form of its salts, some of which are very unstable (Ref 1)
1-Azido-3-iodosobenzene, yell amorphous mass, explodes at 125⁰. Its Formate explodes at 78⁰ (Ref 2)
1-Azido-4-iodosobenzene, explodes on heating to 130⁰ or in the presence of conc nitric or sulfuric acid. Its Chromate salt explodes by friction or by heating to 71⁰. Its Formate salt explodes at 85⁰ (Ref 2)

Other props & methods of prepn are given in the Refs. It is claimed as a fusehead constituent by Ingram (Ref 3)

Refs: 1) Beil 5, (142) 2) Beil 5, (143)
3) L.K. Ingram, USP 2241406 (1941) & CA 35, 5318 (1941)

Mononitriiodiosobenzene, O₂NC₆H₄IO; mw 265.02, N 5.17%. Three isomers are known: 1-Nitro-2-iodosobenzene, orn colored prisms (from chl), mp — decomp at 100⁰. Its salts are unstable on heating (Ref 1)
1-Nitro-3-iodosobenzene, yell solid, mp, decomp on heating at low temp. Its chromate salt, O₂NC₆H₄ICh₁₄, orn pd, explodes on heating to 95⁰ (Ref 2)
1-Nitro-4-iodosobenzene, solid, mp — explodes at 82–83⁰ (Ref 3). Its salts are unstable on heating

Other props & methods of prepn are given in the Refs

Refs: 1) Beil 5, 252 2) Beil 5, 253 & (622)
3) Beil 5, 254, [191] & (624) 4) L.K. Ingram, USP 2241406 (1941) & CA 35, 5318 (1941)

Dinitriiodiosobenzene, (O₂N)₂C₆H₄IO, and Trinitriiodiosobenzene, (O₂N)₃C₆H₂IO, deriva not found in Bell

1-Iodoso-2-chlorobenzene, ClC₆H₄IO; mw 254.5, white-yellow powder, v sl sol in eth, chl, benz or petr eth; somewhat sol in w. Explodes mildly at 83–85⁰. Prep by passing chlorine into o-chloriodobenzene and then treating with dil NaOH. The 3-chloro isomer, bright yellow needles, decomp at 100⁰ & the 4-chloro isomer, bright yellow amorphous mass, decomp at 116–117⁰

Ref: Beil 5, 220 & 221
1-Iodo-3-iodobenzene, IC₆H₄IO; mw 345.9, bright yellow amorphous powder; insol in usual org solvents; explodes mildly at 124° on rapid heating; on slow heating it melts (decomp) at 207°. Prepd by passing chlorine thru a chl soln of m-diiodobenzene
Ref: Beil 5, 225–226

1,3-Di-iodosobenzene, OIC₆H₄IO; mw 361.9, bright yellow amorphous powder, almost insol in all solvents; explodes around 108°. Prepd by passing chlorine thru a glacial HAc soln of m-diiodobenzene
Ref: Beil 5, 226

m-Iodosotoluene, CH₃C₆H₄IO; mw 234.0, amorphous yellow powder, sol in cold glacial HAc. Decomp at 180–85°; explodes at 206–07°. Prepd by passing chlorine thru a soln of m-iodosotoluene in chl. The ortho & para isomers decompose with gas evolution at 170–75° & 175–78°, respectively, but no mention is made of their exploding
Ref: Beil 5, 310, 311 & 313

4-Iodo-3-nitrotoluene, O₂NC₆H₄(CH₃)IO; mw 280.12, N 5.03%, red powder; explodes mildly at 129°. Prepd by passing chlorine thru a chl of 4-iodo-2-nitrotoluene and neutralizing with dil NaOH. The Chromate salt, [O₂NC₆H₅(CH₃)I- (OH)]₂CrO₄, red-orange crystals, explodes at 94°. The Formic acid salt, O₂NC₆H₅(CH₃)I(H₂O)₂, orange-colored powder, explodes mildly at 72°. The Acetic Acid salt, O₂NC₆H₅(CH₃)I(C₂H₃O₂)₂, bright yellow needles, explodes at 200°
Ref: Beil 5, 337

2-Iodo-4-nitrotoluene, O NC H (CH₃)IO, insol powder; explodes at 180–81°. Prepd as above from 2-iodo-4-nitrotoluene
Ref: Beil 5, 338

4-Iodo-1-propylbenzene, CH₃CH₂CH₂C₆H₄IO; mw 262.0; decomp on storage; explodes at 105°. Prepd by passing chlorine thru a soln of 4-iodo-1-propylbenzene in chl & petr eth and then treating with a dil NaOH soln. Its Perchlorate salt, C₉H₇I₁(OH)ClO₄ explodes at 73° and also explodes spontaneously on storage. The Chromate salt also explodes spontaneously
Ref: Beil 5, 392

Iodotoluene and Derivatives
Iodotoluene, CH₃C₆H₄I, is a stable non-explosive compd; so are its mononitro derivatives. Higher nitro derivs or azido derivs were not found in Beil
Ref: Beil 5, 310 & 337

IODOXY COMPOUNDS
As in the case of iodoso compds, the only iodoxy compds reported in the literature are ArIO₂ & not AlkIO₂. Vibration spectra of several ArO₂ compds are given in Ref 2. Individual explosive iodoxy compds are described below. No other refs to explosive iodoxy compds were found in CA 1947–1971

Iodoxybenzene, C₆H₅IO₂; mw 236.02; ndls (from benz), mp – explodes at 236–37°; almost insol in chl, acet & benz; v sl sol in petr eth; sl sol in hot water & glacial acetic acid. It can be prepd by oxidation of iodobenzene with various agents. Iodoxybenzene explodes in the presence of concd H₂SO₄ or phosphorus pentachloride. The Perchlorate salt is expl

Azoiodoxybenzene, N₃C₆H₄IO₂; mw 277.03, N 15.17%. Three isomers are known:
1-Azido-3-iodoxybenzene, brownish ndls (from glacial acetic acid), mp – explodes violently on heating to 157° or by friction (Ref 1)
2-Azido-3-iodoxybenzene, brownish ndls, mp – explodes on heating to 175–80°; sol in water & glacial acetic acid (Ref 2)
3-Azido-4-iodoxybenzene, brn crysts, mp – explodes on heating to 170°, or in contact with concd H₂SO₄ (Ref 3). The Chromate of the 1,4 compd is a very dangerous, deep-red material that explodes at 71° or upon being rubbed lightly. Its Formate (in the form of plates) explodes at 85°. Its Nitrate melts at 102° with decomposition. The iodoxyazidobenzenes are claimed by Ingram (Ref 4) to be useful as fusehead ingredients
Other props & methods of prepn are given in the Refs

Mononitroiodoxybenzene, O₂NOC₆H₄IO₂; mw 281.02, N 4.99%. Three isomers are known:
1-Nitro-2-iodoxybenzene, tablets (from glacial acetic acid), mp — explodes violently at 210°; almost insol in eth, pet eth & benz; sl sol in glacial acet. water & alc (Ref 1)
1-Nitro-3-iodoxybenzene, plates (from w), mp — explodes on heating to 215—18° (Ref 2)
1-Nitro-4-iodoxybenzene, wh plates, mp — explodes at 212—13°; v sl sol in glacial acet ac (Ref 3)
Methods of prep & other props are given in the Refs
Refs: 1) Beil 5, 253 & {622} 2) Beil 5, 253 & {223} 3) Beil 5, 254, [191] & {624}

Dinitroiodoxybenzene, (O₂N)₂C₆H₄IO₂; mw 326.02, N 8.59%. Two isomers are found in Beil

1,3-Dinitro-4-iodoxybenzene, plates or prisms (from w) or prisms (from aq HNO₃), mp — explodes on heating to 140—60°. Can be prep'd by oxidation of 1,3-dinitro-4-iodobenzene (Ref 1)
1,4-Dinitro-2-iodoxybenzene, crys, mp — explodes on heating rapidly at 150°; was prep'd by reaction of HOCl & 1,4-dinitro-2-iodobenzene in glacial acet ac (Ref 2)
Refs: 1) Beil 5, [202] & {642} 2) Beil 5, {642}

Trinitroiodoxybenzene, (O₃N)₃C₆H₂IO₂, not found in Beil

1-iodoxy-2-chlorobenzene, C₆H₄(Cl)IO₂; mw 270.5; needles (from w), explodes at 203°; sl in w, alc & glacial HAc. Prep'd by heating o-chloriodosobenzene in w or alc
Ref: Beil 5, 220
1-iodoxy-3-chlorobenzene, ClC₆H₄IO₂; mw 270.47; colorless crystals, explodes at 233°; sl in w, alc or glacial HAc. Prep'd by heating m-chloriodosobenzene in w
Ref: Beil 5, 220
1-iodoxy-4-chlorobenzene, ClC₆H₄IO₂; mw 270.47; colorless crystals, explodes at 243°; sl in w, alc or glacial HAc. Prep'd by heating p-chloriodosobenzene in w
Ref: Beil 5, 221

1-iodoxy-2,5-dichlorobenzene, Cl₂C₆H₃IO₂; mw 304.91, white needles (from hot w). Explodes mildly at 230°. Prep'd by treating 2,5-Dichloro-1-iodosobenzene with steam
Ref: Beil 5, 222

1-iodoxy-4-bromobenzene, BrC₆H₄IO₂; mw 314.93; leaflets (from glacial HAc), explodes at 240°; sl sol in glacial HAc. Prep'd by heating p-bromoiodosobenzene in w
Ref: Beil 5, 224

1-iodoxy-2,5-dibromobenzene, Br₂C₆H₃IO₂; mw 393.38; white amorphous powder, explodes at 218°; sl sol in hot w; sol in hot glacial HAc. Prep'd by treating 2,5-dibromo-1-iodosobenzene with steam
Ref: Beil 5, 224

1-iodoxy-3-iodobenzene, I₃C₆H₄IO₂; mw 361.93; colorless needles (from boiling w or glacial HAc), explodes at 216—18°. Prep'd by treating m-iodiodosobenzene with steam
Ref: Beil 5, 226

1-iodoxy-4-iodobenzene, I₃C₆H₄IO₂; mw 361.93; small needles (from glacial HAc), explodes around 232°. Prep'd by treating p-iodiodosobenzene with steam
Ref: Beil 5, 227

1,3 Di-iodoxybenzene, IO₂C₆H₄IO₂; mw 393.93; white tablets, explodes very violently at 261° or on impact; v sl sol in w or glacial HAc. Prep'd by heating m-diiodosobenzene with steam
Ref: Beil 5, 226

2-iodoxytoluene, CH₃C₆H₄IO₂; mw 250.05; white crystals, explodes at 210°. Prep'd by heating o-Iodosotoluene in w. Its Hydrofluoride salt, C₆H₄IOF₂, colorless leaflets, explodes mildly at 180°
4-iodoxytoluene, $\text{CH}_3\text{C}_6\text{H}_4\text{IO}_2$; mw 250.05; prepd as above from p-iodosotoluene. It melts with decomp at 229°.

4-iodoxy-1,3-dimethylbenzene, also called asym iodoxy-m-xylene, $\text{(CH}_3)_2\text{C}_6\text{H}_2\text{IO}_2$; mw 264.07; small crystals (from glc HAc) or amorphous powder (from w), explodes at 195°; sol in hot w or HAc. Prepd by passing chlorine thru a soln of asym iodo-m-xylene in pyridine
Ref: Beil 5, 377

5-iodoxy-1,3-dimethylbenzene, also called asym iodoxy-m-xylene, $\text{(CH}_3)_2\text{C}_6\text{H}_2\text{IO}_2$; mw 264.07; leaflets (from w), explodes at 216°. Prepd by steam distilling asym iodo-m-xylene. Datta & Choudhury claim that both the 4 & 5-iodoxy-isomers explode at 193° (Ref 2).

4-iodoxy-1-methyl-2-ethylbenzene, $\text{CH}_3\text{C}_6\text{H}_3\text{(C}_2\text{H}_5\text{)}\text{IO}_2$; mw 278.10; white leaflets, explodes at 229°. Prepd by heating 4-iodoso-1-methyl-2-ethylbenzene in w
Ref: Beil 5, 396

4-iodoxy-1-methyl-3-ethylbenzene, $\text{CH}_3\text{C}_6\text{H}_3\text{(C}_2\text{H}_5\text{)}\text{IO}_2$; mw 278.10; white leaflets, explodes at 229°. Prepd by treating corresponding methyl-ethyl-iodobenzene with Na hypochlorite soln
Ref: Beil 5, 396

4-iodoxy-1-propylbenzene, $\text{CH}_3\text{CH}_2\text{CH}_2\text{C}_6\text{H}_4\text{IO}_2$; mw 278.10; leaflets (from w), explodes at 185–200°. Prepd by storing corresponding iodoso compd in w
Ref: Beil 5, 393

5-iodoxy-1,2,4-trimethylbenzene, $\text{(CH}_3)_3\text{C}_6\text{H}_2\text{IO}_2$; mw 278.10; small needles (from glc HAc), explodes mildly at 212°; sl sol in chl; insol in eth or benz. Prepd by heating the corresponding iodoso compd in w
Ref: Beil 5, 404

2-iodoxy-1,3,5-trimethylbenzene, $\text{(CH}_3)_3\text{C}_6\text{H}_2\text{IO}_2$; mw 278.10; needles (from glc HAc), explodes at 195°; sl sol in glc HAc or alc. Prepd by heating iodomesitylene and treating it with steam
Ref: Beil 5, 409

4-iodoxy-1-tert-butylbenzene, $\text{(CH}_3)_2\text{CC}_6\text{H}_4\text{IO}_2$; mw 292.12; crystalline, explodes at 201°; sol in glc HAc; insol in w. Prepd by reacting p-tert-butyl-iodo-chlorobenzene with Na hypochlorite soln
Ref: Beil 5, 417

4-iodoxy-1-isooamylnbenzene,
$\text{(CH}_3)_2\text{C}_6\text{H}_3\text{CHCH}_2\text{CH}_2\text{C}_6\text{H}_4\text{IO}_2$; mw 306.15; tablets (from w or glc HAc), explodes at 200–03°. Prepd by oxidation of p-iodosooamylnbenzene with a Na hypochlorite soln
Ref: Beil 5, 435

Ion Exchange is defined in the Condensed Chemical Dictionary (Ref 31) as:
"A reversible chemical reaction between a solid (ion exchanger) and a fluid (usually a water solution) by means of which ions may be interchanged from one substance to another. The superficial physical structure of the solid is not affected. The customary procedure is to pass the fluid through a bed of the solid, which is granular and porous, and has only a limited capacity for exchange. The process is essentially a batch type in which the ion exchanger, upon nearing depletion, is regenerated by inexpensive brines, carbonate solutions, etc. Ion exchange occurs extensively in soils.

Ion exchange resins are synthetic resins containing active groups (usually sulfonic, carboxylic, phenol, or substituted amino groups) that give the resin the property of combining with or exchanging ions between the resin and a solution. Thus a resin with active sulfonic groups can be converted to the sodium form and will then exchange its sodium ions with the calcium ions present in hard water. "Amberlite" resins are of this type.

Some specific applications of ion exchange:
water softening; milk softening (substitution
of sodium ions for calcium ions in milk); removal of iron from wine (substitution of hydrogen ions); recovery of chrome from plating solutions; uranium from acid solutions; streptomycin from broths; removal of formic acid from formaldehyde solutions; demineralization of sugar solutions; recovery of valuable metals from wastes; recovery of nicotine from tobacco-dryer gases; catalysis of reaction between butyl alcohol and fatty acids; recovery and separation of radioactive isotopes from atomic fission; chromatography; establishment of mass micro standards; in cigarette filters to remove polonium from smoke.”

The discovery of the phenomenon of ion exchange is attributed to H.M. Thompson & J.T. Way, British agricultural chemists. They reported in 1848 the exchange of Ca and NH₄ ions in soils. No practical application of this discovery was made, however, until the German chemist R. Gans proposed (in about 1910) to apply the ion exchange principle to water softening. The first successful organic ion exchange materials appeared on the market about 1935, and about the same time I.G. Farbenindustrie started a systematic research program.

Since that time, the ion exchange industry, using various resins, has expanded enormously. Among the scientists who have contributed to the ion exchange industry, the following may be mentioned; O. Liebknecht & R. Griessbach of Germany, P. Smit of Holland, B.A. Adams & E.H. Holmes of Great Britain, H.L. Tiger, S. Sussman & A. Mindler of the United States and L. Wiklander of Sweden.

The ion exchange substances used commercially at present (in particular, ion exchange resins) are ionic solids in which one of the ionic species (either an anion or cation) is a highly cross-linked, polymeric, high molecular weight, non-diffusible ion, whose multivalent charge is balanced by relatively small diffusible ions of the opposite charge. These exchangers constitute a class of electrolytes having properties that are in many respects similar to true solutions of electrolytes.

The chief advantage of an ion exchange technique lies in simplicity and rapidity with which various separations or concentrations may be achieved. The application of ion exchange technique in analytical chemistry includes: concentration of dilute solutions, fractionation of ions having similar analytical properties, removal of interfering ions etc. Chromatographic technique is one of the branches of ion exchange technique. The ion exchange technique has found extensive application in the purification of water for laboratory uses, to replace the more expensive distilled water.


Ionic Energy in Propulsion. See Propulsion, Ionic in future Vol


Since the appearance of Vol 4, relatively little has been published on ionization in detonation waves, or ionization in shocked condensed media. However, there is considerable new information on shock ionization of gases. We discuss these phenomena below

Ionisation in a Detonation Wave. Hay et al (Ref 9) obtained high-speed framing camera pictures and time-resolved spectrograms of the detonation of solid explosives in a high vacuum. The salient feature of these observations is the very high velocity, 20 km/sec, of the leading products emitted from the ends of charges of explosives which contain hydrogen. These authors propose two mechanisms which might account for these high-speed emissions. They obviously favor mechanism 1). We quote:

"1. The mechanism of ambipolar diffusion can be invoked to double the speed of the ionized portion of the cloud. In ambipolar diffusion, a pressure gradient in the plasma at the surface of the explosive will tend to accelerate the electron gas at several thousand times the rate for ions. Any tendency for the electrons to move ahead of the ions, however, is quickly balanced by an electric field caused by the separation of charges. The electrons, being highly mobile in comparison to the ions, will quickly achieve a near equilibrium between the influences of the pressure gradient and electric field. However, the same electric field acts on the ions but in the same direction as the pressure gradient. This subjects the ions to a total force away from the surface double that on a neutral species. An order of magnitude calculation shows that a charge separation of even a tenth of a percent creates an electric field which would produce forces on the charged particles many orders of magnitude greater than the pressure gradient. The difference in arrival times of various species (hydrogen, lithium, sodium) in the pink glow is presumably due to the fact that the same pressure and electric field act on both light and heavy species in the detonation "plasma," accelerating the lightest species the most.

2. The mechanism proposed by Johansson and Selbert (Ref 3) which invokes elastic collision between heavy molecules in the detonation products and lighter molecules in the ambient gas. Although this is an attractive hypothesis, it appears doubtful that the lighter molecules actually come from the ambient gas for two reasons: (a) the intensity of the light emitted when these molecules impinge on a surface ought to increase as the number of available molecules increases; however, over a million-fold increase in ambient pressure (10^-7 to 10^-1 torr) no obvious increase in luminosity was observed;
(b) velocities of the sodium atoms or ions, which certainly do not originate in the ambient gas, are very nearly as high as those of the hydrogen atoms, molecules or ions."

The use of ionization pins or pin switches in the measurement of detonation velocity was described in Vol 4, pp D632-638. Their principle of operation is the closure of an electric circuit by the ionization associated with a detonation wave. The latest pin switch techniques are described in Ref 11

Shock Ionization in Condensed Media. Dremen et al have studied shock polarization of both polar & nonpolar liquids. They conclude that these polarizations are ionic in nature. We quote from a recent article by Dremen & Yakushev:

"It has been found in studying polarization of liquid dielectrics in shock waves, that liquids involving polar molecules, such as water, acetone, chlorobenzene, bromobenzene, iodobenzene, nitrobenzene, and others (Ref 5) give polarization signals. When shock waves are propagated over nonpolar liquids such as benzene, carbon tetrachloride, etc, no polarization signals appear (Ref 8). In this connection, an assumption arose that with decomposition of nonpolar molecules of a liquid behind the front of a shock wave into polar fragments, a polarization signal may be obtained from propagation of a second reflected shock wave over a shock-compressed substance (Ref 8). Carbon tetrachloride was the liquid used."

On the basis of their results as well as the results of others, Dremen & Yakushev conclude:

"The electrochemical nature of signals observed for all liquids studied is evidence that the electric conductivity arising on shock compression is of an ionic nature. Hamann and coworkers (Refs 1, 2, 6 & 7) came to the same conclusion when studying the conductivity of water, methanol and acetone in shock waves.

The persistence of transparency on shock compression of water, methanol and dichloroethane within the range of pressures corresponding to good conductivity (Ref 4) is another evidence in favor of the ionic nature of these liquids under the given conditions"


Shock Ionization of Gases. If temperatures behind a shock front are in the range of 15000-20000 K the gas through which the shock is moving will be appreciably ionized. The establishment of ionization equilibrium is the rate-determining step in this process and consequently controls the thickness of the wave front (Ref 6).

The ionization process is quite complex, and to avoid even further complications, the following discussion will be limited entirely to the shock-ionization of monatomic gases. One compelling reason for doing this is that complications due to dissociation (eg of N2 or more complex polyatomic molecules) are thus avoided. Furthermore for a given driver shock, the highest temperatures, and consequently the greatest likelihood of ionization, occur in monatomic gases since such gases cannot "soak-up" energy in internal modes, ie, in rotation or vibration. In what follows A represents an atom of gas (Argon by preference since much of what follows is derived from studies of shocked Argon). The main steps in the process are represented by the following equilibria:

\[
A + A = A^+ + e^- + A \quad (1)
\]

\[
A + \nu = A + e^- \quad (2)
\]

\[
A + e^- = A^+ + 2e^- \quad (3)
\]

Electron avalanching via (3) is the main process for producing A+ ions. However this requires the presence of "priming" electrons which can be
produced by \( A + A \) collisions in the shock (step (1)) or photoelectrically by the absorption of sufficiently energetic radiation to photoionize \( A \) (step (2)). However, under shock conditions where ionization is observed, the best evidence suggests that steps (1) & (2) cannot provide a sufficient number of "priming" electrons for step (3). This discrepancy is still unresolved (Ref 6). Biberman & Yakubov (Ref 4) have suggested that absorption of resonant radiation from the equilibrium region produces many \( A \) atoms that are electronically excited; such atoms are then readily ionized by electron impact and thus lead to electron avalanching.

Equilibrium calculations based on combining the Saha equation (in terms of the internal partition functions of \( A, A^+ \) & \( e^- \)) with the conservation conditions for the shock are fairly involved (Ref 1). We chose instead to derive the following set of equations for an idealized case, which, though they overestimate the degree of ionization and underestimate the temperature, point out the shock and gas parameters that control the ionization of monoatomic gases.

Consider a perfect gas of molecular weight \( M \) which has been shocked and has attained ionization equilibrium with the fraction of atoms ionized designated by \( \alpha \). For a shock pressure \( P \), much greater than ambient pressure, gas density \( \rho \) in the shock and \( \rho_0 \) ahead of the shock and a shock velocity \( U \), the conservation conditions lead to:

\[
P = \rho_0 U^2 (1-\rho_0/\rho) = \rho_0 M U^2
\]

where \( \mu \equiv 1-\rho_0/\rho \), and

\[
\Delta E = \frac{1}{2}P(1/\rho_0-1/\rho) = \frac{1}{2} \mu U^2
\]

combining Eq (1) with the perfect gas law gives

\[
P = \rho_0 \mu U^2 = \rho(1+\alpha)RT/M
\]

or

\[
T = \frac{\rho_0 \mu U^2 M}{(1+\alpha)R} = \frac{\mu(1-\mu)U^2 M/(1+\alpha)R}
\]

If we assume that a negligible amount of energy is used in electronic excitation of the \( A \) atoms, then the incident shock energy \( \Delta E \) is used to ionize some of the \( A \) atoms and to heat the mixture of ionized and neutral atoms. This energy balance is expressed by:

\[
\Delta E - \Delta I = c_v \Delta T
\]

where \( \Delta I \) is the total ionization energy of the gas and \( c_v \) is the const vol heat capacity of the equilibrium gas. From kinetic theory \( c_v = 3/2 \) R/M, which, combined with Eq (2), leads to:

\[
\frac{1}{2} \mu U^2 - \alpha I_1/M = 3/2(1+\alpha)R \Delta T/M
\]

because at equilibrium there are \( (1-\alpha) \) moles \( A, \alpha \) moles \( A^+ \) & \( \alpha \) moles \( e^- \) or a total of \( (1+\alpha) \) moles of "monoatomic" gas. Here \( I_1 \) is the first ionization potential of \( A \), i.e., we are neglecting all but singly ionized species.

Since \( \Delta T = T-T_0 \) and if \( T \gg T_0 \) (\( T \sim 10000 \) to \( 50000^o \)K which is much greater than \( T_0 = 300^o \)K), Eq (6) gives

\[
T = \frac{\mu U^2 M}{3(1+\alpha)R} - \frac{2}{3} \alpha I_1 / (1+\alpha)R
\]

and after rearranging, combining terms, & eliminating \( T \) via Eq (4):

\[
\alpha = \frac{U^2 M}{2I_1} \left[ \mu^2 - 3(1-\mu)\mu \right] = \frac{U^2 M \mu}{2I_1} [4\mu-3]
\]

and

\[
T = \mu(1-\mu)U^2 M/[1+U^2 M \mu(4\mu-3)] R
\]

Eq 8 immediately tells us that the degree of ionization depends strongly on the shock velocity \( U \). For strong shocks \( \mu \approx 0.9 \) (see Fig 6 of Ref 2) for all monoatomic or even polyatomic gases. Thus the only important shock parameter is \( U \). Similarly the only important gas parameters are the molecular weight \( M \) and the first ionization potential \( I_1 \).

It is instructive to examine the noble gases in terms of Eqs 8 & 9. In the series \( He \) to \( Xe \), \( M \) increases and \( I_1 \) decreases, thus at a given \( U \) and a nearly constant \( \mu \), the degree of ionization \( \alpha \) is much greater for \( Xe \) than for \( He \).

In fact \( \alpha \) increases regularly in the series \( He < Ne < Ar < Kr < Xe \). Temperature of the shocked gases also increases in the same manner but not as greatly as \( \alpha \) because the additive term in the denominator of Eq 9 increases progressively from \( He \) to \( Xe \) and tends to counteract the increase in \( M \) in this progression.

For strongly shocked Argon (see Table 1), calculations based on Eq 8 give \( \alpha \)'s that are 30 to 40% higher than the more accurate (partition function) calculations of Reynolds.
<table>
<thead>
<tr>
<th>U (mm/µsec)</th>
<th>ρ/ρ₀ (Ref 2)</th>
<th>u (Ref 2)</th>
<th>μ (Ref 2)</th>
<th>α (Eq 8)</th>
<th>α (Ref 2)</th>
<th>T (Eq 9)</th>
<th>T (Ref 2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>8</td>
<td>10</td>
<td>0.900</td>
<td>0.889</td>
<td>0.454</td>
<td>0.322</td>
<td>19000</td>
<td>23100</td>
</tr>
<tr>
<td></td>
<td>9.5</td>
<td>0.895</td>
<td></td>
<td>0.436</td>
<td></td>
<td>20200</td>
<td></td>
</tr>
<tr>
<td></td>
<td>9.0</td>
<td>0.889</td>
<td></td>
<td>0.410</td>
<td></td>
<td>21500</td>
<td></td>
</tr>
<tr>
<td></td>
<td>8.5</td>
<td>0.882</td>
<td></td>
<td>0.393</td>
<td></td>
<td>22800</td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>10</td>
<td>0.900</td>
<td>0.896</td>
<td>0.574</td>
<td>0.417</td>
<td>22200</td>
<td>25500</td>
</tr>
<tr>
<td></td>
<td>9.5</td>
<td>0.895</td>
<td></td>
<td>0.552</td>
<td></td>
<td>23600</td>
<td></td>
</tr>
<tr>
<td></td>
<td>9.0</td>
<td>0.889</td>
<td></td>
<td>0.518</td>
<td></td>
<td>25200</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>10.5</td>
<td>0.905</td>
<td>0.901</td>
<td>0.736</td>
<td>0.523</td>
<td>23800</td>
<td>28350</td>
</tr>
<tr>
<td></td>
<td>10.0</td>
<td>0.900</td>
<td></td>
<td>0.709</td>
<td></td>
<td>25300</td>
<td></td>
</tr>
<tr>
<td></td>
<td>9.5</td>
<td>0.895</td>
<td></td>
<td>0.681</td>
<td></td>
<td>26900</td>
<td></td>
</tr>
</tbody>
</table>

& Seely (Ref 2). This is to be expected since we neglected the energy used in electronic excitation of Ar atoms and this neglect shows up as increased ionization. Our calculated temperatures (see Table 1) are about 10% lower than Seely’s (Ref 2) if ρ/ρ₀ is the same in both calculations. Presumably this better agreement between temperature and α’s is due to the following compensating effect. Inclusion of electronic excitation would make it appear as an additional negative term in Eq (7) and would tend to reduce T. However, inclusion of electronic excitation makes α smaller. Thus the first right-hand term of Eq (7) becomes larger and the second (negative) term becomes smaller, both of which tend to increase T and counteract the direct effect of including a negative term for electronic excitation.

There are no direct measurements of the degree of ionization of strongly shocked noble gases. Thus the validity of the above discussion can only be tested indirectly by comparing computed shock temperatures (which contain terms in α) with measured shock temperatures.

Shock temps and shock velocities in Ar were measured by the writer (Ref 3) & Zatsepin et al (Ref 7). Velocities and temps of shocks in Kr & Xe were also measured by the writer (Ref 5) and shock temps in Xe by Zatsepin et al (Ref 7). The shock velocity of 8.3 mm/µsec reported for Ar in contact with Comp B (Ref 3) is incorrect. This is an average velocity of the propagation of intense luminosity and/or ionization which changes very little over at least 10 cm travel from the Comp B face except in the region very close to the Comp B. Unfortunately it is the shock velocity in this close-in region that we need since peak temp is achieved within 1 µsec of the shock entry into the Ar. Our best estimate of U for Ar in the close-in region is ~10 mm/µsec. The values of U for Comp B driven Kr & Xe, to be used in Table 2, are also estimates for the close-in region. They are quite tentative since they are based on few data.

Table 2.

<table>
<thead>
<tr>
<th>Gas</th>
<th>U_OBS (mm/µsec)</th>
<th>T_OBS (ºK)</th>
<th>(ρ/ρ₀)CALC</th>
<th>T_CALC (ºK)</th>
<th>T_CALC (ºK)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ar</td>
<td>~10</td>
<td>28500</td>
<td>10.1 (a)</td>
<td>25300 (c)</td>
<td>28700 (d)</td>
</tr>
<tr>
<td>Kr</td>
<td>~7.6</td>
<td>~37500</td>
<td>8.6 (b)</td>
<td>32600 (c)</td>
<td>37600 (d)</td>
</tr>
<tr>
<td>Xe</td>
<td>~6.8</td>
<td>~47500</td>
<td>7.8 (b)</td>
<td>40200</td>
<td>47000 (d)</td>
</tr>
</tbody>
</table>

(a) Ref 2
(b) Assumed to be the same as for Ar (Ref 2) at the corresponding U
(c) According to Eq 9 with α obtained via Eq 8
(d) All α’s arbitrarily reduced by a factor of 1.4
Agreement between experiment and computation is good if one assumes that, as appears to be the case for Ar, all "accurate" α's are 40% smaller than the corresponding α's computed via Eq 8. A presumably exact calculation (no details given) by Zatsepin et al. (Ref 7) gives a shock temp of ~37000°K(?) for a shock in Xe moving at U = 6.8 mm/μsec and ~29000°K for shock in Ar moving at 10 mm/μsec.

Written by J. ROTH


Ions; Action on Explosives. Kallmann & Schrändl (Ref 2) claim that Mercuric Fulminate, Azides & even TNT were initiated by the action of H, Ar & Hg ions. This was not substantiated (Ref 3). Nitrogen Iodide is definitely initiated by α-particles (Refs 1 & 4) but it is atypical, see iodides and Initiation in this Vol


IPA Composition. Recognizing that the hygroscopicity of Black Powder is due largely to the charcoal, a non-hygroscopic ignitor-type pdr was developed. It was found that substituting Tetranitrocubazole (See Vol 2, pC48-L) for charcoal results in a compn, designated IPA Composition, which is only half as hygroscopic as Blk Pdr and can be made only one-fifth as hygroscopic by the use of chemically pure instead of spec grad K nitrate

Ref: S. Livingston, PATR 647 (March 1947)

Ipatieff, V.N. (1867-1952). Vladimir Nikolaevich Ipatieff, of Universal Oil Products Co and Northwestern University, was born in Moscow and attended military schools under the czarist regime. His work carried him to a high position in the Russian army and to recognition as one of the outstanding scientists of Russia.

Although he had been interested in chemistry since boyhood, his studies in organic chemistry began at 27 when he presented a thesis dealing with the addition of hydrogen bromide to unsaturated hydrocarbons, a reaction which soon led to the proof of structure of isoprene. He studied in both Germany and France and on returning to Russia he presented a thesis on the synthesis of isoprene. His work on decomposition of organic compounds at high temperatures with special catalysts, he began about 1900. His work on the dehydration and dehydrogenation of alcohols and their combination in the preparation of butadiene from alcohol has formed the base for much of the synthetic rubber development in the US. He later developed the first high pressure bomb for catalytic reactions, then found promoters which aided the catalysts in their activity. From these discoveries branched his work in hydrogenation (both organic and inorganic), isomerization, polymerization, condensation, cyclization, alkylation, and related reactions.

It was at his brother's home that the Czar had taken refuge and was assassinated at the time of the revolution in Russia. Dr. Ipatieff managed to continue his research under the new regime but this became increasingly difficult and he finally left Russia permanently, abandoning all his financial assets. Through the efforts of Gustav Egloff of Universal Oil Products and Ward Evans of Northwestern University, who arranged for him to carry on work here, he came to the United States in 1931. He was denounced
by the Soviet government for his refusal to return to that country and the honors which had been conferred upon him were withdrawn.

Among his many honors were election to both the American and Russian Academies of Sciences, the Willard Gibbs Medal of the Chicago Section of the ACS, the Berthelot Medal, the Lenin Prize, and honorary doctor’s degrees. Out of the income from a trust fund set up by Dr and Mrs Ipatieff the Ipatieff Prize was established several years ago which would encourage work in the field of catalysis and high pressure.

At both the Universal Oil research laboratory and the Ipatieff Catalytic Laboratory at Northwestern (an extension of the original laboratory set up by Ipatieff) Dr Ipatieff was active almost to the time of his death.

Ref: Anon, Chem & Engr News 30, 5300 (1952)

IRECO Chemicals. IRECO Chemicals was organized in Salt Lake City, Utah, in 1962 from a merger of Intermountain Research and Engineering Company and Mesabi Blasting Agents, Inc., for the purpose of commercializing slurry explosives. Slurry explosives were invented by Dr. Melvin A. Cook and H. Earl Farnam, Jr., in 1956 and represent a major breakthrough in explosives technology, the magnitude of which can be compared with dynamite’s victory over Black Powder in the 1860’s.

Slurry explosives are based upon a system consisting of oxidizing agents and nonexplosive fuel materials. The explosive energy derived from these unique explosives is the result of extremely rapid reduction-oxidation reactions between the fuels and oxidizers upon initiation by a high explosive booster. A slurry explosive can be visualized as a colloidal system which comprises basically two phases, a dispersion phase and a dispersed phase. The fundamental concept which led to the discovery of slurry explosives was that an aqueous oxidizer solution (e.g., ammonium nitrate) could be used as the dispersion medium of a colloidal system to disperse the required fuel (aluminum) and thereby achieve a multiplicity of beneficial results.

A particularly significant feature of the slurry explosives is that they are easily made water resistant by means of a suitable hydrophilic colloid which binds the solid particles of the system together and prevents diffusion of water into or out of the products. Other principal advantages of slurry explosives are substantially increased safety, greater overall economy, more favorable fume characteristics, and ease of varying the energy and density to enable custom designing of the explosive for each specific application to ensure optimum energy/cost ratios. These and many other favorable characteristics have enabled slurry explosives to replace conventional explosives in mining, quarrying, and construction operations around the world.

Prior to its merger into IRECO Chemicals, Intermountain Research and Engineering Company (IRECO—founded in 1958) pioneered the early development of slurry explosives under a contract from the Iron Ore Company of Canada (IOCC) who owned the original Cook and Farnam invention.

Although the original slurries had been licensed by IOCC to major explosives manufacturers around the world, progress in the marketplace for the first two years was slow. In December 1959 IRECO conducted a test at the US Steel Pilotae operation on the Mesabi Iron Range in northern Minnesota at the request of IOCC to promote the use of slurry in open-pit blasting. The results were so spectacular that a group of IRECO scientists quickly founded a manufacturing and sales company called Mesabi Blasting Agents, Inc., or MBA (in 1960), obtained a sublicense from IOCC, and began to manufacture and sell slurries on the Mesabi Iron Range. Within a year MBA had successfully promoted the use of slurry explosives in the mining industry.

The first-generation slurries were called Dense Blasting Agents, or “DBA’s,” and comprised two types: slurries sensitized primarily with coarse “Pelletol” TNT and slurries sensitized with aluminum. Second-generation slurries, designated the IREGEL 300 series, were introduced in 1969 as a replacement for the DBA’s except for DBA-22M, the first and only slurry yet to be completely qualified by the US Military.

The IREGEL 300’s were significantly stronger and more water resistant than their DBA counterparts by virtue of lower water content, ensured continuity of aqueous oxidizer-
salt solution phase, and better thickening. In 1971 IRECO achieved another first with their introduction of IREMITE, a cap-sensitive series of slurry explosives designed for use in small-diameter bore-holes as a replacement for dynamite. These versatile small-diameter products offer distinct safety, fume and performance advantages and are available in a wide range of diameters, densities, and energies and are sufficiently revolutionary in concept to have the same effect in the small-diameter applications as the SMS system (see below) did in large open pit mining.

Yet another economically attractive series of slurries called the IREGEL 600 series was introduced in the US in late 1972. This series is also being manufactured today in Canada, South Africa and Australia.

In order to initiate slurry explosives and other modern blasting agents, IRECO Chemicals developed and patented the Procure booster—a booster designed for maximum safety with a cap-sensitive core protected by a non-cap-sensitive explosive shell comprising cast TNT or composition B. These boosters are characterized by extremely high detonating pressures. Because the booster requirement, as measured, for example, by the minimum booster (MB) required to detonate a blasting agent, varies as the reciprocal of the detonation pressure (ie MB = K/P^2), Procure boosters are very efficient. For instance, the MB with a Procure booster is much smaller than the MB with a gelatin dynamite or low-density (powder) high explosive. "Quickness" (or absence of a transient build-up of detonation) in a booster is another important factor which is possessed almost ideally by the original "Pentomex" (cast pentaite) and the present Procure boosters. Procure boosters are used extensively in blasting operations around the world to initiate all types of modern blasting agents. They are manufactured in a wide variety of sizes and weights to meet individual requirements.

One of the most significant events in explosives history has been the development of the IRECO Chemicals Site-Mixed-Slurry (SMS) system, commonly referred to as the pump truck system, which is today the safest blasting system known. Nonexplosive raw materials are carried by the pump truck from the storage area to the borehole site where they then become an explosive capable of being detonated after placement in the borehole. By adjusting the settings on the control panel the pump truck operator can deliver a wide range of explosive strengths and densities to meet individual blasting requirements.

The first SMS system was established for the Kaiser Steel iron ore operation at Eagle Mountain, California. The first SMS system outside North America was established in early 1965 at Phalaborwa, North Africa, for the Palabora Mining Company. Shortly thereafter the SMS system was established in rapid succession for mining operations throughout the world.

In 1966 fifty percent of the capital stock of IRECO Chemicals was acquired by the Rio Tinto-Zinc Corporation, Ltd. worldwide association of mining and related industrial enterprises such as ore processing, smelting, fabricating metals, and the production of chemicals. Since that time IRECO Chemicals has continued to grow and is today the world leader in the research, manufacture and marketing of slurry explosives with operations in important mining centers worldwide.

Written by M. GARFIELD COOK

Irradiation of Explosives with High Speed Particles was studied by Bowden & Singh. They subjected to irradiation a number of sensitive explosive crystals (such as Pb, Ag & Cd azides, Ag acetylide and nitrogen iodide) by electrons, neutrons, fusion products and X-rays. All these substances were exploded by an intense electron beam but it was shown that this was due to a thermal effect. Fission products exploded nitrogen iodide but in the other substances some changes within the crystals took place but no explosions. The experiments showed that, in general, the activation of a small group of adjacent molecules was not enough to cause explosion.


Iron Acetylide. See Acetylides and Carbides (Inorganic) in Vol 1, p A76-R

Iron Azide. See List of Inorganic Azides in Vol 1, p A543
Iron Picrate. See Picrates

Iron, Powder for Pyrotechnics. Sidorov et al. claim a spark-forming composition for ppg fireworks which forms bright sparks of different colors, containing a thermal mix & a metal powder, eg (%wt): NH₄ClO₄ 55 ± 5, urotropine 14 ± 2, iditol 8 ± 2, metal powder (Fe or steel chips, powdered Al or its alloy with Mg) 23 ± 5 (Ref 2).


Isano Oil. Isano oil, a conjugated triple-bonded glyceride, when heated sufficiently, reacts exothermically with violence. Its uses & chem reactions are described Ref: T.A. Kneeland et al, JAOC 36, 361 (1958) & CA 52, 15096 (1958)

Isazaurolin. (6-Oximino-Δ²-dihydro-1,2,4,5-oxatriazin in Ger). (HO,N):C,NH,NH
O- N=CH

mw 116.08, N 48.27%; mp turns orange ca 85° and puffs off ca 112-113° with loud report. Colorless needles which become yellowish on drying and then yellow-orange on storage. Was prepared by Wieland & Hess (Ref 2) by treating methyl azaurolitrilic acid with concd HCl.

Dissolves in dil mineral acids (colorless) and in alkalis (orange-red). It is a mild explosive. Forms salts, for instance.

Hydrochloride. C₃H₆O₂N₄·HCl; mp (dec 148-150°) prisms (from abs alc with small amt of eth); very sol in w; insol in alc.
Refs: 1) Beil 27, 783 2) H. Wieland & H. Hess, Ber 42, 4188-89 (1909)

Isoamylamine. See under Amylamine in Vol 1, p A395-R

Isoamylpicrate. See iso-Amylpicrate in Vol 1, p A399-R

Isoamylureidoacetyl Azide. See iso-Amylureidoacetyl Azide in Vol 1, p A399-R

2-Isocyanate Benzoyl Azide. See 2-Azidoformylphenylisocyanate in Vol 1, p A638-R

Isocyanic Acid. See Cyanic Acid in Vol 3, p C582-L

Isobutana. See iso-Butane in Vol 2, p B368-L

Isobutanesodiol. See iso-Butanesediol in Vol 2, p B370-L

Isobutanol. See under Butanol and Derivatives in Vol 2, p B372-R

Isobutyl Alcohol. See under Butanol and Derivatives in Vol 2, p B372-R

Isobutyl Peroxide. See Di-iso-butyl Peroxide in Vol 5, p D1201-L

2-Isocyanatobenzoylazide. See 2-Azidoformylphenylisocyanate in Vol 1, p A638-R

Isocyanogen Tetraazide. (N₂)₂C·NN·C(N₃)₂, mw 220.13, N 89.09%, OB to CO₂ -29.1%, mp 89°. Prepd by reacting tetrabromoisocyanogen, Br₂·CN·CBr₂, dissolved in Me₂CO with aq NaN₃. Its use as an initiating explosive is claimed Ref: C.J. Grundman & W.J. Schnabel, USP 2990412 (1961) & CA 65, 25256 (1961)

Isomelamine. See Cyanuramide in Vol 3, p C589-L

Iso-Me-NENA. Designation of N-(β-Nitroxypropyl)-nitrine or 1-Nitramino-2-propanol Nitrate described in Vol 1, p A253-L
Isomers. Compounds having the same empirical formulas but different molecular arrangement, as well as different chemical and physical properties, are known as isomers.

For instance, ortho, meta and para-mononitrotoluenes all have the same empirical formula, C₇H₅NO₂, but the NO₂ group is attached to different carbons of the benzene ring.

Isomers of Trinitrotoluene. Several isomers are known: alpha, beta, gamma, delta, epsilon and zeta (see Trinitrotoluene, described under Toluene and its Derivatives).

The most important of these explosive isomers is the alpha-trinitrotoluene, known as TNT. In crude, commercial TNT, 4 to 5% of impurities are present, consisting mainly of a mixture of beta-, or 2,3,4-, and gamma, or 3,4,6-trinitrotoluenes. These impurities may be removed by treating the crude TNT with an aqueous solution of sodium sulfite. According to Davis (Ref 3) the beta- and gamma-isomers react with lead oxide in alcohol to form lead dinitrocresolates, while alpha-TNT remains unaffected under similar conditions.


Isonitramines are compounds having a general formula, R.N(OH).NO or likely

\[ \text{RN} - \text{NOH} \]

where R stands for a radical, CH₃-, C₆H₅-, CH₃C₆H₅-, etc. They may be prepd by treating ketones or nitroparaffins in alcoholic solutions with nitrous oxide (N₂O₃) in the presence of sodium ethylate.

For discussion of the nomenclature of isonitramines see Ref 3.

Metallic salts of these compounds were proposed by von Herz to be used in detonators & percussion caps. Lead Methylene disisnotramine is specifically mentioned (Refs 1 & 2).

Refs: 1) E. von Herz, BritP 241892 (1924) & CA 20, 3574 (1926) 2) E. von Herz, USP 1625966 (1927) & CA 21, 2065 (1927) 3) Sidgwick (1937) p 455

Isonitraminoacetic Acid. See Nitrosohydroxymethanoacetic Acid under Hydroxylamine Derivatives in this Vol.

α-Isonitraminoacetoacetic Acid, Ethyl Ester. See 2-(Nitrosohydroxylamino)-3-butanone-1-acetic Acid, Ethyl Ester under Hydroxylamine Derivatives in this Vol.

α-Isonitraminoacetopropionic Acid. See 2-(Nitrosohydroxylamino)-1-propanoic Acid under Hydroxylamine Derivatives in this Vol.

Isoälefin Polymers of molecular wt 50000 to 100000, such as Isobutylene polymer, were patented as thickeners for flammable naphtha used in incendiary bomb mixts. Up to 25% may be incorporated. Small amounts of metal soaps may replace part of the polymer. Thermite mixts, Na, K and P may also be included in such an incendiary filler.


Isopicramic Acid. See 2,6-Dinitro-4-aminoanisole in Vol 1, p A243-R

Isoprene (2-Methyl-1,3-butadiene) Peroxide, Polymeric. (No formula given). Resinous substance; decomp slightly above 0°. Reported in Ref 2 as a highly explosive substance. May be prepd by the reaction of isoprene with molecular oxygen.

Refs: 1) Bell, not found 2) K. Bodendorf, ArchPharm 271, 1 (1933) & CA 27, 4472 (1933) 3) W. Kern et al, Makromol Chem 3, 223 (1949) & CA 50, 1431 (1956) 4) Tobolsky & Mesrobian (1954) 31 & 178

Isopropyliden-di-cyclohexyliden-triperxyxyl. One of the Ger names for the "Acetone Compound" of 1,1'-Bis(hydroperoxycyclohexyl)-peroxide described in Vol 2, p B144-R.
Isopurpuric Acid or Picrocyaninic Acid. (or Dicyandinitro-α-phenylhydroxylamine)

\[
\begin{align*}
\text{O}_2\text{N} & \quad \text{CN} \\
\text{HO} & \quad \text{NO}_2 \\
\text{NH} & \quad \text{OH}
\end{align*}
\]

mw 265.14, N 26.42%. It is an unstable compound of dark-yellow color, isolated for the first time by Borsche and Böcker in fairly pure state. It may be prepd by mixing its potassium salt with an ice-cold aqueous solution of phosphoric acid.

The salts of isopurpuric acid are easier to prepare, are more stable and some are explosive, for example:

Potassium Isopurpurate. C₆H₅O₂N₃K, mw 303.24, N 23.10%. Brownish-red crystals; may be prepd by adding slowly, with agitation, a solution of 10g of KCN in 40ml of H₂O to 10g of PA dissolved in 170ml of alcohol. The temperature should be kept below 35°. Stirring is continued for a while after completing the addition of KCN and then the mixture is left to stand. After this, the precipitate is filtered off and crystallized from about 1½ l of hot water.

The salt is fairly sol in w and easily sol in most organic solvents. It is a violent explosive, very sensitive to impact and friction.

It puffs off at 215°

Refs: 1) Beil 15, 61 2) W. Borsche & E. Böcker, Ber 37, 4396 (1904) 3) L. Gody, Traité théorique et pratiques des matières explosives, Namur (Belgium) (1907), p 562 4) Not found in recent CA’s

Iso-Trioil. A product resulting from the purification of crude TNT by recrystallization from nitric acid. This process has been used in Europe on a production scale; however, the Sellite purification process was selected for use in the USA. Isothiol has been found to be a satisfactory substitute for TNT in Dynamite consisting of: RDX 78, TNT 15, cornstarch S & polyisobutylene 2% and for DNT in the M-1 Propellant consisting of: NC 84.5, DNT 9.9, dibutylphthalate 4.9 & diphenylamine 1.0%

Ref: J. Cussen, “Utilization of By-Product

 Iso-Trioil Resulting from Nitric Acid Purification of Crude TNT”, PATR-DB-TR:16-58 (Jan 1959)

Isotrotyl is a yellow material, mp 57-58°, containing about 98% of alpha-TNT and its isomers. It is prep’d by nitrating the so-called “liquid TNT,” containing 16.9-17.4% N with mixed nitric-sulfuric acid. It has been suggested that “isotrotyl” consists of addition compounds of the isomers instead of free isomers. Since the alpha, beta and gamma compounds are practically of the same value as explosives, this low-melting explosive (isotrotyl) was found to be useful as a constituent of blasting explosives, such as Roburite, Bellite etc. It was also used to lower the melting points of other explosives, eg TNX or PA


α-Isoxazole. See under α-Isoxazolecarboxylic Acid below

α-Isoxazolecarboxylic Acid and Its Explosive Derivatives. ON:CH₂:CH₂CO₂H; mw 113.07, N 12.39%; pale yel crystals; mp 149°; bp (loses CO₂ on heating above its mp)

First obtained but not identified by Testoni & Mascareli (Ref 1) from products of reaction between acetylene and fuming nitric acid.

Qulico & Freri (Ref 2) prep’d it in the same manner and assigned to it the above structural formula.

Qulico & Freri (Ref 3) used this acid to prepare the α-isoxazole hydrazide, ON:CH₂:CH₂CO₂HN₂; mw 127.10, N 33.06%, OB to CO₂ -106.8%; mp 141-42°, which on treatment with NaNO₂ + HCl gave the explosive α-isoxazoleazide, ON:CH₂:CH₂CO₂N₂; mw 138.09, N 40.58%, OB to CO₂ -73.9%; mp 37° (decomp); bp (explodes violently on heating above its mp)

Refs: 1) Testoni & Mascareli, Gazz 32 (1), 202 (1902) 2) A. Qulico & M. Freri, Gazz 59, 930-41(1929) & CA 24, 3484 (1930) 3) M. Freri, Gazz 62, 457-63 (1932) & CA 26, 5954 (1932) 4) No further refs found under Isoxazole derivs or Isoxazolecarboxylic acid in CA 1947-1971
Isoxazoleazo Derivatives of Trinitromethane Prepared from Acetylene. In studying the reaction products of nitric acid with acetylene, Quilico isolated an explosive product of yellow color corresponding to the formula $C_6H_6O_2N_6$, mw 246.10, N 34.15%.

He assigned to it tentatively the structural formula $\text{ON:CHCH:CN:(NC(NO_2)_3}$ and named it $\alpha$-isoxazoleazo trinitromethane. The compound melts with decompn at 78\(^\circ\)C, when heated cautiously, but when heated rapidly or struck it explodes with a flame.

Refs: 1) Beil 1(902) 2) A. Quilico, Gaz 62, 503-18 (1932) & CA 26, 5953-4 (1932) 3) No other refs found under Isoxazole derivs in CA 1947-1971

4-(3-Isoxazolyl)-3-furazancarboxylic Acid was prepd in complex sequence of reactions by Quilico & Freri (Ref 1). It melts at 133\(^\circ\)C & decomposes ~190\(^\circ\)C.

Its Ag salt explodes weakly when heated. (No formulas for the acid or its salt are given)

Refs: 1) A. Quilico & M. Freri, Gaz 76, 3 (1946) & CA 41, 381 (1947) 2) No further refs found in CA 1947-1971

I\(_{sp}\). Abbreviation for specific impulse (sometimes called specific thrust) is the propulsive impulse delivered by a propulsion engine per unit weight of propellant. In the US the units of I\(_{sp}\) are lb-sec/lb. It is related to measurable quantities by

$$I_{sp} = \frac{V_e + (P_e - P_0)A}{m_0g}$$

where the subscripts (e) refer to exit quantities, and the subscript (a) refers to ambient quantities; \(V\) = product stream velocity, \(P\) = static pressure, \(A\) = area, \(m\) = rate of mass flow & \(g\) = gravitational acceleration (Ref 1). Typical values of I\(_{sp}\) for "ordinary" systems is 200-270 lb-sec/lb and 270-400 lb-sec/lb for "high-energy" systems (Ref 2). Theoretical calculations of I\(_{sp}\) (based on thermodynamics & thermochemistry) are in good agreement with experimental measurements (Ref 3)


ITALIAN EXPLOSIVES AND RELATED ITEMS

Introduction

During WWII, the Italians had several large military explosives and ammunition plants and some of its mining expls plants manufd military expls. They produced materials of good quality and in quantity nearly sufficient to conduct the war.

List of Principal Italian Factories Manufacturing Explosives and Related Items (Obtained thru the courtesy of Dr Omero Vettori, Director of Cheddite Factory, near Aulla (Massa Carrara) (April 1975)

Note: The firms marked with an asterisk are still in production.

Section A. - Explosives and Propellants

1) B.P.D. (Bombini-Parodi-Delfino), now SNIA VISCOSA; factory in Colleferro, near Rome (Production of TNT, Propellants, Hunting Powder and Missiles)

2) ACNA (Aziende Colori Nazionali Affini); factory in Cengio (TNT)

3) SGEM (Società Generale Esplosivi Munizioni); factories in:

a) Carmignano (TNT, Propellants — for cannons and rifles) (destroyed)

b) Pallerone, near Aulla (Double-Base Propellants) (This factory is now the property of the Italian Army)

c) Villafranca Lunigiana, near Aulla (Propellants for cannons and rifles) (destroyed)

d) Avigliana, near Turin (RDX, PETN, Dynamites and Propellants) (Closed)

4) Bussi e Pratole Peltina (RDX) (destroyed)

5) Unknown Name Firm; factory in Narni (PETN) (This factory is now the property of the Italian Army)

6) SIPE (Società Italiano, Prodotti Esplosivi); factories in:

a) Spilamberto, near Modena (NC, PETN, Fuse, Detonating Fuse, Primer & Dynamite)

b) Gallicano, near Lucca (now called SIPE NOBEL) (Black Powder)

6) SGEM Factory at Orbetello (now SIPE NOBEL) (Dynamite Detonating Fuse, Industrial Explosives, Blasting Caps, Shell Loading, Detonators)
7) Stacchini, Factory in Saliera Apuana, near Aulla (Black Powder)
8) AFE (Azienda Forniture Esplosimenti), Factory in Stazzema, near Viareggio (Black Powder)
9) Polverifico Benedetto Cacciotti, Factory in Viterbo (Black Powder, Industrial Explosives, Safety Fuse)
10) Army Factory: Capua (Protecnico Esercito) (Caps, Primers, Small Arms Ammunition)
11) Nobel SGEM Factory in Taino, near Varese (Detonators, Blasting Caps, Primers, Fuses, Primary Explosives (now closed)
12) Army Factory (Sopletificio Torre Annunziata) (Fuse Production)
13) S.R.C.M. (Società Romana Costruzioni Meccaniche), Factory near Rome (Primary Explosives, Hand Grenades, Caps, Primers)

Section B – Shell Loading Plants:
1) Army Factories in:
   a) Piacenza (exploited in 1940)
   b) Noceto, near Parma (Laboratorio Caricamento Proiettoli Esercito)
   c) Baiano di Spoleto (Laboratorio Caricamento Proiettoli Esercito)
2) Mangiarotti, Factory in Codroipo (Shell Loading, Industrial Explosives, Detonating Fuses)
3) Ditta Fratelli Rovina, Factory in Spilimbergo (Shell Loading and Unloading)
4) Ditta Finocchi, Factory in Rossano Veneto (Shell Loading and Unloading)
5) Ditta Amelotti, Factory in Rivalta Scrivia (Shell Loading) (Closed)
6) Polverifico Stacchini, Factory in Bagni di Tivoli near Rome (Shell Loading and Loading of Propellants)
7) B.P.D. (Bombardi-Pardi-Delfino), now SNIA VISCOSA, Factory at Ceccano (Shell Loading)
8) Sorlini Ing. Antonio, Factory in Ghedi, near Brescia (Shell Loading, Compression of Explosives, Industrial Explosives)
9) Sorlini Luciano, Factory in Carzago della Riviera (Dynamites, Industrial Explosives, Detonating Fuses, PETN)
10) Ditta Vulcana, Factory in Fascia d’oro, near Brescia (Shell Loading, Industrial Explosives, TNT Purification, Compression of Explosives) (Now closed)
11) Ditta Federico Marzan, Factory in Peschiera del Garda, near Verona (now SALMA) (Shell Loading, Shell Manufacture, Mines, Explosives Manufacture) (Now closed)
12) Ditta Simmel, Factory in Castagnole Paese, near Treviso (Shell Loading, Assembly)
13) Ditta La Precisa, Factory in Teano, near Naples (Shell Loading, Primary Explosives, Primers, Caps)
14) Chedite Italia, Factory in Aulla (Massa Carrara) (Shell Loading, Assembly, Detonating Fuses, Industrial Explosives, such as Cheddites)

Section C – Industrial Explosives
1) Dinamite S.p.A. (Società per Azione), Factory in Mereto di Tomba, near Udine (Dynamites, DNT, PETN, Industrial Explosives, Detonating Fuse, Blasting Machines)
2) Pravissani Esplosivi, Factory in Sequals, near Udine (Dynamites and other Industrial Explosives)
3) SES (Società Esplosivi Siciliana), Factory in Latina (Industrial Explosives)

Section D – Small Arms Ammunition and Weapons
1) Beretta Armis listed in Catalog
2) OTO MELARA, Factory in La Spezia (Manufacture of cannons and missiles)
3) Army Arsenals in Naples, Turin and Piacenza
4) BREDIB MECCANICA BRESCIANA in Brescia (Rifles and Pistols)
5) Army Factory in Terni (Cannons and other Weapons)
6) SMI (Società Metallurgica Italiana), Factory in Campotizzoro (Small Arms Ammunition)
7) Leon Beaux (Small Arms Ammunition including 20-mm) (Now closed)
8) Giulio Fiocchi Lecco (Small Arms Ammunition)

Italian explosives, ammunition and weapons (small arms and artillery pieces) have always been considered of very good quality. With the exception of T4 (Cyclonite or RDX) and Tritolita (Cyclotol), which the Italians developed and used before both Great Britain and the USA, there are no high explosives of unusual interest or originality. However, there are several explosives similar to the German Ersatzprengstoelle (Substitute Explosives), which were developed in Italy due to the shortage of aromatic compounds.

In the following pages are listed Italian explosives, some of them obsolete. We are able
to bring the list up to date due to the assistance of Dr Omero Vettori of Aulla (Massa Carrara) and the appearance in 1974 of the monumental work of Dr Camillo Belgrano, “Gli Esplosivi”, contg 695 pages. It is the 2nd, greatly enlarged, edition of his 1952 book (Ref 13). The new edition is listed as Ref 31.

**Alphabetical List of Italian Explosives and Related Items**

**Acapnia** is one of the smokeless propellants, listed in Belgrano (Ref 31, p 581) and by Molina (Ref 1, p 409). See under Polveri da Caccia.

**Acardite** or *asym*-Diphenylurea, described in Vol 1 of Encycl (Ref 24, p A47-R), was developed in Germany under the name of *Akardit*. It has been used, accdg to Belgrano (Ref 31, p 218) as a stabilizer in some Italian smokeless propellants.

**Accenditori** (Lighters) are devices serving to light fuses (mice). In the catalog of Montecatini of 1959, Accenditori “Pirea” Montecatini (p 40) and Accenditori “Dardo” Montecatini (p 41) are described.

**Accenditori in bacchette** (Lighter Rods). Accdg to Belgrano (Ref 31, p 530), it is a fuse of diam 5.5–5.6mm and ca 10cm long filled with a very slow-burning powder contg a finely pulverized mixture of KNO₃ 79.00, beechn carbon 3.75 & sulfur 17.25%.

**Accenditori Elettrici** (Electric Lighters). A brief description is given in Belgrano (Ref 31, pp 514–15). In catalog of Montecatini of 1959, p 31, are briefly described Accenditori elettrici ritardati (Delay Electric Lighters).

**Accenditori Militari** (Military Igniters). The following items are described in TM-9-1985-6 (1953) (Ref 16): 

a) Chemical Delay Igniter (pp 173–74 with Fig 245)
b) Time Delay Igniter (Lead Shear Wire) (pp 174–75 with Fig 246)
c) Friction Delay Igniter—Miccia 40 and 60 (p 175 with Fig 247)
d) 50-Day Clock (pp 175–76 with Fig 248)

**Acido d’argento**. See Azotidrato d’argento

**Acido azotidrico** (Hydrazoic Acid) is briefly described in Ref 31, p 429

**Acido nitrico-sulfurico** (Nitric-Sulfuric Acid) is described in Ref 31, pp 159–64

**ACIDO PICRICO o TRINITROFENOLE**

[Picric Acid, abbrd PA or Trinitrophenol (TNPh)], also called in Ita *Meilinata o Pertita*, HO.C₆H₂(NO₂)₃. Its prepn, props, uses and analytical procedures are described by Belgrano (Ref 17, pp 281–87). Its props are: Density (max) 1.68, Explosion Temperature 310°, Temperature of Explosion 3200°, Heat of Explosion 1040kcal/kg, Volume of gas at 0° & 760mm 675 l/kg, Specific Pressure 8900 atm/kg, Trauul Test 315cc, Detonation Velocity 7100m/s and Impact Sensitivity with 2kg wt 40cm. Straight PA has been used for loading 100mm, 120mm & 149mm Shells and in composite expls, such as MABT, MAT, MBT, Polvere verde & Victorite (Ref 28, p 318).

**Acido di piombo**. See Azotidrato di piombo (Lead Azide, abbrd as LA)

**Acido stignico o Trinitroresorcina** (Styphnic Acid, abbrd as StA), (HO)₂C₆H(NO₂)₃. It is described in Ref 31, pp 453 to 455 and in Ref 28, p 318. Used in the form of its lead salt *Stignato di piombo* (qv), meaning Lead Stypnate, abbrd LSt.

**Afocita**. A blasting expl compn existing in two formulations: 1) AN (Ammonium Nitrate) 58–62, KN (Potassium Nitrate) 28–31, carbon 7–9 & sulfur 2–3%; 2) AN 58–62, KN 31–38, charcoal 3.5–4.5, sulfur 2–3 & moisture 1.5% (Ref 28, p 318). The 1st formulation is also given by Belgrano (Ref 31, p 340)

**Ager** (Explosivi) expls are based on AN (or other inorg nitrates) and aromatic nitro-compounds. Three compns are listed in the catalog of Società Vulcania at Brescia which gives their props & compns with numerical values: Ager C—Trauul test value 320cc & detonation velocity 2800m/s; suitable for use in quarries, with rocks of medium hardness. Ager D—Trauul value 400cc & deton vel 4700m/s; suitable for hard rocks. Ager E—
Trauzl value 450cc & deton vel 3823m/s; suitable for permissible (See also Ref 31, p 336)

**Albite.** A white fusible mixture consisting of AN 60, NGU (Nitroguanidine) 20 & GuN (Guanidine Nitrate) 20%, introduced during WWI by Manuelli & Bernardini and used as bursting chge in some shells (Ref 4, pp 342–44). The compn of Albite used during WWII for loading shells was AN 58.6, NGU 19.1 & GuN 22.3% (Ref 24, Vol 1, p A120-L; Ref 28, p 318 & Ref 31, p 254)

**Alfa.** Sporting propellant similar to Schultz’s smokeless powder (Ref 31, p 581)

**Alti esplosive.** High Explosives (HE’s). See Esplosivi alti

**Alvisi esplosivi.** See Esplosivi Alvisi

**Amatolo,** described in Ref 4, pp 240–41, is similar to Amer or Brit *Amatol* described in Ref 24, Vol 1, pp A158-L to A164-L. Most popular was the mixture “tipo 60/40” (called Esplosivo 60/40), which contd AN 60 & TNT 40%. Its deton vel is 6500m/sec, and it was used cast-loaded in shells. Less powerful (deton vel 4500m/s) was “tipo 80/20”, which was used press-loaded in sea-mines (Ref 4, p 241). Still less powerful was 90/10 mixt with deton vel 2500m/s, used in mining operations (Ref 28, p 319 and Ref 31, p 243) (Compare with MNDT, MST and Nougat)

**Ammonafite.** Mining expl contg AN 76.7, NG 15.0, Colloid Cotton 0.3 & grain flour 8.0% with 0.5% of yellow ocher incorporated (Ref 31, p 83)

**Ammonal.** described in Ref 4, pp 237–40, was similar to Austrian, Brit, German, French and Amer Ammonals described in Ref 24, Vol 1, pp A287-L to A292-R. The Ital Army used during WWII the mixture: AN 71–72, Al powder 22 & tar (bitume) 6–7%, under the name *Nitramite* (Ref 4, p 238). Several formulations used during WWII contd AN 46–64, Al 17–22, TNT 15–30 & carbon 3%. Such a compn was known as *Toluol-ammonial* or *T-ammonal.* TNT (Trinitronaphthalene) was used in some compns in lieu of TNT (Ref 28, p 319 and Ref 31, p 367)

**Ammondinamite.** See under DYNAMITE

**Ammondite.** Same as Ammonite No 1

**Ammongelatina.** See under DYNAMITE

**Ammonite No 1.** A blasting expl consisting of AN 88, DNT (Dinitrotoluene) 3, NG 3, vegetable flour 5 & DPhA (Diphenylamine) 1% (Ref 28, p 319)

**AN—FO (Ammonium Nitrate—Fuel Oil),** called in Italy *NA—OC* (Nitrat ammonico—Olio combustibili). Belgrano (Ref 31, pp 318–19) gives a standard compn consisting of “prilled” AN of density 0.70–0.75,94.4 and combustible oil 5.6%, Trauzl test value 295cc and detonation velocity ca 400m/s

**Anigrina lamellare.** Single-base smokeless, sporting procplnt with NC completely gela-tinized (Ref 31, p 580)

**ANS (Esplosivo).** Same as ASN

**Antiflame (Antiflame or Flame Suppressor) and Refrigerenti (Cooling Agents).** Belgrano (Ref 31, pp 219–20) lists: vaseline, mineral oils, acardite, NGU (Nitroguanidine), Nitronaphthalene, inorganic chlorides, Amm sulfate, oxalates of Amm, Na & K and K tartrate as cooling agents or flame suppressors for Dynamites

**Antigrandine: Esplosivi, Cannoni e Razzi.** (Antihail: Explosives, Cannons and Rockets). As bursting charge explosives for antihail projectiles, TNT and some Cheddites were used in Italy. An antihail cannon is illustrated on p 337 of Ref 31, whereas a rocket is on p 626 *Addn Ref from CA:* L. Peseni, ItalP 518413 (1955) & CA 51, 17170 (1957) (Antihail rockets designed to reach an altitude of 1800m are supplied with BnPdr mixture contg KNO₃ 24, charcoal 5 & sulfur 0.65kg)

**Antigrisou (Esplosivi) o Esplosivi antigrisutosi (Permissible Explosives).** The following are
listed in Ref 31, p 326: Antigrisou N.0 – AN 80.57, DNN (Dinitronaphthalene) 6.36 & AChl (Ammonium Chloride) 13.02%; Antigrisou N.2 – AN 81.49, DNN 11.11 & AChl 7.40%; Antigrisou N.3 – AN 82, TNN (Trinitronaphthalene) 5 & AChl 13%

Requirements for Italian permissible explosives are given by Belgrano (Ref 31, pp 327–29), while tests are described on pp 329–32

Antisanzionite. See ASN

Antonite (Explosivi). Mining expls manufd by the Società Vulcania at Brescia. Two compns are listed in their catalog of 1960; in Ref 24, Vol 1, p A473-L; and in Ref 31, p 317:

Antonite per cava — for work in quarries — consists of AN & TNT in proportion to make O content 5.9% by weight; Trauzl test value 340cc and deten vel 3400m/s
Antonite per galleria — for work in tunnels — consists of AN & TNT in proportion to make O content 3.08%; Trauzl value 415cc and deten vel 4000m/s

Aquila. A smokeless, sporting propInt with completely gelatinized NC (Ref 31, p 580)

Ares. A powdery mining expl based on AN & a combustible (Ref 31, p 317)

Aria liquida e Ossigeno liquido (esplosivi). Liquid Air and Liquid Oxygen Explosives, such as Oxylquiri or LOX have been used in high mountain hydroelectric works by the Ital Corps of Engineers (Genio Militare) (Ref 15, p 38 & Ref 31, p 305)

ARMI PORTATILI MUNIZIONI (Small-Arm Ammunition). The ammunition used during WWII was similar to British and German ammo. The following types are described in TM 9-1985-6 (1953) (Ref 16, pp 65–72): 6.5-mm Ball (p 67, Fig 76); 7.35-mm Ball (p 67, Fig 77); 7.7-mm Ball (p 68, Fig 78); 7.7-mm API (Armor-piercing Incendiary) (Blue Tip) (p 68, Fig 79); 7.7-mm API (Incendiary) (Green Tip) (p 69, Fig 80); 8-mm Ball (p 69, Fig 81); 8-mm AP (Armor-piercing) (p 70, Fig 82); 12.7-mm Tracer (Red Tip) (p 70, Fig 83); 12.7-mm Incendiary (Blue Tip) (p 71, Fig 84); 12.7-mm APIT (Armor-piercing Incendiary Tracer) (White Tip) (p 72, Fig 85); 12.7-mm HE (High Explosive) (p 72, Fig 86)

In “MUNIZIONAMENTO ITALIANO” (Addnl Ref C) under Cartuccie (plural Cartucci) are listed Cartuccie Cal 6.5 per Moschetto (Carbine) Mod 91/38; Cartuccie Cal 7.35 per Fusil (Rifle) e Moschettone (Carbine) Mod 38; Cartuccie Cal 7.65 per Pistola Berretta; Cartuccie Cal 8 per Mitragliatrice (Machine Gun) Mod 35 e Breda Mod 37; Cartuccie Cal 9 per Mitragliatrice Mod 38; Cartuccie Cal 9 per Pistola automatica Berretta Mod 34; Cartuccie Cal 10.35 per Pistola Mod 39; Cartuccie Cal 13.2 per Mitragliera d’Aereo (Aerial Machine Gun)

Actually cartucci are complete rounds of small-arms ammunition, while cartocci are complete rounds of artillery ammunition

Armi subacquee (Underwater Weapons). They include: siluri (submarine torpedoes), bombe torpedini di profondità (depth bombs), mine subacquee (underwater mines), etc. Their teste (warheads) can have as bursting charge (carica esplosiva) Tritolital (TNT 60, RDX 20 & Al 20%); Tritolito (RDX 40 or 60 incorporated in fused TNT 60 or 40%) or Torpex (RDX 44, TNT 38 & Al powder 18%) (Ref 31, p 382)

Artifici de guerra — Military Pyrotechnics. See under PIROTECNIA

ARTIGLIERIA ITALIANA — Italian Artillery

Its history is described in the book of General Carlo Montu (Ref 3). Artillery pieces used during WWI & WWII are described by the late Col J.B. Jarrett in Ref 5a (illustrated)

It would be well to remember that Italian practice is to refer to the caliber in millimeters followed by the tube length in calibers. Thus, 65/17 means a 65-mm bore diam and a tube 17 calibers long

Following Chart is copied in abbreviated form from Ref 5a, p 668
### CHART 1
Breakdown of Important Data Concerning the Most Used Italian Pieces

<table>
<thead>
<tr>
<th>Designation</th>
<th>MV (ft sec)</th>
<th>Range (yds)</th>
<th>Ammunition</th>
</tr>
</thead>
<tbody>
<tr>
<td>.65/17</td>
<td>1,140</td>
<td>7,100</td>
<td>HE, AP-HEC&amp;BC, Hollow chge</td>
</tr>
<tr>
<td>75/13</td>
<td>1,240</td>
<td>9,800</td>
<td>HE, AP-HEC&amp;BC, Shrapnel</td>
</tr>
<tr>
<td>75/18, Model 34</td>
<td>1,300</td>
<td>10,300</td>
<td>HE, AP-HEC&amp;BC, Shrapnel, Hol chge</td>
</tr>
<tr>
<td>(This piece was found on 3 carriage models, and was also used on the Semovente or S.P. M13/40 tank version)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>75/27, Model 11</td>
<td>1,640</td>
<td>11,100</td>
<td>HE, AP-HEC&amp;BC, Shrapnel, Hol chge</td>
</tr>
<tr>
<td>(A modification of this weapon was seen on S.P. carriages)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>75/34</td>
<td>1,650</td>
<td>13,500</td>
<td>HE, AP-HEC&amp;BC, Hollow chge (streamlined shells)</td>
</tr>
<tr>
<td>77/28 (Originally Krupp 1896 design)</td>
<td>1,762</td>
<td>7,300</td>
<td>HE, shrapnel and case shot</td>
</tr>
<tr>
<td>100/17, Model 14</td>
<td>1,400</td>
<td>10,100</td>
<td>HE Shrapnel, Hollow chge</td>
</tr>
<tr>
<td>105/28</td>
<td>1,880</td>
<td>15,000</td>
<td>HE (several patterns flat based and streamlined)</td>
</tr>
<tr>
<td>149/12, Model 14</td>
<td>1,100</td>
<td>7,500</td>
<td>HE and Shrapnel (several patterns of HE)</td>
</tr>
<tr>
<td>149/13, Model 14</td>
<td>1,100</td>
<td>9,600</td>
<td>HE and Shrapnel (several patterns of both)</td>
</tr>
<tr>
<td>149/17</td>
<td>1,660</td>
<td>12,500</td>
<td>HE, Shrapnel (several patterns)</td>
</tr>
<tr>
<td>149/40 (Modern gun)</td>
<td>2,600</td>
<td>23,900</td>
<td>HE (2 patterns)</td>
</tr>
<tr>
<td>152/37 Howitzer</td>
<td>1,300</td>
<td>10,400</td>
<td>HE (several patterns)</td>
</tr>
<tr>
<td>152/37</td>
<td>2,270</td>
<td>21,800</td>
<td>HE (several patterns)</td>
</tr>
<tr>
<td>210/22</td>
<td>1,870</td>
<td>17,400</td>
<td>HE and an AP (likely anti-concrete)</td>
</tr>
<tr>
<td>(Not seen in Africa)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>305/8 (Skoda mortar 1916 and not seen in Africa)</td>
<td>1,300</td>
<td>12,000</td>
<td>HE (base fuzed and Shrapnel)</td>
</tr>
</tbody>
</table>

### Anti-aircraft Artillery (AA Guns)

<table>
<thead>
<tr>
<th>Vertical Range (feet)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2-cm Breda</td>
</tr>
<tr>
<td>2-cm Scotti</td>
</tr>
<tr>
<td>37/54 Breda</td>
</tr>
<tr>
<td>75/27 Krupp</td>
</tr>
<tr>
<td>75/46 Ansoldo, Model 34</td>
</tr>
<tr>
<td>75/50 Skoda</td>
</tr>
<tr>
<td>75/53 French, Model 30</td>
</tr>
<tr>
<td>76/40 Ansoldo</td>
</tr>
<tr>
<td>90/53 Ansoldo</td>
</tr>
</tbody>
</table>
Following addnl info is taken from p 669 of Ref 5a:

CHART II

Antitank (A/T Guns)
47/32 Breda
47/32 Austrian Boehler
47/50 Schneider (French captured)
75/27 Series, using AP, Hollow charge
75/46 AA Dual purpose
90/53 AA Dual purpose
100/17 series, using AP, Hollow charge

Seacoast Guns
57/30
57/43
120/21 Same ammunition
120/25
120/40
149/35
152/32 (two versions)
152/50 Same ammunition
280/9
280/10 Same ammunition
280/11
280/16
305/17
305/50
381/40
420/12

Abbreviations:
AP = Armor-piercing
A/T = Antitank
HE = High Explosive
HEBC = High Explosive, Ballistic Cap
HEC = High Explosive, Capped
Hol chge = Hollow (or Shaped) Charge
PD = Point Detonating Fuze
SD = Self-destroying
SP = Self-propelled

Note: In the opinion of Col G.B. Jarrett, the Italian artillery support to other arms performed poorly during WWII, even though the Italian gunners were usually very brave men. Briefly, their troubles lay in poor design of both carriages and ammunition (Ref 3, pp 663–64) (See also CANNONE)

ASN of Tonegutti, proposed after WWI consisted of AN 70, PETN 20 & DCyDA 10%. It melted at 115° and was suitable for cast-loading shells (Ref 4, p 245 & Ref 24, Vol 1, p A496-L). ASN of Tonegutti, proposed before WWII, consisted of: AN 60, PETN 20, DCyDA 10 & GuN (Guandine Nitrate) 10%. It has a low mp (10^c) due to the presence of DCyDA & GuN. It was used during WWII for loading Naval shells. When used in underwater ammunition, such as torpedoes, depth charges & sea mines, its efficiency was increased by incorporating some Al powder (Ref 28, p 320 and Ref 31, p 181 under Antisanzionate)

Aster. A smokeless, sporting proplnt with NC completely gelatinized (Ref 31, p 580)

Astralita 1 e 2 were Ialt AN+TNT+NG+woodmeal exps manufd during WWI by the Società Dinamite Nobel, Avigliana (Ref 15, p 32). They were used as fillers for trench mortar shells and hand grenades. The expls were similar to German Astralits listed in Ref 24, Vol 1, pp A497-R & A498-L. Belgrano (Ref 31) lists on p 122 the ballistic mortar strength of Astralita antigrisou 3^e classe as 4.75% of Gelatina 92/8 (Sprenggelatine), but gives no compn

Accdg to Ref 28, p 320, Astralites are now used in mining operations

Avigliana 3. See Nitramite

Azotidrato d’argento (Silver Azide–SA) or Acido d’argento, AgN₃, is described in Belgrano (Ref 31, pp 449–51). Trauzl value for 2g sample 22.5cc, Detonation Velocity 5700m/sec, Temperature of Explosion 3545° & Detonation Temperature 297° (Ref 31, pp 447 & 451). Small quantities of SA are used in primers (Ref 28, p 318) (See also Ref 24, Vol 1, p A597-R to A601-R)

AZOTIDRATO DI PIOMBO (Lead Azide–LA) or Acido di piombo, Pb(N₃)₂, is described by Belgrano (Ref 31, pp 439–49). Its props are: Density (max) 4.79, Explosion Temperature
327°, Temperature of Explosion (Flame Temperature at Explosion) 3350°, Specific Volume 310 l/kg, Heat of Formation -364 kcal/kg, Impact Sensitivity with 2 kg wt 8 cm, Trauzl with 10 g sample 115 cc and Detonation Velocity 5300 m/sec. It was manifold by Nobel SGEM at Tiana, near Turino and by Bombrini-Parodi-Delfino SA at Colloferro-Roma. LA nearly entirely replaced MF (Mercuric Fulminate) as an initiating detonating agent of priming comps. It can be used alone but is preferable to use it with LSt (Lead Styphnate) which is more sensitive to flame initiation (Ref 28, p 318) (See also Ref 24, Vol 1, pp A545-L to A563-L)

Balistite (Ballistite). A double-base propellant with high NG content was invented in 1887–1888 by A. Nobel. Several formulations were developed later and used in many countries. The varieties used in Italy are listed in Ref 24, Vol 2, pp B8-R & B9-L. Ballistite al 60, which contd NG 60 & NC 40% with 1–2% stabilizer added, was too erosive for use as a proplnt in guns, but proved to be very suitable for use as a bursting charge in some smaller caliber shells, such as 37/40mm HE and 37/40mm HEP (High-Explosive Armor-Piercing)

Belgrano (Ref 31,) lists on p 205: Ballistite ordinario contains NG 50 & CC (Cotone colloidio) 50%; Ballistite a basso titolo cont NG 42 with CC & other ingredients; Ballistite attenuata cont NG 25, Pircocolloidio 60 & liquid DNT 15%. Every compn containd ca 0.5% DPhA (Diphenylamine) added (See also Ref 4, p 181)

Bassi esplosivi. See Esplosivi bassi

Bersaglio. Target

Bicchere. Projectile's Case or Body. See PROIETTO o PROIETTILE

Blasting Gelatin. See Gomma A

BM (Esplosivi Mangiarotti). A series of mining expls manufd by the Società Mangiarotti, Codroipo (Udine). The following formulations are listed in their catalog of 1960: BM, 1, per galleria — gray powder consisting of TNT, AN & thermite; Trauzl test value 440 cc & deton vel 3550m/sec; used in galleries not contg firedamp

BM, as, per uso a cielo aperto — reddish pdr consisting of AN, with cyclic aliphatic compds & metallic pdr's; Trauzl value 470cc & deton vel 4100m/sec; used in open-cut quarries with rocks of medium hardness

BM, ac, per uso a cielo aperto — brownish-yel pdr consisting of AN & TNT; Trauzl value 450cc & deton vel 3800m/sec; used in open-cut quarries

BM, 57, per uso a cielo aperto — grayish-black pdr based on AN; Trauzl value 370cc & deton vel 2050m/sec; used in open-cut quarries with materials of medium hardness, such as limestone, clay, soil for cement, etc

Super BM, per galleria — green pdr, sl plastic contg aromatic nitrocompounds with org nitrates and plasticizers; Trauzl value 420cc & deton vel 5000m/sec; used for blasting in galleries contg no firedamp

BM, a2 — brown pdr contg AN, TNT & thermite; Trauzl value 440cc & deton vel 3000m/sec; used in open-cut quarries with soft & medium hard rocks; also in agriculture, such as for destruction of tree stumps, etc

Super BM, Cava — gray pdr of density 1.20; compn not given; Trauzl value 510cc & deton vel 4600m/sec; used in quarries for blasting very hard materials such as granite

BM, 2 per galleria — gray pdr contg inorganic nitrate, org nitrocompounds and Al pdr; Trauzl value 450cc & deton vel 4100m/sec; used in gallerias contg no firedamp

Note: BM is mentioned in Ref 31, p 317 w/o giving compn or props

Boceda. Sporting smokeless proplnt with completely gelatinized NC (Ref 31, p 580)

Bomba atomica is briefly described by Belgrano (Ref 31, pp 519–20). General description is given by C.G. Dunkle in Ref 24, Vol 1, pp A499-L to A500-L. Historical development is discussed on p A500-L, under “Atomic (or Nuclear) Energy”

BOMBA (plural Bombe) — Bomb (Bombs), Conventional

Italian bombs of WWII are described in OP 1668 (Ref 8, pp 1–27) and in TM 9-
1985-6/TO 39B-1A-8 (Ref 16, pp 1–27). They were constructed of more than one piece, being assembled by screws or rivets, or welded. They were usually filled thru the base, which was closed by a base plate, attached by screws or rivets. The bombs were generally constructed of sheet steel but in some cases aluminum alloy was used. Demolition bombs were usually constructed of mild steel, while armor-piercing bombs (AP) were made of hardened steel. The anti-personnel (A/P) bombs differed from demolition bombs in construction; in one type (Type F) the filling was enclosed in a sheet container, on the outside of which a steel strip was wound spirally, while in another type (Type Mtr) the filling was contained in a sheet-steel case which was enclosed in a larger container. The space between the two containers was filled with steel fragments.

The majority of bombs contained tail fuzes but quite a number contained nose fuzes. Some medium and large bombs, eg, 100, 250, 500, 800 and 1000 kg, contained both tail and nose fuzes.

The principal bomb filling was cast Tritol (TNT); however, quite a few bombs were filled with Amatol; and, in the case of shaped-charge bombs, with a mixture of T₄ (RDX) 60, TNT 38 & wax 2%.

The bombs were either galvanized or painted to protect them from corrosion. The following colors were used to identify different types:

<table>
<thead>
<tr>
<th>Type of Bomb</th>
<th>Color of Body</th>
<th>Color of Nose</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fragmentation (F)</td>
<td>Blue</td>
<td>Red</td>
</tr>
<tr>
<td>High Explosive (HE)</td>
<td>Grey</td>
<td>Red</td>
</tr>
<tr>
<td>Anti-personnel (A/P)</td>
<td>Black or Blue</td>
<td>Red</td>
</tr>
<tr>
<td>Incendiary (IB)</td>
<td>Reddish-brown</td>
<td>Red</td>
</tr>
<tr>
<td>Gas (G)</td>
<td>Bright Yellow</td>
<td>Red</td>
</tr>
<tr>
<td>Practice</td>
<td>Grey</td>
<td>Grey</td>
</tr>
</tbody>
</table>

Following are Italian terms for bombs and for some grenades:

- **Bomba a mano**
- **Bomba a mano contro i carri armati**
- **Bomba a mano dirompente**
- **Bomba antisommergibile; (bomba da getto; bomba di profondità o bomba torpedine)**
- **Bomba da demolizioni**
- **Bomba da tromboncino**
- **Bomba dirompente**
- **Bomba fumogeno**
- **Bomba incendiaria**
- **Bomba leggera**
- **Bomba luminosa**
- **Bomba perforante**
- **Granata bomba**
- **Bomba da esercitazione**

- Hand grenade
- Anti-tank grenade
- Fragmentation grenade
- Depth bomb or depth charge
- Demolition bomb
- Rifle grenade
- Fragmentation bomb
- Smoke bomb
- Incendiary bomb
- Illuminating bomb
- Flash bomb
- Armor-piercing bomb
- Demolition bomb
- Practice bomb
Table
Listing Names, Dimensions, Explosive Charges and Fuzes Used in Italian Bombs of WWII

<table>
<thead>
<tr>
<th>American Designation of Bombs</th>
<th>Italian Designation of Bombs</th>
<th>Overall Length (inches)</th>
<th>Maximum Diameter (inches)</th>
<th>Type of Filling</th>
<th>Weight of Filling (kg)</th>
<th>Total Weight (kg)</th>
<th>Fuze</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 kg A/P</td>
<td>F</td>
<td>6.0</td>
<td>4.5</td>
<td>TNT</td>
<td>0.380</td>
<td>1.72</td>
<td>Tail, Type K</td>
</tr>
<tr>
<td>2 kg A/P</td>
<td>Mtr</td>
<td>6.0</td>
<td>4.5</td>
<td>TNT</td>
<td>0.220</td>
<td>1.87</td>
<td>Tail, Type K</td>
</tr>
<tr>
<td>3 kg A/P</td>
<td>Mtr</td>
<td>12.1</td>
<td>8.2</td>
<td>TNT</td>
<td>0.17</td>
<td>3</td>
<td>Nose, Type M</td>
</tr>
<tr>
<td>4 kg A/P</td>
<td>Manzolini (Thermos)</td>
<td>12.3</td>
<td>7.3</td>
<td>TNT</td>
<td>0.67</td>
<td>3.68</td>
<td>Manzolini Types I &amp; II</td>
</tr>
<tr>
<td>12 kg A/P</td>
<td>F</td>
<td>32.4</td>
<td>17.3</td>
<td>TNT</td>
<td>1.93</td>
<td>12.20</td>
<td>Nose, Type F</td>
</tr>
<tr>
<td>12 kg A/P</td>
<td>Mtr</td>
<td>3.5</td>
<td>3.5</td>
<td>TNT</td>
<td>1.90</td>
<td>12.88</td>
<td>Nose, Type J</td>
</tr>
<tr>
<td>14 kg Frag</td>
<td>I</td>
<td>22.1</td>
<td>4.2</td>
<td>TNT</td>
<td>2</td>
<td>14</td>
<td>Nose, Type I</td>
</tr>
<tr>
<td>14 kg Frag</td>
<td>II</td>
<td>22.1</td>
<td>4.2</td>
<td>TNT</td>
<td>2</td>
<td>14</td>
<td>Nose, Type I</td>
</tr>
<tr>
<td>24 kg GPHE</td>
<td>--</td>
<td>30.5</td>
<td>6.4</td>
<td>TNT</td>
<td>12</td>
<td>24</td>
<td>Tail, Type N-1</td>
</tr>
<tr>
<td>40 kg GPHE</td>
<td>--</td>
<td>32.3</td>
<td>9.0</td>
<td>TNT</td>
<td>12</td>
<td>40</td>
<td>Tail, Type N-3</td>
</tr>
<tr>
<td>50 kg GPHE</td>
<td>--</td>
<td>40.5</td>
<td>9.9</td>
<td>Amatol</td>
<td>29.20</td>
<td>59.31</td>
<td>Tail, Type C or Y</td>
</tr>
<tr>
<td>100 kg GPHE</td>
<td>--</td>
<td>51.3</td>
<td>10.7</td>
<td>TNT</td>
<td>50.6</td>
<td>100</td>
<td>Tail, Type C or Y</td>
</tr>
<tr>
<td>100 kg GPHE</td>
<td>--</td>
<td>51.3</td>
<td>10.7</td>
<td>Amatol</td>
<td>49.5</td>
<td>100</td>
<td>Tail, Type C or Y</td>
</tr>
<tr>
<td>250 kg GPHE</td>
<td>--</td>
<td>73.8</td>
<td>17.6</td>
<td>TNT</td>
<td>125.7</td>
<td>259.1</td>
<td>Nose, Type A and Tail, Type O</td>
</tr>
<tr>
<td>500 kg GPHE</td>
<td>--</td>
<td>96.6</td>
<td>18.0</td>
<td>TNT</td>
<td>216</td>
<td>508</td>
<td>Nose, Type A and Tail, Type O</td>
</tr>
<tr>
<td>800 kg GPHE</td>
<td>--</td>
<td>127.8</td>
<td>18.0</td>
<td>Unknown</td>
<td>357</td>
<td>821.6</td>
<td>Nose, Type A and Tail, Type O</td>
</tr>
<tr>
<td>1000 kg GPHE</td>
<td>--</td>
<td>140.0</td>
<td>20.5</td>
<td>TNT</td>
<td>Unknown</td>
<td>1000 (appr)</td>
<td>Nose Fuze Only</td>
</tr>
<tr>
<td>500 kg GPHE Time Bomb</td>
<td>--</td>
<td>93.7</td>
<td>18.1</td>
<td>TNT</td>
<td>247.0</td>
<td>Unknown</td>
<td>Nose and Tail Fuzes</td>
</tr>
<tr>
<td>15 kg SAP</td>
<td>--</td>
<td>31.0</td>
<td>4.7</td>
<td>TNT</td>
<td>5.2</td>
<td>15.5</td>
<td>Tail, Type N</td>
</tr>
<tr>
<td>31 kg SAP</td>
<td>--</td>
<td>31.7</td>
<td>6.4</td>
<td>TNT</td>
<td>10.5</td>
<td>31.0</td>
<td>Tail, Type N-2</td>
</tr>
<tr>
<td>100 kg SAP</td>
<td>--</td>
<td>50.5</td>
<td>9.9</td>
<td>Amatol</td>
<td>27.3</td>
<td>109.0</td>
<td>Tail Fuze, Type C-1 or Y-1</td>
</tr>
<tr>
<td>104 kg SAP</td>
<td>--</td>
<td>43.0</td>
<td>10.0</td>
<td>TNT</td>
<td>30</td>
<td>104.0</td>
<td>Tail, Type C</td>
</tr>
<tr>
<td>12 kg Sm</td>
<td>--</td>
<td>47.0</td>
<td>5.2</td>
<td>Sm compn</td>
<td>–</td>
<td>28</td>
<td>Nose Fuze</td>
</tr>
<tr>
<td>0.5 kg Inc</td>
<td>IP</td>
<td>6.1</td>
<td>2.75</td>
<td>Cotton wicks soaked in gasoline Thermite</td>
<td>–</td>
<td>0.5</td>
<td>Tail, Type K</td>
</tr>
<tr>
<td>0.5 kg Inc</td>
<td>IT</td>
<td>5.1</td>
<td>2.5</td>
<td>Thermite</td>
<td>–</td>
<td>0.5</td>
<td>Tail, Type K</td>
</tr>
<tr>
<td>0.5 kg Inc</td>
<td>FI</td>
<td>4.9</td>
<td>2.5</td>
<td>Phosph</td>
<td>–</td>
<td>0.5</td>
<td>Tail, Type K</td>
</tr>
<tr>
<td>1 kg Inc</td>
<td>I &amp; II</td>
<td>6.1</td>
<td>2.7</td>
<td>Mg powd Thermit</td>
<td>0.084</td>
<td>1</td>
<td>Tail, Type K</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(Continued)
<table>
<thead>
<tr>
<th>American Designation of Bombs</th>
<th>Italian Designation of Bombs</th>
<th>Overall Length (inches)</th>
<th>Maximum Diameter (inches)</th>
<th>Type of Filling</th>
<th>Weight of Filling (kg)</th>
<th>Total Weight (kg)</th>
<th>Fuze</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 kg Inc</td>
<td>I</td>
<td>12.2</td>
<td>2.7</td>
<td>Thermit Oil Mg, HgO &amp; MNB</td>
<td>0.321</td>
<td>2.12</td>
<td>Tail, Type K</td>
</tr>
<tr>
<td>2 kg Inc</td>
<td>II</td>
<td>12.2</td>
<td>2.7</td>
<td>-</td>
<td>0.339</td>
<td>Unknown</td>
<td>Tail, Type K</td>
</tr>
<tr>
<td>20 kg Inc</td>
<td></td>
<td>34.0</td>
<td>6.3</td>
<td>Thermit</td>
<td>10.58</td>
<td>20.17</td>
<td>Tail, Type E</td>
</tr>
<tr>
<td>70 kg Inc</td>
<td></td>
<td>47.2</td>
<td>9.9</td>
<td>Thermit</td>
<td>36.6</td>
<td>74.5</td>
<td>Tail, Type E</td>
</tr>
<tr>
<td>100 kg Spec</td>
<td></td>
<td>532.0</td>
<td>10.7</td>
<td>TNT</td>
<td>Unknown</td>
<td>113.0</td>
<td>Nose, Type X &amp; or</td>
</tr>
<tr>
<td>(Combination A/P bomb &amp; bomb container)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Tail, Type Z</td>
</tr>
<tr>
<td>15 kg Gas</td>
<td></td>
<td>31.0</td>
<td>4.7</td>
<td>DPhClAr</td>
<td>3.65</td>
<td>16</td>
<td>Tail Fuze</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>TNT (burster)</td>
<td>1.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>25 kg Gas</td>
<td>Furretta</td>
<td>32.7</td>
<td>6.3</td>
<td>Lacrym</td>
<td>10.0</td>
<td>25</td>
<td>Tail, Type K</td>
</tr>
<tr>
<td></td>
<td>(C40 P)</td>
<td>32.3</td>
<td>9.9</td>
<td>DPhClAr</td>
<td>6.5</td>
<td>47</td>
<td>Tail Fuze</td>
</tr>
<tr>
<td></td>
<td>(A)</td>
<td>32.3</td>
<td>9.9</td>
<td>Burster</td>
<td>13.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>(C40 P)</td>
<td>32.3</td>
<td>9.9</td>
<td>Mustard</td>
<td>14.7</td>
<td>40.6</td>
<td>Tail Fuze</td>
</tr>
<tr>
<td></td>
<td>(B)</td>
<td>32.3</td>
<td>9.8</td>
<td>Burster</td>
<td>0.18</td>
<td></td>
<td></td>
</tr>
<tr>
<td>55 kg Gas</td>
<td></td>
<td>32.3</td>
<td>9.8</td>
<td>Phosgene Burster</td>
<td>20</td>
<td>55</td>
<td>Unknown</td>
</tr>
<tr>
<td></td>
<td></td>
<td>50.2</td>
<td>10.7</td>
<td>DPhClAr</td>
<td>14.3</td>
<td>101.9</td>
<td>Tail Fuze</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Burster</td>
<td>28.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>100 kg Gas</td>
<td>(C100 P)</td>
<td></td>
<td></td>
<td>Mustard</td>
<td>214</td>
<td>264</td>
<td>Nose Fuze</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(appr)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>250 kg Gas</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Nose Fuze</td>
</tr>
<tr>
<td>500 kg Gas</td>
<td>(S00T)</td>
<td>96.6</td>
<td>18.0</td>
<td>DPhClAr</td>
<td>210</td>
<td>298</td>
<td>Nose, Type T</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Inc</td>
<td>Unknown</td>
<td>5.2</td>
<td>Nose, Type S</td>
</tr>
<tr>
<td>5 kg Vento -Marker(a)</td>
<td></td>
<td>17.4</td>
<td>5.2</td>
<td>Sm or</td>
<td>Unknown</td>
<td></td>
<td></td>
</tr>
<tr>
<td>160 kg A/S</td>
<td>(CS)</td>
<td>69.8</td>
<td>13.3</td>
<td>TNT</td>
<td>99</td>
<td>176</td>
<td>Nose &amp; Tail Fuzes</td>
</tr>
<tr>
<td>160 kg HE</td>
<td></td>
<td>62.4</td>
<td>12.6</td>
<td>TNT</td>
<td>Unknown</td>
<td>163.5</td>
<td>Nose Fuze</td>
</tr>
<tr>
<td>3 kg AA</td>
<td>(CV)</td>
<td>13.5</td>
<td>3.2</td>
<td>Amatol</td>
<td>0.40</td>
<td>3.0</td>
<td>Nose, Type I</td>
</tr>
<tr>
<td>20 kg AA</td>
<td>(CV)</td>
<td>30.7</td>
<td>5.5</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Nose, Type I</td>
</tr>
<tr>
<td>3.5 kg HoC</td>
<td></td>
<td>18.8</td>
<td>6.0</td>
<td>RDX 60</td>
<td>22</td>
<td>3.5</td>
<td>Tail Fuze</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>TNT 38</td>
<td>(appr)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Wax 2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5 kg HoC</td>
<td></td>
<td>Unknown</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Tail Fuze</td>
</tr>
<tr>
<td>25 kg HoC</td>
<td></td>
<td>Unknown</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Tail Fuze</td>
</tr>
<tr>
<td>50 kg HoC</td>
<td></td>
<td>Unknown</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Tail Fuze</td>
</tr>
<tr>
<td>100 kg HoC</td>
<td></td>
<td>Unknown</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Tail Fuze</td>
</tr>
<tr>
<td>Bomb Container (b)</td>
<td></td>
<td>5’6”</td>
<td>11</td>
<td>AP or Inc Mg</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Nose, Type L</td>
</tr>
<tr>
<td>Parachute</td>
<td>Martellona</td>
<td>36.9</td>
<td>4.0</td>
<td>(powder)</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Friction Igniter for Nose</td>
</tr>
<tr>
<td>Flare</td>
<td></td>
<td>12.5</td>
<td>1.45</td>
<td>Unknown</td>
<td>Unknown</td>
<td>Unknown</td>
<td></td>
</tr>
<tr>
<td>Parachute</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Flare (cardboard)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Regulations: AA Antiaircraft; AP Armor-piercing; A/P Antipersonnel; appr approximately; A/S Antisubmarine; C, CS and CV Italian designations for bombs; Comp Composition; Design Designation; diam diameter; DPhClAr Diphenylchloroarsine; F Ital designation (See note c); Frag Fragmentation; GP General purpose; GPE General purpose high explosive; HE High Explosive; HoC Hollow (shaped) charge; Inc Incendiary; Lacrym Lacrymator; MNB Mononitrobenzene; Mtr Italian designation (see note d); Phosph Phosphorus; SAP Semi-armor-piercing; Sm Smoke; Spec Special

Notes:

a) The Vento Marker was used in conjunction with chemical bombs to indicate the force and direction of the wind at ground level. Two fillings were used, a smoke filling in the daytime and an incendiary filling for use at night.
b) The Bomb Container was a bomb-shaped hollow body of sheet metal, holding 8 columns of A/P or incendiary bombs.
c) "F" Bomb consisted of a thin steel cylinder surrounded by a tightly coiled spring of rectangular cross section. The object of the spring was to provide shrapnel effect. In fragmentation, the bomb usually broke into pieces about 1" x 0.2" x 0.18".
d) "Mtr" Bomb consisted of two cylinders of sheet metal. The inner cylinder contained the explosives, and the outer cylinder was threaded at the top to take a screwed circular cover. Between the two cylinders were small steel pellets embedded in concrete.

Bomb Fuze — Spolette per bombe

Italian Bomb Fuzes used during WWII were mechanically operated, except for Time Fuzes (Spolette a tempo) used in 500-kg Time Bomb. Most of the fuzes functioned on impact. Normally the fuzes were made of brass, steel or Al and were coated with shellac or varnish to prevent corrosion. Both nose and tail fuzes were used.

Technical Manual TM 9-1985-6 (Ref 16) lists twelve types of MINF (Mechanical Impact Nose Fuzes) (pp 29–39); nineteen MITF (Mechanical Impact Tail Fuzes) (pp 40–49 & 54–55); three MTNF (Mechanical Time Nose Fuzes) (pp 49–50); one ETNF (Electrical Time Nose Fuze) (p 51); one CLDNF (Clockwork Long-Delay Nose Fuze) (p 52); one CLDTF (Clockwork Long-Delay Tail Fuze) (p 53); one Tail Fuze for Hollow Charge Bomb (p 56); one Hydrostatic Tail Fuze—"Grand Daddy" (p 57); one MIND—"Orphan" (p 58); and one Mechanical Anti-Disturbance Fuze—"Manzolini" (pp 59–60)

Italian terms for Fuzes are given under Spoletta (plural Spolette)

Booster Charge. See Carica di rinforzo

Borani — Propellenti liquidi per razzi (Liquid Rocket Propellants) consist of liquid hydrogen with boron or other light element, such as beryllium (Ref 31, p 129)

BOSSOLO (plural Boscoli). Accdg to the "Military Dictionary", listed here as Addn Ref A, it can be translated as Cartridge Case, Shell Case or Shrapnel. This signifies that the term bossoho has several meanings and the same conclusion can be drawn after reading the following:

Belgrano (Ref 31, p 611) describes only boscoli da caccia which are cardboard or plastic cartridge cases used in sporting (hunting) ammunition.

Accdg to "MUNIZIONAMENTO ITALIANO", listed as Addn Ref C, boscoli for small arms' ammunition are cartridge cases, usually made of brass, provided with percussion primers at their base. These boscoli serve as containers for propelling charges (cariche di lancio). In case of fixed or semi-fixed artillery ammunition, such as for granata perforante (armor-piercing shell), bossoho is a brass tube located inside a cartridge case and attached at its perforated base to a percussion primer. The bossoho is loaded thru its upper open end with an easily ignitable mixture, usually Black Powder (polvere nera, abbrd as p.n.), which serves to ignite on striking the percussion primer with a trigger, the propelling charge (carica di lancio) in the cartridge case. This type of bossoho may be called a Powder Tube or an Igniter Tube (Compare with Igniter Tubes described in Ref 24, Vol 1, p A385-L and in Vol 2, p C75-R) (See also under CARTOCCIO)

Burster Charge. See Carica di scoppio
C (Polvere). A smokeless propellant with NC of 12.5% N completely gelatinized with ether-alcohol (Ref 31, p 580). Do not confuse with Miscela C (qv).

C-2 (Polvere) o Cordite Italiana. A double-base propellant produced by Società Dinamite Nobel at Avigliana for use during WWI and WWII. Its compn as given in Ref 31, p 210 is NC 70.5, NG 23.5, vaseline 5.0 & Na bicarbonate 1% (See also Ref 4, p 179).

C-12 (Polvere). Ital solventless propInt: NC 68.0, NG 25.0, Centralite 5.0, vaseline 1.0, Na bicarbonate 0.5 & Amm oxalate 0.5% (Ref 4, p 183).


Cadinite. A mining expl consisting of Na nitrate 56, NG 26, sulfur 10 & carbon (or a cellulosic material) 8% (Ref 8, p 321).

Cannel (Polvere). See Polvere Cannel.

CANNELLO (plural Cannelli). Accdg to Military Dictionary (AddnRef A), it is not a Cannon Primer, but it is rather a Projectile Primer. The following types exist:

- Cannello elettrico – Electric Primer
- Cannello a frizione – Friction Primer
- Cannello fulminante – Igniter
- Cannello fulminante a strappo – Pull Igniter
- Cannello fulminante a pressione – Push Igniter
- Cannello a percussione – Percussion Primer

Accdg to AddnRef C, the cannello is a part of the colpo completo (Complete Round of Ammunition).

CANNONE (plural Cannoni). Accdg to Military Dictionary (AddnRef A), it can be translated as Cannon, Gun or Piece (of Artillery).

They may be subdivided into:

- Cannone ad anima liscia – Smoothbore Cannon
- Cannone ad avancarica – Muzzle-loading Gun
- Cannone antiacera – Antiiaircraft Gun
- Cannone anticarro – Antitank Gun
- Cannone a retrocarica – Breech-loading Cannon
- Cannone a tiro rapido – Rapid-fire Gun

Cannone automatico – Automatic Gun
Cannone contro i carri armati – Antiarmored Vehicle Gun
Cannone da campagna – Field Gun; Field Piece
Cannone da carro armato – Armored Vehicle Gun; Tank Gun
Cannone da costa – Coast Artillery Gun
Cannone da esercitazione – Drill Gun
Cannone da montagna – Mountain Gun
Cannone d’assedio – Siege Gun
Cannone di grosso calibro – Heavy Gun
Cannone di piccolo calibro – Light Gun
Cannone di torre corazzata – Turret Gun
Cannone fuori bordo – Out-board Gun (Avn)
Cannone per fanteria – Infantry Gun
Cannone rigato – Rifled Gun
Cannone suodato – Shielded Gun

CAPSULA (plural Capsule). Capsule or Percussion Cap

Capsula detonante – Blasting Cap or Detonator

Capsule incendive ad accensione elettrica – Electric Igniter Cap consisted, accdg to Piantanida (Ref 4, p 196), of a mixture of Gunpowder cotton (dry) 50, KClO₃ 25 & Sb₂S₃ 25%, which surrounded the Pt-Ir wire, connected to two electrodes.

Capsule incendive a percussione – Percussion Caps for igniting propellant charges consisted, accdg to Piantanida (Ref 4, pp 194–95), of mixtures of MF, KClO₃, Sb₂S₃, glass powder & guncotton. Less expensive mixture contd KClO₃, Sb₂S₃ & guncotton or KClO₃, Pb(SCN)₂ & Pb ferrocyanide. Numerical values and other compns are listed on pp 195–196. These mixtures are probably obsolete.

CARICA (plural Cariche) – Charge

Carica cave o Carica ad effetti concentrati – Hollow or Shaped Charge. In the Catalog of the Società Mangirotti, Codroipo (Udine), published in 1960 are described on p B5 compressed charges of TNT (Tritolo). They are in the shape of inverse funnels with inside cavity to create the Munroe-Neumann Effect action on detonation initiated by a cap. The bottom diameters of chgs are 60 & 75mm and their weights are 80, 100, 120, 160, 230, 300 & 360g. Their intended uses are demolition of...
large blocks of stone & punching holes thru steel plates

A brief, general description of cariche cavi (including an illustration; Fig 77) is given by Belgrano (Ref 31, pp 267–69)

Carica d’inflammazione – Igniting Charge

Carica d’innescamento – Priming or Initiating Charge. Their compositions are listed under Composizioni (o Miccele) innescanti

Carica di lancio – Propelling Charge. See under ESPLOSIVI DA GUERRA. Acdg to AddnlRef C; it is a part of the colpo completo (qv)

Carico di polvere nera – Black Powder Charge

Carica posteriore – Base Charge

Carica preparata – Prepared Charge with Container; Cartridge

Carica di rinforzo – Booster Charge. See under ESPLOSIVI DA GUERRA

Carica di scoppio – Bursting Charge. See under ESPLOSIVI DA GUERRA

Carsoniti. Perchlorate based expls, such as
a) KClO₄ 85 & vaseline 15% and
b) KClO₄ 88 & DNBenzene 12% (Ref 31, pp 362–63)

CARTOCCI (singular Cartoccio) (Propellant Containers or Cartridge Cases). The description given in “MUNIZIONAMENTO ITALIANO” (AddnlRef C) is incomprehensible without illustrations, and for this reason we are including here two Figs from the article of Col Jarrett, listed here as Ref 5a. The 1st Fig is cartoccio a bossolo (Cartridge with Igniter Tube), while the 2nd Fig is cartoccio a sacchetto (Cartridge Bag). The illustrations in Jarrett’s article include cartocci a bossolo for 75mm Shell and also for 105/28, 100/17 and 149/13 Shells

In TM 9-1986-6 (1953), listed here as Ref 16 are described and illustrated under Cartridge Cases, without including Italian names, the cases for several weapons. Their igniter charges range from 10 to 50g of Black Powder

Cartridge Cases for the following weapons are described and illustrated: 75/13-mm Mountain Gun (p 123, Fig 185); 75/18-mm Mountain Howitzer (p 124, Fig 186); 75/27-mm Field Gun (pp 124–25, Fig 187); 75/32-mm Light Field Gun (p 125, Fig 188); 77/28-mm Field Gun (p 126, Fig 189); 100/17-mm Light Field Howitzer (p 126, Fig 190); 105/14-mm Light Field Howitzer (p 127, Fig 191); 105/28-mm (p 127, Fig 192); 149/13-mm Heavy Field Howitzer (p 128, Fig 193); 149/13-mm Field Howitzer (p 129, Fig 194); 152/37-mm Gun (p 130, Fig 195); 380/15-mm Heavy Howitzer (p 131, Fig 196)

CARTOCCI GRANATA. Under the title, “MUNIZIONAMENTO ITALIANO” (Addnl-Ref C) are listed complete rounds (colpi completi) for weapons caliber 20-mm and larger, which for small arms, the corresponding rounds are listed as cartucci under armi portatili

For example Cartocci Granata per Mitragliera (Machine Gun) da 20-mm Mod 35 consists of bossolo e canello (Cartridge Case with Primer), carica di lancio (Propelling Charge), bicchiere (Projectile Body), carica di scoppio (Bursting Charge) and spoletto (Fuze). Bossolo is of brass and is filled with polvere FC 4. Bicchiere is of steel with copper bands and filled with TNT or fieglmatized RDX or PETN

Other cartocci granata described in Addnl-Ref C arc:
Cartocci Granata per Mitragliera da 37/54, Mod 39
Cartocci Granata per Cannone da 47/32
Cartocci Granata per Obici (Howitzers) da 75/13, Mod 32
Cartocci Granata da 75/27 per Cannone, Mod 32
Cartocci Granata per Cannone da 76/40-45, Mod 36
Cartocci Granata per Cannone da 90/53
Cartocci Granata per Obice da 100/17, Mod 14-50
Cartocci Granata per Obice da 100/22
Cartocci Granata per Cannone da 105/28, Mod 32
Cartocci Granata per Obice da 149/19, Mod 32
Cartocci Granata per Cannone da 149/40, Mod 35
Colpi completi (Complete Rounds) per Obice da 210/22
Colpi completi per Mortai da 81-mm
Colpi completi per Mortai da 120-mm

Cartridges from 6.5mm to 12.7mm, incl. are listed under Armi Portatili, Munizioni (Small-Arms Ammunition), while artillery cartridges of caliber 37mm and higher are listed in MUNIZIONAMENTO ITALIANO as cartocci granata and described in TM 9-1985-6 (1953) (Ref 16).

From Ref 16, we are listing here:
37/40-mm HE (p 73, Fig 87); 37/40-mm HE/AP (p 73, Fig 88); 37/45-mm AP (p 74, Fig 89);
37/54-mm AA (p 74, Fig 90); 40/39-mm AA (p 75, Fig 91); 47/32-mm HE, M35 (p 75, Fig 92); 47/32-mm AP, M35 (With Cap) (p 76, Fig 93); 47/32-mm AP, M35 (Without Cap) (p 76, Fig 94); 47/32-mm AP, M39 (p 77, Fig 95); 47/32-mm EP (Hollow Charge) (p 77, Fig 96); 47/32-mm EPS (Hollow Charge) (p 78, Fig 97); 57/43-mm AP (Without Cap) (p 78, Fig 98); 57/43-mm AP (With Cap) (p 79, Fig 99); 65/17-mm HE (p 79, Fig 100); 65/17-mm AP (p 80, Fig 101); 65/17-mm EP (Hollow Charge) (p 80, Fig 102); 65/17-mm EPS (Hollow Charge) (p 81, Fig 103); 70/17-mm HE (p 81, Fig 104); 75-mm HE (p 82, Fig 105); 75/13-mm HE (Light Case) (p 82, Fig 106); 75/13-mm HE, M32 (p 83, Fig 107); 75/27-mm HE, M32 (p 83, Fig 108); 75/32-mm HE (p 84, Fig 109); 75/46-mm HE, M34 (p 84, Fig 110); 75/46-mm HE, M36 (p 85, Fig 111); 75/32-mm ADE-HE (p 85, Fig 112); 75-mm AA (p 86, Fig 113); 75-mm AP (p 86, Fig 114); 75/27-mm AP (p 87, Fig 115); 75/32-mm AP (p 87, Fig 116); 75-mm EP (Hollow Charge) (p 88, Fig 117); 75-mm EPS (Hollow Charge) (p 88, Fig 118); 75/13-mm EP (Hollow Charge) (p 89, Fig 119); 75-mm EPS, M42 (Hollow Charge) (p 89, Fig 120); 75-mm Incendiary (F/C) and (F1/C) (p 90, Fig 121); 76/40-45-mm HE (p 90, Fig 122); 76/40-45-mm HE, M36 (p 91, Fig 123); 77-mm HE (Short) (p 91, Fig 124); 77-mm HE (Long) (p 92, Fig 125); 77-mm AA (p 92, Fig 126); 90/42-53-mm HE (p 93, Fig 127); 90-mm AA (p 93, Fig 128); 90/53-mm AP (p 94, Fig 129); 100-mm HE (p 94, Fig 130); 100/17-mm HE, M32 (p 95, Fig 131); 100-mm ADE-HE (p 95, Fig 132); 100/17-mm ADE-HE (p 96, Fig 133); 100-mm EP (Hollow Charge) (p 96, Fig 134); 100-mm EPS (Hollow Charge) (p 97, Fig 135); 105/28-mm HE (p 97, Fig 136); 105/28-mm HE, M32 (p 98, Fig 137); 105/28-mm ADE, M32 (p 98, Fig 138); 105/28-mm HE (p 99, Fig 139); 105/25-mm EP (Hollow Charge) (p 99, Fig 140); 105/50-mm Hollow Charge, M43 (p 100, Fig 141); 120/21-mm HE (p 100, Fig 142); 120/21-mm HE (Cast Steel) (p 101, Fig 143); 120/25-mm HE (Short) (p 101, Fig 144); 120/25-mm HE (Long) (p 102, Fig 145); 120/25-mm HE (Cast Steel) (p 102, Fig 146); 120/40-mm HE (Cast Steel) (p 103, Fig 147); 120/40-mm HE (Base Fused) (p 103, Fig 148); 120/45-mm HE (p 104, Fig 149); 149/12-mm HE (Short) (p 104, Fig 150); 149/12-mm HE (p 105, Fig 151); 149/12-13-mm HE (Light) (p 105, Fig 152); 149/13-mm HE (p 106, Fig 153); 49/35-mm HE, M32 (p 106, Fig 154); 149/35-mm HE, M32/38 (p 107, Fig 155); 149/12-13-mm HE (Cast Steel) (p 107, Fig 156); 149/13-35-mm HE (British) (p 108, Fig 157); 149/35-mm HE (British) (p 108, Fig 158); 149/35-mm ADE, M32 (p 109, Fig 159); 149/40-mm ADE-HE, M35 (p 109, Fig 160); 152-mm HE (p 110, Fig 161); 152/13-mm HE (Short) (p 110, Fig 162); 152/13-mm HE (Long) (p 111, Fig 163); 152/37-mm HE (p 111, Fig 164); 152/45-50-mm HE (Base-Fused) (p 112, Fig 165); 152/32-45-mm HE (Base-Fused) (p 112, Fig 166); 152/37-mm AP (p 113, Fig 167); 210-mm HE (p 113, Fig 168); 210-mm HE (Cast Steel) (p 114, Fig 169); 210-mm HE (Bomba) (p 114, Fig 170); 210/22-mm HE, M35 (p 115, Fig 171); 260/9-mm HE (p 115, Fig 172); 260/9-mm HE (Cast Steel) (p 116, Fig 173); 305-mm HE (Short) (p 116, Fig 174); 305-mm HE (Long) (p 117, Fig 175); 305-mm HE (Light-Case Long and Short) (p 117, Fig 176); 305-mm HE (British) (Long and Short) (p 118, Fig 177); 305/17-mm HE (p 118, Fig 178); 305/17-mm HE (Bomba) (p 119, Fig 179); 305/17-mm HE (Cast-Steel) (p 119, Fig 180); 305/17-mm HE (One-Piece) (p 120, Fig 181); 380/15-mm HE (Base-Fused) (p 120, Fig 182); 420-mm HE (Short and Long) (p 121, Fig 183); 240 and 400-mm HE (Bomba) ~ for use in Mortar (p 122, Fig 184)
**Abbreviations:** AA — Anti-aircraft; ADE-HE — HE Projectile provided with booster system “Detonatore AD Alto Esplosivo”; AP — Armor-Piercing (Perforanti); API — Armor-Piercing-Incendiary; EP — Hollow Charge with Base Fuze; EPS — Hollow Charge incorporating a nose fuze with a long flash tube leading to the center of expl charge; HE — High Explosive (Alto esplosivo); I — Incendiary (Incendiario)

**Cartocchio a bosso.** Cartridge Case (for Fixed or Semi-Fixed) Rounds (See Illustration under CARTOCCHIO)

**Cartocchio a sacchetto** — Cartridge Bag (for Separate-Loading Rounds) (See Illustration under CARTOCCHIO)

**Cartridge Case.** See CARTOCCHIO and under Bosso

**CARTUCCIA.** Cartridge (plural Cartucce). The following items are listed in Military Dictionary (AddnRef A):

- *Cartuccia a capsula centrale.* Center Fire Cartridge
- *Cartuccia a pallottola perforante.* Armor-Piercing Cartridge
- *Cartuccia a pallottola tracciante.* Tracer Cartridge
- *Cartuccia da caccia.* Sporting Cartridge (Ref 31, pp 595 & 619)
- *Cartuccia di lancio.* Propellant Cartridge

**Cartuccia "Magnum".** Cartridge “Magnum”.

Several varieties are described in Ref 31, pp 594–95

Small-arms Cartridges are listed here under *Armi portatili munizioni*, while Artillery Cartridges, which are actually complete rounds (*colpi completi*) are listed under CARTOCCI GRANATA

**Cava D**
**Cava I**
**Cava extra 2**

Italian explosives listed in Ref 31, p 317 without giving their comps

**Centraliti (Centralites),** invented in 1906 in Germany, have been used, accdg to Belgrano (Ref 31, p 218), in Italy as stabilizer for NC–NG proplnts. Description of Centralities is given in Ref 24, Vol 2, pp C126ff. Centralities have also been used in some Italian single-base proplnts, such as Polvere per mitragliatrice FIAT mod 35, described here under Nitrocellulose a solvente volatile and in Ref 31, p 204

**CG Polvere.** See Polvere CG

**CHEDDITE.** Cheddite (plural Cheddittis) or *Esplosivi Street*. Cheddites, invented in 1897 by E. Street of England, have been patented since 1898 in many countries. Accdg to Dr Omero Vettori of Aulla (Massa Carrara), a subsidiary of the French firm located in Chedde, Haute Savoie, established in 1901 at Saviano, near Livorno, Italy, a plant which is likely to be one of those belonging to the Società Italiana Esplosivo Cheddie with the main office at Torino. This Company, which is now a subsidiary of the S.A. Suisse d’Explosifs, Liestal, near Basel, Switzerland, has also plants located at Torano (Carrara), Borgo Franco (Ivrea) and Cinzano Torinese. Cheddites are expl comps based on chlorates or perchlorates of Amm, K or Na. Other ingredients are combustibles and binder. A detailed description of Cheddites is given in Ref 24, Vol 2, pp C155 to C164

Accdg to Dr Omero Vettori (private communications of 1962 & 1963), the plant of Società Italiana Esplosivo Cheddie at Saviano manufactured at that time: Cheddite OS Extra, Cheddite O Extra, Cheddite O Extra B, Plastigel I and Plastigel II. Their comps and props are given in Table 4, p C159 of Ref 24, Vol 2. There is also listed Cheddite gelatina.

In the book of Belgrano (Ref 31, pp 355–60) are described chlorate and perchlorate Cheddites

Accdg to Ref 28, p 321, Cheddites are too sensitive to mechanical action to be used as bursting charges in shells, but they can be used for loading land mines

(See also Cremonite, Esplosivo 86/14, Esplosivo P, Esplosivo S, Manilante, Polvere cancell and Romeite)

**Cidonite o Ciclotrimetilentritriminoammina.** See T₄ and in Belgrano (Ref 31, pp 255–59)

**Ciclottetrametilentetranitroammina.** Same as Octogene o HMX. It is described in Belgrano (Ref 31, pp 259–60)

**Ciclolt.** See Tritolite
Claramite. A mining expl manufd after WWI by the Dinamificio di Orbetello, Italy, utilizing scrap military propellants, consisted of Balistite (or Cordite) 64, Amm Perchlorate 20, NaNitrate 15 & K dichromate 1% (Ref 14, Vol 3, p C329-R & Ref 28, p 321)

Colpo completo (Shot or Complete Round of Artillery Ammunition). Accdg to "MUNIZIONAMENTO ITALIANO" (AddnRef C), the colpo completo consists of: il proietto (Projectile); la carica di lancio (Propelling Charge); il canneto (Projectile (or Fuze) Primer) and la spoletta (Fuze)

In our opinion, to these must be added il bossolo (Igniter for Propelling Charge)

Commercial (or Industrial) Explosives of Non-permissible Type are listed in Ref 24, Vol 3, pp C438-R to C440-R. Some of them are also listed here in alphabetical order

Commercial (or Industrial) Explosives of Permissible Type are listed in Ref 24, Vol 3, pp C451-L & C454-L. Some of them are also listed here in alphabetical order

Complete Round of Ammunition. See Colpo completo

Composizioni (o Miscale) fungogene (Smoke Compositions). See under PIROTECNIA o ARTIFICI DA GUERRA and in Belgrano (Ref 31, p 630)

Composizioni illuminanti (Illuminating Compositions). See under PIROTECNIA o ARTIFICI DA GUERRA and in Belgrano (Ref 31, pp 629–30)

Composizioni (o Miscale) incendiarie (Incendiary Mixtures). Not found in the books of Molina (Ref 1), Pianiana (Ref 4), Caprio (Ref 11), Belgrano (Ref 13), Giorgio (Ref 26) and Belgrano (Ref 31), but found in the book of Giua (Ref 19, pp 412–14)

Giua, after giving a general description of incendiaries (including Napalm) used during WWII in aerial bombs, lists on p.414 the following incendiaries patented in Italy after WWII:

a) Termite, ItalP 448101 (1949) of Ministero della Difesa Aeronautica and in CA 45, 1770 (1951) (Also Ref 31, p 640) consisted of Al 17, celluloid 20, susgissidio di ferro (Fe2O3) 43, Ba peroxide (BaO2) 11.5, Na silicate (Na2SiO3) 6.5 and a substance which regulates the duration of combustion (such as bitumen, resin or tar) 2%. The mixture could be heated to 100° and compressed to 5000–6000 atm, w/o danger of expln

b) Miscela incendiaria Tonegutti, ItalP 446010 (1949) of Ministero della Difesa Marina e M. Tonegutti and CA 45, 1770 (1951), consisted of K chloride 50, AN 20, Cu sulfate 10 & Mg (or Al) powder 20%

Another Italian Patent listed in CA is:

c) Bombirini Parodi-Delfino, ItalP 430931 (1948) & CA 43, 8682 (1949): TNT (PETN or RDX) 15–60, oxidizers (such as chlorates, nitrates, oxides, perchlorates or peroxides) 20–40 & finely pulverized metals (such as Al, Mg, Zn or Fe) 20–40%

Composizioni (o Miscale) innescenti (Initiating Compositions) or Composizioni primari (Primary Compositions). The following formulations were found in some Italian ammunition captured during WWII and examined at Picatinny Arsenal: a) KClO3 43, Sb2S3 24, MF (Mercuric Fulminate) 24 & abrasive 9%; used in some cartridge cases, such as the 47-mm APRN (Armor-piercing Round Nose) shell b) KClO3 44, Sb2S3 48, MF 6 & abrasive 2%; used as a primer in fuzes of some bombs and in 47-mm APRN shell c) LA (Lead Azide) 55, LS (Lead Styphnate) 44 & binder 1%; used as the upper charge in detonators with base charge of T4 (RDX) in some shells (Ref 28, p 322)

The following typical Italian military initiating compns are listed in Belgrano (Ref 31):

a) MF 15–40, KClO3 20–50, gelatine 0.5–2, Sb2S3 30–35 & Si carbide 10–20% (p 423)

b) MF 13.7, KClO3 41.5, gelatine 0.7, Sb2S3 33.4 & powdered glass 10.7% (p 423)

MF 38, PETN 15, Sb2S3 39, KClO3 7 & K2Cr2O7 1% (p 425)

MF 38, PETN 15, Sb2S3 40, K2Cr2O7 2 & Ba(NO3)2 5%. Gave the best results (p 425)

c) Miscale innescenti inossidabili a base di stifenato di piombo e tetrazene – LS 25–55, Tetracene 1.2–5, Ba(NO3)2 25–45, PbO2 5–10, Sb2S3 9–10, CaS 3–15 & glass powdered 0–5% (p 484)
f) Miscela innescanti inossidabili, found in capsule tipo “Sinoxid” Italiane — LSt 35—40, Tetracene 1—3, Ba(NO₃)₂ 35—40, Sb₂S₃ 10—15, CaSi₂ 3—5 & PbO₂ 3—4%. Some mixts contd ca 2% of carbonamide (p 484)

g) Miscela inossidabili. Eight formulations contig Ba nitrate, LSt, Tetracene & Sb sulfuride are listed on p 485

h) Miscela inossidabili, such as 1) LA 5, Pb(SCN)₂ 25, KCIO₃ 55 & Sb₂S₃ 15%
2) LA 25, KCIO₃ 35, Sb₂S₃ 35 & SiC 5%
3) LSt 35, Sb₂S₃ 5, Ba(NO₃)₂ 40 & CaSi₂ 20% (Ref 31, p 487)

i) Miscelle di innesco non corrosive containenti fosforo rosso (Noncorrosive Mixtures Containing Red Phosphorus) developed betw 1950 & 1960 at the Olin Mathieson Chemical Corp and probably tried in Italy. We are listing two examples — Red P 25, PETN 5 & Ba(NO₃)₂ 70%
2) Red P 17, LSt 25, PETN 5 & Ba(NO₃)₂ 53% (Ref 31, p 488)

j) Mixtures contig chloride or perchlorate of thallium developed in 1958—59 by the Manufacture générale des munitions, France, were probably tried in Italy. Three formulations are listed in Ref 31, p 489

k) Noncorrosive mixts contg nitrate of thallium or/and cesium, developed in 1959 by the Manufacture générale des munitions. Two formulations are listed in Ref 31, p 489

l) Noncorrosive mixts contg LSt, Tetracene with added 0.1 to 5% “Aerogel”, developed in 1957 by the Olin Mathieson Chemical Corp and probably tried in Italy. Several formulations are listed in Ref 31, p 490

m) Mixtures contg thiocyanate of lead or mercury developed in France and probably tried in Italy. Five formulations are listed in Ref 31, p 491

Giorgio (Ref 26, p 163) lists several mixtures, among them Cu (Pb or Hg) thiocyanate 30—40, K chloride 50—55, Sb sulfide 0—10 & powdered glass 0—5%

Composizioni (o Miscelle) a luci colorate (Colored Lights Compositions). See under PIROTECNIA o ARTIFICI DA GUERRA and in Belgrano (Ref 31, p 627)

Composizioni di scoppio (Bursting Charge Compositions). See Esplosivi di scoppio

Composizioni (o Miscelle) per tracciatori (Tracer Compositions). See under PIROTECNIA o ARTIFICI DA GUERRA

Cordite Italiana. See C₂ (Polvere)

Cotone colloccio. (Collodion Cotton, abbrd as CC). See under Nitrocellulose

Cotone fulminante. (Guncotton, abbrd as GC). See Fulmicotone under Nitrocellulose

Cremonite. A Cheddite-type expl mixture proposed in 1902 by U. Alvisi — Amm Perchlorate 48.85 & Amm Picate 51.15% (Molina, Ref 1, p 200)

Cresiltite. Mixture of TN-m-Cr (Trinitrometacresole) 60 & PA (Picric Acid) 40%, used in Italy for loading large-caliber shells. It was developed in France under the name of Crèyllite No 2 (Ref 28, p 323 & Ref 31, p 292) (See also Ecristale)

Demolition Charges. Accdg to OrdnSergeant (Ref 4a, p 18), Italian military demolition expls of WWII were plastic, such as the one consisting of RDX 67.2, NG 16.3, Al 12.2 & wax 4.1% (adds to 99.8%). It was of gray color and the charge shaped like a pancake, ball or hollow cylinder. Another charge consisted of RDX, NG & wax and was of yellow color. The 3rd chge was light-brown in color and consisted of RDX with a desensitizing agent. Mg shavings (1—3 oz per 1 lb of HE) were added to all above demolition chges to increase their incendiary effect

Demolition Fuse, Instantaneous is described under Fuse (Miccia)

Detonatori. See under INNESCAMENTO

DINAMITI (Dynamites)

Accdg to Giua (Ref 19, pp 338—45), Italian Dynamites may be subdivided into: 1, Dinamiti a base inerti (Dynamites with inert base) and 11, Dinamiti a base attiva (Dynamites with Active Base)

1. Dynamiti a base inerti may be subdivided into:
a) **Tipo I** – Nitroglicerina (NG) 70–75% & kieselguhr 30–25%

b) **Tipo II** – NG 50 & kieselguhr 50%

c) **Dinamite nera** (Black Dynamite) – NG 45–55 & coke 55–45%

d) **Dinamite al carbonio di legno** (Charcoal Dynamite) – NG 90 & charcoal 10%

e) **Dinamite rossa** (Red Dynamite) – NG 68 & tripoli 32%

f) **Wetter-dinamite** (Permissible Dynamite) – NG 35–40, kieselguhr 14–10 & Mg sulfate 32–50%

II. **Dinamiti a base attiva** may be subdivided into:

g) **Gelatina gomma o Gelatina esplodente** (Blasting Gelatin) – NG 92–93 & CC (Collodion Cotton) 8–7%

h) **Gelatina espressiva da guerra** (Military Blasting Gelatin) – NG 86,4, CC 9,6 & camphor 4,0% (Suitable for use as bursting charge)

i) **Gelatina dinamite** (Gelatin Dynamite or Gelatin) – NG 67–86, CC 3–5,5, KN (Potassium Nitrate) 5–25 & wood flour 2–10%

j) **Ammon dinamite** (Gelatina 65%) – NG 63, CC 2, AN 30 & wood flour 5%

k) **Gelatina dinamite incongelabile o antigelo** (Nonfreezing Gelatin Dynamite) – NG 20–55, CC 1–2, Nitrotoluenes 8–21, SN (Sodium Nitrate) with AN 25–60 & cereal flour 1–8%

l) **Gelatina 40%** – NG 40, drip oil (liq DNT (Dinitrotoluene)) 10, SN 44 & cereal flour 6%

m) **Ammon-gelatina I** – NG (gelatinized with CC) 40, AN 45, SN 5 & wood flour 10%

n) **Ammon-gelatina II** – NG (gelatinized with CC) 20, AN 75 & wood flour 5%

o) **Gelatina Vender o Dinamite incongelabile, Vender** (Gelatin of Vender or Nonfreezing Dynamite of Vender). A series of exprs invented by Venzo Vender and manufd before WWII at the Dinamitificio di Cengio. They were based on NG mixed with 10–30% of Dinitromonoformin or Dinitromononacetin and other usual ingredients of Dynamites. It was claimed that these exprs remained plastic at temps as low as −20°C

p) **Dinamite No 1, Non-gelatinizzate** – NG 70–74 & wood flour 30–26%

q) **Dinamite No 2, Non-gelatinizzate** – NG 35–48, SN and/or PN 52–39 & cereal flour 12–17%

r) **Unknown Name Dynamite** – NG 20–25, AN 20–25, SN and/or KN 30–35 & charcoal 20%

Some Dynamites were used by Italians during WWII as demolition charges

Dynamites and other exprs permitted for use in gaseous and/or dusty coal mines are “Esplosivi antigirusouti”, “Esplosivi ammissibili” or “Esplosivi di sicurezza” (Ref 28, p 324)

Belgrano (Ref 31, p 170– lists on Table 26 the following Dynamites as the principal exprs used in Italy: GDI, GDI, GD2, GDM, GEO,
Gomma A and Gomma B. All of them are listed, including their properties, under ESPLOSIVI DA MINA in Table 1. In addition to these, Dynamite GEOM is listed there

**AddnlRef**: L. Avogadro (of Montecatini, Avigliana, Torino), AnnChim (Roma) 49, 352– 57 (1959) & CA 53, 15567 (1959) (Sensitization of Dynamite Gelatin by 1–5% of inert substances of high or moderate hardness such as pumice, quartz, Al₂O₃, hematite, pyrite, etc can raise deton velocity to as high as 6800 m/sec)

**Dinamon.** Accdg to Belgrano, 1st Edn (1952) (Ref 13, p 163), Dinamon consisted of AN 69, KCIO₄ 8, TNT 20 & Al 3%. It was listed in Ref 24, Vol 3, p C440-L

**Dinamon 1B.** Accdg to Belgrano, 2nd Edn (1974) (Ref 31, p 171), it is a mining expr cong NG 3–6%, the rest being AN, TNT, woodmeal and some other ingredients

**Echos or Escho.** Accdg to Molina (Ref 1, p 342), Esposivo Echos – AN 75, Si (95% pure) 16, Al pdr 2 & dried horse dung (called “Ipposino”) 7% – was used by the Italians for military purposes. Silicon can be replaced by ferro-silicon. Belgrano (Ref 31) lists it as Echo on p 316 but erroneously gives AN content as 25 instead of 75%

**Ecrasite.** This name is derived from **Ekrasit**, an Austrian military expr developed in 1892 and used until 1908 when all of its available supply was blown up in Kiev, Russia. Its compn was kept secret and originally was supposed to be Ammonium Trinitrocrelate. Discussion on this subject is given in Ref 24, Vol 5, pp E8-R & E9-L
Erasite 60/40. Same as Creasite or Cresylite No 2

Esanitrofenilamina, Exil o p-Dipierilamina. Hexanitrophenylamine, 
(O₂N)₃C₆H₂.NH.C₆H₂(NO₂)₂, described, under Diphenylamine, in Ref 24, Vol 5, pp D1434-Rff. Accdg to Ref 28, p 325 it was used as a HE either alone or inmixts with TNT (See also Ref 1, p 399)

Belgrano (Ref 31, pp 375–77) describes it as Exil and states that its mixture with TNT & AI has been used in underwater arms (armi subacque) 

Esanitrosorbit (Hexanitrosorbitol). An expl compd first prepd and examined during WWII in Italy by A. Tetramanzi & N. Arnaldi and reported in Atti della Accademia delle Scienze di Torino, Classe di Scienze Fisiche, Matematiche e Naturali 77, 278–81 (1942) & CA 38, 3841 (1944). This compd, O₉N.O.CH₂(CH(ONO₂))₄.CH₂.O.NO₂, was obtd by nitrating sorbitol of high purity by means of fuming nitric acid below 0°, followed by gradual addn of sulfuric acid at below minus 15°. The product obtd, in 97% yield, was (after crystn from alcohol) in the form of plates of density 1.58, melting at 55°. Its props are reported as follows: Heat of combustion 1465 kcal/kg; Heat of formation 135 kcal/mole; Heat of explosion 1500 kcal/kg; Detonation Velocity 7230 m/sec and Sensitiveness to Impact and Stability — similar to NG. It was not considered a satisfactory replacement for NG as a gelatinizer for NC because of physical changes which occurred in the mixture. Its use for military purposes was not reported (Ref 28, p 325)

Escho. See Echos

Esplosivi alla nitroglicerina. See under DINAMITI

Esplosivi Alvisi. Perchlorate-based, Chedidite type, expls patented by Ugo Alvisi beginning in 1899. They were superior to chlorates-based expls. The following are described by Molina (Ref 1, pp 199-200): Manilamite, Polvere Cannel, Cremonite and Kratite

Esplosivi ammissibile. Permissible Explosives. See Esplosivi antigrisutosi

Esplosivi alti (High Explosives) o Esplosivi dirompenti (Brisant Explosives). To these belong Esplosivi da guerra, such as Trinitol (TNT), Pentrite (PETN), T₄ (RDX), Melinite (FA), Tetrile (Tetryl), etc and Esplosivi da mina, such as listed in Tables I & III and under DINAMITI, Items a, b, c, d, e, g, h, i, j, k, l, m, p & q. Also Gelignite, Gomma A, Gomma B, etc.


Esplosivi antigrisutosi o Esplosivi ammissibili (Antifiredamp Explosives or Permissible Explosives). See under Esplosivi da mina, Tables II & IV and under DINAMITI, Item f. Also Grisounite and Grisoutina

Esplosivi bassi (Low Explosives). To these belong weak Esplosivi da mina (such as Esplosivi antigrisutosi), DINAMITI with low NG content and Esplosivi di lancio o Esplosivi propellenti

Esplosivi deflagranti. See Esplosivi di lancio

ESPLOSIVI DA GUERRA o ESPLOSIVI MILITARI. To these belong: Esplosivi innescenti o primari (Initiating or Priming Explosives); Esplosivi di lancio o Propellenti (Propellants); Esplosivi di rinforzo (Booster Explosives) and Esplosivi di scoppio (Bursting Explosives). They are described separately below

Esplosivi innescenti o primari. (Initiating or Primary Explosives).

Under this title Belgrano (Ref 31, pp 405ff) describes:
1) Innescenti primari (Priming Explosives), which include Fulminato di mercurio (MF) (pp 406–428); Fulminato organico, CHO.CNOH (p 428); Fulminato di argento (AgF) (pp 428–429); Azotidrati (Azides), which includes Acido azotidrico, HN₃ (Hydrazoic Acid) (p 429); Azotidrato di sodio, NaN₃ (SA) (pp 430–38); Azotidrato di piombo, PbN₆ (LA) (pp 438–49); Azotidrato d’argento, AgN₃ (pp 449–51); Azotidrato di ammonio, NH₄N₃ (p 451);
Acido stifico or Trinitroresorcin, (HO)$_2$C$_6$H(NO$_2$)$_3$ (pp 453–55); Dinitroresorcin, (HO)$_2$C$_6$H$_2$(NO$_2$)$_2$ (p 456); Stilfratto di piombo (Lead Stypheate) (LS) (pp 456–63); Guaninitrosoammina o Tetracene (Tetracene), C$_6$H$_5$N$_1$O (p 464–67); Solfoeisancato di piombo (Lead Thiocyanate), Pb(CNS)$_2$ (pp 467–68); Diazidinitrofenolo o DDNP, C$_6$H$_2$N$_4$O$_9$ (pp 471–74); Tritriydroxocianato di piombo, Pb(C$_6$HN$_3$O$_3$Pb) (p 475–76); Nitrato di diazobenzene, C$_6$H$_5$N$_2$O$_2$ (pp 476–77); Solfo di azoto, N$_4$S$_4$ (pp 477–78); Esmatidinetrprofessionalminio o HMDT, C$_6$H$_2$N$_2$O$_6$ (pp 478–79); Cianurazide, C$_3$N$_3$(N$_2$)$_3$ (pp 481–82); Acidio diazotriazolcarbonio,

$$\text{H}_2\text{N} \rightarrow \text{N} \rightarrow \text{C}_3\text{CO}_2\text{H} \quad \text{(pp 482–83)}$$

Principal Italian Initiating Compositions (or Mixtures) are described under Composizioni (o Miscele) innescenti

**Esplosivi da lancio, Esplosivi deflagranti o Propellent (Propellants).**

To these belong the following items described in Belgrano (Ref 31): Polvere nera da guerra (p 342), Balistite (p 205), under Polveri senza fumo (pp 201ff), C$_2$ (Polvere) (p 210), and Propellenti per razzi (p 128). They are described here separately. To these must be added Filit (qv), which was not described in Belgrano (See also POLVERI ITALIANI DA LANCIO SENZA FUMO)

**ESPLOSIV (o POLVERI) DA MINA.** Mining Explosives

Many Italian Mining Explosives are listed under COMMERCIAL OR INDUSTRIAL EXPLOSIVES in Ref 24, Vol 3, on pp C438-R to C440-R (Nonpermissible Explosives) and on pp C451-L & R and C454-L (Permissible Explosives). Some of them are listed in this section

Belgrano (1952) (Ref 13) lists in Tables on pp 280ff 164 formulations, while in the new edn of Belgrano (1974) (Ref 31) there are listed in Tables 53 to 62 incl, 259 Mining Explosives, some of them French. The tables give, besides compns, the following properties: Trauzl (Lead Block Expansion), Distanza colpo (Gap), and Velocità detonaione (Detonation Velocity) values

The tables which follow list expls selected from Belgrano’s books

**TABLE I**

**ESPLOSIVI DA MINA GELATINOSE CON NITROGLICERINA**

(Mining Explosives with Nitroglycerol)

<table>
<thead>
<tr>
<th>Composition (%) and Some Properties</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>Gorina</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>GDI</td>
<td>GD2</td>
<td>GDI</td>
<td>GDIM</td>
<td>GEOM</td>
<td>GEO</td>
<td>B</td>
<td>A</td>
<td>GDM</td>
<td></td>
</tr>
<tr>
<td>Nitroglycerin</td>
<td>43.2</td>
<td>48.3</td>
<td>7.0</td>
<td>60.0</td>
<td>38.0</td>
<td>57.0</td>
<td>71.0</td>
<td>40.75</td>
<td>77.5</td>
<td>82.5</td>
</tr>
<tr>
<td>Colloid Cotton</td>
<td>2.3</td>
<td>2.7</td>
<td>0.8</td>
<td>3.5</td>
<td>2.3</td>
<td>3.5</td>
<td>5.0</td>
<td>1.5</td>
<td>5.0</td>
<td>5.5</td>
</tr>
<tr>
<td>Am Nitrate</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>Am Perchloreate</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>Woodflour</td>
<td>7.0</td>
<td>5.8</td>
<td>1.0</td>
<td>5.2</td>
<td>−</td>
<td>3.5</td>
<td>5.0</td>
<td>−</td>
<td>5.0</td>
<td>3.0</td>
</tr>
<tr>
<td>Dinitrotoluene</td>
<td>−</td>
<td>−</td>
<td>10.0</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>Triinitrotoluene</td>
<td>−</td>
<td>−</td>
<td>5.0</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>Na Nitrate</td>
<td>45.5</td>
<td>42.7</td>
<td>32.2</td>
<td>30.5</td>
<td>6.0</td>
<td>7.0</td>
<td>−</td>
<td>12.0</td>
<td>8.5</td>
<td>−</td>
</tr>
<tr>
<td>K Nitrate</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>Ca Silicide</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>Oil</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>PETN</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>Ocher, red (Hematite)</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>Ocher, yellow (Limonite)</td>
<td>1.0</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>Na Carbonate</td>
<td>1.0</td>
<td>0.5</td>
<td>−</td>
<td>0.5</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>0.5</td>
<td>0.5</td>
<td>−</td>
</tr>
<tr>
<td>Ca Carbonate</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>−</td>
</tr>
</tbody>
</table>

| Trauzl Test, cc                     | 340 | 355 | 430 | 440 | 475 | 500 | 505 | 525 | 540 | 560 | 630 | 475 |
| Gap Test, cm                       | 14 | 15 | 6 | 20 | 21 | 23 | 26 | 25 | 29 | 30 | 35 | − |
| Veloc of deton, m/sec               | 5000 | 5200 | 4700 | 6000 | 5900 | 5400 | 5900 | 7000 | 6700 | 6900 | 7200 | − |
### TABLE II
**ESPLOSIVI DA MINA POLVERULENTI CON NITROGLICERINA**
(Pulverulent Mining Explosives with Nitroglycerol)  

<table>
<thead>
<tr>
<th>Composition (%) and Some Properties</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
<th>12</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitroglycerin</td>
<td>14.0</td>
<td>11.0</td>
<td>10.0</td>
<td>4.0</td>
<td>4.0</td>
<td>8.0</td>
<td>12.4</td>
<td>12.76</td>
<td>4.0</td>
<td>12.0</td>
<td>11.7</td>
<td>9.0</td>
</tr>
<tr>
<td>Dinitrotoluene</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>1.5</td>
<td>-</td>
<td>0.5</td>
<td>0.3</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Trinitrotoluene</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>10.0</td>
<td>0.3</td>
<td>-</td>
<td>20.0</td>
<td>-</td>
</tr>
<tr>
<td>Collodion Cotton</td>
<td>0.3</td>
<td>-</td>
<td>0.25</td>
<td>1.5</td>
<td>0.1</td>
<td>0.5</td>
<td>0.3</td>
<td>0.24</td>
<td>-</td>
<td>0.3</td>
<td>0.3</td>
<td>-</td>
</tr>
<tr>
<td>Woodflour</td>
<td>-</td>
<td>6.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Cereal flour</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>5.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>3.0</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Am Picrate</td>
<td>65.7</td>
<td>51.0</td>
<td>89.75</td>
<td>82.0</td>
<td>89.4</td>
<td>81.0</td>
<td>79.0</td>
<td>83.0</td>
<td>76.0</td>
<td>78.7</td>
<td>76.0</td>
<td>70.7</td>
</tr>
<tr>
<td>Na Nitrate</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>K Nitrate</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Dinitronaphthalene</td>
<td>-</td>
<td>10.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>8.0</td>
<td>4.0</td>
<td>10.0</td>
<td>-</td>
<td>10.0</td>
<td>-</td>
</tr>
<tr>
<td>K Perchlorate</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Na Chloride</td>
<td>20.0</td>
<td>22.0</td>
<td>-</td>
<td>11.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>T4 (RDX)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>6.5</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>2.0</td>
<td>2.0</td>
</tr>
<tr>
<td>Oil</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>5.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Ca Silicide</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

Traulz Test, cc  | 270 | 265 | 300 | 305 | 320 | 375 | 390 | 390 | 400 | 420 | 430 | 450 |
Gap Test, cm     | 2  | 5  | 20  | 3  | 12  | 6  | 3  | 4  | 11  | 1  | 1  | 6  |
Veloc of deton, m/sec | 2120 | 2300 | 2038 | 1710 | 2500 | 3200 | 2715 | 3200 | 2348 | 2960 | 2350 | 2912 |
Abbreviations:    | CM - Coal Mines | G&OM - Galleries & Open Mines | OM - Open Mines Only |

Note: Composition No 3 is called Grisoutina

### TABLE III
**ESPLOSIVI DA MINA POLVERULENTI SENZA NITROGLICERINA**
(Pulverulent Mining Explosives Without Nitroglycerol)  

<table>
<thead>
<tr>
<th>Composition (%) and Some Properties</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
<th>12</th>
<th>13</th>
<th>14</th>
</tr>
</thead>
<tbody>
<tr>
<td>Am Nitrate</td>
<td>85.5</td>
<td>77.0</td>
<td>40.0</td>
<td>78.0</td>
<td>82.0</td>
<td>80.0</td>
<td>79.5</td>
<td>70.0</td>
<td>84.5</td>
<td>90.0</td>
<td>-</td>
<td>79.0</td>
<td>70.0</td>
<td>63.0</td>
</tr>
<tr>
<td>Dinitronaphthalene</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Dinitrotoluene</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>1.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Trinitrotoluene</td>
<td>4.5</td>
<td>15.0</td>
<td>-</td>
<td>16.0</td>
<td>14.0</td>
<td>20.0</td>
<td>10.0</td>
<td>10.0</td>
<td>-</td>
<td>8.0</td>
<td>90.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Am Perchlorate</td>
<td>-</td>
<td>-</td>
<td>25.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>K Perchlorate</td>
<td>-</td>
<td>-</td>
<td>29.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>K Nitrate</td>
<td>-</td>
<td>-</td>
<td>29.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Woodflour</td>
<td>10.0</td>
<td>6.5</td>
<td>4.0</td>
<td>-</td>
<td>3.0</td>
<td>-</td>
<td>2.5</td>
<td>-</td>
<td>-</td>
<td>2.0</td>
<td>-</td>
<td>1.0</td>
<td>-</td>
<td>3.0</td>
</tr>
<tr>
<td>Oil</td>
<td>-</td>
<td>-</td>
<td>2.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Paraffin</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Na Nitrate</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>4.5</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>8.0</td>
<td>15.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>K Bichromate</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>4.5</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Carbon (pulverized)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>1.5</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>1.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Charcoal (pulverized)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>PETN or RDX</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>10.0</td>
<td>20.0</td>
<td>30.0</td>
<td>21.0</td>
<td>-</td>
</tr>
<tr>
<td>Ca Carbonate</td>
<td>-</td>
<td>-</td>
<td>1.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Other (Hematite)</td>
<td>-</td>
<td>-</td>
<td>0.5</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

Traulz Test, cc  | 330 | 330 | 350 | 400 | 420 | 430 | 450 | 450 | 360 | 395 | 375 | 395 | 400 | 425 | 450 | 480 |
Gap Test, cm     | 1  | 2  | 3  | 6  | 6  | 7  | 7  | 2  | 3  | 1  | 4  | 5  | 3  | 6  | 6  | 6  |
Veloc of Deton, m/sec | 1900 | 2300 | 2400 | 3600 | 3700 | 4300 | 3500 | 1600 | 3100 | 2300 | 7000 | 2850 | 2100 | 3000 | 3000 | 3000 |
TABLE IV
ESPLOSIVI DA MINA CON RESIDUATI DI POLVERI DI LANCI0
(Mining Explosives Containing Surplus Propellants)

<table>
<thead>
<tr>
<th>Composition (%) and Some Properties</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
<th>12</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ballistite</td>
<td>45</td>
<td>12</td>
<td></td>
<td>60</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td></td>
<td>50</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Polvere B</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>15</td>
</tr>
<tr>
<td>Polvere C</td>
<td></td>
<td></td>
<td></td>
<td>54</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>50</td>
<td></td>
<td></td>
<td>54</td>
</tr>
<tr>
<td>Polvere Dupont</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>10</td>
<td>20</td>
<td>40</td>
<td></td>
</tr>
<tr>
<td>TNT</td>
<td></td>
<td>10</td>
<td></td>
<td>10</td>
<td></td>
<td>4</td>
<td>10</td>
<td>10</td>
<td></td>
<td></td>
<td></td>
<td>10</td>
</tr>
<tr>
<td>Am Nitrate</td>
<td>55</td>
<td>64</td>
<td></td>
<td>70</td>
<td>80</td>
<td>55</td>
<td></td>
<td>60</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>K Nitrate</td>
<td></td>
<td></td>
<td>40</td>
<td></td>
<td></td>
<td></td>
<td>5</td>
<td>40</td>
<td>40</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Na Nitrate</td>
<td></td>
<td>9</td>
<td>26</td>
<td></td>
<td></td>
<td></td>
<td>20</td>
<td></td>
<td></td>
<td></td>
<td>26</td>
<td></td>
</tr>
<tr>
<td>Am Chlorate</td>
<td></td>
<td>20</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>K Perchlorate</td>
<td></td>
<td></td>
<td></td>
<td>5.5</td>
<td>10</td>
<td>5</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Woodflour</td>
<td></td>
<td></td>
<td></td>
<td>4.5</td>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ca Silicide</td>
<td></td>
<td>5</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>5</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Trauzl Test, cc</td>
<td>430</td>
<td>425</td>
<td>445</td>
<td>315</td>
<td>320</td>
<td>330</td>
<td>350</td>
<td>350</td>
<td>375</td>
<td>410</td>
<td>435</td>
<td>435</td>
</tr>
<tr>
<td>Gap Test, cm</td>
<td></td>
<td>1</td>
<td>7</td>
<td>4</td>
<td>4</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>3</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>Veloc of Deton, m/sec</td>
<td></td>
<td></td>
<td>3300</td>
<td>3200</td>
<td>1350</td>
<td>1500</td>
<td>1320</td>
<td>2500</td>
<td>2328</td>
<td>2900</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note: After WWII considerable amounts of left-over propellants (both of American & Italian origin) were left unused in Italy. Some of these propellants were used to prepare mining explosives. The usual procedure was to grind a propellant to pass a sieve with 16 openings per sq cm (US Std Sieve No 12) and to mix it with an oxidizer and other ingredients ground to pass a sieve with 20 openings per sq cm (US Std Sieve No 14).

Black Powder is still used in mining and its compn: K nitrate 60–72, carbon 14–21 & sulfur 13–18% is listed in Belgrano (Ref 31, p 342) as Polvere nera da mina

AddnRefs from CA:
A) S. Custodero, ItaIP 522944 (1955) & CA 53, 3698 (1959) [Synthetic resin-base expls are prepd by incorp of oxidizers into a liquid polymer which is then solidified by addn of a suitable catalyst. Thus, to 20% of a mixt contg 100 parts “Araldit D” and 9–10ps “951” (an aliphatic polyamine), 80% NaClO3 was added with slow stirring to obtain a homogeneous paste. After it was shaped into the desired form by means of a press, it hardened on standing for 14 hours]

B) A. Simoncini, ItaIP 515805 (1955) & CA 52, 15910 (1958) [Tannin contg expl mixts and flammable powders are prepd in the following examples: 1) Powdered 76 parts KClO3, mixed with 24 parts powd chestnut wood extract (f) (as a paste), dried and ground; 2) Powd 38 pts KNO3 mixed with 16ps (f) and another 38ps KNO3 mixed with 8ps powd S, moisten, dried and ground]

C) T. Seguitti, IndMineraria (Roma) 16 (6), 289–98 (1965) & CA 64, 7960 (1966) [Explosive mixts contg AN and combustible oil for general mining uses is described in detail]

D) T. Seguitti, IndMineraria (Roma) 16 (8), 413–22 (1965) & CA 64, 7960–61 (1966) [Explosive mixts contg AN and combustible oil for underground uses are described in detail]

Esplosivi di rinforzo (Booster Explosives).
The following expls have been used: Acido picrico (Ref 31, p 281); T₄ (RDX) (Ref 31, p 263); Pentrite (PETN) (Ref 31, p 176) and Tetritile o Tetrite (Tetryl) (Ref 31, p 272)

Illustration of an Italian Booster is given in Ref 16, Fig 38 on p 28

Esplosivi di scoppio (Bursting Explosives or Fillers).
Following are the principal High Explosives (Esplosivi alti) used as fillers of projectiles, by
themselves, or in compns: Amatolo (TNT 60 & AN 40%) (Ref 31, p 246); Ammonal (Ref 31, p 367); ANS o ASN (Ref 31, p 181); Exil (Hexanitrodiphenylamine) (Ref 31, p 275); Octogene o HMX (Ciclotetrametilentetranitro- 
ammina) (Ref 31, p 259); Pentrite o Tetrinitro- 
pentaeritrite (PETN) (Ref 31, pp 176ff); Pen- 
trite paraffina, 90/10 (Ref 31, p 182); Pentoliti 
(PETN 20–80 & TNT 80–20%) (Ref 31, p 182); 
T₄, Esogene o Trimetilentriammina (RDX, 
Cyclonite or Hexogen) (Ref 31, pp 255ff); 
Tetritol (Tetryl 75 & TNT 25%) (Ref 31, p 246); 
Tritolite, Tritolo o Trinitrotoluene (Ref 31, p 
233); Tritolital (TNT 60, RDX 20 & Al 20%) 
(Ref 31, p 263); Tritolite (RDX 60, TNT 39 
& beeswax 1%) (Ref 31, p 246) and Xilite o 
Trinitrometaxilolo (Ref 31, pp 248–49) (See 
also the book of E. Brandimarte, “Cariche di 
Scoppio”, Ref 29) 

The following Bursting Explosives patented in Italy after WWII are listed in CA: 
a) Montecatini Società Generale, ItalP 433633 
(1948) & CA 44, 1709 (1950) [Cast expl prepared 
by incorporating RDX (or PETN) 75 with Nitro-
isobutylglycerol Triacetate 25% at 80°C] 
b) Polversifici Giovanni Stacchini SA, ItalPats 
433636 (1948); 445206 (1949); 445601, 
445602 & 445603 (1949) and CA 44, 1709 
(1950); 45, 1770 & 3160 (1951) [Comps of 
RDX (or PETN) 5–65, TNT 30–90 with pow-
dered metal (such as Si, B, Mg, Cu, Fe, Al or 
Zn) 5–30%] 
c) Direzione Superiore del Servizio Tecnico di 
Artiglieria a Roma, ItalP 450103 (1949) & CA 
44, 11098 (1950) (RDX 86–92 mixed with 
caster oil 8–14% and pressed in the form of 
beads) 
d) Bombrini-Parodi-Delfino S.p.A., FrP 1424216 
(1966) (Ital) & CA 65, 10420 (1966) [Plastic 
explosives of high power suitable as bursting 
dges can be prep as follows: Into molds 
or bomb bodies are introduced a granulated 
mixt of NC and HE’s, followed by the addn of 
a solvent contg a nitrate ester, a stabilizer, a 
plasticizer and sometimes an accelerating agent. 
The compn of the final expl may be: NC 5–36, 
nitrate ester 33–40, HE 20–60, plasticizer 1.5– 
2.5, stabilizer 0.5–0.6 and accelerating agent 
0.4–0.5 pt. The granulated compn is prep'd 
by mixing the NC, the HE, a plasticizer (di-
ethylphthalate) and a stabilizer (Centralite) 

with a NC solvent (acetone). The paste obt'd 
is impregnated in the mold with NG and the 
DEt ph in a solvent, centrifugd, and gelled at 
50°C] 
e) E. Ravelli, ItalP 648270 (1962) & CA 65, 
5440 (1965) [Explosives of high density suitable 
for loading proj's are obt'd by mixing T₄ (RDX) 
and/or Pentrite (PETN) with high density ni-
trates. For example, compressed mixt of PETN 
25 with Pb(NO₃)₂ 75% gave a loading d 2.5 
to 2.7; mixt of PETN 29 with Ba(NO₃)₂ 71% 
or PETN 24, Ba(NO₃)₂ 74 & Zn stearate 2% 
gave d 2.2–2.3 

Explosivi “slurry”, developed after WWII in US, 
were not manuf’d, nor used as of 1974 when 
the book of Belgrano was published (Ref 31, p 320) 

Explosivo 60/40. Amatol contg 60% AN & 
40% TNT. See under Amatolo 

Explosivo 86/14. Cheddite-type expl consisting 
of Amm Perchlorate 86 & paraffin 14% (Ref 4, p 
230) 

Explosivo ASN. See ASN o Antisanzionite 

Explosivo FNP. See FNP (Explosivo) 

Explosivo MABT. See MABT 

Explosivo MAT. See MAT (Picratol) 

Explosivo MBT. See MBT (Explosivo) 

Explosivo MNDT. See Siperite 

Explosivo MST. See Nougat 

Explosivo “P”. Blasting expl consisting of Amm 
Perchlorate 61, Na nitrate 30, paraffin 8 & 
vaseline 1% (Ref 4, p 230). A similar expl, 
called Explosivo speciale P consisted of Amm 
Perchlorate 53, Na Nitrate 35 & paraffin 12% 
(Ref 28, p 331). It is called in Ref 1, p 202 
Cheddite speciale P 

Explosivo plastico. A plastic expl suitable for 
military use consists of RDX 85–89, petrolatum 
12–10, plastic binder (Trioplasta molle) 0.5–2 
& glycerophthalic acid 0.5–1%. Its Al-contg
modification was: RDX 64–75, petrolatum 10–12, Al powder 25–10, plastic binder 0.5–2 & glycerophthalic acid 0.5–1%

Ref: Dinamite Nobel SA, Milano, ItalP 427535 (1947) & 439931 (1948); CA 43, 4768 (1949) & 44, 6130 (1950) (See also “T₄ plastico”)

Esplosivo “S”. Cheddite-type blasting expl consisting of NaClorate 90, paraffin 7 & vaseline 3% (Ref 1, p 205 & Ref 28, p 331)

Esplosivo S20. Military expl consisting of AN 79, TNT 20 & woodflour 1%. It was manufd at Società Stacchini. Its props are similar to those of French Explosif du type N n°U, which consisted of AN 78.7 & TNT 21.3%. Their props are listed on p 332 of Ref 28. See also L. Mènard & A. LeRoux, MP 34, 201 (1952)

Esplosivo speciale P. See under Esplosivo P

Exil. See Espanitrodifenilammina

Exogene o Trimetilenitritroamina (RDX) See T₄

F₂. A single-base smokeless proplnt listed by Belgrano without giving its compn (Ref 31, p 581)

FB (Polvere). See Polvere FB

FC 4 (Polvere). See Polvere FC 4

Filitte. Ballistite proplnt consisting of NC 50 & NG 50% with 0.5–1% aniline added as stabilizer. It was manufd in the form of cords. Not used now because it badly corrodes gun barrels (Ref 24, Vol 6, p F24-R). Not listed by Belgrano

FNP (Esplosivo). High Explosive compn consisting of AN 75, PETN 20 & wax 5%. Was used during WWII press-loaded in some projectiles (Ref 28, p 332)

Fulmicotone (Guncotton). See under Nitrocellulose

Fulminato d’argento. Silver Fulminato, AgCNO₃, is described by Belgrano (Ref 31, pp 428–29). It was used in small quantities by the Marina Italiana for special detonators (p 429) (See also Ref 24, Vol 6, pp F223-R to F224-R and Ref 28, p 332)

Fulminato di mercurio. Mercuric Fulminate, Hg(CNO)₂, is described in Belgrano (Ref 31, pp 406–27). It was manufd to Dr Omero Vettori, during WWII by the Nobel Società Generale di Esplosivi e Munizioni at Tiana and by the Società Anonima Bombolini-Parodi-Delfino at Colloferro-Roma. It has been used in admixture with KClO₄ & Sb₂S₃ with or without powdered glass. Some post WWII formulations are listed here under Composizioni innescanti and in Belgrano, p 423. See also Ref 24, Vol 6, pp F217-L to F230–R and Ref 28, p 333

Fucchì artificiali. See under PIROTECNIA o ARTIFIZI DA GUERRA

Fuse (Miccia). Most important fuses are:

Miccia a lenta combustione – Slow Fuse, such as Safety Fuse. The core of Italian fuse consisted of K nitrate 60, beech charcoal 25, hemp charcoal 5 & Fe oxide 10% (Ref 31, p 523). Other formulations are BKPDrs, such as K nitrate 70 or 77, S 12 or 10 & carbon 18 or 13% (Ref 31, p 524)

Miccia detonante – Detonating Cord or Cordeau. These may have cores of compressed PA (Picric Acid) - deton vel 5000m/s; core of TNT - deton vel 4500m/s; fuse of PETN prep by a special method and MF (Mercuric Fulminate) fuse prep by a special method (Ref 31, p 528)

In OrdnSergeant (Ref 4a, p 18) is described a WWII Instantaneous Fuse for Initiation of Demolition Charges which contd a core of MF 79.4 with wax 20.6%. When initiated by a cap, the fuse functioned at the rate of about 20000f/s (6100m/s). If initiated by flame it will burn but its use as slow burning fuse was not recommended

GDI, GDII, GD2 and GDIM. Mining Explosives. Their compns and props are listed under Esplosivi da Mina in Table I

Gelatina 40%. See item i, under DINAMITI

Gelatina 65%. See Ammon dinamite, item j, under DINAMITI

Gelatina 92/8. Same as Gomma A

Gelatina da guerra. Accdg to Piantanida (Ref 4, p 257), it consisted of NG 86, CC (Colloid Cotton) 10 & camphor 4%
**Gelatina dinamite.** See item i, under DINAMITI

**Gelatina dinamite incongelabile.** See item k, under DINAMITI

**Gelatina esplodente o Gelatina gomma.** See item g, under DINAMITI

**Gelatina esplosiva da guerra.** See item h, under DINAMITI

**Gelatina gomma.** See Gelatina esplodente, item g, under DINAMITI

**Gelatina P₁.** Gelatinous expl used during WWII: DNT 17, CC (Collodion Cotton) 0.5, Amm perchlorate 36, Na nitrate 25, TNT 3.3 & PETN 18% (Add to 99.8%); Trauzl value 460cc & gap 0cm (Belgrano, Ref 31, p 232)

**Gelatina P₂.** Gelatinous expl used during WWII: DNT 22.5, CC 0.5, Amm perchlorate 36, Na nitrate 25 & TNT 3.75% (adds to 87.75%); Trauzl value 400cc & gap 0cm (Ref 31, p 232)

**Gelatina P₃.** Gelatinous expl used during WWII: DNT 17, CC 0.5, Amm nitrate 42, K perchlorate 16.5, Na nitrate 18 & TNT 6%; Trauzl value 350cc & gap 0cm (Ref 31, p 232)

**Gelatina Vender.** See item o, under DINAMITI

**Gelatinizzanti (Gelatinizers).** Belgrano (Ref 31, pp 218–19) lists: Centralities, phthalides, alkylphthalates (such as ethyl & butyl), diphenyldurethane and ethylphenylurethane

**Geligniti.** Blasting expls originating in England. The formation given by Molina (Ref 1, p 291) is NG 59, CC 4, K nitrate 29 & woodflour 8%. Another weaker formulation called *Gelignite d'ammonio* contd NG 29.3, CC 0.7 & Amm nitrate 70%. The stronger formulation was used by Italians as a bursting chge in some Land Mines (Ref 28, p 334)

**GEO and GEOM.** Mining Explosives. Their comps and props are given in Table I, under ESPLOSIVI DA GUERRA

**Gomma A and Gomma B.** See Table I, under ESPLOSIVI DA GUERRA

**Gomme incongelabili.** Accord to Molina (Ref 1, p 303), these gelatinous, nonfreezing expls were manufd after WWI by the Dinamifico di Orbetello. Their comps were: NG 72 to 65, CC 6 to 5, TNT 7 to 8 & Amm nitrate 15 to 22%

**GRANATA (Grenade).** Although this term is also applied by the Italians to a shell (like German Grenate), it is preferred to use the term *proiettile for shell or projectile*, as it is in Munizioniamento ITALIANO (AddnlRef C). We are applying the term granata to Hand Grenade (*Granata a mano*) and Mortar Grenade (*Granata a mortaio*), but Italians did not use Rifle Grenade (*Granata da fucile*) during WWII (See also CARTOCCI GRANATA and under Hand and Mortar Grenades)

**Grisou (Esplosivi).** See Antigrisou (Esplosivi)

**Grisoudinamite.** Permissible expl contg AN 88, NG 10, CC 0.5 & woodflour 1.5% (Ref 31, p 326)

**Grisounite contd 95.5% AN & 4.5% TNN (Trinitronaphthalene).** (Ref 31, p 326)

**Grisounite gomma contd 70% AN, 29.5% NG & 0.5% CC (Ref 31, p 326)

**Grisounite roccia or Rock Grisonite: AN 91.5 & TNN 8.5% (Ref 31, p 326)

**Grisoutina.** See item 3, in Table II, under ESPLOSIVI DA MINA

**Grisoutina all' contd 11.76% NG, 0.24% CC & 88.00% AN (Ref 31, p 324)

**Grisoutina all: 13.2% NG, 0.52% CC & 86.55% AN (adds to 100.27%) (Ref 31, p 324)

**Grisoutiti o Dinamiti senza fiamma (Flameless Dynamites).** They are described by Molina (Ref 1, pp 296ff). One example is given by Belgrano on p 326 of Ref 31: NG 44, NC 12 & MgSO₄,7H₂O 44%. The sulfate served as a cooling agent for the gases of explns

**Guanammon.** An experimental castable expl mixture of AN 85 & dicyandiamide 15% (Ref 4, p 244)
Hand and Mortar Grenade (Granata a mano e Granata da mortaio). Italian HE-A/P (high
explos-antipersonnel) grenades used during WWII
were almost totally of the "offensive" type.
Although the loading factor was usually low, the
grenade bodies were not adapted for maximum
fragmentation. The antitank (A/T) hand gre-
nades were adaptations of the A/P grenades with
an addnl charge. There was no evidence of the
use of shaped charges in A/T grenades. Like-
wise, chemical grenades were made of adapted
A/P grenades. No rifle grenades were used,
but instead there was used a small mortar,
serving as a grenade projector. The grenades
were of the impact type, being provided with an
"all-ways" fuze which armed in flight
The following types are described in TM 9-
1985-6 (1953) (Ref 16): Breda Hand Grenades
M35, M40 & M42 (pp 155–56, Fig 226);
Hand Grenade OTO, M35 (pp 155–157, Fig
227); Hand Grenade SRCM, M35 (p 158, Fig
228); PCR Grenade (p 158, Fig 229); "L" Type
A/T Hand Grenade (p 160, Fig 230); Breda
Mortar Grenade (pp 160–61, Fig 231); Incendiary
Bottle Grenade (p 162, Fig 232); Smoke Hand
Grenades (p 162, no Fig); Breda Drill Grenade
(p 162, Fig 233); and SRCM Practice Grenade
(p 163, Fig 234)

Hexocire (Esplosivi). RDX–Beeswax Explosives.
The following formulations are listed in Belgrano
(Ref 31, p 264): Hexocire 95/5, Hexocire
graffitata 98/2/1 and 95/5/0.5

Hexocire-Aluminum. RDX 80, beeswax 5 &
Al 15%; deton vel 8350m/sec (Ref 31, p 264)

Hexogene o T4. RDX or Cyclonite. Deton vel
8520m/sec at d 1.71 (Ref 31, p 264)

Hexogene-Nylon, 95/5. Deton vel 8600m/sec
at d 1.73 (Ref 31, p 264)

Hexaliti. RDX/TNT Explosives, such as 30/70,
50/50, 60/40, 70/30, 80/20 & 90/10. Their
deton vels are betw 7240 & 8200m/sec at d
1.6 (Ref 31, p 264)

High Explosives (HE's). See Esplosivi alti

Idrazina o Diammide, H2N.NH2, Hydrazine.
See Belgrano (Ref 31, pp 127–28) and in this
Vol of Encycl, p H190-L

Idrazina nitrat, N2H4.HNO3, Hydrazine Ni-
trate (Ref 31, p 128) and in this Vol of Encycl,
p H 196-R

Idroliti. Ammonium nitrate expls contg water,
which serves to lower mp of AN. They were
manufd by the Societá Dinamite Nobel at Avi-
giana. The example cited by Plantanida (Ref
4, p 246) contd AN 70, Hexogene (T4) 20,
paraffin 3 & water 7%. This expl was insensi-
tive to shock (Ref 28, p 334)

Igniter Composition for Propellants, reported in
OrdnSergeant (Ref 4a, p 17), was BkPdr of
compr K nitrate 74.4, charcoal 16.5 & S 9.1%.
It was initiated by a primer cap contg MF 28.4,
KClO3 34.4, Sb2S3 35.6 & ground glass 1.6%

Igniters, Military. See Accenditori militari

Imperialite. Blasting expl consisting of AN 85–
90 & Al scales 15–10%. Its inventor Imperiali
was killed in a violent expn during mixing of
ingredients (Ref 28, p 334 & Ref 31, p 316)

Incendiarie (Miscela). See Composizioni (o
Miscela) Incendiarie

INNESCAMENTO (Priming or Initiating of
Explosion)
Under this title, Belgrano (Ref 31, p 499ff)
describes the following items:
a) Capsule da mina ordinarie o Detonatori
normali (Blasting Caps or Ordinary Detonators)
(pp 501–505)
b) Principali prove da eseguire sui detonatori
(Principal Tests of Performance of Detonators)
(pp 505–507)
c) Detonatori secondari (Compound Detonators
(p 507)
d) Detonatori o Inneschi elettrici (Electric Deton-
ators or Primers (pp 508–510)
e) Esplositori per l'accensione dei detonatori
elettrici (Exploders for Igniting Electric Deton-
ators) (pp 511–514)
f) Accenditorni elettrici (Electric Igniters) (pp 514-515)
g) Principal Types of Detonators and Their Characteristics are listed in Table 73, pp 516-17. Twelve types are Italian, the rest are W. German, Austrian, USA and French

Giorgio (Ref 26), under the title "Mezzi d’ innesco di uso militare" (Initiating Devices for Military Use), describes the following items:
Detonatori comuni (Ordinary Detonators) (pp 192-94); Detonatori elettrici instantanei (Instant Electric Detonators) (pp 195-98); Detonatori elettrici a ritardo ordinario (Electric Detonators with Ordinary Delay) (pp 198-99); Detonatori elettrici a microritardo (Electric Detonators with Micro Delay) (pp 199-200); Miccia a lenta combustione (Slow Fuse) (pp 189-92); Miccia detonante (Detonating Fuse) (pp 200-202); Miccia a rapida combustione (p 202); Cordone di accensione, called in US Igniter Cord and in Italy Pirofis (pp 202-203)

Accdg. to OrdnSergeant (Ref 44, p 18), Italian Detonators of WWII contd as a primary charge a mixt of LST (Lead Styrphate), LA (Lead Azide) and, as a secondary (base) charge, PETN (Pentaerythritol Trinitrate)

Aadin. Ref from CA:
[Discussion on types of detonators, including metal filaments, is given. Some detonation parameters for T₄ (RDX), Trinitol (TNT), NG, MeNO₃ and Tritolite (TDX-TNT mixt) are included]

Kratiti e Nitrokratiti. Series of expl mixtures developed by U. Alvisi, which consisted of Ammonium Perchlorate with NG and/or NC (Ref 1, p 200 & Ref 31, p 363)

Land Mines. See under Mines and Traps

Luci colorate (Miscela). See Composizioni (o Miscela) a luci colorate in Belgrano (Ref 31, p 627) and under PIROTECNIA o ARTIFICI DA GUERRA

M₂₄, M₆₆, M₈₈ Polveri al nitrometriolo. See under Polvere al nitrometriolo

MABT. High Explosive compn consisting of PA, TNT & DNPPhenol. Can be prep’d by mixing MAT with MBT. Used as bursting charge (Ref 11, Vol 2, p 119)

Manlianiite. A Chedite-type expl consisting of Anm Perchlorate 72, carbon 14.7 & sulfur 13.3%. It was proposed by U. Alvisi (Ref 1, p 199)

MAT (Picratol). A castable HE mixture of PA 60 & TNT 40%. It is a yel-buff solid which on heating becomes plastic at 55° and melts at 85°. Its props are: Ballistic Strength 103% TNT; Brisance by Sand Test 44g sand crushed vs 43g for TNT; Explosion Temperature 240-280° in 5sec; Impact Sensitivity with 2kg Weight 13 inches vs 14 inches for TNT; Rôle Bullet Sensitivity, 10% detonations from impact of a .30 cal bullet fixed at a distance of 90 feet; Stability, stable in storage; Velocity of Detonation 7100m/sec at density 1.62 vs 6900m/sec for TNT. Used for cast-loading medium caliber shells (Ref 11, Vol 2, p 111 & Ref 28, p 335)

MBT (Explosivo). A castable HE mixture of PA 60 & DNPPhenol 40%. It is a yel solid which becomes plastic at 68° and melts at 77°. It is less brisan and less powerful than TNT and requires a strong initiator for detonation; used for loading medium caliber shells. Its mixture with MAT is known as MABT (Ref 11, Vol 2, p 119 & Ref 28, p 336) [Comp with French DD (Explosif) and Japanese Chaoyaku]

Melinita (Picric Acid), described here as Acido picrico

Metriolo (Nitrometriolo) o Nitropentaglicerina, H₃C.C(CH₂ONO₂)₃, (Metriol Trinitrate, Mtterr.NT). A slightly turbid, heavy explosive oil developed before WWII by the SA Bombini Parodi-Delfino (BPD) and manufacturerd at the Colloferrona-Roma Plant. It was obtd in 92% yield by nitratr trimethylolmethylenh in a batch process using mixed nitric-sulfuric acid of zero water. The process is similar to that used for prep of NG, except that high ratios of acids are used. A detailed description of prep is
given in PATR 2510 (1958), p Ger 113-L

Mettriol Trinitrate has been used in smokeless propellant, such as in Polvere BPD described by Caprio (Ref 11, Vol 2, p 154). By itself, MtrTN does not gelatinize NC unless at temps of at least 110°. However, when ca 8% of acetyl-metriolo (triacetate of metriol), H₃C.C(CH₂.OCH₃)₃, is added, gelatinization takes place at 80°. The cool Polvere BPD described by Caprio (Ref 11, Vol 1, p 156) contd: MtrTN 59, NC 33, acetyl-metriol 5.5 & Centralite 2.5%.

Advantages of MtrTN are low volatility, low calorific value (which means less erosion of gun barrels) and a good degree of flashlessness. The Italian Navy was greatly impressed by the considerably reduced flash in “Metriolo propellenti” (Ref 28, p 336).

Mezze d’innesc (Initiating Devices). See under INNESCAMIENTO

Mezze d’innesci di uso militare (Initiating Devices for Military Use). See under INNESCAMIENTO

Miscia (Fuse). See under INNESCAMIENTO

Mines and Traps. Italian antipersonnel (A/P) land mines of WWII were operated by pressure or trip wires. They were often difficult to detect, especially those operated by pressure in which only a portion of the lid or igniter appeared above the ground. Bakelite and wood were used in the construction of some mines to prevent detection by magnetic instruments.

Anti-tank (A/T) mines varied in appearance, being tubular, rectangular, or circular in shape. Some were made of bakelite or wood.

Improvised mines seem to have been the Italian specialty in the field. They used the majority of such mines in Abyssinia, since the supply of standard mines was apparently limited. Most of these mines were of wooden construction and used blocks of TNT for the bursting charge. Sometimes shells and shrapnel were used for mines.

The following items are described in TM 9-1985-6 (1953) (Ref 16): Picket A/P Mine (p 165, no Fig); B4 A/P Mine (pp 165–66, Fig 236); Bakelite and Wooden A/P 1-lb Mine (p 166 & p 237); Ratchet Mine (Railroad Mine) (p 167, Fig 238); Railway Mine (pp 167–68, Fig 239); Wooden Box A/T Mine (p 168, Fig 240); Four-Igniter A/T Mine (pp 168–69, Fig 241); A/T Mine B-2 (pp 169–70, Fig 242); A/T Mines V-3 and V-5 (p 170, Fig 243); Pignione Type I & II Bakelite A/T Mine (pp 171–72, Fig 244).

Note: Although the title of the Section includes the word “Traps”, no description of such item is given.

Miscela C o PE. A plastic expl consisting of RDX and a plasticizer; Miscela C₂ – RDX, CC & plasticizer is water-resistant; Miscela C₃ – RDX, Tetryl, CC & plasticizer; Miscela C₄ – Plastic expl consisting of RDX 91 with Polysobutylene & other ingredients (Ref 31, p 263).

Miscuglio nitrico-sulfurico per nitrazione. (Mixture of nitric-sulfuric Acids for Nitration). As an example, Belgrano (Ref 31) describes on pp 159–162, prepn and analysis of acids used in nitration of glycerol.

MNDT. See Siperit

MST. See Nougat

MTX. A mixture of French origin: Mellinite (PA) 55, TNT 35 & Xilite (TN-meta-Xylene) 10% (Ref 31, p 249).

NAC. See Polvere NAC

NA–OC (Nitrato ammonico-olio combustibile). See AN–FO

Nitram X. One of the mining expls, listed in Ref 31, p 171

Nitramite o Avigliana 3. An Ammonal-type expl mixture consisting of AN 71–72, Al 22 & bitumen pitch (or paraffin) 6–7%. Used as a bursting charbe in all types of ammunition (Ref 4, p 238; Ref 28, p 337 & Ref 31, p 316).

Nitroammonico o Nitratio d’ammonio (Ammonium Nitrate, abbrd as AN). It is described in Ref 31, p 306–16. Used in a wide variety of military and commercial expls, such as: Afocite, Albite, Amanoto, Ammonal, Ammonite No 1, ANS or ASN, Astralite, Cremonite, Dinamiti,
Dinamon, Echos or Escho, Esplosivi da guerra, Esplosivi da mina, Esplosivo S20, FNP Esplosivo, Gellignite, Gomma, Idrolita, Imperialista, MST or Nougat, PAN, PANA, PNP, Polvere “Cannel”, Romite, Sabulite, Schneiderite, Sipere, Solfito, Umbrite, Vibrate, Virrite and others (Ref 28, p 337)

Nitrate X Galleria. One of the mining expls listed by Belgrano (Ref 31, p 171) without giving its composition

**NITROCELLULOSE o NITROCOTONE.** There are three Italian basic types for use in Polveri senza fumo (Smokeless Propellants):

A. **Cotone collodio** (Collodion Cotton), which contains 11.2 to 12.3% N

B. **Pirocolloidio** (Pyrocotton), which contains 12.5 to 12.7% N

C. **Fulmicotone o Cotone fulminante.** (Gun-cotton, abbrd as GC), which contains 13.2 to 13.4% N (Ref 31, p 188)

All NC’s may be divided into the following groups:

a) **Nitrocellulose a solvente volatile** (NC used with ether-alcohol solvent) is usually Pirocolloidio. As an example of its use is the Polvere per mitragliatrice FIAT mod 35 (Propellant for machine gun FIAT, Mod 35), which consists of NC 97, Centralite 2 & DPhA (Diphenylamine) 1% (Ref 31, p 203)

b) **Nitrocellulose a solvente fisso** (NC used with fixed solvent) is usually Cotone collodio of N content ca 12% and Pirocolloidio. As a fixed solvent, NG or its mixture with NGc (Nitroglycol) is used. Polveri NAC (qv) is an example of using Cotone collodio and Balistite attennata (qv) is an example of using Pirocolloidio (Ref 31, pp 205–06). Other examples of propblnts using NC with fixed solvents are Polveri FB, FC, al nitrodigicol, al nitrometriolo, and al trinitroanisol are listed in Ref 31, pp 208–10 and also here under Polveri

c) **Nitrocellulose a solvente misto** (NC used with mixed solvent) is usually a mixture of Pirocolloidio & Fulmicotone. Mixtures of NG with ether-alcohol or with acetone are used as solvents. Examples of these propblnts are Polvere C2 and Solenite ordinaria (Ref 31, p 212)

**Nitrocratiti.** See under Kratiti and in Ref 31, p 363

**Nitrodigicolie, Nitroeterolo o Dinitrodiiten-glico (Diethylenegycol Dinitrate, abbrd as DEGDN), O_2NO.CH_2.O.CH_2.ONO_2, is described in Belgrano (Ref 31, pp 136–39). Colori expl liq used as a component of propnt, and composite expls. The Italian Army Specification Requirements listed in the pamphlet “Capitolato Tecnico Generale per la Fornitura di Esplosivi Propellenti”, MD Esercito (1951), CTF 28, are as follows: N content 14.10%, alkalinity (as %Na_2CO_3) 0.0023; and stability by 120° German test, 30 mins for red coloration of indicator paper (Ref 28, p 337)

Two composite expls contg DEGDN are listed in Ref 31, pp 139–40: a) Esplosivo da mina — DEGDN 36.5, CC 1.5, PETN 10.0, AN 28.0 & Na nitrate 24.0%  b) Esplosivo plastico — DEGDN 32 & RDX 68%, experimentally used by Ital Genio Militare (Corps of Engineers)

Two Polveri al nitrodigicol are listed in Ref 31, p 208: a) A 860cal/g-DEGDN 68, Nitroacetylcellulose 30 & Centralite 2% b) A 730cal/g-DEGDN 27.0, Nitroacetylcellulose 63.5, acetylcellulose 5.0 & Centralite 5%

**Nitrogelatina.** Same as Gelatina dinamite, described as item i, under DINAMITI

**NITROGLICERINA, Trinitroglicerina o Olio esplosivo (Nitroglycerol, abbrd as NG),**

(O_2NO)_3C.CH(OONO_2),CH_2(OONO_2). An only expl liquid first prepb by Ascanio Sobrero in Univ of Torino, Italy. A detailed description of its prep, props, analysis and uses is given by Belgrano (Ref 31, pp 140ff)

The Italian Army Specification Requirements as of 1951 were as follows: Titolo d’azoto (N content) 18.36%; Reazione (Reaction) neutral to phenolphthalein; and Stabilità al saggio Abel a 80°C (Stability by Abel Test at 80°C), 14 minutes (Ref 28, p 338)

Used in many military and commercial expls and propblnts, such as Balistite & other double-base propblnts, Dinamitt, Esplosivi da mina, Gellignite, Gomma, Pentritite & others
Nitroglicol e Dinitroglicol [Ethylene glycol Dinitrate (EGDN) or Nitroglicol (NGc)], O₂NO₂.H₂C.CH₂ONO₂. An oily expl liquid, first prep’d in 1870 by Henry. Its prepns, props and analysis are described by Belgrano (Ref 31, pp 133–36).

Used in many countries as an additive to NG to prevent its freezing. Such expls are known as Dinamiti and Gelatine incongelabili.

Nitroguanidina (Nitroguanidine, abbrd as NGu), H₂N.C(NH)NH.NO₂, wh ndls, mp 246–47°, first prep’d in 1874 by Jousselain. Its prepns, props, uses and analysis are described by Belgrano (Ref 31, pp 251 to 254). NGu has been used as a component of military and commercial expls and as a cooling agent for smokeless propnts. Four mining expls contg 15 to 28% are listed in Ref 31, p 253 and in Ref 28, p 338. Expl compns Albite and Umprite were used during WWII as bursting charges (Ref 28, p 338).

Nitrometriolo (Metriol Trinitrate). See under Metriolo.

Nitronaftiti. Accdg to Piantanida (Ref 4, p 236), they are mixts of RDX 75–80% with MN-Naphthalene 25–20% manuf’d by the SA Dinamite Nobel at Avigliana. The mixt softens at 75–80° but does not melt; is press-loaded, while soft, in ammunition (Ref 28, p 339).

Accdg to Belgrano (Ref 31, p 26), Nitronaftiti are plastic mixts of RDX & DNNaphthalene, fusing at 70–80°. They are more powerful than TNT.

Nobelita Galleria. A mining expl consisting of NC 37, KCIO₄ 34, NaNO₃ 24, DNT 3 & mineral oil 2%; temp of expln 2800°C; burns in air at ca 130° (Ref 31, p 362).

Nougat o MST (Esplosivo). An Amatol-type expl consisting of AN 49, TNT 44 & DNanaphthalene 7% (Ref 11, Vol 2, p 93 & Ref 31, p 297). It has been used cast-loaded in the following shells: 120/21mm HE, 120/21mm HE (cast steel); 149/35mm HE, 149/35mm HE (one piece); 210mm HE (bomba), 305/17mm HE (cast steel) and 305/17mm HE (one piece) (Ref 28, p 339).

NTP. A military expl mixture consisting of AN 77, RDX 20 & paraffin 3%; deton vel 5850m/sec at d 1.62 (Ref 11, Vol 2, p 114 & Ref 31, p 266).

NX. Mixture of AN 77 & Xilite (TN-m-xylene) 23% (Ref 31, p 249).

Obice. Howitzer.

Olio da combustione. Fuel Oil.

Olio esplosivo. One of the names for NG (Nitroglycerol).

Olio di sgocciolamento (Drip Oil in the USA and Tröpföl in Germany). A crude, commercial, black, oily substance resulting from the two-stage nitration of toluene in the manuf of TNT. Its main constituents are isomers of DNT with small quantities of isomers of TNT and some other aromatic nitrocompounds. It has been used as an antifreeze addn to Dynamites, such as in Dinamiti incongelabili; in some AN expls and as a gelatinizer of NC used in some propnts. Due to the fact that its composition is variable, and it contains some sulfuric acid derivatives, its use in military expls is not recommended.

Another way to utilize “drip oil” is to separate it from purified DNT, and nitrate the oil to TNT, thus making the manuf in three stages. The resulting product is nearly as powerful as TNT but of inferior stability (Ref 28, p 339).

A typical commercial explosive is given on p 340 of Ref 28: Olio di sgocciolamento 16, TNT 7, AN 45, Na nitrate 15 K perchlorate 16.5 & CC 0.5%

Ossonitu. A liquid, Sprengel-type expl, prep’d in situ by mixing ca 58 parts of PA (or 28 pts of MNB) with concd nitric acid, just before use (Ref 31, p 379). Accdg to Ref 28, p 340, some liquid N peroxide is also added to PA and nitric acid.

Oxilliquite. It is one of the esplosivi all’aria liquida (Liquid Air Explosive) prep’d in situ by pouring liquid air (or liquid oxygen) into a mixture of 3 parts of carbon, impregnated with 2 parts of petroleum (Ref 31, p 305).
Oxilite. A liquid air expl prep’d in situ by pouring liquid air (or liquid oxygen) into a mixture of fossil flour, previously impregnated with petroleum (Ref 31, p 305)

NOTE: Accdg to Belgrano both mixts (Oxilique & Oxilite) have been used in high mountains for construction of hydroelectric plants, for construction of the Sempione tunnel and for work by Genio Militare (Corps of Engineers) (Ref 31, p 305)

P₁, P₂ & P₃. See Gelatina P₁, P₂ & P₃

PA (American). Abbrn for Picric Acid

PA (Italian). Mixtures of Pentrite (PETN) with Acetato di pentaoeritrile (Pentaerythritol Tetranitrate or PETA in proportions of 65/35 & 75/25. They were developed by BPD (Bombini-Parodi-Delfino SA, Colloferro-Roma). They melt at 85–88°C, have the same power as TNT and are fairly insensitive to shock. PA mixts were proposed for cast-loading shells, but were seldom used because PA is more expensive than TNT. Accdg to Plantanida (Ref 4, p 232), PA was also used in compressed form as a secondary charge in detonators. See also Ref 28, p 340 & Ref 31, p 182)

Palla – Solid Shot; Ball

Palla dum dum – Dumdum Bullet

Palle – Ball Ammunition

PALLOTTOLA – Bullet; Rifle Bullet; Shot

Pallottola esplodente – Explosive Bullet; Dumdum Bullet

Pallottola perforante – Armor-piercing Bullet

Pallottola a revestimento d’acciaio – Steel-jacketed Bullet

Pallottola tracciante – Tracer Bullet

PAN. Mixt of PETN 42, PETA 23 & AN 35% developed by BPD for press-loading ammunition (Ref 4, p 232; Ref 28, p 340 & Ref 31, p 182)

PANA. HE mixt of PETN 32, PETA 24, AN 35 and Al powder 9% developed by BPD for press-loading ammunition. It is more powerful than Ammonal, but also more expensive (Ref 4, p 232; Ref 28, p 340 & Ref 31, p 182)

PE (Amer). Abbrn for Pentaerythritol

PE (Italian). Designation of Miscela C (qv). Do not confuse with C (Polvere)

Pentaerythritol Tetranitrate (PETN). See PENTRITE

Pentolite. See Pentritolo and Pentrol

Pentiniti. Powerful mixts of PETN 80–85 & NG 20–15% invented in 1929 by the late Dr A. Stettbacher of Switzerland. They are described by E. Plantanida in MAF 14, 458 (1935). Belgrano (Ref 31) gives on p 182 its compn but does not list its props

PENTRITE o Tetranitrito di Pentaeritrile (Pentaerythritol Tetranitrate, abbrd as PETN), C(CH₂ONO₂)₄. Its prep, props, uses and analysis are described by Belgrano (Ref 31, p 176–183). Its props given on p 181 of Ref 31 are as follows: Density (max) 1.62, Explosion Temperature 195°C, Flame Temperature on Explosion (Temperature Developed on Explosion) 3600°C, Heat of Explosion 1400 kcal/kg, Specific Pressure 10500 atm/kg, Volume of Gas at 0°C & 760mm 780 l/kg, Traufl Test 500cc, Impact Sensitivity with 2kg Weight 30cm and Detonation Velocity 8400m/sec. Giua (Ref 19) gave some slightly different values, such as: Explosion Temperature 190°C, Flame Temperature 4050°C, Traufl Test 540cc and Impact Sensitivity with 2kg Wt 38cm (listed also on p 258 of Ref 31). Straight, compressed, PETN was used during WWII as a bursting charge in small caliber shells (such as 20mm) and as a booster or base charge in detonators. PETN, phlegmatized by 8–10% wax (or paraffin) and dyed blue, was used as bursting charge in shells larger than 20mm and as demolition charges (Ref 28, p 341). PETN was also used in composite expts, such as ASN, Esplosivi da guerra, Esplosivi da mina, FNP Esplosivo, Idrolita, Nitronaftita, PA, PAN, PANA, Pentiniti, Pentritolo, Pentrol, FNP Esplosivo, Unknown Name Explosives and others
Pentritolo o Pentrol (Pentolite). Mixtures of PETN & TNT in various proportions. Accdg to Caprio (Ref 11, Vol 2, p 115), mixts of TNT with PETN 40–60% were used under the name of Pentrol, by the Genio Militare (Corps of Engineers) in demolition charges. Accdg to Ref 28, pp 341–42, mixts of PETN 50–80 and TNT 50–20% were used as early as 1934 as bursting charges in land mines, demolition charges, and in some underwater ammunition, such as depth charges. Some Pentritols were dyed red and some of them further phlegmatized by immersing in molten paraffin or by incorporating some MNT (Mononitrotoluene) in the mixture. Accdg to Belgrano (Ref 31, p 182), mixts with 40–60% PETN, called Pentrol, have been used in granite (shells or grenades), razzi (rockets), proiettli anticarro (A/T projectiles) or for cariche cave (shaped charges)

Pertite. See ACIDO PICRICO

PETN. See Pentrite

Picramide. See Trinitroanilino

Picrato ammonico o Picrato di ammonio (Ammonium Picrate or AP), (O₂N)₃C₆H₂ONH₄. Its prep and props are in Belgrano (Ref 31, pp 288–89). Straight AP has been used in the US, under the name Explosivo D, while in Italy it was used in composite expls Cremonite (qv) & Picratol

Picratol. Mixture of Ammonium Picrate 52 & TNT 48% listed in Belgrano (Ref 31, p 246) as for use in semi-armor piercing projectiles. Its power by Trauzl test is 380cc vs 285cc for TNT

Piombite. Accdg to Caprio (Ref 11, Vol 2, p 96) and Belgrano (Ref 31, p 351), it is an expl mixture consisting of Lead nitrate 76, Tritonaphthalene 16, Calcium silicide 5 & vaseline 3%. Trauzl Test value 155cc vs 285cc for TNT. It was used in mining expls.

Accdg to Molina (Ref 1, p 358), the following Piombite was used during WWI as a bursting charge in cast-iron shells of medium caliber: Pb nitrate 75, Ca silicide 16, Pb carbonate (basic) 6.5, TNaphthalene 1.5 & vaseline oil 1% (See also Ref 28, p 342)

Pirocollodio. See under Nitrocellolose

PIROTECNIA o ARTIFIZI DA GUERRA
(Military Pyrotechnics). Colonel and now General Attilio Izzo in "Pirotecnia e Guochi Artificiali" (Pyrotechnics and Fireworks), listed as Ref 12, describes various Italian military pyrotechnic devices and gives on pp 204–238 numerous compositions. More recent book is that of Dr Camillo Belgrano (Ref 31) who on pp 623–35 lists, under the title "Pirotecnia", many comps

A. Composizioni a luce gialla per stelle (Comps for yellow light stars) a) NaNO₃ 74, S 6 & Al 18% b) KNO₃ 44, Na₂CO₃ 5, pulverin (meal powder) 23 & shellac 6% (Izzo, p 204)

B. Comp a luce gialla per segnalazioni (Comps for yellow light signals) a) NaNO₃ 55.5 & polyvinyl chloride 44.5% b) KNO₃ 62.5, NaNO₃ 10.5, S 23 & charcoal 4% (Izzo, p 204)

C. Comp a luce gialla per torre (Yed light compn for torches) KClO₄ 18, Na oxalate 9, Ba(NO₃)₂ 60, S 4 & shellac 9% (Izzo, p 205)

D. Comp a luce rossa per stelle (Comps for red light stars) a) KClO₃ 40, Sr(NO₃)₂ 40, charcoal 6, lampblack 6 & tar 8% b) KClO₃ 60, Sr(NO₃)₂ 25 & shellac 15% (Izzo, p 207)

E. Comp a luce rossa per segnalazioni (Comps for red light signals) a) KClO₃ 30, Sr(NO₃)₂ 44, S 18, charcoal 2 & Sb₂S₃ 6% b) KClO₄ 54, Sr(NO₃)₂ 36 & shellac 10% (Izzo, p 207)

F. Composizioni per traccianti (Tracer Comps). In the book of Izzo (Ref 12, pp 210, 213 & 235) are listed a) Red (rosso) -colored tracer – SrO₂ 40, Mg (powder) 40 & Sr oxalate 20% b) Green (verde) tracer – BaO₂ 72.5, Mg 15, Ba oxalate 5 & gumlac 7.5% c) Yellow (giallo) tracer – KNO₃ 50, red arsenic 30 & sulfur 20%

In the book of Belgrano (Ref 31, p 633) are listed under the title traccianti a) Red tracer – Sr nitrate 69, Mg 25 & binder 6%; b) Green tracer – Ba nitrate 55, Mg 35 & binder 10%; c) Tracer – KClO₃ 58, SrCO₃ 22 & shellac 20%

A typical Italian tracer compn examined during WWII at Picatinny Arsenal contd Ba nitrate 63.0, Mg pdr 34.3 & binder-fuel 2.7%. It was used in 47-mm AP (Armor-piercing) Shell (Ref 28, p 322)

G. Comp a luce verde per stelle (Comps for green light stars) a) KClO₃ 30, Ba(NO₃)₂ 53, shellac 15 & lampblack 2%; b) Ba(NO₃)₂ 70, shellac 17 & milk sugar 13% (Izzo, p 211)

H. Comp a luce verde per segnalazioni (Compns
for green light signals) KClO₃ 31, Ba(NO₃)₂ 52, S 10 & charcoal 6% (Izzo, p 211)

I. Comp a luce verde per traccianti (Green light tracer compn) BaO₂ 72.5, Ba oxide 5, Mg 15 & gumlac 7.5% (Izzo, p 213)

J. Comp a luce azzura per stelle (Compn for blue light star) a) KClO₃ 70, ammoniacal Cu sulfate 15 & shellac 15%; b) KClO₃ 50, Paris green 20, Ba(NO₃)₂ 17, shellac 10 & gum arabic or dextrin 3% (Izzo, p 213)

K. Comp a luce azzura per segnalazioni (Compn for blue light signals) a) KClO₃ 54.5, ammoniacal Cu sulfate 27.5 & charcoal 18%; b) KClO₃ 53, ammoniacal Cu nitrate 26, S 5 & charcoal 16% (Izzo, p 214)

L. Comp a luce violetta per stelle (Compn for violet light star) KClO₃ 58.5, SrCl₂ 14.5, CuCO₃ 10, gumlac 7 & S 10% (Izzo, p 215)

M. Comp a luce bianche per stelle (Compn for white light star) a) KNO₃ 65, S 19 & Sb₂S₃ 16%; b) KNO₃ 59, S 30 & powder (meal 11%); c) KNO₃ 60, charcoal 8, petunia 8, iron filings (paraffined) 16 & gum arabic or dextrin 8% (p 216) and d) KClO₄ 61, Al 31 & lycopodium powder 8% (p 218)

M₁. Comp per stelle (Star compn). Belgrano, pp 632–33 lists the following compns: a) NaNO₃ 74, S 8 & AI 18%; b) NaNO₃ 56, S 21, charcoal 14, Na bicarbonate 4.5 & SrSO₄ 4.5% and c) Ba nitrate 50, KClO₃ 35 & gumlac 15%.

N. Comp a luce bianche per segnalazioni (Compn for white light signals) a) KNO₃ 52, S 22, powder (meal 22 & Sb₂S₃ 4%); b) KNO₃ 61, S 19, powder (meal 5 & Sb₂S₃ 15% (Izzo, p 218)

O. Comp per miscelle sibilanti (Compn for whistling mixtures) a) K picrate 70 & NaNO₃ 30%; b) Gallic acid 50 & KClO₃ 50% (Izzo, p 226)

P. Composizione per fumata grigia (Compn for gray smoke) KNO₃ 50, lampblack 12.5, charcoal 12.5, colophony 12.5 & As₂S₂ 12.5%. The mix is packed in cylinders 10cm in diameter and 25cm long. For its initiation a mixture of KNO₃ 66, S 13, powder (meal 11 & Sb₂S₃ 11% is used (Izzo, p 231)

Q. Composizioni per miscelle fumogene a fumo bianco (Compn for mixt producing white smoke). Formula used by Italians during WWI: NaNO₃ 25, mineral oil 15, sawdust 50 & water 10%. It was initiated by a mixture of NaNO₃ 47.5, KNO₃ 5 & sawdust 47.5% (Izzo, p 233). Belgrano, p 631 gave CuCl₄ 50, Zn dust 25, ZnO 20 &

fossil meal 5%

R. Miscela all’esclorotano (Mixtures with hexachloroethane). There are two mixt proposed by Izzo and three mixts by the Società ACNA. They contd 45 to 52.5 hexachloroethane, 25 to 31.5 Zn dust, 0 to 2.0% ZnO and some other ingredients (Izzo, p 234). Belgrano, p 631, gives HClE₈ 50, Zn 25, ZnO 20 & fossil meal 5%

S. Composizione a tetrachlorotano (Compn with tetrachloroethane): TeCl₄ 40, ZnO 20, Zn dust 15, Ca silicate 15 & NaClO₃ 10% (Izzo, p 234)

T. Composizioni per miscelle fumogene colorate (Compn for colored smoke mixts). Good results were claimed to be obttd with the following: Coloring substance (dye) 36, KClO₃ 28, milk sugar 28, sulfur 4 & Ammonium chloride 4% (Izzo, p 231)

U. Composizioni a fumo giallo (Compn for yellow colored smoke) a) KNO₃ 27, As₂S₂ 27, S 27 & Sb₂S₃ 19%; b) Redalgar (As₂S₃) 30, S 20 & KNO₃ 50%; used in traces for rockets; c) Chrysoidine 30, Auramine (yellow) 10, KClO₃ 35 & milk sugar 25% (Izzo, p 235). Belgrano, p 632 gave Chrysoidine 9–30, Auramine yellow 34–10, KClO₃ 33–35 & lattusio (milk sugar) 24–25%

V. Composizione a fumo arancione (Compn for orange-colored smoke) Chrysoidine 45, KClO₃ 25, milk sugar 30 & mineral coal flour 5% (Izzo, p 235). Belgrano, p 632, gives HClE 20, Auramine 14, Red of p-Nitroaniline 23, KClO₃ 25, MgCO₃ 4 & lattusio (milk sugar) 14%

W. Composizioni a fumo rosso (Red smoke compn) a) Red of para-Nitroaniline 60, KClO₃ 20 & milk sugar 20%; b) Red of p-Nitroaniline 26, Rhodamine B 53 & KClO₃ 21% (Izzo, p 236). Belgrano, p 631, gives Rhodamine 70, KClO₃ 15 & milk sugar 25%; also HClE 30, Rhodamine 34, KClO₃ 18, milk sugar 14 & MgClO₄ 4%

X. Composizioni a fumo verde (Green smoke compn) a) Auramine yellow O 15, Indigo 26, KClO₃ 33 & milk sugar 26%; b) Auramine yellow O 10, Indigo 20, KClO₃ 30, milk sugar 20 & malachite green oxide 20% (Izzo, p 236). Belgrano, p 632, gives HClE 31, Green of Malachite 35, KClO₃ 16, milk sugar 14 & MgClO₄ 4%

sugar 15-20 & Methylene Blue 20 or 0%
Z. Composizioni a fumo nero (Black smoke comps) a) Hexachloroethane 50, anthracene 15, naphthalene 15 & Mg powder 20% (Izzo, p 236 & Belgrano, p 631); b) HCIE 65, anthracent 6, α-naphthol 17 & Al powder 12%. They were used in Italian smoke bombs (bomba fumogena) initiated by a mixture of KNO₃ 61, S 17.5, As₂S₂ 17.5 & gum arabic or dextrin 4% (Izzo, p 236) and c) KClO₃ 31, naphthalene 11, red P 11 & tar 47%; d) KClO 45, naphthalene 40 & charcoal 15% (Izzo, p 237)
Z₁. Composizione a fumo viola (Violet smoke comp) 1-methylaminonitroquinone 18, 1,4-
diamino-2,3-dihydroanthraquinone 26, KClO₃ 30.2, S 11.8 & Na bicarbonate 14% (Belgrano, p 632)

Accdg to Gen Izzo (Ref 12, p 200), the principal Italian factories manuf pyrotechnic comps and items were, as of 1950, as follows: Polveri-fichi Stacchini di Roma, la ditta Camocini di Como, la Muggia, i, la Società ACNA, etc

The following Italian Military Pyrotechnic smoke comps used in colored smoke signal devices for air-to-air liaison were described by J. Goldenson & C.E. Danner in Chem & Engg News 26, 1976-78 (1948):
Red — KClO 26, lactose 27, Sudan IV 27, Rhodamine B 14, Auramine 5 & sucrose 1%
Yellow — KClO₃ 30, “Auramine O” 65 & kieselguhr 5%
Green — KClO₃ 27, lactose 25, dimethylanilinoazobenzene 21 & 1,4-di-p-toluidinoanthraquinone 27%
Black — KClO₃ 60, naphthalene 20 & α-MN-Naphthalene 20% (Ref 28, pp 342-43)

Following is addnl information given in the book of Belgrano (Ref 31)
I. Miscela a luci colorate (p 627), calc'd in parts: a) White — KNO₃ 70, pulvemin 30, S 30 & Sb₂S₃ 5 pts; b) Red — Sr nitrate 45, KClO₃ 30, S 18, Sb₂S₃ 6 & carbon 2 pts; c) Green — KNO₃ 50, KClO₃ 30 & 10 pts; d) Yellow — KNO₃ 60, NaNO₃ 10, S 22 & carbon 4 pts; e) Violet — KClO₃ 42, Sr nitrate 18, S 30, calomel 5 & CuCO₃ 5 pts and f) Blue — KClO₃ 50, Cu nitrate 25, carbon 15 & S 5 pts
II. Miscelle illuminanti compresse (Ref 31, pp 629-30)
a) Mg 43, Na nitrate 47 & binder (laminac) 10%. For its initiation use 75 pts of above mixture with 25 pts of Black Powder. When compressed at 150 atm, its luminosity is 600000 candles
b) Mg 55, Na nitrate 33, Laminac 6 & Ba stearate 6%; luminosity 600000 candles
c) Mg 35, Al 5, Ba nitrate 52, Na oxide 5.5 % & oil 2.5%; luminosity 540000 candles
III. Miscela innescente per fumogeni (Initiating mixture for smoke comps) KNO₃ 61, red arsenic 17.5, S 17.5 & gum arabic 4% (Ref 31, p 632)

PNP Esplosivo. A HE consisting of PETN 20, AN 77, & wax 3% has been used press-loaded as a bursting charge in various projectiles (Ref 26, p 160 & Ref 31, p 182)
PNT. Belgrano (Ref 31) lists it on p 267 without giving its compn
Polvere alla fitale — centralite. See Polvere FC
Polvere alla nitrocellulose. See Polvere NAC
Polvere BPD (Belgrano-Parodi-Delfino Powder). See under Metriolo (Nitrometriolo)
Polvere bruna o Polvere cioccolata (Brown Powder or Chocolate Powder) contd KNO₃ 79, incompletely carbonized wood as powder 18 & sulfur 3%; used in some cannons prior to the invention of smokeless propints (Ref 1, p 147-8 & Ref 28, p 343)
Polvere C. Same as Cordite Italiana
Polvere C₂. See C₂ (Polvere)
Polvere C-12. See C-12 (Polvere)
Polvere “Cannel”. A Cheddar-type expl contg Amm Perchlorate 80 & Cannel coal from Scotland 20% was proposed in 1900 by U. Alvisi as a blasting expl (Ref 1, p 200)
Polvere CG-13. A Ballistite-type expl contg NG 25, NC (N 12.5%) 60 & DNT (solid) 15% (Ref 11, Vol 2, p 139 and Ref 31, p 210)
Polvere CG-14. A Ballistite-type expl contg NG 25, NC (N 12.5%) 60, DNT (solid) 10 & Centralite 5% (Ref 4, p 183 and Ref 31, p 210)
Polvere FB o Polvere allo fitalato di butilo.
There are 860 and 960 calorie-types, which contd., respectively: NG 32 & 31, NC (12% N) 57 & 61, butyl phthalate 9 & 5.5 and Centralite 2 & 2.5%; used in rapid-fire cannons (Ref 11, Vol 2, p 148 & Ref 31, pp 206 & 208)

Polvere FC o Polvere alla fitalide-centralite.
There are 860 & 960 calorie-types, which contd., respectively: NG 28 & 32, NC (12% N) 64 & 62.5, phthalide, C₆H₄.CH₂.O.CO 4 & 3.5, Centralite 3 & 1.5 and vaseline 1 & 0.5%; used in rapid-fire cannons (Ref 11, Vol 2, p 148 & Ref 31, p 206)

Polvere per mitragliatrice FIAT mod 35 e cal 8
(Propellant for Machinegun FIAT, Mod 35, Cal 8). See "Polvere italiana per carucce cal 8, per mitragliatrici", under POLVERI DA LANCIO SENZA FUMO, item A. Polveri alla sola nitrocellulosa o Polveri a una base

Polvere NAC o Polvere alla nitroacellullosa
(Propellant with Nitrocellulose). There are 860 & 960 calorie-types, which contd., respectively: NG 27 & 32, Nitrocellulose (N 11.2%) 66 & 63, Centralite 7 & 4 and DPHA 0 & 1%; used in cal 20 & 37mm cannons (Ref 11, p 147 & Ref 31, p 206)

POLVERE NERA (Black Powder) o Polvere da fuoco (Fire Powder) (Abbreviated as BkPdr)

The latest Italian book describing Black Powder is that of Belgrano (Ref 31, pp 341—50). Black Powders are subdivided into three types: 1) Polvere da guerra (Military Type) KNO₃ 74—75, carbon 15—16 & sulfur 10%; 2) Polvere da caccia (Hunting or Sporting Type) KNO₃ 75—78, carbon 12—15 & sulfur 9—12%; 3) Polvere da mina (Mining or Blasting Type) KNO₃ 60—72, carbon 14—21 & sulfur 13—18%

The following average props of BkPdr are listed by Belgrano on p 349: Density (max) 1.7; Temperature of Explosive Decomposition 300⁰C; Temperature of Explosion 2700⁰C; Heat of Explosion 740cal/g, Specific Pressure 3250 atm/kg; Volume of Gas Developed on Explosion, calc'd at STP 325 liters/kg; Trauzl Test value 30—40cc; Impact Sensitivity with 2kg Weight 70cm; and Velocity of Detonation 1400m/sec (See also Ref 28, p 345)

The principal Italian uses of Military BkPdrs are as igniters in percussion primers. For example, the Igniter of the 105/28mm cartridge case contd. 25—30g of BkPdr & ca 5g of lead foil, serving as a decoppering agent (Ref 28, p 345)

Polvere al nitrodiglicol (Propellant with DEGCdN).

Two types are described in Belgrano (Ref 31, p 208), 860 cal & 730 cal of comps, respectively: DEGCdN 68 & 27, Nitrocellulose 30 & 63, acetylcellulose 0 & 5 and Centralite 2 & 4.5%

Polvere al nitrometriolo (Metrol Trinitrate Propellant). Three types are described in Belgrano (Ref 31, p 210), M4, M6 & M8 with comps, respectively: Metrol Trinitrate 55.5, 57.5 & 59%; NC 40, 36 & 33%; acetylmetrol 2.5, 4.5 & 5.9%; and Centralite 2, 2 & 2.5%

Polvere al trinitroanisolo—Stabilite. Two formulations are listed in Belgrano (Ref 31, p 210):

a) TNAns (Trinitroanisole) 40 & NC 60% b) TNAns 34, NC 60, NG 5 & water 1%

Polvere verde (Green Powder). A Cheddite-type expl: KNO₃ 66.7, PA 19.0 & K₂Fe(CN)₆ 14.3%; was used in blasting operations (Ref 28, p 346).

Belgrano (Ref 31, p 356) lists this expl, but gives for KNO₃ 39%, which is evidently in error

POLVERI DA LANCIO SENZA FUMO (Smokeless Propellants) o Polveri colloidalni (See also Explosivi da lancio o Deflagranti)

Giorgio (Ref 26, pp 204—08) subdivides them as follows:

A. Polveri alla sola nitrocellulosa a solvente volatili o Polveri a una base (Single Base Propellants with Volatile Solvent). To these belong a proplnt listed by Giorgio, p 205 as Polvere italiana per cartucce cal 8 per mitragliatrici, which contd.: NC (12.8% N) 97, Centralite 2 & DPHA ( Diphenylamine) 1%. It is in the form of tubes 0.75—01 x 2mm, graphited. The same proplnt is listed by Belgrano (Ref 31, p 204) as "Polvere per mitragliatrice FIAT mod 35 e cal 8" (Propellant for Machinegun FIAT Mod 35, Caliber 8)

B. Polveri a doppia base (con NGL) [Double-Base Propellants with NG (Nitroglycerin)] are divided into:
a) Polvere a doppia base senza solvente volatile (Double-Base Propellants Without Volatile Solvent), called by Belgrano Polvere a solvenente falso (Propellants with Fixed Solvent). To these belong Balistite normale e ordinaria (with 50% NG); Balistite a basso titolo (with 42% NG) (Ref 26, p 205); Polvere FB, Polvere FC & Polvere NAC (Ref 31, p 206)
b) Polvere a doppia base con solvente volatile (Double-Base Propellants With Volatile Solvent), called by Belgrano (Ref 31, p 211) Polvere a solvenente misto (Propellants with Mixed Solvent). To these belong Polvere C₂ [described here as C₂ (Polvere)], also known as Cordite italiana and Solenite (qv) (Ref 26, p 206)

C. Polveri a più componenti (Propellants with Several Components). Giorgio (Ref 26) lists on p 207 among them some propellants which are listed by Belgrano on p 206 under Polveri a solvenente falso. One of them, Polvere alla Centralite, listed by Giorgio on p 207, was not found in Belgrano. Its compn is NG 24, NC 69 & Centralite I 7%

Under the title “La preparazione delle polveri colloidali”, Giorgio described on pp 209–10 the prep of single-base and double-base propulnts

Under the title “Fabricazione delle polveri a solvenente falso”, Belgrano described on pp 206–07 methods of prep of double-base propulnts with fixed solvent

Under the title “Analisi sulle polveri di lancio”, Belgrano described on pp 213–14 analytical procedures used for single-base and double-base propulnts contg as stabilizers DPhA, Centralite, etc

Polveri senza fumo. See POLVERI DA LANCIO SENZA FUMO

PROIETTO o PROIETTILE (Projectile)

Accdg to “MUNIZIONAMENTO ITALIANO” (Addnl Ref C), the terms proietto o proiettile are used referring to any heavy object which is fired from a gun and produces a destructive effect at a distance. This object may be solid (like an AP Bullet) or hollow (like HE Projectile) loaded with a bursting charge expl (carica di scoppio) which is detonated by an initiation train

Accdg to TM 9-1985-6 (1953) (Ref 16, p 61), Italian projs used during WWII were very similar to British, French and Japanese. Most field equipment was semi-fixed, with the smaller caliber and AA (Antiaircraft) guns using fixed ammunition. When the semi-fixed type of ammunition was used, the propellant charge was divided into a number of parts which were enclosed in silk bags. Igniters of BkPdr were also included. The complete charge was enclosed in the cartridge case and closed by means of a cardboard cup. In the base of the cases were fitted percussion primers of standard design

In guns using bag ammunition, an igniter which had an extension piece was fitted into the vent hole. For initiation of propellant charges, the types of tubes used were friction, percussion, and electric percussion

The Italians designated their projectiles by the caliber, the specific equipment in which they were used and the model numbers of the projectiles. In most cases the model number indicated the year in which the item was designed

Accdg to MUNIZIONAMENTO ITALIANO (Addnl Ref C), Proiettili Italiani may be divided into: Proiettili di piccoli calibri (Small caliber projectiles, inclusive to 100-mm); Proiettili di medi calibri (Medium caliber projectile, from 102 to 210-mm) and Proiettili di grosso calibri (Large caliber projectiles, above 210-mm)

Projectiles may be classified also accdg to their effect on the target (bersaglio), as follows:
a) Proiettili ad azione esplosiva. Projs filled with HE’s, which when initiated by the fuze blow up either during flight or on reaching the target (Granate mine, Granate bombe, Granate torpedini and Granate scoppianti) (Mine-, Bomb-, Torpedo- and Explosive Shells)
b) Proiettili ad azione proiettiva (Antipersonnel or Projecting Shells), which on bursting scatter splinters or pellets over more or less wide range and are designed to hit personnel or equipment. They may be called antipersonnel or fragmentation (Granate ordinarie, Granate di ghisa acciaiosa, Granate dirompenti o a tempo e Granate a frattura prestabilita) (Ordinary-, Steel-pig, Demolition-, Time- or Predetermined Time Burst Shells)
c) Proiettili speciali (Special Shells). These include incendiari (Incendiary), fumogeni (Smoke), nebbiogeni (fog), illuminanti (Illuminating) e proiettili carichi di aggressivi chimici (CW Shells)
d) Proiettili perforanti (Armor-Piercing Pro-
jectiles, abbrd AP). They contain a small expl charge encased in specially treated steel. Used against tanks, ships and fortifications
e) **Proiettili semiperforanti** (Semi-penetrating Shells) contain a larger expl chge and, in addn to penetrating effect, they have an expl effect
f) **Scatola a mitraglia** (Case Shot or Canister) consists of sheetmetal cylinders contg pellets cemented by resinous or similar binder. Used in close-range combat
g) Hollow (Shaped) Charge Projectiles (Cariche cava) are not described in AddnRef C

There were also inert projectiles used for training purposes

Italian WWII projectiles are described and illustrated under “ITALIAN PROJECTILES AND CASES” in TM 9-1985-6 (1953) (Ref 16) and in “MUNIZIONAMENTO ITALIANO” (AddnRef C), also as complete rounds. We are describing rounds for small arms under Armi portatili, munizioni, while artillery ammunition is described and illustrated under CARTOCCI GRANATA, where many Hollow Charge (Carica cava) Shells are listed

**PROJECTILE FUZES** (Spolette per proiettili).

Accdg to MUNIZIONAMENTO ITALIANO (AddnRef C), projectile fuzes may be divided by their location into *spolette anteriori* (Nose Fuzes) and *spolette posteriori* (Base Fuzes); and in terms of operation into *instantaneo* (Instantaneous), *ordinario* (Regular) and *ritardato* (Delay)

A number of fuzes used before WWII are known, but it is preferable to list here fuzes of WWII, which are described and illustrated in TM 9-1985-6 (1953) (Ref 16). The fuzes are divided into:
a) Nose Percussion Fuzes (Field Equipment), *spolette percussioni*
b) Nose Percussion Fuzes (Naval and Coastal Defense)
c) Time Fuzes (Field Equipment), *spoletta a tempo – Combustion Type* and *spoletta meccanica a tempo – Mechanical Type*
d) Time Fuze (Naval and Coastal Defense), *OT – Nose Time Combustion Type* and *OMT – Nose Mechanical Time Type*
e) Time and Percussion Fuze ADE

Following are the fuzes described in Ref 5a:
- Percussion Nose Fuze for 65/17 HE Projectile (p 134, Fig 201); Percussion Nose Fuzes M10 (Guerriore) PC and M (Guerriore) MEGC (p 135, Fig 202); Percussion Nose Fuze 1, M35 (p 136, Fig 203); Percussion Nose Fuze 1, M38 (p 137, Fig 204); Percussion Nose Fuze 1, M32/38 (p 137, Fig 205); Percussion Nose Fuze M40 with Detonator M40-Ordinary (p 138, Fig 206); Ditto – Instantaneous (p 138, Fig 207); Percussion Nose Fuze for 37/40 AP (p 139, Fig 208); Percussion Nose Fuze M39 (p 140, Fig 209); Percussion Nose Fuze OK BO SC 41 (p 141, Fig 210); Percussion Nose Fuze M16 (OK 25912) (p 141, Fig 211); Percussion Nose Fuze O BO 34/37 (p 142, Fig 212); Percussion Nose Fuze for 37/54 HE Shell (p 143, Fig 213); Nose Time Fuze M900/14 (p 144, Fig 214); Nose Time Fuze OT 32 (p 145, Fig 215); Time and Percussion Nose Fuze ADE M99 (p 146, Fig 216); Nose Time Fuze M06/17 (p 147, Fig 217); Time and Percussion Nose Fuzes ADE M06 (p 147, Fig 218); Time and Percussion Nose Fuze ADE M12 (p 148, Fig 219); Time and Percussion Nose Fuze ADE M36 (p 149, Fig 220); Time and Percussion Nose Fuze ADE M32 (p 150, Fig 221); Nose Time Fuze OT 33 (p 151, Fig 222); Mechanical Time Nose Fuze M36 (p 152, Fig 223); Base Fuze for 47/32 AP (p 153, Fig 224); Base Fuze for 100/17 Hollow Charge (Carica cava) (pp 153–54, Fig 225)

**Propellenti (Propellants).** See Esplosivi da lancio

**Propellenti per razzi (Rocket Propellants).** A brief description is given by Belgrano (Ref 31, pp 128–29). They are divided into *liquidi e solidi*. Liquids are divided into *monopropellenti* and *bipropellenti*, and monopropellenti are subdivided into three groups:

1) Those contg in its molecule both combustible and oxidizer, such as nitromethane, nitroethane, ethyl nitrate, NG, etc

2) Compounds which easily undergo exothermic decmpn, such as hydrazine, acetylene, ethylene, etc

3) Synthetic mixtures of oxidizers with combustibles

Propellants used during WWII were usually liquid bipropellants, consisting of gasoline, aromatic amines and alcohol as combustibles with oxidants, such as liquid oxygen, hydrogen
peroxide or fuming nitric acid

Among solid rocket propellants, Belgrano mentions the mixtures of Na Nitrate & Amm Picrate with thermoplastic resin of urea-aldehyde type

Belgrano also mentions liquid propellant mixtures of hydrogen with light substances such as boranes of lithium or beryllium.

It is not shown by Belgrano which of the above propellants were used by Italians

Addnt Refs from CA:
A. M. Pallotta of Bombrini-Parodi-Delfino S.p.A., Societa per Azioni, ItalP 640004 (1962) & CA 58, 11163 (1963) [Inhibition of Solid NC-NG base Rocket Propellants and/or sticking them to their container can be achieved by a mixt compatible with the ingredients of the propln, eg, 2-Nitrophenylamine — it consists of a substance having ≥1 reactive HS groups, another having ≥1 unsat monomers with 1 or 2 double bonds, and a substance or a combination of substances which promotes the co-polymerization. The adhesion is stable betw -30° & +50°. For example, a NC-NG based propln contg 2-NDPhA and plasticizers was centered in a container of steel or Al. In the empty space, a mixt of 100 parts by wt of a polysulfide, HSO(C$_2$H$_4$OCH$_2$OC$_2$H$_4$)$_{23}$(C$_3$H$_4$- OCH$_2$OC$_2$H$_4$SH), 64 ps Me methacrylate, 20 ps TiO$_2$, 1 p vinyl acetate, and 2 ps acrylonitrile was introduced. To this, 6 ps cuminhydroperoxide and 0.4 p diphenyldiurea were added. In <3 hours at RT, complete polymerization followed resulting in perfect inhibition and sticking of the grains]

B) R.M. Corelli (Univ Roma), Missili 4 (4), 13–24 (1962) & CA 60, 15672 (1964) [Development of high-energy chemical propellants which were classified into two groups: 1st group, consisting of liquid, solid, and hybrid (liquid, solid or lathegoric) proplnts, is based on oxidn-redn reactions; 2nd group, based on the metastability of the substance, is the so-called free-radical group of high-energy fuels, primarily H, O, N, NH$_3$, Me and Et. The liq proplnts include O$_2$, O$_3$, F, boranes, dimethylhydrazine, and derivs of F & B. The solid proplnts include thermoplastics (such as polyvinyl chloride), & polyethylene; the thermosetting compds (such as polyurethane, epoxy & polyester resins); and elastomers (such as polyisobutylene & Thiokol).

The free-radical group of fuels offers specific impulses much higher than conventional proplnts, but handling problems appear to be insurmountable]

C) G. Partel, Rivlng 15 (10), 969–76 (1965) & CA 64, 14015 (1966) [Nitric oxide, NO, tested in 28 expts as an oxidant in rocket proplnts appeared to be unsuitable for that purpose, because it required a long reaction time inside the rocket engine. However, it could be used as regulator, when nitric acid is the oxidant]

D) D. Perfumo (of Bombrini-Parodi-Delfino S.p.A.), ItalP 696714 (1965) & CA 66, 16784 (1966) [In the manuf of solid rocket proplnts of the extrud type, the acton of plasticizers on the primary proplnt is accelerated by the addn of small amts of N-alkylformamidns (such as HCONMe$_2$). Processing temps can be low and primary plasticizers (such as alkylphthalates) can be omitted. In a typical prep a proplnt contg NC 61.5, NG 4, Nitroglycol 3, Amm Perchlorate 30.5 & Ethylcentralitate 1%, was treated with a liq contg NG 62, NGc 34, Ethylcentralitate 1, and HCONMe$_2$ 3%]

E) Mario Giaccio (Univ of Pescara), Quad Merco1 7 (1), 63–87 (1968) & CA 71, 146 (1969) [Historical notes on solid proplnts for rockets, from Chinese craftsmen to large-scale industry are given. Modern proplnts (fuels and oxidizers) are included and future developments are discussed]

F) G.B. Guarise & G.A. Menin (Univ of Padova), AttiMemAccadPatarinaSciLettArti 1967–1968 (Pub 1969) 80 (Pt 2), 399–415 [Chem & mech characteristics of oxygen gas-JP-4 proplnt mixts were detd in an expnl rocket engine. Details of expts and results, such as specific impulse & thrust limit are given]

Randite. A Cheddite-type expl contg KClO$_3$, MnBenzene 15 & MnO$_2$ 43% (Ref 31, p 356)

Razzi (Rockets). No description is given by Belgrano except for razzo antigradino (Antithal Rocket). Its outside view is on p 626

RDX. See T$_4$

Romite. A Cheddite-type expl used for loading land mines: KClO$_3$ 39, AN 49 & naphthalene or paraffin 12% (Ref 28, p 346 & Ref 31, p 356)
Sabulite. Originally Sabulite, developed in Belgium and used as permissible (SPG) Dynamite, condt AN 54, K nitrate 22, TNT 6, Amm chloride 13 & Ca silicic 5%. The Italians modified it to AN 78, TNT 8 & Ca silicic 14% for use during WW1 as a bursting charge in some ammunition (Ref 1, p 342). Then they replaced TNT (probably due to its shortage) by TNN (Trinitronaphthalene) and developed, acdg to Piantanida (Ref 4, p 246), the following comps for use as bursting charges: a) Sabulite normale – AN 60, Na nitrate 18, TNN 8 & Ca silicic 14% and b) Sabulite extra – AN 65, TNN 10 & Ca silicic 25%

All these comps are listed in Ref 28, p 346 and, in addition, there is Sabulite 18 which condt AN 42, Na nitrate 40 & TNN 18%

Belgrano (Ref 31) stated on p 329 that Sabulite is a Belgian expl type SPG (sécurité-grisou-poussière), while on p 316 he stated that Sabulite consists of AN, Ca silicic & TNT with a Trauzl value of 2550c
(See also Piantanida, Ref 4, p 246)

Saggio (Test), plural Saggi. The following stability tests are described in Belgrano (Ref 31): Saggio Abel (pp 64–7), Saggio Angeli (p 67) and Saggio di Bergmann-Junk o Saggio Tedesco (p 199)

Schneiderite. An expl mixture of AN 87.4 & DNN (Dinitronaphthalene) which originated in France and used during WWI as a bursting charge. It was adopted in Italy and in Russia. It was also used as a demolition charge. Its mixture with TNT is known as Esplosivo MST, described here as Nougat (Ref 4, p 241; Ref 28, p 346 & Ref 31, p 316)

Sebomite. A Cheddite-type expl: KCIO3 90, MNT 5 & tallow (sego) 5% (Ref 1, p 194)

Securite. A Dynamite which used to be manufd at Dinamitificio di Segni: NG 52 & 30 and NC 48 & 70% (Ref 1, p 298). Belgrano (Ref 31, p 317) lists another Securite consisting of AN & m-DNBz (meta-Dinitrobenzene)

Sedulite. A Cheddite-type expl: KCIO3 85, MnO2 5 & vaseline or paraffin 10% (Ref 1, p 198)

Seismic Explosives. See under Tutamite

Selenite Italiana. NG 36.0, Guncotton 30.5, Collodion Cotton 30.5 & mineral oil 3.0% (Ref 4, p 180)

Sevrante A. An expl mixt of French origin consisting of Amm Perchlorate 31, PETN 48, DNT 18 & Al 3%; density 1.55 & deton vel 5000 m/sec (Ref 31, p 363)

Sevrante B. Amm Perchlorate 42, PETN 42 & DNT 16%; density 1.50 & deton vel 4500 m/sec (Ref 31, p 363)

Siperite o MNDT (Esplosivo). Acdg to Molina (Ref 1, p 342), Siperite was an expl compn consisting of AN 82–87 & DNN 18–13 parts, to which 20pe of TNN was added. Piantanida (Ref 4, p 242) gives: AN 72.8, TNT 16.7 & DNN 10.5%. Belgrano (Ref 31, p 297) gives its compn as AN 73, DNN 10 & TNT 17%. Acdg to Ref 28, p 347, it was used as a bursting charge for some projectiles

Small-Arms Ammunition. See ARMI PORTATA TILI MUNIZIONI

Solenite italiana o ordinaria. Acdg to Piantanida (Ref 4, p 180), it consisted of NG 36.0, fulmicotone 30.5, cotone collodio 30.5 & olio minerale 3%. Same compn is given by Belgrano (Ref 31, p 212) except that he lists pirocollodio instead of cotone collodio. It was used as a rifle proplnt for Fucile Mod 91 & Mannlicher (See also in this Vol of Encycl, p H152-R)

Solfitre. Mixture of AN 83–88 & sulfur 17–12% proposed in 1930 by Dr Pannoncini. It is listed in Ref 28, p 347 and in Ref 31, p 317. Its use is not indicated

Solfoclianato di piombo (Lead Thiocyanate). Pb(CNS)2 is described by Belgrano (Ref 31, pp 467–68). It was used in many initiating mixtures such as KCIO3 50–55, Pb(CNS)2 ca 25, Sb2S3 15–18 & Pb(N3)2 ca 5%

Solfituro di antimonio (Antimony Sulfide), Sb2S3, has been used in many initiating mixtures contg
MF (Mercuric Fulminate) with KClO₃, such as listed in two tables on p 423 of Belgrano (Ref 31).

**Spoletta** (plural Spoletto). Fuze (Fuzes). See under Bomb Fuzes and under PROJECTILE FUZES

**Stiffnato di piombo** (Lead Stypnate, abbrd as LSt) or Trinitrotoreocrinato di piombo), C₆H(NO₂)₃O₂Pb is described in Belgrano (Ref 31, pp 456–64). It has been manufd by the Società Nobel and used as an igniting charge for LA Lead Azide) in detonators and as an ingredient in priming compns (Ref 28, p 350).

**T₄**, **Esogene o Ciclotrimetilentrinitoamina** (RDX, Cyclotrimethyltrinitramine, Hexogene or Cyclonite). Its prepn, props, and analytical procedures are described by Belgrano (Ref 31, pp 255–59).

Its props listed in Ref 31, p 259 are Density (max) 1.69, Explosion Temperature 250°C, Flame Temperature at Explosion (Temperature of Explosion) 3380°C, Heat of Explosion 1390kcal/kg, Volume of gas at 0°C & 760mm 890 l/kg, Impact Sensitivity with 2kg Weight 30cm, Detonation Velocity 8400m/sec, and Trauzl Test Value 470cc. Giuta (Ref 19) gave some slightly different values, such as Density (max) 1.70, Explosion Temperature 290°C, Volume of Gas at 0°C & 760mm 908 l/kg, Impact Sensitivity with 2kg Wt 42cm, Detonation Velocity 8380 m/sec and Trauzl Test 520cc (listed also on p 258 of Ref 31).

T₄ by itself is too sensitive to be used as a bursting charge in projectiles, but it is suitable for use in detonators & in blasting caps. When so used, T₄ is dissolved in cold acetone and precipitated by adding cold water to the soln. After filtering, the T₄ is dried in an aluminum-jacketed rotating drum thru which air preheated to 120°C is driven. Then the dry particles are coated with 1% graphite to facilitate their press-loading in detonators.

In a slightly desensitized form, as for example 95/5—T₄/wax or paraffin, it is used as a booster charge in ammunition (Ref 28, p 348). Belgrano (Ref 31, p 261) calls such expl T₄ flemmatizzato and says that it is colored pink (rosa) in Italy. Other flemmatizzati listed by Belgrano are NGU, DNT, mineral oil, castor oil, acetanilide, MNN and synthetic resins.

Mixtures of RDX with TNT, called Tritoliti o Hexoliti (qv), and with TNT & Al, called Tri-tolital (qv), are used as bursting charges. Other mixts contg RDX are Hexocire, Hexocire-Aluminum, Hexogene-Nylon and Idrolitii are listed here.

A mixture consisting of AN 73.4, RDX 22 & wax 4.6% was used, accdg to Ref 28, p 348, during WWII, as a bursting chge for some armor-piercing long-nose shells. A similar mixture called NTP (qv) is listed in Ref 31, p 266, as one of the Miscela betliche polverulente a base di T₄.

Addnl Refs from CA:

A) Firmo Rag. G. Mangiarotti, ItalP 528927 (1955) & CA 53, 3598 (1959) [Purification of a mixture of T₄ (RDX) and Tritol (TNT) can be achieved by dissolving it at 45°C in acetone, cooling to 25°C and, after decantation, filtration, centrifugation and treatment with C or decolorizing earth, crystallizing in a vertical crystallizer. The crystals of T₄ which separate are continuously filtered out and TNT is separated from the mother liquor by pouring it into water or by evaporating the solvent.]

B) G. Dini & G. Manfredi, ItalP 558782 (1957) & CA 53, 22956 (1959) [Details of conventional method of prepn of T₄ (Cyclonite or RDX) from hexamethylenetetramine (treatment with excess nitric acid), are claimed which make it possible to recover this excess directly in high concn.]

**T₄ plastico** (Plastic RDX). One such is RDX 89 & vaseline 11%. By adding 10–20% of Al powder, the T₄ plastico con allumino is obtNd (Ref 31, p 262).

In Ref 28, p 348, is listed a mixture contg RDX 79.5, DEGDN (with 0.3–0.4% of Colloidion Cotton) 17.5 & vaseline 3%. This mixt is known as “Italian” and supersedes the so-called “American” mixture consisting of RDX 89 & petrolatum 11%, which is listed under Esplosivo plastico. The Italian mixt was used in land mines and in antitank bombs.

Other RDX plastic expls are Miscela C, Miscela C₃ and Miscela C₄. They are listed here and in Ref 31, p 263.

**T-Ammonial or Toluolummonal.** See under Ammonial.
Termite (Thermite). See under Composizioni (o Miscele) Incendiarie and in Belgrano (Ref 31, p 640) and Giua (Ref 19, p 414)

Tetralite. See TETRILE

Tetranitroanilina (Tetranitroaniline or TeNA). It is described in Belgrano (Ref 31, pp 271–72). Used in cryst form in some detonators and in mixts with TNT, TNN & DNBz

Tetranitropentaeritrite. See PENTRITE

Tetrazene o Guanilnitrosoammina (Tetrazene or Guanynitrosoammine) is described by Belgrano (Ref 31, pp 464–67). Used in Composizioni (o Miscele) innescanti, items e, f, g and l

TETRILE, Tetranitromedianilina, Trinitrofenilmetilnitrosoammine o Tetralite (Tetryl, Amer and CE, British). It is described in Belgrano (Ref 31, pp 272–75). He lists its uses in Gt-Britain and the USA but says nothing about its uses in Italy. In Ref 28, p 349 it is stated that it is used as a component of some expl mixtures, as with TNT, such as Tetritol

Tetritol (Tetrytol). Accdg to Belgrano (Ref 31, p 246), the mixture used in Italy consisted of Tetryl 75 & TNT 25% and was colored yellow. It was used for cast-loading in demolition charges

Tipo I and II Dinamite. See under DINAMITE

Titan. One of Ital mining expls. It is listed without giving its compn by Belgrano (Ref 31, p 317)

TNT. See TRITOLO

Tolite. See Tritolo

Tolulammonal. See under Ammonal

Torpex. Accdg to Belgrano (Ref 31, p 265), it is a mixt consisting of RDX 44, TNT 38 & Al powder 18%. Trauzl Test value is 490ec. It was used for loading warheads of torpedoes, depth bombs & sea mines

Tovex (Esplosivi “slurry”). Expl mixts contg AN, Na nitrate, TNT, with or w/o Al and ca 15% of water were developed in the USA after WWII. Four formulations of Tovex are listed by Belgrano (Ref 31, p 322), who states on p 320 that slurry expls are not yet manufd, nor used in Italy

Tracianti (Tracers). See under PIROTECNIA

Trialen 105. An expl mixture similar to Torpex and Tritolital. It contd TNT 50, RDX 25 & Al 25%. Used in torpedoes and other underwater ordnance (Ref 28, p 349)

Trilite. See TRITOLO

Trinitroanilina o Picramide (TNA) is an explosive equal in power to PA (Picric Acid). Its props are described in Belgrano (Ref 31, pp 270–71) but uses are not indicated

Trinitroanisolo (TNAns) is described in Belgrano (Ref 31, pp 300–02). It was not produced in Italy industrially but studies were made in regard to its use for mining purposes. One of such mixts consisted of AN 80 & TNAns 20%. Incorporation of TNT in this mixt increased its brisance. It was used during WWII for military purposes by Japan, Germany & France (Ref 28, p 350)

Trinitrocresolo (TNCrs). Its isomer, Trinitrometacresolo (Trinitro-m-nethylphenol) is described in Belgrano (Ref 31, pp 291–92). Was used in Italy in mixture: TNCrs 60 & PA 40%, under the name Gesilite (qv), for military purposes

Trinitrofenilmetilnitrosoammina. See Tetrole

Trinitrofenolo. See ACIDO PICRICO

Trinitrofluorogluconato di piombo (Trinitrofluorogluconate of Lead or Trinitro-1,3,5-trihydroxynate of Lead) is described in Belgrano (Ref 31, pp 475–76). Used in inititating comps and is considered by Belgrano to be better than LSt (Lead Stypnate)

Trinitroglicerina. See NITROGLYCERINA

Trinitrometacresolo. See under Trinitrocresolo
Trinitrometila. See Xilite

Trinitrotetrolina (TNN) is described in Belgrano (Ref 31, pp 298–99). It was used during WWI in mixture with TNT called Trinitrol (qv); also in some “esplosivi di sicurezza” (Safety Explosives) such as TNN 27, AN 27 & Na nitrate 58% (See also Sabulti)

The higher deriv, Tetranitrotetrolina (TeNN) has not been used on account of difficulty of manuf (Ref 31, p 299)

Trinitroresorcina. See Acido stfinico

Trinitroresorcinito di piombo. See Stfinato di piombo (Lead Stynphate, abbrd LSi)

Trinitrotolueno. See Tritolo

Triperossido di esamatiletetramino o Esamatilenetriperossidiamma. Italian names for Hexamethyleneperoxidediamine (HMTD) which is described in this Vol of Encycl, p H83-L Belgrano (Ref 31) describes it on pp 478–79 without indicating its uses in Italy

Tritolito. Accdg to Ref 28, p 351 & Ref 31, p 263, it is a castable explosive used during WWII by Italians and Germans for loading underwater weapons, such as torpedoes, depth charges & sea mines. It consisted of TNT 60, RDX 20 & Al 20%. Its Trauzl test value is 400cc, deton vel 7400m/sec and brisance by Ital steel plate test (prova della piastra di acciaio) 21mm

Tritolito o Ciclotol. Accdg to Ref 28, p 351, it is a castable mixt of TNT 50 & RDX 50% proposed in 1930 by M. Tonegutt for loading underwater weapons, such as torpedoes, sea mines & depth charges. Its props are accdg to Ref 31, p 265: cast density 1.65, Trauzl value 460cc, deton vel 6900m/sec and brisance by Ital steel plate test 23mm. The comp was changed during WWII to TNT 60 & RDX 40%. Its props are cast density 1.62, Trauzl value 440cc, deton vel 6600m/sec and brisance 21mm, but the 50/50 formulation, being stronger, was used for loading aerial bombs and antitank shells (Ref 31, p 265)

The formulation RDX 60, TNT 39 & wax 1% is similar to Amer Composition B, which contd 1% paraffin (instead of wax). Its props are Trauzl test value 470cc and deton vel 6950m/sec. Was used in aerial bombs, armor-piercing shells, torpedoes and mines (Ref 31, p265)

Accdg to Ref 28, p 351, the following mixts were used during WWII for loading shaped charge shells and mines:

a) 50/50–RDX/TNT; b) 60/38/2–RDX/TNT/Wax (dyed red); c) 25/50/25–RDX/TNT/Al (Trialen 105) and d) 20/60/20–RDX/TNT/Al

The simplest method to prepare blends of RDX & TNT is to dissolve them in acetone and precipitate the powder by adding cold water

TRITOLIN (Trotite, Trilite, Tolite, Trinitrotoholo) (TNT or Trinitrotoluene). It is described in Belgrano (Ref 31, pp 233–47). TNT is manufd in Italy in two grades: 1) A.P. (alto punto) with solidification (setting) point above 80° and 2) B.P. (basso punto) solidifying between 77 & 78°

The following values for props of TNT are listed in Ref 31, p 241: Density (max) 1.58, Explosion Temperature 240°, Flame Temperature at Explosion 2800°, Heat of Explosion 980kcal/kg, Detonation Velocity at d 1.5 6700m/sec, Volume of Gas at 0° & 760mm 680liters/kg, Specific Pressure 8100atm/kg, Trauzl Test Value 285cc and Impact Sensitivity with 2kg Weight 90cm. Giua (Ref 19) gave some slightly different values such as density (max) 1.59, Explosion Temperature 295°, Volume of Gas 690liters/kg, Trauzl Test 310cc & Impact Sensitivity at 2kg Wt 60cm (listed also on p 258 of Ref 31)

Accdg to Ref 31, p 236, the grade of TNT solidifying above 80° was used straight in detonators and in some projectiles, while the lower grade (77–78°) was used in mixtures such as Amatol.

Accdg to Ref 28, p 352, straight TNT has been used by the Italians to a much lesser extent than by the Americans or British. Pressed TNT is used as a booster in some bombs and as a main filler for some demolition charges. Granular TNT is used as the bursting chge of hand grenades, mortar shells and in mines. Cast TNT is used as a bursting charge in some bombs, 2 to 1000kg in weight, and for artillery shells, 57 to 420mm

TNT has also been used extensively in compo-
site expls, such as Amatol, Ammonal, Dinamite, Esplosivi da guerra, Esplosivi da mina, Esplosivo S20, Gomma inorganica, MABT, MAT, MST o Nougat, Pentritolo, Pentrol, Picratol, Siperite or MNDT, Tetritol, Trialen 105, Tritolital, Tritonite, Tritonal and Tritriminal

Analytical procedures and tests used in Italy are described in Ref 31, pp 243–45

**Tritonal.** Composite explosive consisting of TNT 80 & Al powder 20%; Trauzl Test Value 350cc; used for cast-loading light case underwater ammunition (Ref 31, p 246)

**Tritriminal of WWI.** Composite expl of 1 part TNN (Trinitronaphthalene) and 2 parts TNT; used during WWI for cast-loading some ammunition (Ref 31, p 299)

**Trotol.** See Tritolo

**Tutamite.** One of the Italian “Esplosivi per prospezioni geosismiche” (Explosives for Geo-seismic Prospecting) is listed by Belgrano (Ref 31, p 334) together with other Italian seismic expls *Sismite* and *Geo-Mon A*, without giving their compositions

Tutamite is also listed in Catalog of Montecatinni SGIMC, Milano (1959), p 30. Its props are given but no compn.*

*Geo-Mon A* is listed in the same catalog, p 19 with props but no compn. Other Esplosivi per prospezioni geosismiche listed in Montecatini catalog are *Geo-Din A* and *Geo-Din B* (p 17)

In catalog of Vulcanica SPA, Brescia, are listed three seismic expls: 1) *Vulcanica DB*, based on TNT & nitric salts; Trauzl value 320cc, Deton Vel 2600m/sec at d 1.1 and Impact Test with 2kg Wt > 90cm; 2) *Vulcanica DBS*, based on TNT, RDX & AN; Trauzl value 370cc, Deton Vel 5600m/sec at d 1.25 and Impact with 2kg Wt > 90cm; 3) *Sismite A*, based on TNT, Tetritol and nitrates; Trauzl value 400cc, Deton Vel 5000m/sec at d 1.4 & Impact Test 70cm

[Italian catalogs were obtd thru the courtesy of Dr Omero Vettori of Auola (Massa Carrara)]

**Umbrite.** Two formulations are listed by Belgrano (Ref 31, p 254): Umbrite A = NGu 48.4, AN 37.3 & ferrosilicon 14.3%; Trauzl value 260cc and Umbrite B = NGu 45.1, AN 41.4 & ferrosilicon 13.5%; Trauzl value 340cc. Accdg to Ref 28, p 352 they were used as bursting chges in some projectiles

**Universal Italiana.** Acdg to Belgrano (Ref 31, p 580), it is one of the *polveri da caccia* [Hunting (Sporting) Propellants], which was based on completely gelatinized NC. Molina (Ref 1, p 412) gives its compn as NC 80–82, DNT 12–10, DEDPhU (Diethylidiphenylurea) 4–4, vaseline 3–3 & volatiles 1–1%

**UNKNOWN NAME EXPLOSIVES.**

The following expls are extracted from Chemical Abstracts, A.M. Tonegutti, FrP 833720 (1938) & CA 33, 3590 (1939):

a) A castable bursting type expl – AN 70, PETN 20 & cyanoguanidine 10%
b) A press-loaded expl for use as a bursting chge for 47mm AP Shells – AN 73.4, RDX 22 & wax 4.6%
c) A press-loaded expl for use as a booster in ammunition – RDX 95 & wax (dyed red) 5%
d) A castable expl for use as a bursting chge in 500-kg bomb – PETN 65 & PETA (Penta-erythritol Tetraacetate) 35%. It is comparable to TNT in sensitivity to shock, but is about 25% less brisant

**VE₄.** One of the mining expls manufd by Vulcanica SPA, Brescia, which is based on aromatic nitroderivatives and AN. Its Trauzl test value is 335cc & Deton Vel 3550m/sec at d 1.02. Suitable for use in quarries with rock not too hard (Vulcanica Catalog of 1960, p not indicated)

Belgrano (Ref 31) lists VE₄ on p 317 without giving its composition

**Vibrite.** Blasting expl contg AN 78, TNNaphthalene 8 & Ca silicide 14% (Ref 1, p 342 & Ref 28, p 353) (Compare with Sabulite)

**Victorite.** Cheddite-type expl consisting of KClO₃ 40.0, PA 53.2, K (Na or Ba) nitrate 4.8 & carbon 2% (Ref 28, p 353)

**Virite.** Black Powder-type expl consisting of K nitrate 33–38, AN 35–40, sulfur 4–5, carbon 10–12 & Amm oxalate 9–12% (Ref 28, p 353 & Ref 31, p 340)
Vulcania DB and DBS. Seismic prospecting expls listed here under Tutanite

Vulcanite P. A mining expl suitable for blasting tunnels thru hard rock. It is based on inorg nitrates, nitroaromatic comps and nitric diethers. Its Trauzl test value is 515cc, Deton Vel 5100m/sec at d 1.15 & Impact with 2kg Wt 80cm [Catalog of Vulcania SPA, Brescia (1960)]

Vulcanite PR. A mining expl suitable for tunnel work thru hard rock. It is based on inorg nitrates & nitroaromatic comps. Its Trauzl value is 510cc, Deton Vel 5600m/sec at d 1.15 & Impact 70cm [Catalog of Vulcania SPA, Brescia (1960)]

Vulcan Powder. It is listed in Belgrano (Ref 31, p 339) as cocont NG 30.0, Na nitrate 52.5, carbon 10.5 & S 7.0% without indicating which country used it

Xilito o Trinitrometilxirole (TNX or Trinitromethylbenzene). It is very briefly described in Belgrano (Ref 31, pp 248–49). It was used during WWI in mixtures with AN, such as in NX, contd AN 77 & xilito 23%. Another mixture was MTX which contd PA 55, TNT 35 & xilito 10%

References for Italian Explosives, Ammunition and Weapons:
1) R. Molina, "Esplodenti e Modo di Fabricarli, Hoepli, Milano (1930)
2) M. Giua, "Lezioni di Esplosivi", 2 Vols, Ratterio, Torino (1932–33)
3) General Carlo Montu, "Storia dell’ Artiglieria Italiana", Vol 1 (1935) [Reviewed in MAF 14, 557 (1935)]
4) E. Pantanilda, "Chimica degli Esplosivi e dei Gas di Guerra", Regia, Accademia Navale, Livorno (1940)
4a) Lt Col G.B. Jarrett, OrdnanceSergeant, Aug 1943, p 18
5) Anon, "Data on Foreign Explosives", US Chemical Warfare Service Field Lab Memo 4-6-2 (1944), US Office of Technical Services PB Report 11544
6) J.D. Parsons, "Visit to Italian Explosives Factories", PB Rept 12663 (1945)
9) A. Mangini, "Quaderni di Chimica Industriale 14; Esplosivi", Patron, Bologna (1947), pp 147–235
12) A. Izzo, "Protezione e Fuochi Artificiali", Hoepli, Milano (1950)
13) C. Belgrano, "Gli Esplosivi", Hoepli, Milano (1952)
14) Ministero, de la Difesa Marina, Editor, "Guida per l’Esecuzione delle Prove Pratiche o della Analisi sugli Esplosivi", Roma (1952)
15) A. Izzo, "Manuale del Minatore Esplosivistico" (Fochino), Hoepli, Milano (1953)
18) B.T. Fedoroff et al, "Italian Explosives and Ammunition", Pictatiny Arsenal Manuscript, Unpublished (1955)
22) R.P. Antonelli, "Encyclopedia of Explosives", OTIA, Ordnance Liaison Group, Dunham, North Carolina (1960), pp 150–51 (Italian Terms)
26) C. Giorgio, "Tecnica degli Esplosivi", Vol 1, Del Bianco, Udine (1964)
28) O.E.

Additional References:
A) US War Department “Military Dictionary”, TM 30-2569, English-Italian and Italian-English, Washington, DC (1943)
B) “Military Italian-Russian Dictionary”, GosizdatNostri i NazioniSlovarey, Moscow (1953)
C) “MUNIZIONAMENTO ITALIANO” (1941) sent by Dr Omero Vettore, Aulla (MC) in May, 1973
D) Picatinny Arsenal Technical Reports (PATR) on examination of Italian Ammunition picked up at the front or captured:
   a) C.G. Scheibner, PATR 903 (1938) (Cal .50 Expl Shell)
   b) A.B. Schilling, 1241 (1943) [47-mm AP, HE (LN) CRA]
   c) Ibid, 1256 (1943) [47-mm AP, HE (MB) CRA]
   d) R.M. Dennis, 1315 (1943) [47-mm HE (Tritolo) CRA]
   e) A.B. Schilling, 1327 (1943) (Stick-Type Land Mine)
   f) R.M. Dennis, 1332 (1944) (37-mm LE Shell CRA)
   g) Ibid, 1344 (1944) (81-mm HE Mortar Shell CRA)
   h) A.B. Schilling, 1348 (1944) (Variable Pressure Land Mine)
   i) Ibid, 1349 (1944) (20-mm AP, HE Shell Complete Round)
   j) Ibid, 1362 (1944) (81-mm HC, HE Mortar Shell CRA)
   k) Ibid, 1381 (1944) (Pressure Igniter and Detonation Block)
   l) F.G. Haverlack, 1515 (1945) (120/45-mm HE Shell)
m) A.B. Schilling, 1539 (1945) (Offensive Hand Grenade)
   n) F.G. Haverlack, 1549 (1945) (149/13-mm HE Shell)
o) A.B. Schilling, PAMR 105 (1956) (Hand and Smoke Grenades with Friction Type Igniter)
E) J.D. Parsons, “Visit to Italian Explosives Factories” (1945) (US Office of Technical Services, PB Rept 12663)
F) Anon, “Allied and Enemy Explosives”, Aberdeen Proving Ground, Maryland (1946)

Abbreviations: CRA — Complete Round of Ammunition; AP, HE, LN — Armor-Piercing, High Explosive, Long Nose; MB — Monoblock; HC — High Capacity; LE — Low Explosive

Italian Catalogs, furnished thru the courtesy of Dr Omero Vettore of Aulla (Massa Carrara):
I) Catalog of “Esplosivi ed Accessori da Mina”, Montecatini SGIMC, Milano (1959). Described are the properties and uses of mining expls designated as Gomma A, Gomma Agel, Gomma B.M., G.E.O.M., G.D. 1°M.T., G.D. 1°M.B., G.D.S., Gelignite S.A., Semigel A, Dinamone 1°, Dinamon S, Cava M, Grisoutina 13.20%, Grisoutina 10%, Tionite, Geo-Din A, Geo-Din B, Geo-Mon A and Tutamite. No compositions are given. Also are described initiating devices, nonelectric and electric
II) Catalog of “Esplosivi Mangiarotti”, Codroipo (Udine) (1960). Described are the properties, uses and testing of mining explosives designated as BM.1, BM.2, BM.3, BM.5, BM.57, Super BM, BM.a2, Super BM.Cava, BM.2 per Galleria. Compns are given without numerical values
III) Catalog of “Vulcania SPA. Polveri per Mina, Caccia, Agricultura, Difesa”, Brescia (1960). Described are the properties and uses of mining expls: Antonites, Alger C, Alger D, Alger E, VE4, Vulcania D.B., Vulcania D.B.S., Vulcanite P, Vulcanite PR and Sismite A. Compns are given without numerical values
IV) Catalog of “Sorlini Esplosivi SPA, Brescia” (1961). Described are properties and uses of mining expls: Martia Alpha, Martia Beta, Martia Gamma, Martia Delta 45, Martia Eta S. No compositions are given
V) Catalog of SMI (Società Metallurgica Italiana) SpA “Catalogo Munizioni Militari”. At their plant near Campo Tizzoro (Province of Pistoia) they manufacture Small Arms Cartridges, while at Forina di Barga (Prov of Lucca) Artillery Cartridges are manufactured. The Co also manufactures steel cartridges for Recoilless Rifles (Bossoli Acciaio per Cannoni Senza Rinculo), Percussion & Electric Primers (Cannelli a Percussione ed Elettrici) and other metallic items.

VI) Professor L. Musciarelli, “Storia delle Armi” (History of Weapons) which includes History of Beretta Factory and Catalog of Pistols, Rifles & Shotguns. Published by Fabbrica d’Armi Pietro Beretta, Milano.

Ivory Nut (Corajo, Coroso or Vegetable Ivory). A hard white substance obtained from the tagus nut, such as Phytelephas Macrocarpa or negrito palm, which grows in Ecuador.

It is composed of cellulosic bodies and other hydrocarbons.

In powdered form, it is used as a “fuel” component of dynamites and other blasting explosives. According to Marshall (Ref 1), such material should not contain more than 5% of moisture and 3% ash; its solubility in 95% alcohol should not exceed 3%.

Ivory nut meal used in US dynamites is usually either fine (No 1), which has about 5% remaining on No 20 sieve and 30% passing No 100 sieve, or coarse (No 2) with 5% remaining on No 10 and 5% passing No 60. The percentage of ash in both cases is below 5% and moisture below 10%. Absorption value: 30-35%

for the fine and 25% for the coarse. Densities are 0.80 to 0.90 for No 1 and 0.5 to 0.7 for No 2. Oxygen balance to CO₂ is about -115% for the dried material.


Ivory Nut, Nitrated. No specs or procedures for nitration were found. Its use in the following Dynamite compn is claimed by Stine (Ref 1): NG 35, NC 1, Nitrated ivory nut 5%, NaNO₃ 52 and Woodmead 7 parts. The nitrated ivory nut is insol in NC and its use as a substitute for an equal amt of NC produces an explosive which is more easily detonated.

Nitrated Ivory Nut is also claimed (Ref 2) as an explosive by itself or a constituent of primary or secondary charges containing other explosives, such as MF, LA, TNT etc.

Refs: 1) C.M. Stine, USP 114330 (1915) & CA 9, 2313 (1915) 2) E. Marks, BritP 124303 (1919) & CA 14, 2713 (1920)

Izod Test. See Charpy and Izod Tests in Vol 2, p C154-L

J (Poudre de chasse). A French smokeless sporting propellant invented by Bruneau and manufactured before WWI. It contained NC 83 & ammonium bichromate 17%. It replaced poudre pyroxylin type S
Refs: 1) Daniel (1902), p 385  2) Pascal (1930), p 228

Jablonsky’s Propellant. Two samples of propellant powder were submitted by Dr Frans Jablonsky for tests as to suitability for use in small arms ammo. The powder was stated to contain NC recovered from scrap movie picture film & other chemicals intended to increase the oxygen content. Chem analysis showed the samples to have the following comp:

<table>
<thead>
<tr>
<th>Composition, %</th>
<th>Sample No 1</th>
<th>Sample No 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitrocellulose</td>
<td>76.26</td>
<td>74.08</td>
</tr>
<tr>
<td>Ammonium Picrate</td>
<td>13.82</td>
<td>10.89</td>
</tr>
<tr>
<td>Potassium Chlorate</td>
<td>5.40</td>
<td>7.75</td>
</tr>
<tr>
<td>Camphor</td>
<td>3.46</td>
<td>3.54</td>
</tr>
<tr>
<td>Ethyl Acetate</td>
<td>0.86</td>
<td>3.74</td>
</tr>
<tr>
<td>Red org material</td>
<td>0.20</td>
<td>—</td>
</tr>
</tbody>
</table>

Tests showed the propellants to be of relatively low stability, unduly volatile & hygroscopic. Ballistic tests gave results indicating the propellants to have less ballistic potential than Pyrocotton, and to deposit unburned residue in the rifle barrel. The propellants, represented by the two samples, were considered to be of little promise for use in military ammo

Jacket. A cylinder of steel covering & strengthening the breech end of a gun or howitzer tube. It also refers to the water jacket on some machine guns
Refs: OrdTechTerm (1962), p 168-L

Jacket, Bullet. A metal shell surrounding a metal core, the combination comprising a bullet for small arms. The jacket is either composed of, or coated with, a relatively soft metal, such as gilding metal, which engages the rifling in the bore, causing rotation of the bullet
Refs: OrdTechTerm (1962), p 168-L

Jacket, Coolant. The metal enclosure that consists of an inner wall (liner) and an outer metal wall (outer case), with a coolant passage between them, occasionally held apart by spacers or coolant helices. Term also applied to the double metal walls of a regeneratively-cooled thrust-chamber assembly

The main component of a modern liquid-propellant rocket thrust-chamber assembly is called the coolant jacket. There are many jacket designs but, in nearly all cases, the jacket structure extends thru the throat region as well as the combustion chamber

Jacketed Projectile. In order to take advantage of the increased striking energy made possible by the use of a smaller caliber, higher vel, perforating missile, designers have made use of the principle of the subprojectile as shown in the Fig. Here the jacket around the smaller tungsten carbide core, or subprojectile enables the missile to derive full benefit of the propellant powder gases. Further, since the jacket or carrier is made of Al, the lighter over-all wt of the complete projectile coupled with the use of high potential propellants results in a higher muzzle vel. Unfortunately, the decrease in over-all wt, while helping to give greater muzzle vel, also results in a lower ballistic coefficient and, therefore, a higher rate of loss of the muzzle vel. Since the jacket is rigidly attached to the core and does not fall off until contact with the target, the HVAP (hypervelocity armor-piercing)

IDEALIZED PROJECTILE TYPES

![Diagram of projectile types](attachment:projectile_types.png)
is also known as the composite rigid or compo-
rigid type.
Ref: ArmamentEngg (1954), pp 208–09

Jacque Stability Test. Jacqué, director of the
Spanish plant Cantabrica at Galdécano, near
Bilbao, proposed a test for NC which was simple,
rapid, and did not require complicated apparatus.
It is not advisable to use this test when it is
known in advance that the NC is of low stability
Procedure. A NC sample of 2-3g, previously
dried in a loosely covered weighing dish or
crystallizer to constant wt, is transferred to an
oven maintained at 130–40° and left there for
2 hrs. It is cooled in a vacuum desiccator and
reweighed. The sample is again heated at 140°
and weighed at intervals of ½–2 hrs until it
decomposes, as indicated by an abnormal loss of
wt. After termination of the test, it is ad-
visable to wash the NC with distilled water,
add a few drops of 0.1N KMnO₄ soln and de-
termine the amt of HNO₃, either by the nitron
or diphenylamine method
A somewhat analogous test is described by
Sy [(JACS 25, 549 (1903)] and by Berkhout
[SS 17, 33 (1922)]. See also Heat Tests in
Vol 1 of Encycl, p XV
Refs: 1) M. Jacqué, SS 1, 395 (1906) & CA
2, 1346 (1908) 2) Reilly (1938), p 85

Janinite Powders. Austrian mining expats patented
by Jahn many years ago. They contained NaNO₃
70–75, sulfur 12–10, lignite 17–10, KClO₃
0.4–2, Na Picrate 0–3, PA 0.3–0 & NaCO₃
0.3–0%
Refs: 1) Cundill (1889) in MP 6, 19 (1893)
2) Daniel (1902), 387–88

Jalnias Powders. The following compns for
Cannon Powder – KNO₃ 75, Na Picrate 3,
sulfur 10, coal 10 & KClO₃ 2% and
Rifle Powder – KNO₃ 15, Na Picrate 8, sulfur
10, coal 15 & KClO₃ 2% are found in Ref
Ref: Pérez Ara (1945), p 223

JANAF. Abbr for Joint Army-Navy-Air Force
(often pronounced as one word)
Ref: OrdTechTerm (1962), p 168-L

Jonite. A large grain, black blasting powder
impregnated with NG. It was used to a limited
extent for excavation work on the Isthmus of
Corinth in Greece
Ref: Daniel (1902), p 386

JANNAF. Abbr for Joint Army-Navy-NASA-
Air Force (often pronounced as one word)
established in 1969. Its activities are described in
Ref
Ref: Expls & Pyrots 6 (2), 1973

Jannopoulos Test. A stability test for expls which
is a modification of Abel's Test (See Vol 1
of Encycl, p A2-L) and of Guttmann's Test
(See Vol 6 of Encycl, p G196-L). The finely
ground sample is previously warmed to 35–38°
for four days before testing. The test is not
applicable to double-base propints contg NG
Ref: Reilly (1938), p 78

Janovsky Test. Color reactions produced by
 treating aromatic nitro compds with hot &
cold alcoholic NaOH or KOH and acetic
NaOH solns. A table of color reactions of some
nitro compds is given in Ref 2. See
Janovsky's Reagents under COLOR REAC-
TIONS and COLOR REAGENTS in Vol 3 of
Encycl, p C405-R
Refs: 1) J.V. Janovsky, Ber 24, 971 (1891)
2) G.D. Parks & A.C. Farthing, JChemSoc
1948, 1278

JAPANESE EXPLOSIVES, AMMUNITION
AND WEAPONS

Accdg to Ref 5, p 360, the Japanese used during
WWII a greater variety of HE's (High Expls) and
ammunition than other nations at the time (except
probably the Germans and Italians). This was due
to a shortage of raw materials necessary for the
manuf of the best expls. Although they used many
standard HE's (such as TNT, Picric Acid, PETN,
RDX and Tetryl), every other available expl sub-
stance was also used as an ingredient of expl compns.
Among these substances may be mentioned the fol-
lowing: TNAnS (Trinitroanisole), TNPhT (Trinitro-
phenotolene), HNDPhA (Hexanitrodiphenylamine),
GuN (Guanidine Nitrate), DNN (Dinitronaphtha-
lene), DNBz (Dinitrobenzene), AP (Ammonium Picrate), as well as a number of commercial exps, particularly Dynamites. As nonexplosive ingredients, the materials used included: inorganic nitrates, chlorates & perchlorates; aluminum, silicon carbide, ferrosilicon, petroleum, woodpulp, powdered coal & charcoal, tar, waxes, oils and their mixtures.

The Japanese Initiating Agents were similar to those used by the Germans, except that Tetracene was not used and fewer mixtures were employed in ammunition. MF (Mercuric Fulminate) was used in many primary mixtures, while LA (Lead Azide) was used in fewer mixtures. It does not appear that LST (Lead Styphnate) was used in compns contg nonexp ingredients. As a base ingredient of detonators, PETN was mainly used and to a lesser extent, Tetryn.

As to Japanese ammunition itself, one can point out the great variety of improvised devices, such as Mines & Depth Charges made from wooden boxes, burlap bags with rubber lining, & oil drums; Grenades made of pottery, glass & gas pipes; Booby Traps made of tin cans; and Bangalore Torpedoes made of bamboo tubes.

According to Refs 2 & 3, the Japanese Explosive Ordnance used during WWII was subdivided into those used for the Army (Rikugun) and those for the Navy (Kaijgun).

**System of Designation**

1. Type Number. Item of Ordnance, as well as most other items of military equipment, were given a type number indicating the year the article was finally adopted for service use. Until the reign of the present Emperor (Showa era, started in 1926), items were designated by the year of the era. Now, however, the year of the Japanese Empire (Japanese year 2600 corresponds to our 1940) could be used. For items introduced up to the year 2600, the last two numbers are used in the designation. Thus Type 99 means that the item was adopted in 2599 (our 1939). The year 2600 may be represented as Type 100 or Type 0, while the years 2601, 2602, etc are represented as Type 1, Type 2, etc.

2. Exptl Ordn items were marked with the year of the Showa era during which the experiment was authorized.

3. Ordin items authorized in the eras preceding the Showa era: namely, Taisho (1912–1926) and Meiji (1867–1912), were designated by the era and the year of the era. Type II (Taisho)=1922; Type 41 (Meiji)=1908

4. The term *Modif* usually indicated a change in basic design, while the term Modification represented minor changes in design or a change in the expl filling.

Additional explanation is given in the introduction to each Section, such as Bombs, Land Mines, Grenades, Projectiles, Rockets, Mortars, Fuzes, Firing Devices and Sabotage Devices.

The Japanese explosives and related items are listed alphabetically in the pages which follow.

**Alphabetical List of Japanese Explosives and Related Items**

- **Aerial Torpedo.** Kural

- **Akatsuki.** An Ammonium Nitrate expl contg: AN 73–75, NG gel 5–6, cellulose 3–5 & other ingredients 14–9%. Detonation Velocity 3490m/s at density 1.0; Detonation Pressure (experimental) 42.3kbar, while the calculated value is 33kbar.


- **A(ko) or Type A (Explosive).** See Otsu-B

- **Aluminum.** See Arumiyumu

- **Ammonai — Ammonal.** Compr and specific uses are not known (Ref 5, p 361)

- **Ammonium Chlorate,** used in “Brown Powder” of compr: Amm chlorate 51.5, Ba nitrate 34.5, TNNaphthalene with oil 8.2 & woodpulp 5.0% (adds to 99.2%). Found as a booster in bombs for demolition (Ref 1, p 32)

- **Ammonium Nitrate.** See Ammon Shosanen

- **Ammonium Perchlorate** was used in Karito (Army Expl), Kaishikuyaku (Army Expl) and in Type 88 (Navy Expl) & Type 4 (Navy Expl)

- **Ammonium Picrate.** See Picurinsan Ammonia. Used in Type 1 Explosives (Navy Expls)

- **Ammon Shosanen.** Ammonium Nitrate (AN)
Used in a number of commercial and military expl comps, such as: Akatsuki, Ammonāru, Ammonyaku, Anboyaku, Angayaku, “E” (Explosive), Koi, Kiri Nos 1, 2 & 3, Ko-Shoan Bakuγu, Shin-Ki, Shinγyoryoku, Shin-Toku-Shoan, Shoan-Bakuγu, Shoan Bakuγu Nos 104 & 201, Shoanyaku, Shobenyaku, Shonγyaku, Shotoγu, Type 4 Mk 2, and others (Ref 5, p 361)

There have also been conducted research on the use of AN in Composite Propellants for Rocket Engines (Ref 5, p 361)

Ammonyaku. Army expl compn consisting of AN & charcoal. Japanese documents cite it as “substitute powder”. It was never recovered by Americans (Ref 1, p 29; Ref 5, p 361 and Ref 6, p 8-3)

AMMUNITION (Danyaku)

Introduction

Ammunition and Weapons used by the Army (Rikugun) were in most cases different from those used by the Navy (Kaigun). For this reason it is necessary to indicate whether the item was used by the Army or the Navy

Accdg to OPNAV 30-3M (1945) (Ref 1, p 123), Japanese Army Ammunition of WWII was divided into the following groups:

a) Small Arms Ammunition — items smaller than caliber 20-mm

b) Aircraft Cannon Ammunition — items of caliber 20-mm and over

c) Medium and Large Caliber Gun Ammunition — items of 20-mm and over

d) Mortar Ammunition

e) Fuzes

Under the title “Japanese Ammunition—Explanatory Notes”, TM 9-1985-5 (1953) (Ref 3, p IV), the following information is given:

The Japanese Army Terminology of WWII for projectiles conformed fairly well with US custom. The abbrn AP indicated a projectile intended for piercing heavy armor, such as armor plate of thickness equal to or greater than the caliber of the projectile. These projs had a small HE bursting charge. Abbrn AP-HE indicated a solid-nosed proj in general similar to AP but designed for much lighter penetration.

They contd larger HE charges than AP and smaller than standard HE projs

There were “old” and “new” color systems for marking the Army projectiles, and they are briefly explained on pp 266–69 of Ref 3

The following abbrns are used in this Section on AMMUNITION:

AAMG — Antiaircraft Machine Gun
AC — Aircraft Cannon
ACMG — Aircraft Machine
Amno — Ammunition
AP — Armor-Piercing
AP-HE — Armor-Piercing—High Explosive
API — Armor-Piercing Incendiary
APT — Armor-Piercing Tracer
ATk — Anti-Tank
BkPdr — Black Powder
HE — High Explosive
HEAT — High-Explosive Antitank (Shaped Charge)
HEI — High-Explosive Incendiary
HEIT — High-Explosive Incendiary Tracer
HET — High-Explosive Tracer
HMG — Heavy Machine Gun
How — Howitzer
1 or Incend — Incendiary
IT — Incendiary Tracer
LMG — Light Machine Gun
MG — Machine Gun
MK — Mark
Mod — Modification (or Modified)
Proj — Projectile
Propnt — Propellant
SD — Self-Destroying
Tk — Tank
WP — White Phosphorus

Ammunition, Army (Danyaku Rikugun)

Accdg to TM 9-1985-5 (1953) (Ref 3, p 265), such ammunition and weapons were generally copies of German and French designs or were developed following their customs. In comparison with items used by other countries in the years preceding WWII, the Japanese ammo and weapons appear to be outmoded. This is particularly true in considering Small Arms (Shōkahei) because there is little evidence that they were standardized. The weapon used prior to 1930 was 6.5mm, but shortly thereafter it was superseded by cal 7.7mm, but the change was not complete and many 6.5mm
items were used during WWII

The foreign influence in design was particularly evident after 1939 when Aircraft Cannons (Kôkûki taihô) of German and Italian design were copied. The largest Aircraft Cannon devised was 120mm, but nothing larger than 57mm was ever put in service use

Most Infantry Artillery Weapons (Hôheihe) were characterized by their immobility, as very few of them were designed for rapid motor transport. The standard Field Piece (Yahô) was 75mm, although 105mm and 150mm were frequently encountered.

One outstanding characteristic of Japanese armament was the large variety of type and sizes of Mortars (Kyûhô) in use. They were used not only as Infantry (Hôhei) support weapons but also as Artillery Pieces (Hô). They ranged in size from the 50-mm Grenade Discharger (Tekidantô) to the 320-mm “Spigot” Mortar (“Spigot” Kyûhô)

The standard Antiaircraft Gun (Kôshôhô) was a 75-mm Gun, but there was also an 88-mm AA Gun which was one of their most effective weapons and a 105-mm AA Gun. The Japanese also designed a 150-mm AA Gun for the defense of the home islands, but this was used only in the last few months of WWII

The newest trend in research and development in ammunition was along the line of Rockets (Roketto). Very few types were used during the war, but there were many experimental models of Antitank (Taisensha) and Artillery (Hôhei) Rockets, ranging in size from 75mm to 60cm.

Research was also conducted on Smoothbore (Kakkô no) and Recoiless (Hihandô) Weapons (Mid-Handô), but none of them was developed during the war beyond the experimental stage

The following Items of Japanese Army Ammunition are described, including pictures, in TM 9-1985-5 (1953) (Ref 3, pages indicated under each item):

- Type 38 6.5-mm Ammo for Rifle, Carbine and Light MG; used single-base graphited proplnt (pp 270-71)
- Type 99 7.7-mm (Rimless) and Type 92 7.7-mm (Semi-rimmed) for MG’s (pp 274-75)
- 7.92-mm Aircraft MG Ammo (p 277)
- 8-mm Pistol and 9-mm Pistol Ammo (pp 277-78)
- 12.7-mm AC Cannon Proj contd PETN & Incend chge (pp 278-79)
- Type 97 & Type 98 20-mm Ammo contd single-base proplnt; no info about filling of proj. Used in A/Tk and AA-A/Tk Guns (p 280)
- 20-mm HE Tracer, Type 98 HET and Type 100 HET Projectiles contd RDX and Tracer compr (pp 281-82)
- 20-mm HE Incendiary and 20-mm HEI (Ma 201) Projs contd RDX or PETN and Incendiary (pp 282-84)
- 20-mm HEIT Ammo for use in Ho-1 & Ho-3 AC Cannons. Its proj contd RDX, Incendiary & Tracer (p 284)
- Type 100 Mod 2 20-mm IT, SD Ammo for use in Type 98 AC-A/Tk Gun. Its proj contd Incend and Tracer comprs but there is no HE except in Booster (p 285)
- 20-mm APT Ammo for Type 97 A/Tk, Type 98 AA-A/Tk, Ho-1 & Ho-3 Guns. Its proj contd 7.0g of Tracer compr (pp 285-86)
- Ho-5 20-mm AP Ammo for use in Army AC Machine Gun. Its proj was solid steel shot and proplnt consisted of graphited cylindrical grains of single-base pdt (p 286)
- Type 2 and Type 2 Mod 20-mm HEI Ammo. The proj contd Cyclonite and Incend compr (p 287)
- Type 4 20-mm HEI (Ma 202) Ammo. Its proj contd PETN and Incend compr contg Ba nitrate, Al pdt, Mg pdr & wax (pp 287-88)
- Type 2 20-mm APT Ammo. Its proj contd 7.0g of Tracer compr (pp 288-89)
- 37-mm HEI Ammo for use in Ho-203 AC Cannon. Its proj contd Cyclonite & Incend compr; proplnt consists of 59.8g of flat 4-mm squares poured loosely into brass case and sealed in with cardboard disc (pp 289-90)
- 37-mm HEI Ammo for use in Ho-204 AC Cannon. Its proj contd Cyclonite/Wax and Incend compr; proplnt — 75.1g of the same type as above (pp 290-91)
- 37-mm Army Ammunition for eight A/Tk and Tk Guns listed on pp 293-97 used the same type of projs filled with PA (Picric Acid), but different cartridge cases. They were for the following guns:

Type 11th Year Infantry Gun — 1.85oz of graphited proplnt in the form of flakes; they were poured into the case and sealed with a cardboard disc (pp 293-95)
Type 94 Tank Gun - 2.7oz of graphited propellant in the form of rectangular flakes (pp 293 & 295)
Type 94 A/Tk, Type 98 Tank and Type 100 Tank Guns - 4.3oz of graphited propellant in the form of unitubular grains contd in a silk bag (pp 293 & 295)
Type 97 A/Tk Gun - 5.0oz of graphited propellant in the form of short, cylindrical grains contd in a silk bag (p 293)
- Type 94 37-mm HE Ammo had the propellant as above and the forward 1/3 of the projectile cavity filled with PA, while the after 2/3 of the cavity filled with cast TNT. Used in all 37-mm Guns (p 295)
- Type 94 37-mm AP-HE Ammo had the propellant as above and the proj filled with PA; used in many A/Tk & Tank Guns (p 295)
- Type 1 37-mm HE Ammo had the propellant as above and the proj filled with PA; used in Type 1 Tank & A/Tk Guns and also in Type 94 A/T Gun (p 296)
- Ho 301 40-mm HE Ammo for use in Ho-301 40-mm AC Cannon; had proj filled with TNT, contd in forward part a pellet of PA. The propellant chg (0.4oz) consisted of small greenish-gray, square flakes (ca 1mm in size) enclosed in a silk bag (pp 296-97)
- Type 1 47-mm HE Ammo for use in Type 1 A/Tk and Type 1 Tank Guns. The filling (87g) of its proj consisted of two preformed paper-wrapped blocks taped together and wrapped. The forward block consisted of two pellets of PA - a ring pellet around the gane and a solid pellet behind the gane. The after block was one cast piece of TNT. Its propellant chg (398g) consisted of single-perforated cylindrical grains of a graphited double-base smokeless propellant contg NC 60.0, NG 34.5, Etcentralite 3.0 & Diphenylformamide 2.5% (pp 297-98)
- Type 1 47-mm AP Ammo for use in Type 1 A/Tk and Tank Guns. Its proj contd 18g of 90/10-RDX/paraffin and a Tracer compn. Used 398g of the same double-base propellant as in HE Ammo (p 299)
- Type 90 5.7-cm (57-mm) HE Ammo for Type 90 and Type 97 Tank Guns. Its steel proj was filled with 250g of TNT and its propellant was 113g of Mk 1 square grain (med) (p 300)
- Type 90 5.7-cm (57-mm) Ammo with Substitute Cast Iron Projectile was used in Type 90 and Type 97 Tank Guns. The proj was filled with Black Powder and its propellant was the same as in above HE Ammo (p 301)
- Type 92 5.7-cm (57-mm) AP Ammo was used in Type 90 & Type 97 Tank Guns. Its proj contd two sections of expl, individually wrapped in paper; upper section was pressed PA/Wax, while lower section was cast TNT. ProjInt was Mk 1 square grain (med) (p 302)
- Ho-401 5.7-cm (57-mm) Ammo for use in Ho-401 AC Cannon; no data are given for its propellant and for filling of proj (p 303)
- Type 92 7-cm (70-mm) HE Ammo for use in Type 92 Infantry Gun (Howitzer). Its proj contd 590g of cast TNT (or pressured RDX/AN mixture). ProjInt (50g) was in square flakes contd in four pads made by sewing portions of the projInt betw two dark blue silk discs. The sections were not uniform in size, holding 5.4, 8.9, 16.8 & 18.7g of projInt each. The 5th disc, light blue in color, contd 3.1g of BkPdr serving as an ignition chge. The cartridge cases were of two designs; the newer type designated “B” (OTSU) was one-piece and slipped off the proj to change the projInt chge; an older design, presumably “A” had a threaded base which could be unscrewed to vary the chge (pp 304-05)

Note: Its variety for use in Type 94 Tank Gun is shown on p 308
- Type 92 7-cm (70-mm) Ammo with Substitute Projectile for use in Type 92 Infantry Gun (Howitzer). Its proj was filled with BkPdr and its projInt arrangement was the same as in 7-cm HE Ammo (p 305)
- Type 3 7-cm (70-mm) Hollow Charge Ammo for use in Type 92 Infantry Gun (Howitzer) as an A/Tk weapon. Its proj contd a Ho chge of cast TNT/RDX mixture wrapped in varnished paper; projInt - same as in Type 92 7-cm HE Ammo (pp 305-06)
- Type 95 7-cm (70-mm) Illuminating Ammo for use in Type 92 Infantry Gun (Howitzer). Filling of proj consisted of ejection chge (BkPdr) and Illuminating chge, probably consisting of mixt of Mg & Al pdr with Ba nitrate; projInt - same as in Type 92 7-cm HE Ammo (pp 306-07)
Type 95 7-cm (70-mm) AP Ammo for use in Type 94 Tank Gun contains proj filled with 170g of preformed, paper wrapped HE’s: Ōshiyaku (qv) in forward portion and TNT in rear portion; projInt was 120g of 5-mm square grain (medium) (p 309)
7-cm (75-mm) Army Ammunition. There were ten or more 75-mm (designated as 7-cm) guns in use. Although the weapons varied considerably in design, length of bore, and employment, the bore diam was held constant to 75-mm and the projs were designed to be interchangeable for numerous guns. In this case identical projs could be assembled with different sizes of cartridge cases and propol charges to fit the various weapons.

Ammunition for Navy guns designated “8-cm” have a bore diam 3 inches (76.2mm) and are not interchangeable with Army ammunition.

Following are the 75-mm Army guns belonging to this group, as listed on p 311:

Type 41 Mountain (Regimental)
Type 94 Mountain Gun
Type 38 Field Gun
Type 41 Cavalry Gun
Type 38 Improved Field Gun
Type 95 Field Gun
Type 11 Year Field or AA Gun
Type 90 Field Gun
Type 1 Self-Propelled Gun
Type 88 Field Gun
Type 88 Field or AA (Special) Gun

Propellent cases for these guns are illustrated on pp 312–15 and complete rounds are listed in Table on pp 316–17.

Following projectiles were used in the above guns:

7-cm (75-mm) HE-AA Proj filled with Önayaku (qv) was used in Type 11-Year Field AA Gun (p 318)

Type 90 7-cm (75-mm) HE-AA Long-Pointed Proj was filled with cast TNT (standard) and as alternate, PA. Used in Type 11-Year & Type 88 Guns (pp 319–20)

Type 90 7-cm (75-mm) HE, Long Pointed Projectile was filled with cast TNT (standard) or Heineiyaku (qv) as alternate. Used in Type 38 Field Gun Group (four weapons), Type 88 Field AA (Special) Gun, Type 90 Field Gun and Type 94 Mountain Gun (p 321)

Type 94 7-cm (75-mm) HE Proj was filled with TNT, while Angayaku and Heineiyaku were used as alternates; used in the same guns as above (pp 321–22)

7-cm (75-mm) Design “A” HE Proj contd 625g of Önayaku (qv); was used in Type 41 and Type 94 Mountain Guns and in Type 38 Field Gun Group (p 323)

7-cm (75-mm) Design “B” HE Proj contd Önayaku filling and was used in the same guns as above (p 324)

Type 98 Modified 7-cm (75-mm) HE Proj was filled with cast TNT and used in Type 41 Mountain Gun and Type 38 Field Gun Group (pp 325–26)

Type 90 7-cm (75-mm) HE Semisteel Proj was filled with cast TNT; was used in Types 41, 94, 38 & 90 Guns (pp 326–27)

Type 97 7-cm (75-mm) HE Semisteel Proj was filled with cast TNT; used in Type 41 Mountain Gun (pp 327–28)

Type 90 “A” 7-cm (75-mm) Substitute Proj was filled with BkPdr; used in Types 41, 94, 38 & 90 Guns (p 328)

Type 2 7-cm (75-mm) Hollow-Charge Proj contd 500g of cast 60/40–TNT/RDX wrapped in varnished paper; used in Type 94 and Type 41 Mountain Guns (pp 328–29)

Type 95 7-cm (75-mm) AP-HE Proj contd 460g of Önayaku; used in Types 41, 38, 90, 94 & 88 Guns (pp 330–31)

Type 1 7-cm (75-mm) AP Proj contd 53g of 90/10–RDX/Paraffin, coated with graphite and packed in an Al foil wrapper; used in Types 94, 41, 90 & 38 Guns (pp 331–32)

Type 38 7-cm (75-mm) Shrapnel Proj contd 100g of BkPdr as bursting chge together with lead balls; used in Types 41, 98, 38 & 90 Guns (pp 332–33)

Type 90 7-cm (75-mm) Shrapnel Proj contd 100g of BkPdr as bursting chge and ca 272 lead balls; used in the same Guns as above (pp 333–34)

7-cm (75-mm) Smoke (WP) Proj contd BkPdr as bursting chge and WP as chemical agent; no data for weapons (p 335)

Type 90 7-cm (75-mm):Smoke (WP) Proj contd 100g of Önayaku as bursting chge and 700g WP; used in Types 41, 94, 38 & 90 Guns. Its screening capacity was 20 meters high & wide and duration 1–2 minutes (pp 336–37)

Type 90 7-cm (75-mm) Incendiary Proj contd 20g of BkPdr as expelling chge and 50g as incendiary chge; used in the same guns as above (p 337)

Type 90 7-cm (75-mm) Illuminating Proj contd BkPdr as expelling chge and 250g of illuminating chge (probably a mixture of Al, Mg & Ba nitrate); used in the same guns as above (pp 338–39)
Type 11-Year 7-cm (75-mm) Target Proj contd folded parachute and shrouds in cardboard cylinder. After propelling the proj, the parachute became ejected by a small chge of expl, located in Time Fuze. Then the descending parachute and shrouds served as targets for AA practice; used in Type 11-Year AA Gun and in Type 88 Field AA Gun (p 339)
7-cm (75-mm) Liquid Incendiary Proj contd 40g of granular PA in paper container as bursting chge and soln of WP in CS₂ (300g) with rubber pellets (320g); used in Types 41, 94, 38 & 90 Guns (pp 339-40)
7-cm (75-mm) Vomit-Gas Proj contd 460g of 70/30-TNT/Naphthalene as bursting chge and 170g of Diphenylcyanarsine as liquid chemical agent; used in Type 41 Regimental or Mountain Gun (p 341)
Type 100 8-cm (88-mm) HE-AA Ammo with Long-Pointed Proj which was filled with 900g of TNT; was propelled by 2330g of No 16 cylindrical proplnt; used in Type 99 AA Gun (pp 342-43)
9-cm (90-mm) HE Ammo with Proj filled with 590g of crude TNT; used in an antiquated weapon classified as mortar; no info about proplnt used (p 343)
9-cm (90-mm) HE Ammo with Semisteel Proj which is filled with 650g of TNT; used in the same type of weapon as bove; no info about proplnt used (pp 343-44)
Army 10-cm (105-mm) Ammunition
There were three 105-mm Howitzers and four 105-mm Guns, which are designated 10-cm by the Japanese. The projs were in most cases interchangeable. All weapons used semifixed ammo except the Type 14-Year AA Gun which used fixed ammo (p 344)
Following list (extracted from p 346) gives types and weights of propellant charges used in 105-mm Weapons:
Type 14-Year Howitzer - 430g in 3 increments of Mk 1 square grain or Mk 3 strip
Type 91 Howitzer - 1135g in 4 increments of Mk 2 square grain
Type 14-Year Improved Howitzer - 692g in 3 increments of Mk 2 square grain
Type 92 Gun - 4000g in 2 increments of Mark 3 strip
Type 14-Year Gun - 2535g in 2 increments of Mark 3 strip
Type 38 Gun - 1740g in 1 increment of Mk 2 strip
Type 14-Year AA Gun - 3075g in 1 increment of Mk 3 strip
Illustrations for cartridge cases used in the above weapons are on pp 347 to 350 and following are the projectiles used in 105-mm weapons:
Type 91 10-cm (105-mm) HE Proj is filled with 2300g of cast TNT; used in Type 91 How and Type 38, 92 & 14-Year Field Guns (p 351)
Type 91 10-cm (105-mm) HE, Long-Pointed Nose Proj was filled with cast TNT and “White Composition” of AN, RDX & Gun; used in Type 91 How, Types 38, 92 & 14-Year Field Guns and 14-Year AA Gun (with fixed Ammo) (pp 352-53)
12-cm (120-mm) Shrapnel Amm with Proj contg BkPdr as ejecting chge and 539 lead balls packed in a resin matrix. The proplnt was in two increments each encased in a silk bag with a small igniter chge sewed to the bottom of each; the 1st increment was NC in 5/16” square flakes, while the 2nd was NC in 1/16” square flakes; used in Type 38 How (pp 353-54)
12-cm (120-mm) AP-HE Amm with Proj filled with PA; used the same proplnt and in the same weapon as for shrapnel ammo (pp 354-55)
15-cm (150-mm) Army Ammunition
There were three 150-mm Howitzers and four 150-mm Guns, which varied greatly in design from a very short-range How to long-range Field Guns. The projs were not as interchangeable as in 75-mm & 105-mm weapons. Usually there was one type of proj for use in Hows and another type for the Guns. All the weapons used semifixed ammo with the exception of Type 88 Gun which used a bag charge
Following list (extracted from p 357) gives types and weights of proplnt chges used in 150-mm Weapons:
Type 4-Year How - 2260g in 5 increments of Mk 2 square grain
Type 38 How - 825g in 2 increments of Mk 1 square grains
Type 96 How - 2930g in 5 increments of Mk 2 square grains
Type 45 Gun
Type 7-Year Gun
Type 90 Gun
Type 89 Gun - 9700g in 2 increments of Mk 4 strip
Illustrations of cartridge cases used 150-mm wea-
pons are given on pp 358–61 and the following projectiles used in these weapons:

Type 92 15-cm (150-mm) HE Proj was filled with Angayaku (qv); used in Type 4-Year, Type 96 & Type 38 Hows (p 362)

Type 93 15-cm (150-mm) HE Proj was filled with cast TNT; used in Type 89, Type 45, Type 7-Year and Type 90 Guns (p 363)

Type 92 15-cm (150-mm) HE, Long-Pointed Proj was filled with cast TNT; used in Type 4-Year, Type 96 & Type 38 Hows (p 364)

Type 93 15-cm (150-mm) HE, Long-Pointed Proj was filled with cast TNT; used in Type 89, Type 45, Type 7-Year & Type 90 Guns (p 365)

Type 95 15-cm (150-mm) AP-HE Proj was filled with two preformed blocks of high grade TNT; used in Type 4-Year, Type 96 & Type 96 Hows and in Type 45, Type 7-Year, Type 90 & Type 89 (bag charge) Guns (p 366)

15-cm (150-mm) AP-HE Proj was filled with PA in two preformed, paper-wrapped bags; used in the same guns and Hows as above (p 367)

Type 13 15-cm (150-mm) Smoke (WP) Proj contd PA as bursting chge and WP cast in a brass cylinder which, surrounded by wax, was fitted below PA chge; used in Type 4-Year, Type 96 & Type 38 Hows (pp 367–68)

30-cm (305-mm) AP-HE Proj; no data for its filling (p 369)

30-cm (305-mm) AP-HE Proj; no data for its filling (p 370)

Army Rockets (Ref 3, pp 371–72) — See under ROCKETS

Army Mortars (Ref 3, pp 372–90) — See under MORTARS

Army Projectile Fuzes (Ref 3, pp 391–426) — See under FUZES

**Ammunition, Navy (Dairyaku Kaigun)**

Accdg to Ref 1, p 170, Japanese Navy Ammunition of WWII was divided into the following groups:

a) Small Arms Ammunition — items caliber 7.7-mm, 7.9-mm, 13-mm and 13.2-mm

b) Aircraft Cannon Ammunition — items caliber 20-mm and 30-mm

c) Anti-aircraft “Automatic Weapons” Ammunition — items caliber 25-mm and 40-mm

d) Large Caliber Ammunition — items of caliber 5-cm (50-mm) and over

Under the title "Japanese Ammunition—Explanatory Notes", TM 9-1985-5 (1953) (Ref 3, p IV), the following information is given:

The Japanese Navy Terminology of WWII for projectiles was highly irregular and cumbersome. Complete and accurate identification of proj required identification of the gun, descriptive nomenclature of the proj, and mark (or type) and modification number. For this reason an arbitrary system of nomenclature is used in TM 9-1985-5 (Ref 3). All Navy proj with a relatively heavy HE charge (including light AP types) are designated TSUJODAN, which may be translated either "Ordinary Projectile" or "Common Projectile". In TM 9-1985-5, the term "Common" is reserved for light penetrating type of projectile (solid nose, base fuzed), while for proj having point-detonating fuzes, the US designation HE is used.

Color system of marking projectiles for all sizes over 40-mm is explained on pp 427–31 of Ref 3, and proj of 40-mm and below on pp 432–33.

The following Items of Japanese Navy Ammunition are described, including the illustrations, in TM 9-1985-5 (1953) (Ref 3, pages indicated under each Item):

7.7-mm Aircraft Machine Gun Ammunition with Ball AP, Incend, Tracer & HE Projectiles; no data for proplnt; used in Type 92 & Type 97 ACMG's and in Type 92 AAMG (pp 434–35)

13-mm ACMG Ammunition with Tracer, HE (PETN)T and Incend (WP) Projs; no data for proplnt; used in Type 2 ACMG (pp 436–37)

13.2-mm AC and AAMG Ammo with Tracer, AP & HE(PETN) Incend (WP) Projs; no data for proplnt; used in Type 3 ACMG & Type 93 AAMG (pp 438-39)

**Naval 20-mm Ammunition** was divided into the following:

Type 99 20-mm Ammo (p 440) existed as Mk I and Mk II and was copied from the Swiss Oerlikon design. The proplnt consisted of graphited single-base, single-perforated cylindrical grains, 13.6g for Mk I and 21.4g for Mk II. Both Mk's used the same types of projectiles, which were as follows:

20-mm HE Proj contd as filling 50/50–Pentolite (p 441)

20-mm HE Incend Proj contd as filling TNT and WP (p 442)
20-mm HETracer Proj contd Pentolite and tracer compn (p 443)
30-mm Type 2 & Type 5 HE Incid Projs contd 3.56g of Pentolite and 19.94g of WP (p 452)
20-mm HET, Self-Destroying Proj contd Pentolite and Tracer compn. The proj was similar to the HET type except for its self-destroying feature. This was accomplished by ignition of BkPdr train leading from the tracer thru the hole drilled thru the septum of the proj and HE filling to the base of the gaine (p 444)
30-mm Type 5 HETracer Proj contd in upper cavity Pentolite, while Tracer was in lower cavity (p 453)
20-mm AP Incend Proj contd Incend consisting of NC 77.5, Na nitrate 11.3 & Al pdr 11.2% which was loaded at the base. Upon impact the rear end of proj ruptured and the heat generated ignited the Incend compn. There was no fuze (p 444)  
30-mm Type 5 Tracer Proj contd nothing but Tracer (p 454)
20-mm Practice Proj contd no filling; was used as the first round to clear the gun bore (p 455)  
40-mm Naval Ammunition was used in 40-mm Navy's Vickers-Armstrong Guns. Its proplnt consisted of 95.9g cylindrical length of smokeless pdr (p 455) and the following projectiles were used:
40-mm AP Proj contd 23g of cast TNT (p 456)
40-mm HE AA Proj contd 70g of cast TNT (p 457)
40-mm Tracer Proj contd nothing but Tracer compn (p 458)
5-cm (47-mm) Complete Round and Common Projectile. Its proplnt consisted of 67g unperforated, cylindrical, amber-colored sticks of double-base proplnt (5C2 Type 5-Year) enclosed in a compartmented bag of heavy brownish silk, placed in brass or steel cartridge. Its proj was 50g of loose granular BkPdr. Used in Short 5-cm Gun, mounted on wooden-spoked wheels as a field piece (pp 458–59)
8-cm (76.2-mm) (3-inch) Ammunition for use in various 8-cm/40 Guns included:
8-cm (76.2-mm) Complete Round (Semifixed) used 900g of 203C (Type 89) proplnt which consisted of unperforated 1/16–1/32 by 12.75 inches cylindrical sticks of double-base pdr enclosed in heavy paper and placed in brass cartridge (p 460); used one of the following projectiles:
8-cm (Ordinary Mk 2 Mod 2) HE Proj contd 0.71 lb of Shimose (cast PA) in single block sealed in a waxed paper (p 461)
8-cm Shrapnel Proj contd BkPdr as ejecting chge and shrapnel balls (p 462)
8-cm (Ordinary Mk 3 Mod 1) HE Proj contd 0.9 lb of Shimose (cast PA) (p 464)
8-cm AP Type 1 Special Common Proj contd 0.41 lb Type 91 Expl (Trinitroanisole) (p 465)
8-cm Time Practice Proj filled with either smoke compd or spotting dye (p 466)
10-cm (100-mm) (3.9-inch) Complete Round (Fixed) contd 12.7 lb of double-base proplnt
(Type 91 Mk 2) in the form of flat strips (pp 467–68). It was used with:

10-cm (100-mm) HE Proj which contd 3.13 lb of Type 92 Expl (cast TNT). Used in Type 98 10-cm/65 & 10-cm/50 Dual Purpose Guns (p 468)

12-cm (120-mm) (4.7-inch) Complete Round (Fixed) contd 1.1 lb Type 89 Proplnt consisting of unperforated cylindrical stick 1/32 by 6.7 inches of double-base pdr. Used in Short 12-cm Gun, 12 Calibers Long (p 469). Its projectile was: 12-cm/Short HE Proj contd 5.5 lb of Type 91 Expl (Trinitrotoluene) (p 470)

12-cm (120-mm) (4.7-inch) Complete Round (Semi-fixed) contd in brass cartridge 11.88 lb of Type 13 Propellant consisting of unperforated cylindrical sticks of double-base pdr. Used in Type 11-Year 12-cm/45 Gun (Low Angle) with the following projectiles:

12-cm HE (Ordinary Mk 3) Proj contd 3.61 lb of cast PA (Shimose) (pp 472–73)

12-cm HE (Ordinary Mk 3 Mod 1) Proj contd 3.61 lb of Shimose (p 474)

12-cm HE (Ordinary Mk 4) Proj contd Shimose (p 475)

12-cm Common Projectile contd Shimose (p 476)

12-cm Illuminating Proj contd 0.5 oz of BkPdr as expelling chge and illuminant consisting of Ba nitrate 35.6, K nitrate 10.4, Mg 38.0, wax 13.9, carbon 0.8 & sulfur 1.3%. Compn of first fire compd was: K nitrate 63.2, wax 5.5, glass 6.1 & carbon 13.1%. Illuminant was located in 10 cardboard cylinders arranged in two layers of 5 cylinders, each contg 5.4 oz of illuminant. The proj was used in Type 11-Year and Type 3-Year 12-cm/45 Guns (pp 477–78)

12-cm (120-mm) (4.7-inch) Complete Round (Fixed) contd in brass cartridge 12.0 lb of Type 13 double-base proplnt in the form of cylindrical sticks. Used in Type 10-Year 12-cm/45 Dual Purpose Gun with the following projectiles:

12-cm HE (Ordinary Mod 1) Proj contd 3.96 lb Type 91 Expl (pp 481–82)

12-cm HE (Ordinary Mod 2) Proj contd 3.46 lb of Shimose (p 482)

12-cm Incendiary Shrapnel contd 1.19 lb of Shimose and Incendiary charge consisting of WP & 48 steel pellets, total wt 8.55 lbs (pp 482–83)

12.7-cm (127-mm) (5-inch) Complete Round (Fixed) contd in brass cartridge 8.87 lb of Type 13 Double-Base Propellant consisting of graphited unperforated cylindrical sticks 0.079 by 15.75 inches packed in a compartmented bag of heavy silk. Used in Type 88 & Type 89 12.7-cm/40 Dual Purpose Guns with the following projectiles:

12.7-cm HE Proj was filled with 3.94 lb of Shimose (p 485)

12.7-cm Incendiary Shrapnel (Type 3 Ordinary) Proj contd 162g Shimose, 2.3g Tetryl, 0.13g LA, 35g BkPdr and Incendiary charge, which consisted of 43 steel pellets filled with the following compn: Mg pdr 54, Ba oxide 26, rubber 15, Fe oxide 1.5 & sulfur 3% (adds to 99.5%) (p 486)

14-cm (140-mm) (5.5-inch) Propellant and Powder Tanks.

Under the term “powder tanks” sealed waterproof cylinders were used for storing propellant charges. Two types were recovered by Americans: one was a heavy tank (26.4 lbs) made of cast steel and bronze; the other was of sheet steel and Al (15.4 lbs). Both tanks were well lacquered inside and had the same internal dimensions – 6.2 by 33.4 inches

Propellant chge of 14-cm Bag Ammo consisted of 24.51 lbs of double-base proplnt 37C which consisted of graphited unperforated cylindrical sticks 0.15 by 28 inches, enclosed in a heavy silk bag. Used in 3-Year 40-cm/50 Gun (Low Angle) with the following projectiles:

14-cm HE Proj contd 7.9 lb Shimose (p 489)

14-cm HE (Type 0, Ordinary) Proj contd 6.60 lb Shimose (p 490)

14-cm (Ordinary Mk 1) Common Proj contd 5.94 lb Shimose (p 491)

14-cm (Ordinary Mod 1) Common Capped Proj contd 4.80 lb Shimose (p 492)

14-cm Illuminating Proj contd: primary ejection chge (100g BkPdr), secondary ejection chge (41g BkPdr); initiating pellet, relay train and delay were BkPdr; ignition compd for the illuminant (mixt of K nitrate, Fe oxide, Al, S & wax); illuminant (mixt of Ba nitrate, Mg & wax). Total weight of filling 31 lbs (p 493)

14-cm Smoke Proj was filled presumably with WP (p 494)

14-cm (140-mm) (5.5-inch) Cartridge Case (Semi-fixed) was of brass and contd 6860g of Type 13 double-base proplnt consisting of graphited unperforated cylindrical sticks 1/8 by 19% inches enclosed in a heavy silk bag. No
projs used with this case are listed (p 495).

15-cm (152-mm) (6-inch) Complete Round (Semifixed) used 18.94 lb 37DC Double-Base
Propln made of graphited unperforated cylindrical sticks 1/8 by 26 inches enclosed in heavy silk bag packed in a brass cartridge. Used in Type 41 (Meiji) 15-cm/40 Gun (Low Angle)
with the following projectiles:

15-cm Common (Ordinary Mod 1) Proj contd
Shimose (p 497)
15-cm HE (Ordinary Type 0) Proj contd 6.8 lb
Shimose (p 498)
15-cm Common (Ordinary Mk 4) Proj contd
5.9 lb Shimose (p 499)
15-cm Incendiary Shrapnel Proj — no data for
filling (p 500)

15.5-cm (155-mm) (6.1-inch) Ammunition was
used in 15.5-cm Gun (Bag). No data for propln
used but the following projectiles are listed:

15.5-cm HE (Ordinary Type 0) Proj contd
7.5 lb Shimose (p 502)
15.5-cm AP (Type 91) Proj contd 6.84 lb Tri-
nitroanisole (p 503)

15.5-cm Illuminating Proj was filled with the
same substance as 14-cm Illuminating Proj (p
504).

20-cm/Short (202-mm) (8-inch) Complete
Round (Semifixed) was used in 20-cm Short
Gun. Its propln was double-base 10C3 (Type
89) consisting of unperforated cylindrical sticks
1/32 by 8½—9 inches, enclosed in two com-
partmented heavy silk bags packed in brass
 cartridge. The inner bag contd 2.1 lb of propln,
whereas the outer bag had 2.3 lb. The following
projs were used:

20-cm/Short HE (Ordinary Mk 1) Proj contd
28.50 lbs Trinitroanisole in three cast blocks
(p 506)
20-cm AP, Type 91 Proj contd 17.38 lb TNAns
in a preformed block (p 507)
20-cm HE (Ordinary Type 0) Proj contd 21.34
lb of TNAns (p 508)

36-cm (365-mm) (14-inch) AP Proj used in
36-cm/45 Gun (Bag); probable filling of proj
— TNAns (p 510)

40-cm (406-mm) (16-inch) Mk 5 AP Proj was
used in 40-cm/45 Gun (Bag); probable filling —
Trinitroanisole (p 511)

Note: Under Navy Ammunition are also de-
scribed Rockets (Ref 3, pp 512—14); Mortars
(pp 515—17) and Projectile Fuzes (pp 514—43),
but they are described here separately under
the corresponding letters in alphabetical order

Refs for AMMUNITION:

1) Anon, "Handbook of Japanese Explosive
Ordinance", OPNAV 30-3M (1945), pp 123—57
(Army Ammo) & 170—91 (Navy Ammo)
2) Anon, "Japanese Explosive Ordnance",
& E.J. Hoffschmidt, Eds, "Japanese Combat
Weapons of World War II", WE Inc, Old Green-
wich, Conn, Vol 2 (1968)

Amphibian (Water) Mines. See under MINES

Anbenyaku or Shōbenyaku. HE mixture con-
sisting of AN 55 & Dinitrobenzene 45% used
during WWII as bursting charge of some projectiles.
The name Shōbenyaku was used at Nanman Ar-
senal, Manchuria. An exp1 mixt by the same
name consisted of AN, TNBz & Tetryl

A blasting exp1, also called Anbenyaku, con-
sisting of AN 71.7, NG 8.0, Collod Cotton 0.3
and powdered seaweed 20% was proposed in
1948 by T. Watanabe, JapanP 176113 (1948)
& CA 45, 4930 (1951) (Ref 5, p 361)

Angayaku. Several HE compns were known.
One of them listed in Ref 1, p 27 consisted of
AN 75 & RDX 25%. It was cast in TNT sur-
rond for filling bombs. The compn listed in
Ref 6, p 8-3: RDX 85 & wax 15% was used in
AP Projectile while compn: RDX 41, PETN
50 & wax 8% was used in Machinegun Bullets

In addn to these, the following compns are
listed in Ref 5, p 362: a) AN 84 & RDX 16%;
b) AN 51, RDX 15 & GuN (Guadine Nitrate)
34% and c) AN 48, RDX 20 & GuN 32%

All of the above compns were white in color,
nontoxic and comparable in performance to
Aramatol. It was claimed that compns contg
GuN had low coefficients of shrinkage and,
therefore, could be poured in large casts in a
single pour

Another Angayaku compn considered to be
similar to Brit & US Torpex consisted of AN
43.2, GuN 28.8, RDX 8.0 & Al powder 20% 
was manufd at the Sankanoichi Factory of the
Tokyo Army Arsenal No 2. It was used to a
limited extent in various types of underwater
exp1s (Ref 5, pp 361—62)

As an example of use of Angayaku as a
filler may be cited Type 94 7-cm (75-mm)
HE Proj described and illustrated in Ref 3, p 321

**Armored Cars.** See under Combat Vehicles

**Armor-Piercing High-Explosive (AP-HE) Projectile.** Hakoryudan Tekkodan

**Armor-Piercing (AP) Projectile.** Hakodan or Tekkodan

**Arsenals (Kōshō or Zōneishō) and Powder Factories (Kayaku seizōsho)**

The following Arsenals are listed in Ref 1, p 6: Tokyo, Osaķa, Kure, Toyokawa, Yoko-suka, Maizuro and Sasebo. Tokyo and Osaka were the Army Arsenals, while the other five were the Navy Arsenals. The following Arsenals are listed in the book of Tantum & Hoffschildt, listed here as Ref 7, p 38: Kokura, Tokyo, Nagoya, Heijo, Osaka, Chioda and South Arsenal. In the same book the Tokyo Explosives Plant is listed

**ARTILLERY (Hōhei) AND ITS WEAPONS (Hō)**

In the book *Tantum and Hoffschildt* (Ref 7), Section 3, pp 80–116, the following weapons are described and illustrated:

- 37-mm Infantry Gun, Mod 11 (1922) (pp 80–1)
- 37-mm Gun, Mod 94 (1934) (pp 84–5)
- 47-mm Antitank Gun, Mod 1 (1941) (pp 84–5)
- 70-mm Battalion Howitzer, Mod 92 (1932) (pp 86–7)
- 75-mm Field Gun, Mod 38 (1905) (Improved) (pp 88–9)
- 75-mm Regimental Gun, Mod 41 (1908) (pp 90–1)
- 75-mm Antiaircraft Gun, Mod 88 (1928) (p 92)
- 75-mm Mobile Field AA Gun, Mod 88 (1928) (p 93)

*Note:* On p 94 are given illustrations of the following Japanese Antiaircraft Cannons:

- 75-mm Heavy AA Cannon, M88 (1928) — the Standard Army; 76-mm AA Cannon, M3 (1914);
- 105-mm Army AA Cannon M14 (1925) and
- 127-mm Twin Mount, Dual-Purpose Cannon, M98 (1929)

Table on p 95 lists “Japanese 75-mm Artillery Ammunition.” This includes Models of Projectiles, Their Bursting Charges, Standard Fuzes and Models of Guns:

- 75-mm Field Gun, Mod 90 (1930) (p 96)
- 75-mm Mountain (Pack) Gun, Mod 94 (1934) (pp 97–8)
- 75-mm Field Gun, Mod 95 (1935) (pp 99–100)
- 8-cm (7.62) High Angle Gun, Type 3 (p 101)
- 88-mm Antiaircraft Gun, Type 99 (Year ?) (p 104)
- 105-mm Howitzer, Mod 91 (1931) (pp 103–04)
- 105-mm Gun, Mod 92 (1932) (pp 105–06)
- 120-mm, 45 Calibers, Naval Dual Purpose Gun, Type 10 (p 107)
- 120-mm, 45 Calibers, 11-Year Type Gun (p 108)
- 140-mm Seacoast Gun, Type 3 (Year ?) (p 109)

*Note:* Table on p 110 lists “Characteristics of Principal Japanese Weapons”:

- 150-mm Howitzer, Mod 4 (1915) (pp 111–12)
- 150-mm Howitzer, Mod 96 (1936) (pp 113–14)
- 300-mm Short Howitzer, Type 7 (Year ?) (p 115)
- 105-mm Field Gun, Type 14 (Year ?) (p 115a)

(See also Mortar and Rocket)

**Artillery Ammunition.** See under AMMUNITION

**Aruminyūmu.** Aluminum. It was used in powdered form in many Japanese Navy comps, such as Otsu B, Type 2, Type 4 & Type 92, listed in Ref 1, pp 32–33. It was also contd in one of the Army expls — Angayaku (qv)

**B4 (Incendiary) or Type 2 Navy Explosive.** A light-gray castable mixture of Trinitroanisole 60 to 70 & Al 40 to 30%. The props of the 60/40 mixture were: density 1.90; Brisance by Cu cylinder crusher test 82% (PA=100%); Expln Temp 300°; Impact Sensitivity with 5kg wt 17cm maximum with no expls; Friction Sensitivity with 60kg pressure — no expls; and Power by Ballistic Mortar 64% of PA. It was used in Submarine Gun Incendiary Shells (Ref 5, p 362)

*“Baka” Piloted Rocket Bomb.** See under BOMBS and in Ref 2, pp 116–18

**Bakkan**  Blasting Cap

**Bakubō.** Detonator

**Bakuchiku.** Squib

**Bakudan.** See BOMB
Bakufun. Detonating Explosive or Exploding Powder. A light-gray to tan pdr consisting of MF (Mercuric Fulminate) 28.8, K chloride 37.7, Antimony trisulfide 31.5 & abrasive 2%. Accdg to documents, Mkis I & III are used in ammunition primes while Mk 2 is used in fuze primes (Ref 1, p 25 & Ref 5, pp 362–63). See also Raibun

Bakuhaki. Explosdr; Blasting Machine

Bakuhatsu. Explosion, Blast or Detonation

Bakuhatsu kōryoku. Blast Effect

Bakuhatsu sei. Blasting Gelatin

Bakuhayaku. Detonating Explosive; Brisant Explosive

Bakuhuyō bakudan. General Purpose Bombs; Demolition Bomb

Bakurai. Depth Charge

Bakurai tōshahō. Mine Thrower; Y-Gun

Bakuretsu jūdan. Explosive Bullet

Bakuyaku. Explosive Charge; Powder Charge

Bakuyakytō. Bangalore Torpedo

Balloons, Paper. Ingenious paper balloons were launched in 1944 against the West Coast of the US and Canada. This is described in Vol 2 of Encycl on p B11-L, under “Balloons and Airships and Their Application in War”

Bangalore Torpedo. Bakuyakuto or Hakaitō. See under Demolition Tubes

Barisutaito. Ballistite

Biransei gasu. Blister Gas

Black Powder. Kokushokuyaku (Black Color Explosive) or Yuenyaku (Nonsmokeless Powder); Gunpowder — Kayaku. Black, loose-powdery material consisting of K nitrate, sulfur & charcoal. It was used during WWII for the follow-
ing items: a) Army 20-mm Ammo as a temporary measure; b) Army ejc. charges for 70-mm Barrage Mortar, Shrapnel Shells & Pyrotechnics; c) Army Delays & Relays in Bomb & Projectile Fuzes; d) Army Delays and Igniters in Pyrotechnics; e) Navy ejc. charges for Illuminating Shells & Pyrotechnics; f) Navy Delays & Relays in Bomb & Projectile Fuzes and g) Navy Delays & Igniters for Pyrotechnics (Ref 1, pp 27, 30 & 33)

Blast. Bakuhatsu

Blasting Cap. Bakkan

Bōgyō. Sabotage

Bōgyn. Defense

Bōgyo jirai. Antitank Mine

Bokūyō kaki. Antiaircraft Weapon

BOMBS (Bakudan) (Excluding Flares and Rocket Bombs)

Introduction

Accdg to TM 9-1985-4 (1953) (Ref 2, p 1), the Japanese Army (Rikugun) and Navy (Kaijun) had during WWII separate Air Forces (Kūgun), each of which employed its own distinct bombs and fuzes (Shinkan). For the most part the Army and Navy bombs and fuzes cannot be used interchangeably. Special adapters have been developed, however, which allowed some flexibility of this rule

Until the reign of the present Emperor (Showa era, which started in 1926), items were designated by the year era. Now, however, the year of the Japanese Empire—2600, which corresponds to our 1940, may be used. For items introduced up to the year 2600, the last two numbers are used in the designation. Thus, Type 99 means the item was adopted in 2599 or our 1939. The year 2600 may be represented as Type 100 or Type 0, and the years 2601, 2602, etc as Types 1, 2, etc

Exptl ordnance items are assigned the type numbers indicating the year of Showa era during which the experiment was standardized

Ordnance items standardized in the eras preceding the Showa era; namely, Taisho (1912–
1926) and Meiji (1867–1912) are designated by the era and the year. Type II (Taisho) = 1922 and Type 41 (Meiji) = 1908

Some items developed for a special purpose were designated by a Mark number. The term Model indicates a change in basic design, while the term Modification represents minor changes in design or a change in explosive

Abbreviations are the same as indicated under AMMUNITION

Bombs, Army (Bakudan Rikugun)

The following Army Bombs are listed in Ref 2 on pages indicated under items:
Type 92 15-kg HE contd cast PA or TNT (pp 4–5);
Type 99 30-kg HE contd 48/52–RDX/TNT (p 5);
Type 94 50-kg HE, Type 3 100-kg HE, Type 94 Modified HE, Type 1 50-kg HE, Type 1 100-kg HE, Type 1 250-kg, Type 92 250-kg and Type 92 500-kg HE Bombs were filled with cast PA (pp 6–9);
Type 3 100-kg and Type 3 250-kg Skimming Bombs contd paper-wrapped cast PA blocks sealed with TNT (pp 10–11);
Type 4 100-kg, 250-kg & 500-kg Antishipping Bomb contd paper-wrapped cast PA blocks sealed with TNT (pp 12–13);
1-kg and 5-kg Thermit Incendiary Bombs contd Thermit as main chge and first fire chge (pp 14–16);
Type 97 12-kg Thermit consisted of thin steel body contd two BKpdr chges and three Thermit filled Mg fire pots (pp 16–18);
Type 97 50-kg and Type 100-kg Incendiary Bombs consisted of thin-steel body filled with 400–450 rubber bags impregnated with soin of phosphorus in CS2. The HE chge in the nose and central burster tube was PA (pp 18–20);
Type 100 50-kg Smoke Bomb consisted of thin-steel body filled with FS smoke compn (chlorosulfonic acid 41, sulfur trioxide 54 & sulfuric acid 5%). The HE chge in the nose and in the burster was PA (p 21);
Type 92 Gas Bomb contd 50/50–Lewisiite/ Mustard Gas (p 22);
Type 97 15-kg Concrete Bomb was a light sheet steel cylinder with central exploder tube filled with cast PA which was surrounded by steel pellets set in concrete (p 26);
Type 94 10-kg Substitute Bomb consisted of a thick concrete case surrounding a steel central tube burster filled with loose BKpdr. The tube was surrounded by steel pellets set in concrete (p 27);
Type 1 30-kg Substitute Bomb consisted of a light sheet steel cylinder surrounding a central steel burster tube contg in forward part BKpdr chge and after part HE chge. The space betw the tube and outer casing was filled with steel pellets set in concrete (pp 27–28);
Type 95 4-kg Practice Bomb housed an ampoule contg a smoke compn (p 29)
Cluster Bombs: Type 2 1/3-kg, Type 3 1/2-kg and Type 2 1/2-kg contg TNT 58–60 & RDX 42–40% fillers (pp 29–32). Containers for these bombs are described on pp 33–36;
1-kg Aircraft Missile consisted of a spherical compressed paper container from which a 3-inch high tubular neck of compressed cardboard projected. A wooden plug at the base of this neck housed a friction pull igniter. A central burster chge consisted of granular BKpdr in a silk bag. Surrounding this were 32 cylindrical sheet-metal pellets contg a LE (Low Explosive) chge of K nitrate 55.7, sulfur 16.7, Al 14.6 & antimony sulfide 13.0%. Each pellet had a 1/2-inch safety fuse which was in contact with burster chge. For its operation the cord of the igniter was pulled and the missile was thrown from the plane. After a short delay the BKpdr burster chge exploded rupturing the paper body, scattering the expl pellets and simultaneously igniting the fuse of the pellets. After a short delay the pellets were detonated. This missile was used in air-to-air bombing (pp 36–7 & Fig 31);
50-kg & 100-kg Pamphlet Containers consisted of cardboard case contg a small burster chge and paper pamphlets (p 38);
Miscellaneous Army Bombs — Illustrations without description of several bombs are given on pp 39–40

A shorter description, with illustrations, of Japanese Army Bombs is given in Ref 1, pp 66–80

Bombs, Navy (Bakudan Kaigan)

Introduction

Accdg to Ref 2, p 41, Navy bombs were divided into the following classes: “Land” bombs specially designed for use against land targets; “Ordinary” were designed for use against ships. They included both GP (General Purpose) and SAP (Semi-Armor-Piercing); “Special” are designed for special purposes and
each class is indicated by a mark number;
“Smoke” were used for concealment purposes;
“Practice” were used for practice bombing;
“Target-marker bombs were used as target-
marking beacons;
“Training” bombs were used for training in
handling;
“Dummy” bombs were used for training and
practice bombing.

Individual Navy bombs in these main general
classes were given the Type number (such as
Type 97) which disclosed the year that the
bomb was adopted for service use. In the “land”
and “ordinary” classes, the 1st bomb of a given
wt class was assigned a Type number but
was merely indicated by the wt number. Sub-
sequent designs of the same wt were assigned
Type numbers. Thus there was a No 6 land
bomb and a Type 97 No 6 bomb. The No
indicated the wt in units of tens of kg. Thus a
No 6 bomb weighed 60-kg, a No 25 weighed
250-kg, etc. For “special” bomb the Mk num-
ber was given. The term Model usually meant
a different design of the bomb in the same
general class, while Modification meant a minor
change in the design or filling.

Type 97 No 6 Land Bomb had a sheet steel
body filled with PA or type 98 Explosive (p 45);
Type 2 No 6 Land Bomb had sheet steel body
filled with five 7-kg HE bombs and contd
central burster tube filled with HE (pp 46–7);
No 25 Land Bomb and Type 96 No 25 Land
Bomb had steel body filled with PA or Type
98 Expl (pp 47–9);
No 80 Land Bomb had steel body filled with
PA or Type 98 Expl (p 49);
Type 99 No 6 Ordinary Bomb had steel body
filled with PA or Type 98 Expl (p 50);
Type 99 No 25, also No 80, Ordinary Bombs
Model 1 were filled with Type 91 Expl (pp
50–1 & 53);
Type 2 No 50 Ordinary Bomb Model 1 was
filled with Type 98 Expl (p 52);
No 3, No 6 & No 25 Ordinary Bombs Model 2
were filled with PA (pp 54–5);
No 50 Ordinary Bomb Model 2 was filled with
Type 98 Expl (p 56);
No 6 Mk 1, Type 1 No 6 Mk 1 and Type 4
No 6 Mk 1 Bombs were filled with mustard
gas thickened with methacrylate & polyvinyl
alcohol (pp 56–7);

Type 4 No 6 Mk 1 Bomb was designed to be
filled with mustard gas or with any suitable
CWA (Chemical Warfare Agent) (p 58);
Type 99 No 6 Mk 2, Type 99 No 6 MK 2
Modification 1, Type 1 No 25 Mk 2 Model 1
and Type 1 No 25 Mk 2 Model 1 Modification
1 Bombs were filled with Type 98 Expl (pp
58–63);
Type 99 No 3 Mk 3 Bomb consisted of a sheet
metal cylinder contg a canister with 168 WP-
filled steel pellets. PA chgs were in tail cone
and central burster (pp 63–4);
Type 3 No 6 Mk 3 Model 1 Bomb contd three
cylindrical steel canisters, each contg 87 WP-
filled cylindrical steel pellets; each canister had
central exploder filled with Type 98 Expl (pp
64–6);
Type 2 No 25 Mk 3 Model 1 Bomb contd an
incendiary chge consisting of Ba nitrate 35.8,
Al 13.6, Mg 10.3, Fe oxide & synthetic rubber
shavings 13.1%; two igniter chges (Ba nitrate
75.2, Al 24.2 & oil 2%) were located in the nose
piece and in the tail cone (pp 66–7);
Type 3 No 25 Mk 4 Modification 1 Bomb was
of AP design and contd a small chge of Type
91 Expl used in all Navy AP ammunition (pp
68–9);
Type 99 No 80 Mk 5 Bomb was of forged steel
and contd a large chge of Type 91 Expl (pp
69–70)
Type 2 No 8 Mk 5 and No 150 Mk 5 Bombs
were of AP design and used small chges of
Type 91 Expl (pp 70–1);
Type 98 No 7 Mk 6 Model 1 Bomb contd four
Thermite-filled electron fire pots. A central
channel filled with quick match ran thru the
length of the bomb. A BkPdr chge was located in
the circular recess of the nose piece (pp 71–72);
Type 98 No 7 Mk Model 2 Bomb contd a
central Thermite core surrounded by a solidified
kerosene, petrol, alcohol, soap mixture. A BkPdr
burster chge was located in the nose. A copper
tube contg quickmatch was located inside
Thermite tube (pp 73–4);
Type 1 No 7 Mk 6 Model 3 Modification Bomb
consisted of a sheet steel cylindrical body contg
182 cylindrical pellets arranged around a central
cardboard tube contg gray igniter mixture (Ba
nitrate 75, Al 24.5, oil 0.3 & moisture 0.2%);
The pellets (which contd Ba nitrate 35, ferric
oxide 28, Al 18 & synthetic rubber shavings
Brown Powder — *Kasshokuyaku* (Undercarbonized Black Powder). Known only from documents but was never recovered by US investigators. Use unknown (Ref 1, p 30)

**Brown Powder** (Navy). This name was assigned in Ref 1, p 32 to the mixture: Amm chlorate 51.5, Ba nitrate 34.5, TNM (Trinitronaphthalene) & oil 8.2 & wood pulp 5.0%. Substance dangerous to burn in large quantity. It was found as a booster in preparing bombs for demolition

**B-Fu**. Arm, Weapon

**Bullet. Dangan**

**Bunri yakutō. Separate-loading Ammunition**

**Burst. Haretsu**

**Burst. Sakuyaku**

**Bursting Charge. Sakuyaku**

**Buryoku. Military Force**

**Caliber. Kökel**

**Cannon. Taihō, Kahō, Kanōhō**

**Cap (explosive). Bakkan**

**Carbine. Kijū**

**Carlit.** Accdg to a Pamphlet of “The Japan Carlit Co, Ltd”, located before WWII at Chiyoda-kuki, Tokyo, *Carlit* was invented in 1917 by O. Carlson and manufd in Japan since 1919. It was an Amm perchlorate expl contg ferrosilicon & woodmeal. It was used in the same way and for the same purposes as Dynamites. Properties of Carlit are described in the Pamphlet

In Vol 2 of Encycl, p C68 is described a Japanese Army expl *Carlit or Karitto* consisting of Amm perchlorate 66, Si carbide 16, woodpulp & petroleum 6%. The same compn was given for *Karitto* in OPNAV 30-3M (1945), p 29, listed here as Ref 1. Its Navy name was *Type 88 Explosive* (qv) (See also under Karitto)

**Carriage (Arty). Hōga**
Cartridge (Fixed Ammo). Danyakuhō

Cartridge Bag. Yakunō

Cartridge Case. Yakkyō

Cast (Expl Filling). Chūiten (bakuhatsumu)

Cast Iron HE Projectile. Chutetsu ryūdan

Cavity (in Projectile). Shinkō

Chakatsuyaku. 2,4,6-Trinitrotoluene (TNT), also called Sanshōkī toruōru or Type 92 Explosive (Navy), H₃C₆C₄H₂(NO₂)₃; it yeI to buff expl, d (cryst) 1.65, mp 80.6°C; Brisanse by Sand Test – 48.00 g sand crushed; Expln Temp – decomp at 475°C in 5 sec; Impact Sensitivity, Bur Mines, 2kg wt 100cm², PicArsn App 14–15 inches; Power – 100% TNT, Rate of Deton 6900m/s at d 1.6. Used by the Japanese Navy straight in some 25 & 40mm Shells, while the Army used it in Grenades, Mines, Demolition Charges and a few Bombs

Also used in the following composite explosives: A(ko), Chanayaku, Chaoyaku, Chauyaku, Nigotanoyaku Mk 2, Otsu-B or Type A, Pentoriru, Seigata (Army) or Type 97H (Navy), Shoanyaku, Šōtoyaku, Tanoyaku, Type 92, Type 97H and others (Ref 1, p 26; Ref 5, p 363 & Ref 8, p 350)

Chakuhatsudan. Percussion Shell

Chakuhatsu ryūsandan. Percussion Shrapnel

Chakuhatsu shikan. Percussion Fuze, Impact Fuze

Chanayaku. A yeI castable composite expl consisting of TNT 70 & DNN (Dinitronaphthalene) 30%. Used for loading some Army Artillery Projs (Ref 1, p 26 & Ref 5, p 364)

Chaoyaku. A Lt-yel mixture of PA (Picric Acid) 75 & TNT 25%, which melts below 120°C and explodes at below 350°C, brisanse and detonation velocity are lower than those for PA and it is less sensitive. Press/loaded (or rarely cast) in some Army Bombs (Ref 1, p 26 & Ref 5, p 364)

Chaoyaku. A castable mixture of RDX (Cyclonite) 50 & TNT 50%, corresponding to Amer Cyclotol. Used for loading some Artillery Projectiles. Other RDX/TNT mixtures are listed under Nigotanoyaku Mk 2 (Ref 5, p 364)

Chikka (Army) or Chikka namari (Navy). Lead Azide (LA), Pb(N₃)₂, creamy white to buff or very light gray pdr; detonates w/o melting at ca 350°C; decomp by acetate soln. Most common initiator for detonators and fuzes, especially where a BkPdr relay is present (Ref 1, p 25 & Ref 5, p 364)

Chitai suru. Delay

Chūjō kayaku. Cordite

Chūkei. Relay

Chūki no. Hollow

Chūtai kikan. Recoil Mechanism

Chūtai sōchi. Recoil-operated

Coal. Sekitan

Coast Artillery (Weapons). Kaiganhō

Combat. Sentō

Combat Ammunition. Shōkōri danyaku

Combat Vehicles (Sentō jidōsha) and Their Weapons (Heiki). In the book of Tantum & Hoffschmidt (Ref 7, Section 4, pp 117–135) are described and illustrated the following items:

Tankette, Mod 2592 (1932) (pp 116–17)

Tankette, Mod 2594 (1934) (p 118)

Tankette, Mod 2597 (1937) (p 119)

Light Tank, Mod 2593 (1933) (p 120)

Light Tank, Mod 2595 (1935) (p 121)

Medium Tank, Mod 2589A (1929) (p 122)

Medium Tank, Mod 2589B (1929) (p 123)

Medium Tank, Mod 2594 (1934) (pp 124–25)

Medium Tank, Mod 2597 (1937) (Special) (p 126)

Medium Tank, Mod 2597 (1937) (p 127)

Amphibious Tank (p 128)
Combat Weapons are described in the book of W.H. Tantum IV & E.J. Hoffschmidt, Editors, entitled: “Second World War COMBAT VEHICLES Japanese”, Published by WE Inc., Old Greenwich, Connecticut, Vol 2 (1968), listed as Ref 7 at the end of this Section. Included among others are:

- **Infantry**: Small Arms Weapons, Pistols, Rifles and Machine Guns (pp 39–78)
- **Artillery**: Antitank, Antiaircraft, Light and Heavy Field Guns (pp 79–115a)
- **Combat Vehicles**: Tanks, Self-Propelled Guns, etc (pp 116–54)
- **Miscellaneous Weapons**: Rockets, Mortars, Mines, Grenades, Telescopes and Ammunition (pp 155–92)

**Combination Fuze.** Fukudo shikan

**Combustion.** Nenshō

**Combustion Chamber.** Nenshōshitsu

**Complete Round.** Danyakutō (Fixed Ammo)

**Component.** Kōseibunshi

**Cordite.** Chūjō kayaku

**Cyclonite or RDX.** Shouyaku

**Cyclotol (50/50–RDX/TNT).** Chauyaku

**Dagger.** Tanken

**Dainamaito.** Dynamite NG (Nitroglycerol) contg composite expls from buff to brown color. Accdg to documents there were diatomaceous earth contg, NC (Nitro- cellulose) contg — Gelatin Dynamites (See Geogel). “Faint Smoke” Gelatin Dynamite contd borax or salt. Nonfreezing Dynamites contd NGc (Nitroglycerol) or Dinitroglycerol. Ammonium Nitrate Dynamites contd AN (such as Akatsuki). There is also Semigelatine Dynamite, such as Enoke No 2 Dynamite (Ref 1, p 29). See paper of K. Sassa & I. Ito in Kögyo-Kayaku-Kyōkaishi 32, No 6 (Nov–Dec 1971) translated by Mrs Geti Saad at the BurMines in 1972 (See also under Additional References at the end of this Section)

**Danchaku.** Impact (of Projectile)

**Dandō.** Trajectory

**Dandōgaku.** Ballistics

**Dangan.** Projectile, Shell, Bullet

**Dantai.** Body (of Projectile or Bomb)

**Dantei.** Base of Projectile

**Dantei shikan.** Base Fuze; Tail Fuze

**Dantō.** Nose of Projectile or Bomb; Point of Projectile

**Dantō bakkan.** Point Detonating Fuze

**Danyaku.** Ammunition

**Danyakutō.** Cartridge; Complete Round

**Danyōku.** Fin of Bomb

**Denkayaku.** Booster Charge. See under Gaine

**Denki bakkan.** Electric Primer

**Denki raikan.** Electric Cap

**Denki tenkaçu.** Electric Blasting Machine; Exploder

**Deflagration.** Totunen; Bakunen
Delay. Chitai suru

Delayed-Action Bomb. Chidō bakudan

Delayed-Action Fuze; Delay Fuze. Chidō shinkan

Delayed-Action Mine. Chidō jirai

DEMOLITION CHARGES (Hakaiyaku) and EQUIPMENT (Sōgu)

Following information is given in Ref 1, pp 28–29 & 233–36 and in Ref 2, pp 30 & 252–55

Japanese equipment consisted of:

A. Safety Fuses and Safety Fuse Igniters. Several types of fuses with rate of burning of 30, 32 and 45 seconds per foot were used (p 234). The Igniters were: Trigger Type (Fig 381 on p 233); Pull Type (Fig 382 on p 234) and Type 99 Demolition Tube Igniter (p 233)

B. Blasting Caps. Three types of non-electric and four types of electric caps were known (p 234)

C. Detonating Cord, Type 97, described on p 234, condt a core of PETN (Shoeiyaku), surrounded by cotton, hemp and waterproof layers. Diameter 1/4 inch and Detonation Velocity 6000m/sec

D. Demolition Explosives consisted of the following:

a) Plastic Explosive Koshitsu (qv) (Army) consisted of RDX (Shouyaku) 80 & vegetable oil 20%, made into rolls 4 inches long, weighing 4 oz each. Issued for field use in three rolls packed in a paper package (Ref 1, pp 28 & 234)

b) Prepared Charges. Three main (standard) Army demolition blocks, 2x2x1-inch size were used: 1) PA (Shimose bakuyaku); 2) Haishokuyaku (qv) and 3) Nigonanyaku (Tanoyaku) (50/50–TNT/RDX) (pp 28–30 & 234 of Ref 1 and p 230 of Ref 2); 4) Combined Demolition Blocks – two, three or four TNT and PA blocks were provided with a hole to receive the detonator (p 235 of Ref 1); 5) Demolition Cans were rectangular made of zinc. Three sizes are described in Ref 1, p 234: 1-kg Can was filled with cast, wrapped PA; and 30-kg Can contd 30kg of PA charges, rounded or square

c) Substitute Army Demolition Explosives. The following are listed in Ref 1: Entoyaku (qv) (p 28), Enmayaku (qv) (p 28), Dainamaito (qv) (p 29), Shoanbakuyaku (qv) (p 29), Karito (qv) (p 29) and Ammonyaku (qv) (p 29)

E. Demolition Clocks. The following types are described and illustrated in Ref 2, pp 252–55:

a) Twenty-Four Hour Demolition Clock (p 253; Fig 191, upper)

b) Type 99 Long-Delay Demolition Clock (p 253; Fig 191, lower)

c) Seven and One-Half Day Demolition Clock (p 255, no Fig)

d) Type 92 Seven-Day Demolition Clock (p 255; Fig 192)

F. Demolition Block, shown in Fig 211, p 159 of Ref 1, consisted of a cast-iron cylinder (4¾x6 inches or 4¾x4¾ inches) conta blocks of PA. The body was mounted on a stick 21 inches long. When the device was fired from an Army Mortar, two Pull Igniters in the charge ignited a delay train of 7 seconds

DEMOLITION TUBES. Under this title are described in Ref 1, pp 236–37 several demolition items in the form of tubes, such as:

a) Bangalore Torpedo. The model described in Ref 1, p 236 looks different from Model 99 (1939) described in Ref 2, pp 215–16 and in Ref 7, p 186. This torpedo consisted of a steel tube with shoulders welded to both ends. One end was internally threaded to take the igniter locking collar while the other end was threaded externally to take the pointed nose cover. The igniter system consisted of two pull igniters screwed into an igniter holder which fitted into the igniter locking collar. The rings of the igniters were connected by lines to the lanyard holder. The igniters were simple match compn pull igniters, friction type, with a BKPb delay of 8 seconds, an initant, and a base edge of Tetryl. Each torpedo unit had outside diameter of 1-13/32 inches, total length 51 feet, weight 10 lbs, filled with 3 lbs of 36.4/63.6–TNT/RDX as Bursting Charge. To prepare the
torpedo for use, the storage caps were removed from the required number of tubes and were screwed end for end together. Then the pointed nose cover was screwed onto one end, while at the other end the igniter locking collar was screwed into it. After placing the torpedo on the ground or barbed wires, etc., the safety pin of igniter was removed and the firing lanyard pulled sharply (See Fig 162, p 216 of Ref 2 and description and Fig on p 186 of Ref 7).

The bangalores were used primarily for destruction of wire entanglements, but also to clear passages thru mine fields and destroy railroad tracks. They may be rigged as booby traps (Compare with US Models described in Vol 2 of Encyl, pp B16-B17)

b) **Bamboo Tube**, filled with explosive and fitted with a pull igniter is shown in Fig 386 on p 237 of Ref 1.

c) **Finned Bangalore Tube**, shown in Fig 387, p 237 of Ref 1 and in Fig 299, p 376 of Ref 2, was similar to a regular Bangalore, except that it was made of two pieces with fins attached to the after-body. This section was made to fit into the barrel of the Type 98 Projectile Discharger. A combination Instantaneous-Short Delay Nose Fuze was fitted into the tube. Diameter of tube 1-15/16 & length 78-3/8 inches; total weight 17 lbs, 14 oz.

d) **Type 99 Demolition Tube** was 44 inches long and weighed 35kg. The fuze was actuated by a pull cord and had 7 seconds delay (Ref 1, p 237).

e) **Obstacle Demolition Tube** was over 35 feet long and weighed 102kg. Its construction was similar to those above but the fuze was actuated by electrical means (Ref 1, p 237).

**Denkayaku.** Booster Charge. See under Gaine

**Denki bakkan.** Electric Primer

**Denki tengaku.** Electric Blasting Machine; Exploder

**Density.** Mitsudo

**Density of Loading.** Sōten hijū

**Depth Bomb; Depth Charge** (Bakurai). Accordg to Ref 1, p 61, the depth chge of Japanese Navy, like its mines but unlike its torpedoes, are obsolete in design.

The following types were used during WWII:

Type 88 Depth Charge was a metallic barrel 30.5 inches long and 17.7 inches in diam. This size was standard for all known depth charges. It was filled with 327 lbs of cast FA (Shimose). It had “filling end” and “pistol end”. The hydrostatic pistol was operated by water pressure and could be set to fire at depths of 82 or 148 ft by varying the opposing spring pressure (p 62 of Ref 1).

Type 91 Model 1 Modification 1 Depth Charge was similar to above and contd 220 lbs of Type 88 (qv) Expl. Operated at depths of 82 or 164 ft of water (p 62);

Type 95 Depth Charge (Fig 33) was the regular issue charge until the appearance of the Type 2. It contd 220 lbs of Type 88 Expl. Slow ships dropped the chge to slow down its rate of descent so that the ship could escape the danger area before the chge exploded. The hydrostatic pistol (Fig 34 on p 62) could be set for a depth of 98 ft with parachute or 197 ft w/o one;

Type 95 Mod 1 contd 325 lbs of Type 97 or 98 Expl (p 62);

Type 2 Depth Charge (Fig 35 on p 63) was loaded with 240 lbs of Type 1 Expl. This depth chge and its hydrostatic pistol (Fig 36 on p 63) were almost direct copies of British models. The pistol could be set for depths of 98, 197, 292, 390 and 480 feet (pp 62-3);

Type 2 Mod 1 Depth Charge contd 357 lbs of Type 98 Expl or Type 97 as an alternate filling (p 63);

Type 2 Mod 2 Depth Charge was filled with either Type 1 or 4 Expl (p 63);

50 Kilogram Army Depth Charge (Fig 37 on p 64) was 20 inches long, 13 inches in diameter and carried a chge of approx 75 lbs of TNT. The firing mechanism (instead of hydrostatic pistol) was housed in a well at one end of the case. It was probably used on “suicide boats” (qv);

Type 3 120 Kilogram Army Depth Charge (Experimental) was 24¼ inches long and 15¼ inches in diameter, filled with 200 lbs of HE (no data). It was fired by a pull igniter instead...
of a hydrostatic pistol. The charge (Fig 38 on p 63 of Ref 1) was rigged on Army “suicide crash boats”. The charge was also used as a Mine and could be fired electrically by means of Demolition Charges planted beside it (Ref 1, p 223)

**Destruction.** Hakai

**Detonating Cap.** Raikan

**Detonating Charge; Detonating Explosive.** Kibakusai

**Detonating Cord.** Dōbakusaku

**Detonating Fuze.** Shinkan; Bakkan

**Detonation.** Kibaku; Bakuhatsu

**Detonator.** Kibakuyaku

**Device.** Dōgu, Sōchi, Shikake

**Dōbakusaku.** Detonating Cord

**Dōkaraikan.** Nonelectric Cap

**Dōkasaku.** Safety Fuse; Powder Train

**Dōkasen.** Explosive Train; Powder Train

**Dokuen.** Poisonous Smoke

**Dokugasu.** Poisonous Gas

**Double-Base Propellant.** See Nitoroguriserin meynayaku under PROPELLANTS

**Drill Ammunition.** Giseidan

**Drill Bomb.** Gibakudan

**Drop Test.** Rakka shiken

**Dud.** Fuhatsudan

**Dumdum Bullet.** Damudamu dan

**Dummy Ammunition.** Renshūyō danyaku

**Dummy Projectile.** Giseidan

**Duralumin.** Jururumin

**Dynamite.** Dainamaito (qv)

“E” (Explosive). A light-yellow castable explosive mixture of Trinitroanisole 60 & AN 40%; cast density 1.60; Power by Ballistic Mortar 108% (PA=100%). No info about its uses (Ref 5, p 364)

**Eikadan.** Time Shell

**Eika shikan.** Time Fuze

**Eikōdan.** Tracer Bullet

**Eikō danyaku.** Tracer Ammunition

**Ekka.** Liquid Fire

**Electric Current.** Denryū

**Electric Firing Device Type 3 for Proximity Fuzing of Bombs is described in Ref 2, pp 188–89 with Fig 137

**Electric Primer.** Denki bakkan

**Electric Wire.** Densen

**Electron.** Denshi

**Electron Bomb.** Elekutoron shōdan

**Emmaku.** Smoke Screen

**Emmaku hōshaki.** Smoke Projector

**Enka.** Signal Rocket

**Enkapikurin.** Chloropicrin

**Enka shingō.** Pyrotechnic Signal

**Ennyaku.** A light-yellow explosive mixture of K chlorate 80, MNT (Mononitrotoluene) 15 & castor oil 5%; extremely sensitive to mechanical action. It was used by the Army as Substitute
Demolition Charge and as Substitute Main Charge for Hand Grenades and Mortars (Ref 1, p 28 & Ref 5, p 364)

Enoki Dynamite No 2. See under Dainamaito

Enshūyō bakudan. Practice Bomb

Enshūyō dangan. Practice Shell

Enshūyō jirai. Practice Land Mine

Enshūyō kirai. Practice Sea Mine

Enshūyō tekidan. Practice Grenade

Entai. Trench; Shelter

Entoyaku. A light-yellow expl mixture of K chlorideate 80, DNT (Dinitrotoluene) 16 & castor oil 4%; used for the same items as Ennayaku (Ref 1, p 28)

Equipment. Sōgu

Erosion (of Gun Barrel). Fushoku

Exploder. Bakuhaki

Explosion. Haretta; Bakuhanatsu

Explosive. Bakukatsubutsu; Bakuyaku. See under individual name: Akatsuki, Angayaku, Chakatsuyaku, etc and under “Unknown Name Explosives”

Explosive Bullet. Bakuretsu shōjūdan

Explosive Coal. Food Can and Toothpaste. See under Sabotage Devices

Explosive Train (in Fuze). Dōkasen

Exterior Ballistcs. Tōgai dandōgaku

Factory. Kōjō

Field Gun; Field Piece. Yasenhō

Filler. Sakuyaku

Finned Bangalore Torpedo. See under DEMOLITION TUBES

Fireworks. Enka

Firing (Discharge of a Firearm). Hassha

Firing Devices. See under Igniters or Firing Devices

Firing Pin (Striker). Gekishin

Firing Pin of Fuze. Kakki

Fixed Ammunition or Fixed Round. Kanzen yakutō

Fixed Gun. Koteijū

FLARES (Shōmei) and Flare Bombs (Shomeidan)

The following Japanese Army Flares (excluding Parachute Types) are described in Ref 1, pp 238–43:

Flares for 50-mm Type 10th Year Flare Discharger (p 238 & Fig 388 on p 239) had heavy cardboard cylindrical bodies, 6-1/8 inches long, filled with various color flare compositions. A propint container was attached at the base; 50-mm 10th Year Type “A” Flare (p 238 & Fig 389 on p 239) was a heavy cardboard cylinder 6¼ inches long, which contd a filling of an incendiary type which burned with an intense white flame. It was fired from the Type 89 Grenade Discharger. A cylindrical proplnt container was attached at the base; 81-mm Mortar Signal Flare. Its Green Type was a light iron cylindrical case 3 inches in diam & 7-3/8 inches long loaded with green flare compn. Attached to the base was a proplnt container (p 239); 81-mm Mortar Parachute Smoke Flare was similar to above but contd smoke producing compn (p 239); Flare Signal Cartridges were cardboard cylinders 3¾"x1/16" filled with green, red or white flame producing compns. They were fired from the Type 97 Very Pistol (p 239); Flare Signal Rocket Mk 1 Flare consisted of cardboard cylinder attached to a stick, to the end of which was fixed a 6½ ft length of rope. A pull tape at the base of the rocket case
covered a length of fuse which was lit to fire the rocket (p 240);
Type 93 4-cm Signal Flare was in two types:
a) trailing White Stars and b) Green Comet.
They are described on p 240 and illustrated in
Fig 391;
Safety Fuse Flare, 3¾ inches long and 1¼
inches in diam, was housed in a cylindrical paper
container closed at the base with a wooden plug
to which a short length of safety fuse was
attached. Above the plug were small expelling &
ignition chgs which blew the flare and its
red paper parachute out of the container. The
flare burned for 23 seconds. It could be fired
from the 30-mm Rifle Grenade Discharger (p
240 & Fig 392)
The following Japanese Army Parachute
Flares are described in Ref 1, pp 250–51 and
in Ref 2, pp 22–5 with Figs 18, 19 & 20:
Type 90 Small Model Parachute Flare was a
sheet-steel cylinder (with conical nose) 26¾
inches long & 2-7/8 inches in diam contg 4
lbs of an Illuminating Compn of Ba chlorate
75.2 & gum 24.8% and a BkPdr ignition chge.
The flare burned for 2-2/3 minutes with an
intense greenish-white flame. A detailed
description is given on pp 22–3 with Fig 18;
Type 1 12-kg Parachute Flare was a sheet steel
cylindrical (with conical nose) case 37 inches
long & 4-3/16 inches in diam filled with 15
lbs of an illuminant which was ignited by BkPdr
chge. Compn of illuminant was as indicated
under illuminants in the text. The Illuminant
burned for 1½ to 3 mins with a greenish-white
flame. A detailed description is given on pp
24–5 with Fig 19;
Type 3 Parachute Flare was a sheet steel cylin-
drical (with rounded nose) case 42¾ inches long
& 6¾ inches in diam filled with 15 lbs of illu-
minant, such as described in the text under
I's and a BkPdr Igniter. A detailed description
is given on pp 25–6 with Fig 20
The following Japanese Navy Flares (ex-
cluding Parachute Types) are described in Ref
1, pp 241–43 and some of them in Ref 2, pp
98–100 and 106–08:
Navy Hand Signal Flare was cardboard tube 12¼
inches long filled with a flare mixture and fixed
on a wood handle. The flare was ignited by a
pull igniter wire which extended down the tube
(p 241, Fig 393 of Ref 1);
Illuminating Flare Bomb was of thin iron, conical
in shape with a hemispherical nose welded at
the bottom. Length 22¾ inches, diam at top
6¼ inches. Its filler was ignited by a pull
igniter, resulting in a brilliant greenish-yellow
light (p 241, Fig 394 of Ref 1);
Type 96 Floating Landing Flare was a sheet
metal flare bomb with four sheet metal fins
and a brass nose. Total length 16¾ inches,
diam 5 inches. Its flare mixt filler was ignited
by a pull igniter. When dropped from an air-
plane it floated in water while producing an
extremely bright light to signal landing spots
during the night (p 241, Fig 395 of Ref 1 and
p 98 of Ref 2 with Fig 74);
Navy Ground Signal Flares were cylindrical
cardboard containers 3¾ inches long 1¼ inches
in diam, filled with red, white or green compns.
Used to signal aircraft from the ground (p 242,
Fig 396 of Ref 1);
Type 94 Float Flare, called in Ref 2, p 107
Type 94 Float Light, was tinplate cylinder, 12
inches long & 2¾ inches in diam, with a hemi-
spherical nose. A lead in the nose and a buoy-
cyancy chamber at the base section kept the flare
straight in the water. The body had water
inlet holes in the nose and the side seated by
tear-off strip and a removable end cap. Before
dropping from a plane, the tear-strip and end
cap were removed to allow the water, after
impact, to enter thru the nose inlet holes, thus
wetting Ca carbide in the forward end. This
generated acetylene gas. Simultaneously, water
entered thru the side inlet hole to wet Ca
phosphide gas in the after end of the flare,
thus generating phosphine. When acetylene
passed thru the tail opening, it was ignited by
the phosphine which spontaneously inflames on
contact with the air (p 242, Fig 397 of Ref 1
and with more detailed illustrations on p 106
of Ref 2);
Type 94 Mod 1 Float Flare was a larger flare
than Type 94. Length 20¾ inches & diam 4¾
inches (p 243, Fig 398 of Ref 1);
Type 94 Mod 2 Float Flare was a smaller version
of Type 94. Length 9¾ inches & diam 2-3/8
inches (p 243, Fig 399 of Ref 1 and in Ref 2,
pp 108–09, Fig 81);
Type 0 Model 1 Float Flare was similar in
construction and operation to Type 94. Length
13¾ inches & diam 2-7/8 inches (p 243, Fig
400 of Ref 1 and in Ref 2, pp 108–09, Fig 81); Type 94 Experimental Float Light, described in Ref 2, p 107 and illustrated in Fig 80 of p 106, consisted of tin plate cylinder 21/4 inches long and 43/4 inches in diam. It contd Ca carbide & Ca phosphite. Its operation was exactly the same as the Type 94 Float Flare described above; Navy Float Ball was a large cardboard sphere 9 inches in diam with a dome top covering a paper-covered BkPdr Fuse and a dome bottom contg BkPdr chge in a celluloid cap. The main container carried a cluster of 12 flares attached to a large green tissue parachute. The flare cluster was bedded in a quantity of cottonseed which filled most of the lower half of the sphere. A pull wire permitted removal of the top dome and allowed access to the BkPdr fuse for ignition (Ref 1, p 247, Fig 409).

The following Japanese Navy Parachute Flares are described in Ref 1, pp 251–54 and Ref 2, pp 100–05 with Figs 76, 77, 78 & 79: 5-Kilogram Parachute Flare Model 2 Modification 1 consisted of a light sheet steel cylindrical body 25-1/8 inches long & 3-1/8 inches in diam, contg 5/8 lbs of Illuminant (qv under 1’s), a Pull Igniter and a Parachute. The unit burned for 1½ mins giving brilliant white light. Detailed description is on pp 100–01 with Fig 76 in Ref 2; Type 0 Parachute Flares Model 1 and Model 1 Modification 1, each consisted of sheet steel cylinders 42½ inches long & 6½ inches in diam, contg 66 lbs of Illuminant, a Fuze and a Parachute. They burned for 3-2/3 mins with a bright white light (Ref 1, p 252, Figs 421 & 422). Detailed description is on pp 102–03 with Fig 77 in Ref 2; Type 0 Parachute Flares Model 2 and Model 3, Modification 1 consisted of sheet steel cylindrical bodies 35½ & 39½ inches long and 6½ inches in diam each. Each contd 66 lbs of Illuminant, a Fuze and a Parachute. The flares shed a bright white light during 3-2/3 seconds (Ref 1, p 253, Figs 425 & 426). Detailed description is on pp 103–04 with Fig 78 in Ref 2; Experimental Model 11 Parachute Flares consisted of sheet steel cylindrical bodies 43½ inches long & 9½ inches in diam, provided with fins. One of the models had conical nose while the other had ogival nose. Each unit contd 68 lbs of Illuminant, a Fuze and a Parachute. The flare burned with bright white light during 4-2/3 mins (Ref 1, pp 253–54, Figs 427 & 428). Detailed description is on pp 105 & 107, with Fig 79 in Ref 2; Type 94 Float Light is described as Type 94 Float Flare in Ref 1, p 242 and also in this Section, under FLARES

Floating Flare (Fuhyō shōmei). See under FLARES

Float Lights, Type 94, Type 94 Experimental, Type 94 Model 2 and Type 0 Model 1, described in Ref 2 on pp 107–09 with Figs 80 & 81 as Float Lights are described here as “Float Flares” under FLARES

Floating Mine (Fuhyō suirai). See under MINE, SEA

Floating Smoke Items. See under “Smoke Floats”

Fosugen. Phosgene

Fragmentation Bomb (Sairetsu tōka bakudan). See under BOMB

Fragmentation Grenade (Sairetsu teryūdan). See under GRENADE

Frangible Grenade. Kaembin

Friction Primer. Monkan

Fuel Oil. Nenryōyu

Fuhatsudan. Dud

Fuhatsu suru. Misfire

Furyū suirai. Drifting Mine

Fuse (Kayōhen or Fyūzo). See under Demolition Equipment

Fuse, Safety. Dōkasaku

Fusetsu kirai. Anchored Mine

Fusatsu suirai. Submarine Mine. See MINE, SEA
FUZES (Shinkan)

Japanese Fuzes are divided into Bomb Fuzes and Projectile Fuzes which, in turn, are subdivided into Army and Navy Fuzes.

A. Bomb Fuzes are described in OPNAV 30-3M (1945), listed here as Ref 1, pp 80–88 (Army Bomb Fuzes) and pp 110–21 (Navy Bomb Fuzes). More complete description is given in TM 9-1985-4 (1953), listed here as Ref 3, pp 123–54 (Army Bomb Fuzes) and pp 155–87 (Navy Bomb Fuzes).

Individual fuzes of each service are generally interchangeable for use in bombs of that service, but not interchangeable for use in ordnance of the other service.

The following Army Bomb Fuzes are listed in Ref 2 on pages indicated below:

Type 93 Instantaneous—Short Delay A-2(a) (pp 123–24)
Type 12-Year Instantaneous A-2(b) (pp 124–25)
Type 99 Instantaneous—Short Delay A-2(c) (pp 126–27)
Type 1 Instantaneous A-2(d) (pp 127–28)
Type 92 Nose Fuze A-4(a) (p 129)
Fuzes for Type 3 Bomb A-6(a) and A-6(b) (pp 129–32)
Fuze for Type 2 Bomb A-7(a) (pp 132–33)
Type 4 Two-Second Delay Fuzes A-8(a) & A-8(b) (pp 134–36)
Type 12-Year Tail Fuze B-1(a) (pp 137–38)
Type 1 15-Second Delay Fuze B-1(b) (pp 138–39)
Type 92 Tail Fuze B-4(a) (p 140)
B-4(a) Tail Fuze (p 141)
Experimental 3.5-Second Delay Fuze B-7(a) (pp 141–43)
Type 4 Five-Second Delay Fuze B-8(a) (pp 143–44)
Type 1 Long-Delay Fuze C-3(a) (pp 144–47)
Aerial-Burst Fuzes D-1(a) & D-1(b) (pp 147–49)
Type 1 Aerial-Burst D-5(a) & Combination Fuze D-5(b) (pp 149–51)
Type 1 Antiwithdrawal Fuze E-1(a) (pp 151–53)
Remote Control Radio Fuze (p 153)

The following are Navy Bomb Fuzes described in Ref 2 on pp indicated below:

A-1(a), A-1(b), and A-1(e) Nose Fuzes (pp 154–55)
Type 97 Mk 2 Nose Fuze Model 2 A-3(a) (pp 156–57)
Type 1 Nose Fuze Model 2 A-3(b) (pp 157–59)
Type 2 Nose Initiator A-3(c) (pp 159–60)
Type 97 Mk 2 Nose Fuze Model 1 A-3(d) (pp 160–62)
Type 3 Nose Initiator A-3(e) (pp 162–63)
Type 2 No 50 Ordinary Bomb Model 1 Fuze A-3(f) (pp 163–64)
A-3(g) Bomb Fuze (pp 165–66)
A-5(a) Nose Fuze (pp 166–67)
Type 99 No 25 Ordinary Bomb Fuze B-2(a) (pp 168–69)
Type 99 No 80 Mk 5 Bomb Fuze B-2(b) (pp 169–70)
Type 15 Tail Fuzes Model 2 & Model 1 B-3(a) & B-3(b) (pp 170–72)
B-5(b) and B-5(c) Tail Fuzes (pp 172–75)
Type 97 Rail Initiator B-6(a) (p 176)
B-9(a) and B-10(a) Tail Fuzes (pp 177–79)
Type 99 Special Bomb Tail Fuze C-1(a) (pp 179–80)
Type 99 Special Bomb Nose Fuze C-2(a) (pp 181–82)
D-2(a), D-2(b) and D-2(c) Aerial Burst Tail Fuzes (pp 182–85)
D-3(a) Aerial Burst Nose Fuze (pp 186–87)
Fuze for Type 0 Parachute Flare, Model 1 D-4(a) (pp 187–88)
Type 3 Electric Firing Device for Proximity Fuze of Bombs (pp 188–89)


The following Army Projectile Fuzes are listed in Ref 3 on pp indicated below:

Type 93, Type 100, Type 2 and Type 2 Modified Small Instantaneous Fuzes (pp 391–94)
Type 4 Super Detonating Fuze (p 395)
Ho 301 Impact Fuze (p 396)
Type 88 Fuzes: Small Instantaneous, Short-Delay and Instantaneous (pp 397–400)
Types 90, 93 & 100 Instantaneous Short-Delay (pp 401–04)
Finned Bangalore Torpedo Fuze (pp 404–05)
Type 98 Interior Fuze (pp 405–06)
Type 89 Small Time Fuze (pp 406–07)
Type 89 Powder Time Fuze (pp 408–09)
Auxiliary Detonating Fuze (pp 409–10)
Type 3rd & 5th-Year Combination Powder Time and Impact Fuzes (pp 410–12)
Type 100 Mechanical Time & Impact Fuze (pp 412–13)
Type 2 Combination Powder Time & Impact Fuze (pp 414–15)
Type 94 Small Delay Base Fuze (pp 415–16)
Type 92 Small Short Delay Base Fuze (pp 416–17)
Small Mk 2 & Mk 1 Base Detonating Fuze (pp 417–19)
Medium Mk I Impact Base Fuze (p 420)
Type 88 Small (Howitzer-Mortar) Base Fuze (pp 421–22)
Type 88 Small (Gun) Base Fuze (p 422)
Type 95 Large Mk 2 Mod I Base Fuze (pp 423–24)
Type 95 Medium Base Fuze (pp 425–26)

**Fuze Cap.** Shinkambō

**Fuze Detonator.** Shikan no kibakuzai

**Kyūzo.** Fuse

**GAINES** (Fukutō) and **BOOSTERS** (Denkayaku)

Generally speaking Gaine is a container (sheath) for a Booster. More detailed info about Boosters is given in Vol 2 of Encycl, pp B243-R to B246-R and about Gaines in Vol 6, pp G6-L to G7-L.

Japanese Gaines are divided into Army Gaines (Ref 1, pp 88–9 and Ref 2, p 190) and Navy Gaines (Ref 1, pp 121–22 with Table and Ref 2, pp 191–92).

Accdg to Ref 1, p 88 and Ref 2, p 190, Japanese Army Fuzes utilized a characteristic type of Gaine which differed greatly from that used in Navy Fuzes. Gaines used in Army Bombs are shown in Fig 107, p 88 of Ref 1 and in Fig 138 of Ref 2. They were of brass construction and were screwed into the Fuze.

There were three Types, of which Types I & II were used in the Nose, while Type II was used in the Tail Fuzes. Types I and III were ignited by the flash from the Primer in the Fuze, while Type II Gaine was pierced directly with the firing pin (because it was used in Fuzes having no internal Primer).

Army Boosters were either contained in a brass cup threaded into the Fuze around the Gaine, or in a paper cylinder housed in the main expl chge of the Bomb.

Accdg to Ref 1, p 25, PA under the name Oshokayaku was the main Booster Charge, while Tetryl (Meiayaku) and RDX (Shouyaku) were used in Sub-Boosters (p 26).

Accdg to Ref 1, p 121 and Ref 2; p 191, Japanese Navy Fuzes utilized a characteristic Gaine which incorporated within itself the entire expl train. The Gaines were similar in external appearance but varied in size and internal construction. They were generally made of brass, cadmium plated and lacquered. More recently some Gaines were made of steel.

A Table listing Navy Gaines used during WWII is given in Ref 1, p 121, while on p 122 are given external views of six Gaines. PA (Oshokayaku) was used as a main Booster Charge (p 31), Tetryl (Meiayaku) as Sub-Booster, and as Auxiliary Booster 70/30 TANAnisole/HDNPPhA (H2 Kongo–H2 Mixture), abbr for Type 98 Expl (p 31).

Following Navy Gaines are listed and illustrated in Ref 2 on pages indicated:

Type 97 Land Bomb Gaine “A” and Type 99 Ordinary Bomb Gaine “A”; length 4½ inches, depth 1½ inches. Explosive Train: Primer compn in upper plug; BkPdr delay (0.03 sec) over flash pdr relay in middle plug; Lead Azide detonator over Tetryl in lower plug and PA Booster (p 192 & Fig 140)

Type 97 Land Bomb Gaine “B” and Type 99 Ordinary Bomb Gaine “C”. Same size and Expl Trains, but their delays were 0.2 & 0.1 sec, respectively (p 193, Fig 141)

Type 15 Ordinary Bomb Gaine. Same size and Expl Train as above, but its delay was selective from fractional to 1.5 secs (p 194, Fig 142)

Type 92 Land Bomb Gaine Modification 2. Same size but it was Instantaneous and its Expl Train was: MF (Mercuric Fulminate) over Tetryl in an inverted copper cup in upper plug; no delay chge; Tetryl pellet in lower plug and PA Booster (p 195, Fig 143)

Type 96 Land Bomb Gaine. Same size as above, and Expl Train same as in Types 97, 99 & 15 with delay chge absent (p 196, Fig 144)

Type 4 Gaine for Skipping Bomb. Same size as above with Expl Train: Primer compn in upper plug; circular BkPdr Delay Train (10 to 11 secs) and Relay in middle plug; Primer compn over LA Detonator over Tetryl pellet in lower plug; PA Booster (p 197, Fig 145)

Type 1 Mk 2 Bomb Gaine “A”. Same size and
Train as above, but delay was 3.5±1.0 sec (p 198, Fig 146)

Type 0 Mk 5 Bomb Gaine, length 6½ & depth 1½ inches. Same Expl Train as above but delay was 0.2 sec (p 199, Fig 147)

Electric Bomb Gaine, Instantaneous, length 5-3/8 & depth 1-3/8 inches. Expl Train: The Electric Blasting Cap fitted into hollowed-out cavity in the PA Booster. Leads from the cap ran up to a female plug which was attached by a special adapter to the standard Gaine body (p 199, Fig 148)

Type 2 Small Model Bomb Gaine Model 1, Instantaneous. Length 3, depth 13/16 inches. Expl Train: MF cap over Tetryl in upper plug; PA Booster (p 200, Fig 149)

Practice Bomb Gaine, Instantaneous. Length 3-1/8, depth 11/16 inches. Expl Train: MF in upper plug; Tetryl Booster (p 200, Fig 150) (See also Magazines)

Gasu. Gas, Chemical Agent
Gasubakudan. Gas Bomb
Gasudan. Gas Shell, Chemical Shell
Gasu tekidan. Gas Grenade
Gekishin or Gekkei. Firing Pin
Glide Bomb. Kakku bakudan
Glycerin. Guriserin
Gomu. Rubber
Graphite. Sekiboku
Grapnel. Hikkake-ikari

GRENADERS (Tekidan). There are Hand Grenades (Teryūdan), Rifle Grenades (Jūyō tekidan) and Mortar Grenades (Kyūhō tekidan)

Accdg to Ref 2, p 225, the Japanese Armed Forces developed both before and during WWII a fairly extensive line of Grenades. This type of Ordnance primarily used by ground Forces was developed by the Army, but was also used by the Navy Ground Defense Units. Many Grenades were improvised and some of them, such as of glass construction were dangerous to use. Rifle (as well as Mortar) Grenades were often modified Hand Grenades.

The following Grenades are described in Refs 1 & 2 on pages indicated under each item:

Type 91 Hand, Mortar or Rifle Grenade had a cast iron cylindrical body 3¼ inches long & 2 inches in diam. It had 50 serrated segments and contd 65g of powdered TNT. Its delay was 3 secs (Ref 1, p 195 & Ref 2, p 225 with Fig 169 on p 224)

Type 97 Hand Grenade had a cylindrical, serrated, cast iron or aluminum body, 4 by 2 inches, contg powdered TNT. It was similar to Type 91 except that the base of Type 97 was solid and could not take a propelling charge which is required for conversion to Rifle or Mortar Type. Therefore, it could only be used as a Hand Grenade Delay 4–5 seconds (Ref 1, p 195 & Ref 2, p 226 with Fig 170)

Type 99 Hand Grenade had a cast-steel cylindrical body 3½ by 2 inches with smooth surface, filled with cast PA. Delay 4–5 secs (Ref 2, p 227 with Fig 171). In Ref 1, pp 195–96, this Grenade was subdivided into Type “A” (Kiska Type) (Fig 299) and Type “B” (Fig 300)

Type 4 Pottery Hand Grenade had spherical terra cotta body 3 inches in diam with wall 7/16 inch thick. It contd 3.5 oz of Type 88 Explosive and had 4–5 secs delay (Ref 2, p 228 with Fig 172 & Ref 1, p 197)

Type 23 Full Type Hand Grenade had a cylindrical cast-steel body 3½ by 2 inches, with five transverse depressions, used instead of serrations. It contd 39.5g of granular TNT and had a 5½-sec delay (Ref 2, p 229 with Fig 173 & Ref 1, p 196)

Type 98 Stick Hand Grenade had cylindrical, cast steel body, 7½ inches long and 1-5/16 inches in diam, with smooth surface. It contd 3 oz of cast PA (Shimose) and had a 4–5-sec delay (Ref 2, p 230 with Fig 174 & Ref 1, p 196)

Type 3 Conical Antitank Hand Grenade, called in Ref 1, p 196, "Type 3 Conical Hand-Thrown Mine", existed in "large" and "small" sizes. The basic principles of construction were the same in both cases, but variations occurred in measurements, weights and the expl charges. Each Grenade consisted of a cone-shaped expl chge (Type 94 for large and Pentolite for small), a metal cone and a wooden ring base, all contd in a
silk bag. A fuze was inserted and a grass or hemp tail attached to the apex of the chge. The expl chge was cast in the form of a truncated cone, thus making the Grenade Hollow (Shaped)-Charge Type. It was thrown against tanks.

Large type had body 6% inches long and diam at the base 4-3/8 inches, while smaller type had 5-7/8 & 4 inches, respectively. Length of the tail was 14 inches (Ref 2, pp 230–32 with Fig 175 and Ref 1, p 196)

Sling Hand Grenade had a cast steel cylindrical body which had a metal ring attached to its tapered wooden base plug covered with reinforcing cloth. Overall length 5-7/8, diam 1-13/16, filling 1 oz of TNT and delay 4 or 5 secs. Its fuze was friction-igniter type located in the nose. Grenade could be thrown either by hand or by a sling attached to the base ring (Ref 2, p 232 with Fig 176 & Ref 1, p 198)

1/2-kg Incendiary Hand or Mortar Grenade had brass cylindrical body, 5.6 inches long & 2.0 inches diam filled with WP (white phosphorus) and fuze. For projecting it from Mortar, Model 89, a steel propellant container was screwed into the base of the body. Delay was 4 to 5 secs. (Ref 2, pp 233–34 with Fig 177)

Incendiary Stick Grenade had light steel cylindrical body with hemispherical ends and wooden handle threaded into the base. Length of body 13.2 inches, diam 2.1 inches and length of handle 5.3 inches. The body was filled with 41 rubber pellets, each impregnated with a soin of WP in carbon disulfide. The pellets were scattered by means of a small central burster chge. It is possible that the Grenade was filled sometimes with a SP smoke filling (Ref 2, pp 234–35 with Fig 178 and Ref 1, p 200)

Molotov Cocktail, called “Frangible Incendiary Grenade” in Ref 1, p 200. Its body consisted of a Japanese beer bottle into the top of which was tightly fitted an “all-way” action fuze. The bottle, 11% inches long & 2-1/3 inches in diam was filled with ca 12 oz of an inflammable benzene-type liquid (Ref 2, pp 235–36 with Fig 179)

Phosphorus-Ignited Molotov Cocktail used a bottle 9% inches long & 2-1/8 inches diam, which contd 9.5 oz of 15% soin of polymethylmethacrylate in benzene and sealed with a crown type cap. An adjustable rubber harness held a flat circular glass igniter capsule, contg 1 oz of RP (red phosphorus) to the base. The capsule was well protected in the carton (which held the Grenade) by several layers of cardboard and sawdust. Before launching the bottle, a rip cord was pulled to open the carton and the igniter capsule was secured to the base of the bottle by the rubber harness. When the bottle was thrown with force against a hard target it scattered and the friction between the pieces of RP caused instant ignition of inflammable material. The viscous nature of the filling prevented undue splashing and insured adherence to the target (Ref 2, pp 236–38 with Fig 180 & Ref 1, p 201)

Hydrocyanic Acid (Seison) Frangible Grenades were round glass bowls, 3.9 inches in diam contg 12.2 fl oz of Cu-stabilized ca 80% strong HCNaq soln which acts as a very strong systemic poison. There were two types differing in minor details (Ref 2, pp 238–39 with Fig 181 & Ref 1, p 201)

Frangible Smoke Grenade consisted of a clear glass flat bottomed round flask 2.6 inches in diam, contg 4.1 oz of titanium and silicon tetrachloride. It was closed with rubber stopper and crown cap. Being thrown against a hard object, the bottle broke, releasing the liquid which in contact with air produced a smoke screen (Ref 1, p 201 with Fig 315 and Ref 2, p 239)

40-mm Hollow-Charge Rifle Grenade was light metal cylinder, 1.58 inches in diam, with a semiconical shaped head. Overall length 7.98 inches. It was made in two parts, threaded together and fitted with a ballistics cap and cone to give a hollow-charge effect. The forward part contd 3.81 oz of 50/50–RDX/TNT cast around the cone, while the after part had a rifled collar near the base and contd the Fuze and Expl Train. The Grenade was fired from a cup launcher attached to the standard 6.5-mm rifle (Ref 2, pp 239–40 with Fig 182)

30-mm Hollow-Charge Rifle Grenade was a smaller version of the above: overall length 6.25 inches, diam 1.18 inches and wt of HE chge 1.75 oz (Ref 2, p 241)

Model 3 Modification 1 Rifle Grenade was similar to the Type 99 Hand Grenade (known as “Kiska”), described above in Ref 2, p 227) with a tail assembly added. It contd 3 oz of TNT as Main Charge and was intended to be
fired from the spigot type Grenade Launcher (Ref 2, pp 241–42 with Fig 183)

Smoke Rifle Grenade had a light-metal cylindrical body, 2 inches in diam with fins. It contd a mixture of hexachloroethane 56.2, Zn pdr 27.6, Zn chloride 2.9 & Zn oxide 13.4% (adds to 100.1%). It was used with a special adapter which fitted over the end of a rifle barrel (Ref 2, pp 243–44 with Fig 184)

Small Incendiary Rifle Grenade had a light-metal cylindrical body 1-7/8 inches in diam filled with WP (white phosphorus). It was launched from a spigot type Rifle Grenade Launcher (Ref 2, pp 244–45 with Fig 185)

Improvised Grenades. Accdg to Ref 1, p 202 and Ref 2, p 225 a large variety of Grenades was made from such items as small ammunition, small bombs & shells, pipes, paper and wood. This was probably due to a shortage of materials. Most improvised items were ineffective and dangerous to use

The following improvised Grenades are listed in Ref 1, pp 202–03:

a) Gas pipe, closed at both ends, filled with an expl and fitted with a .22 cartridge case and safety fuse as igniter
b) 25-mm shell case filled with an expl and fitted with a pull igniter or a length of fuse
c) Heavy paper or braided cord container filled with an expl and fitted with fuse
d) 1/2-kg Amry bombs removed from their containers, armed and thrown by hand (Fig 318)

e) A very effective conical Grenade was constructed from the tail section of the Navy 30-kg Practice Bomb. The tail cone was removed and the space around the burster tube was filled with granular PA. Then a grass or hemp cord was attached and the B-6(a) Practice Bomb Tail Fuze fitted to the cone. After arming the Fuze by hand, the bomb was thrown (Fig 319 on p 203)

The following Grenades are described and excellently illustrated in the book of W.H. Tantum IV and E.J. Hoffschmidt, “Second World War Japanese Combat Weapons”, Vol 2 (1968), WE Inc, Old Greenwich, Conn 06870, listed as Ref 7:

Hand Grenade Model 91 (1931) (p 173 with 2 Figs)
Hand Grenade Model 97 (1937) (p 174 with 2 Figs)
Hand Grenade Model 99 (1939) (Kiska) (p 175

with Fig)
Conical Antitank Hollow-Charge Hand Grenade (p 176 with 2 Figs)
Pottery Hand Grenade (p 177 with 3 Figs)
High Explosive Rifle Grenade Model 3 (p 177 with Fig)
Hand Stick Grenade, High Explosive (p 178 with 2 Figs)
Hand Stick Grenade, Incendiary (p 178 with 2 Figs)

Grenade Discharger (Tekidantō) and Grenade Launcher (Rifle) (Tekidanji). A Grenade can be launched by hand, by rifle, by mortar or by special Grenade Launchers or Dischargers. These items were usually modified rifles or mortars. There were also special adapters which fitted barrels of rifles. Most Rifle Grenades could be fired directly from rifles

The following special devices for launching grenades are listed in Ref 1:

50-mm Type 10th Year Grenade Discharger was the forerunner of the “Knee Mortar” (p 158)
Type 89 Grenade Discharger, known as the “Knee Mortar” had a rifled bore and a provision for varying the range with fixed angle of elevation (pp 158–59)

The following devices are described in the book of Tantum & Hoffschmidt, listed as Ref 7:

50-mm Grenade Discharger Model 10 (1921) is the same as listed but not described above and in Ref 1, p 158. It was smooth-bore and weighed 5½ lbs. It could be carried by one soldier. It was used to discharge Model 91 Hand Grenade pyrotechnic signals and smoke shells. Range 175 yds (p 155 with Fig)
50-mm Grenade Discharger Model 89 (1929) was rifled-bore and weighed 10 lbs. Range for Model 91 Hand Grenade was 200 yds. It was also used for pyrotechnic signals, incendiary shells, etc (pp 156–57 with several Figs)
Rifle Grenade Launchers. The following types are described and illustrated on p 179 of Ref 7:
1) Type 2 or Cup Type Launcher which was patterned after the German Launcher fitted over the front sight of the rifle and had a short rifled barrel (See Fig). It fired both the 30-mm and 40-mm Hollow Charge Rifle Grenades, which could penetrate 3-7/8 inches of mild steel plate
2) Type 100 or Kiska Type Launcher, shown in
Fig, consisted of a tube which could be attached to either 6.5-mm or 7.7-mm Rifle. Ordinary ball ammo was used to launch the Grenade (the expanding gas from the fired cartridge was utilized to expel the grenade from the launcher), a feature which enabled the rifle to be carried with the launcher attached and ready for use as either a rifle or as a Grenade Launcher. The Type 99 Kiska Grenade was the only type used with the Launcher. Maximum range was 100 yds (p 179)

3) Spigot Type Grenade Launcher consisted of a rifled barrel threaded to an adapter. It was attached to either the 6.5-mm or 7.7-mm Rifle at the rear of the front sight. It was used to fire Type 91, Type 3 HE, and several types of Smoke and Incendiary Grenades (pp 179-80 with 3 Figs)

Gun (Hō) and Howitzer (Ryūdampō). See under ARTILLERY AND ITS WEAPONS

Gun Barrel. Hōshin

Gunpowder. Menkayaku

Gunpowder. Kayaku or Black Powder. Koku-shokuyaku, also called Yuennaku (Nonsmokeless Powder)

Gyokei suirai or Gyorai. Torpedo

Gyorai. Torpedo

Gyorai jitsuyō tobu. Torpedo Warhead

H2 Kongo (H2 Mixture) or Type 98 (Explosive). Lemon-yel expl comps consisting of TNAnS (Trinitroanisole) 60-70 & HNDPhA (Hexanitrodiphenylamine) 40-30%

The 60/40 mixt was cast-loaded to a density of 1.65; its mp 65-70°, Expln Temp 264° & higher; Brisance, by Copper Cylinder Crusher Test 100% (PA=100%); Power by Ballistic Mortar 96% (PA=100); Deton Velocity by Dautriche Method 7050m/sec; Impact Sensitivity with 15kg Weight 14cm (max wt for no explns); Friction Sensitivity 60kg (max pressure between two rubbing surfaces for no explns). Used, under the name Type 98, in Navy Bombs, Depth Charges, Sea Mines and Torpedoes in place of PA (Picric Acid)

The 70/30 mixt had Expln Temp 237-46°; Brisance by Sand Test 102% TNT; Power by Ballistic Mortar 109% TNT; Rifle Bullet Test 20% detonations; and Impact Sensitivity by PicArsn Test 12.5 inches (TNT 14). It was reported to be toxic, not very stable, and re-active with metals in the presence of moisture. Used pressed by Army and Navy as an Auxiliary Booster in some Bombs and Shells (Ref 1, pp 31 & 32 and Ref 5, p 365)

Haensosan bakuyaku. See Type 88 (Ko)

Haishokuyaku (Gray Powder). A dark-gray, non-toxic expl compn: No 1 Mixture consisted of Amm Perchlorate 76.9, RDX 17.0, silicon carbide 1.3 & paraffin 4.8%; relative Power & Brisance greater than for PA, but it is less sensitive to impact and friction. Used in Army Standard Demolition Blocks; No 2 Mixture consisted of Amm Perchlorate 48, GuN (Guandine Nitrate) 20, RDX 25 & paraffin 5%. It was a commercial mining expl, and also used in Army Demolition Blocks (Ref 1, p 28 & Ref 5, pp 365-66)

Hakaitō. Bangalore Torpedo. See under DE-MOLITION TUBES

Hakai yaku. Demolition Charge

Hakō bakudan. Armor-Piercing Bomb

Hakōdan. AP Projectile

Hakō ryūdan. AP-HE Projectile

Hakū or Ryūkyu. Mortar

Hakugekine. Trench Mortar; Infantry Mortar

Hakurin. White Phosphorus; WP

Hand Grenade. Teryūdan. See under GRENADE

Haretzu. Burst, Explosion

Hasai bōryaku. Sabotage

Hassha. Discharge (of a Firearm)
Hashadani. Projectile

Hatsuen or Hatsuendo. Smoke Bomb or Shell

Hatsuendo. Smoke Grenade

Hatsumoto. Smoke Pot; Smoke Candle

Hatsuendo. Smoke Agent

HE (High Explosive) Projectile. Ryūdan

Heavy Artillery. Jūhōhei

Heavy Gun. Jūhō

Heavy Machine Gun. Juki

Heiki. Ordnance, Weapon, Ammuniiton

Heikibu. Ordnance Department (Japan)

Heiki seisōsha. Ordnance Factory

Heikishō. Ordnance Depot (Japan)

Heineyaku. Trinitrophenetole (TNPhnt) or Ethyl Picate, C₂H₅.O.C₆H₂(NO₂)₃; mw 257.16, N 16.34%; yellowish crystals, mp 78; less powerful, less brisant and less sensitive to impact than TNaNisole; called Type 91 Expl by the Japanese Navy. Deton Vel of TNPhnt 6880 m/sec vs 7640 for TNaNisole [Blatt, OARDC Rpt 1944 (1944)]

Accdg to Ref 3, p 321, Heineyaku was used as an alternate HE filler (instead of TNT) in Type 90 7-cm (75-mm) HE Long-Pointed Projectile and in Type 94 7-cm (75-mm) High Explosive Projectile (Not listed in Ref 1)

It is listed as Keineyaku in Ref 5, p 366 and is stated that it can be used as a Bursting Charge in Projectiles as a Booster Charge, either alone or in mixtures with PA

Heishi. Soldier

Herikoputā. Helicopter

High Explosive. Kökyū bakuyaku

High Explosive Shell. Kökyū ryūdan or Jiraidan

Hikoki. Airplane

Hikōsen. Airship; Dirigible

Hō. Gun; Artillery

Hōbi. Breech (Arty)

Hōdan. Shell

Hōhei. Infantry

Hōhei. Artillery

Hōkō. Muzzle (Arty)

Hollow (Shaped) Charge [Ta(dan)] Japanese Ammunition (Senkōryūdan)

In the paper by A.J. Dere, “Adaptation of the Hollow Charge Principle of Explosives”, published in Ordnance Sergeant, Vol 10, p 13 (Oct 1945), it was stated that Japanese used during WWII, the following Hollow Charge (HoC) items:

75-mm HoC Projectile for Type 41 Regimental or Mountain Gun

HoC Rifle Grenade, diam 2.7 inches, overall length 7.08 inches, which could be fired from a 6.5-mm Infantry Rifle, was modified and several of them were enclosed in a large bomb casing. This ensemble could be dropped on airfields to damage grounded aircraft

Accdg to TM 9-1985-4, listed here as Ref 2, the Army Type 2 1/3-kg Cluster Bomb conté Hollow Charge of 50/50-RDX/TNT (pp 29–30 with Fig 25). There were also Type 3 Conical Antitank Hand Grenade with HoC (pp 230–31 with Fig 175) and 30 & 40-mm HoC Rifle Grenades (pp 239–41 with Fig 182)

In TM 9-1985-5, listed here as Ref 3, the following HoC projectiles are described:

Type 3 7-cm (70-mm) Army HoC Projectile (pp 305–06 with Fig 238)

Type 2 7-cm (75-mm) Army HoC Projectile (pp 328–29 with Fig 257)

In the book of W.H. Tantum IV & E.J. Hoffschmidt, “Japanese Combat Weapons”, WE Inc, Old Greenwich, Conn (1968), listed here as Ref 7, are described the following HoC items:
a) Conical Antitank HoC Hand Grenade (p 176 with Fig). Their construction is identical with
Grenade described in Ref 2, pp 230–31
b) Antitank “Lunge Mine”. This interesting device described on p 184 (with Figs) of Ref 7, consists of a conical-shaped hollow charge encased in a steel container, which was attached at its apex to a long wooden handle. Three legs, equally spaced around the wide base of the cone provided proper stand-off distance. A well in the apex of the cne contained the detonator which is activated by a striker. To operate the device the soldier removed the safety pin, and then using bayonet tactics, lunged forward striking the base of the cone squarely against the tank. When the legs of the mine struck the tank, the long handle was driven forward breaking the shear pin, and the striker was driven into the detonator, initiating explosion of the cne which could penetrate steel plates 4 to 6 inches thick. This mine was considered as a “suicide weapon” (See also Ref 2, pp 208–09 with Fig 155)

Hōshaki. Projector; Discharger

Hōshayaku. Propellant

Hōshin. Gun Barrel; Gun Tube

Howitzer. Ryūdampō. See under ARTILLERY

Hydrocyanic Acid. Seisan

Hyoteki. Target

IGNITER (Tenkayaku or Tengaku) or FIRING DEVICE (Hakkasochi). This includes: Igniting Primer (Tenka bakkan); Igniting Fuze (Tenka shinkan) and Firing Mechanism (Gekihatsu kikan). Igniters are used for igniting propellant charges, pyrotechnics, some fuzes, demolition charges and some special items. The simplest igniters are safety matches and squibs which are used to ignite safety fuses

In Ref 1, pp 233–34 are described devices to ignite demolition charges which are briefly described here under DEMOLITION CHARGES AND EQUIPMENT

In Ref 2 are described the following items:
Type 3 Electric Firing Device developed by the Navy experimentally for Proximity Fuzes (pp 188–89 with Fig 137)

Igniter System for Bangalore Torpedo is described under Demolition Tubes and in Ref 2, pp 215–17 with Fig 162

Mechanical Pull Igniter (pp 246–47 with Fig 186)
Friction Pull Igniters were: Red Type (length 2½ inches) and Black Type (length 3-1/16 inches) (p 248 with Fig 187)

Waterproof Safety Fuse Igniter (pp 248–49 with Fig 188)

Trigger-Type Safety Fuse Igniter (pp 249–50 with Fig 189)

Booby-Trap Firing Device (pp 250–51 with Fig 190)

Time Firing Device Mk 1 (pp 256–57 with Fig 193)

Chemical Delay Firing Device (pp 257–58 with Fig 194 of Ref 2)

Firing Devices used in Land and Sea Mines are listed under MINES

Illuminating Compositions (Shōmei henso).

The following compositions are listed in Refs 2 & 3:

Illuminant for Type 90 Small Model Parachute Flare consisted of Ba chlorate 75.2 & gum 24.8%. The Flare, contg 4 lbs of comp, burned for 2-2/3 mins with an intense greenish-white light (pp 22–24)

Illuminant for Type 1 12-kg Parachute Flare consisted of Ba nitrate 77, Al 8.8, Mg 4.4, S 2.2 & paraffin 4.5 parts. The Flare, contg 27 lbs of comp, burned up to 3 mins with a greenish-white light (pp 24–25)

Illuminant for 5-kg Parachute Flare Model 2 Modification 1 consisted of Ba nitrate 55.5, Al 11.0, Mg 18.0 & wax 9.1 parts. The flare, contg 5½ lbs of comp, burned for 1½ mins with brilliant white flame (pp 100–101 of Ref 2)

Illuminant for 14-cm (140-mm) and 15.5-cm (155-mm) consisted of Ba nitrate, Mg & wax (pp 493 & 504 of Ref 3)

Illuminating Projectile (Shell) (Shōmeidan).

The following projectiles are listed in Ref 3:

Type 95 7-cm (70-mm) Illuminating Projectile (pp 306–07 with Fig 239)

Type 90 7-cm (75-mm) Illuminating Projectile (pp 338–39 with Fig 265)

12-cm (120-mm) Illuminating Projectile (pp 477–78 with Fig 389)

14-cm (140-mm) Illuminating Projectile (p 493 with Fig 404)
15.5-cm (155-mm) Illuminating Projectile (p 504 with Fig 415 of Ref 3)

**Impact.** Shōgeki

**Impact Fuze.** Shōgeki or Chakuhatsu shinkan

**Incendiaries.** (Shōsei).

Accdg to Ref 2, the following incendiaries were used by the Japanese during WWII:

a) Thermite as in Army 1-kg and 5-kg Incend Bombs (pp 15–18)

b) Rubber bungs (1 by 1 inch) impregnated with soin of phosphorus in CS₂ as in Army 50 & 100 kg Incend Bombs (pp 18–20)

c) WP (white phosphorus) filled steel pellets as in Navy Type 99 No 3 Bomb (pp 63–64) and Type 3 No 6 Mk 3 Model 1 (pp 64–65)

d) Steel cylinders contg an Incend mixture of Ba nitrate 35.8, ferric oxide 27.2, Al 13.6, Mg 10.3 & synthetic shavings 13.1%, as in Type No 25 Mk 3 Model 1 (pp 66–68)

e) Thermite-filled electron fireposts as in Type 98 No 7 Mk 6 Model 1 (pp 71–72)

f) Central thermite core contg by solidified kerosene, petrol, alcohol mixture as in Model 2 of above Bomb (pp 72–73)

g) Cylindrical Incend pellets (contg incend mixture of Ba nitrate 35, ferric oxide 28, Al 18 & remainder synthetic rubber resembling Thiokol arranged around central cardboard tube contg igniter mixt (Ba nitrate 75.0, Al 24.5, oil 0.3 & moisture 0.2%) as in Type No 7 Mk 6 Model 3 Modification 1 Bomb (pp 74–75)

h) Incend Shrapnel consisting of short lengths of steel pipe filled with WP, as in Type 4 No 25 Mk 29 (pp 87–88)

i) White Phosphorus as in 1/2-kg Incend Hand or Mortar Grenade (pp 233–34) and Small Incend Rifle Grenade (pp 244–45)

j) Any inflammable benzene-type liquids, as in Molotov Cocktail (pp 235–38)

k) Thermite as in Metal Incendiary Cylinders shown on p 261 of Ref 2

l) Mixture of K chloride, sulfur, ground coal (or sugar), iron filings and wax to form an “Incendiary Brick”, one of the Sabotage Devices (p 262)

m) Mixture of Ba nitrate 30.4, paraffin 19.4, Mg 11.3, Al 11.1, rosin 10.9, ferrosferric oxide 9.1, NC 4.4 & gritty siliceous material 2.6% to form an “Incendiary Soap”, one of the Sabotage items (p 263)

The following Incendiaries are listed in Ref 3 for Army Projectiles:

a) Unknown Incendiaries in 20-mm HEIncend Projectiles on pp 282–84; 20-mm HEIncend-Tracer (p 284); 20-mm Incend (Self-Destroying) (p 285); Type 2 & Type 2 Modified 20-mm HEI (p 286); 37-mm HEI Projs (pp 289–91); Type 90 7-cm (75-mm) Incend (p 337)

b) Incendiary consisting of Ba nitrate, Al, Mg & wax, as in Type 4 20-mm HEI (Ma 202) Proj (pp 287–88)

c) Solution of WP and rubber pellets in CS₂ as in Army 7-cm (75-mm) Liquid Incend Proj (pp 339–40)

The following Incendiaries are listed in Ref 3 for Navy Projectiles:

d) WP in Al canister surrounded by graphited flake NC (p 442)

e) WP as in 25-mm HEI Proj (pp 447–48); Type 2 and Type 5 HEI (p 452); 12-cm (120-mm) IncendShrapnel (pp 482–83)

f) Steel pellets filled with a dry mixture of Mg 54, Ba oxide 26, rubber 15, ferric oxide 1.5 & sulfur 3% as in 12.7-cm (127-mm) Incend-Shrapnel (pp 485–86)

g) Unknown Incendiary as in 15-cm (152-mm) IncendShrapnel (p 500 of Ref 3)

The following Incendiaries used in Japanese Mortars are listed in Ref 3:

h) Incendiary Mixture of K nitrate 47.7, Al 21.7, S 19.9, Sb trisulfide 6.1 & wax 2.8% as in Type 89 50-mm IncendMortar (pp 375–76)

i) WP, carbon disulfide and rubber pellets as in Type 94 90-mm IncendMortar (pp 386–87)

**Incendiary Bomb (Tōka shūidan).** See under BOMBS

**Incendiary Cylinder Mk 1 “A” Large.** See Ref 1, p 249

**Incendiary Grenade.** (Shōiyō tekidan). See under GRENADES

**Incendiary Projectile or Shell (Shōidan).** See under ARTILLERY AMMUNITION

**Indicator (Shijiki).** See Markers or Indicators

**Infantry.** Hōhei
INFANTRY WEAPONS (Hohei heiki). Under this title are described and illustrated on pp 39–78 of the book of Tantum & Hoffschmidt, listed here as Ref 7, the following weapons:

9-mm Revolver, Mod 28 (1898) (p 39)
8-mm Automatic Pistol "Nambu" (pp 40–41)
7-mm Baby "Nambu" (p 41)
8-mm Automatic Pistol, Mod 14 (1925) (pp 42–43)
8-mm Automatic Pistol, Mod 94 (1934) (pp 44–45)
35-mm Pyrotechnic Pistol, Mod 10 (p 46)
Triple Barrel Signed Pistol (p 47)
6.5-mm Rifle, Mod 38 (1905) (p 48)
6.5-mm Carbine, Mod 38 (1905) (p 49)
6.5-mm Cavalry Carbine, Mod 44 (1911) (p 50)
7.7-mm Paratrooper Rifles, Mod 99 (1939) and Mod 2 (1942) (pp 51–53)
6.5-mm Sniper's Rifle, Mod 97 (1937) (p 54)
8-mm Submachine Gun, Type 100 (1940) (p 55)
8-mm Paratrooper's Submachine Gun, Type 100 (1940) (p 56)
6.5-mm Tank Machine Gun, Mod 91 (1931) (p 57)
6.5-mm Heavy Machine Gun, Mod 3 (1914) (pp 60–61)
6.5-mm Light Machine Gun, Mod 96 (1936) (pp 62–63)
7.7-mm Light Machine Gun, Mod 99 (1939) (p 64)
7.7-mm Paratrooper's Machine Gun, Mod 99 (1939) (p 65)
7.7-mm Tank Machine Gun, Mod 97 (1937) (pp 66–67)
7.7-mm Heavy Machine Gun, Mod 92 (1932) (pp 68–69)
7.7-mm Lewis Machine Gun, Mod 92 (1932) (pp 70–71)
13-mm AA Machine Gun, Mod 93 (1933) (pp 72–73)
20-mm Automatic AA/AT Cannon, Mod 98 (1938) (pp 74–75)
20-mm Automatic AT
20-mm Automatic AT Rifle, Mod 97 (1937) (pp 76–77)
25-mm Dual & Triple Automatic AA/AT Cannon, Mod 96 (1936) (p 78)

Initiator. Kökyū tenkayaku; Kibakosochi

Instantaneous Fuze. Shumpatsu shinkan

Interior Ballistics. Tōnai dandōgaku

Iperitto gasu. Mustard Gas
Iron. Tetsu
Jakusōyaku. Reduced Charge
Jet. Funshitsu
Jibaku. Self-Destroying
Jidō. Automatic Gun
Jidō. Automatic Rifle
Jippō. Ball Ammunition
Jirai. Land Mine
Jiraidan. High Explosive Shell
Jirai fusetsu. Mine Field
Jissei kairai. Magnetic Mine
Jitsudan. Live Ammunition
Jūbi kikan. Breech Mechanism
Jūhō. Heavy Gun
Jūhōheiti. Heavy Artillery
Jūken. Bayonet
Jūki or Jūkikanjū. Heavy Machine Gun
Jūkō. Muzzle (SA)
Jūsensha. Heavy Tank
Jūshin. Rifle Barrel
Jūyō tekidan. Rifle Grenade
Kainbin. Frangible Grenade
Kāen hasshaki. Flame Thrower
Kagakudan toshaki. Chemical Mortar
Kagaku heiki or Kaheizai. Chemical Agent, CWA
Kagakusei tekidan. Chemical Grenade
Kagkusei tokagan. Chemical Bomb
Kagakusha. Chemist, Scientist
Kagō. Chemical Composition
Kahō. Gun (General)
Kaiganhō. Coast Artillery Guns
Kaigun. Navy
Kakki. Firing Pin
Kakkōhō. Smooth Bore Gun
Kanashō or Kanō. Cannon (as opposed to Howitzer (called Ryūdanpō))
Kanzan yakutō. Fixed Ammo, Fixed Round
Karitto. Army explosive, originated as commercial expl Calit (qv). Its Navy modification is Type 88 Explosive (qv)
Kasen. Pyrotechnics
Kasen hashaki. Pyrotechnic Projector
Kashokuyaku. Brown Powder. Acdg to documents it was "Undercarbonized Black Powder". Use unknown, but presumably as a substitute for BkPdr (Ref 1, p 30 & Ref 5, p 366)
Keihō. Light Artillery
Keikikanju. Light Machine Gun
Keineyaku. Acdg to Ref 3, p 321 it is Heineiyaku (qv)
Ken. Bayonet
Koyaki. Gelatine Dynamite
Kibaku. Detonation
Kibakuyaku. Initiating Composition. See Bakufun (Ref 1, p 25) and Raibun (Ref 1, p 25)
Kibakuzai. Primer Charge. See Chikkaen (Army) or Chikka namari (Navy), meaning LA (Lead Azide) and also Raisan suigin (Navy) for MF (Mercuric Fulminate) (Ref 5, pp 366 & 369)
Kido shiki kahō. Mobile Artillery
Kihei. Cavalry
Kihō. Carbine
Kikanhō. Machine Cannon
Kikanju. Machine Gun
Kirai. Mine (Sea)
Kiri Nos 1, 2 & 3. Gelatin Dynamites Nos 1, 2 & 3
Ko (Explosive). Japanese Type Explosives, such as Type 88 (Ko)
Kōchū. Bore (Gun)
Kōdan. Flare, Signal Rocket, Star Shell
Kōdō. Gallery of Mine
Kōjō. Factory, Shop
Kōjōhō. Siege Gun
Kōkakuhō. Dual Purpose Gun (Navy)
Kōkei. Caliber
Kōkūki. Aircraft
Kokushokuyaku. Black Powder or Yuenyaku. Nonsmokeless Powder. Composition is not given. Used by the Army as a Main Charge in 20-mm Machine Gun Ammo, as an ejector charge in 70-mm Mortar Shells, Shrapnel Shells & Pyrotechnics; in Delays, Relays & Igniters for Bomb & Projectile Fuzes; and as a "Substitute Charge" for some Bombs, Grenades and Projectiles (Ref 1, p 27 & Ref 5, p 366)
Also used by the Navy as an ejector charge for Illuminating Shells and Pyrotechnics (Ref 1, p 33)
Kōkyū bakuyaku. High Explosive
Kōkyū ryūdan. HE Shell
Kōseisendan. Semi Steel Cast Projectile
Kōsen. Groove, Rifling
Koshahō. Anti-aircraft Gun

Koshitsu (Plastic) or Oshitsuyaku (?). A brown, putty-like expl compn consisting of RDX 80 to 85 & oil 20 to 15%. Brisance by Sand Test 108% TNT; Power by Ballistic Mortar 125% TNT; Detonation Velocity 7400m/sec vs 6900 for TNT; Impact & Friction Sensitivity — same order as for TNT; retains plasticity betw 0° (32°F) & 37.8°(100°F), becoming brittle below 0° and starting to exude oil above 37.8°. Used in Demolition Cartridges 4 inches long and 1.5 inches in diam and Hollow (Shaped) Demolition Cartridges (being 21% more effective than TNT Shaped Charges) (Ref 5, p 367)

Accdg to Ref 1, p 28, the term Koshitsu (meaning "plastic") may be a part of designation. Its compn was RDX 80 & vegetable oil 20%, and it was used in Demolition Rolls. There seems to be another plastic expl contg RDX which was a mixture with NG

Kō-Shuanbakuyaku. Ammonium Nitrate Dynamite
Kōtetsuban. Armor Plate
Kōchū haretusu. Air Blast
Kōchū hatsuendan. Smoke Bomb
Kūgun. Air Force
Kūrai. Aerial Torpedo
Kyoden. Heavy Shell; Heavy Bomb
Kyüḥō. Artillery Mortar
Kyüḥōdan. Artillery Mortar Shell
Kyūnen Kasaka. Quickmatch
LA. Abbrrn for Lead Azide (See Chikkaen)

Land Mine. See under MINES

Lead (Pb). Namari
Lead Azide (LA). See Chikkaen (Army) & Chikka Namari (Navy)

Liquid Fire. Ekka

Livens Projector. Ribensu shiki

Load or Charge of Explosive. Sōyaku

Long-Range Gun. Enkyori shagekibō

L-Shoan. Ammonium Nitrate Dynamite

Machine Cannon. Kikanbō

Machine Gun. Kikanjū

Magazines. There are several items called magazines. One for rifles and pistols is Dansō (in Japan), one for storage is Danyakako and one is a cylindrical item which can be fitted to any Fuze that takes a standard Gaine (qv). Magazines are used to initiate Low Explosives such as BkPdr but never used in a HE filled Bomb

No Army Magazine Gains are listed in Refs 1 & 2, but Navy Magazines are listed in both Refs 1 & 2. In Ref 1 are listed on p 121 Type 98, Mk 6, Model 1 and Model 2 Bomb Gaine Magazines with 0.03-sec Delay and instantaneous, with Fig 185 on p 122. The same Magazines are listed in Ref 2 on p 201 with more detailed illustrations (p 201)

Magnesium Bomb. Maguneshīmu shōidan

Magnetic Mine. Jihoku

Markers or Indicators (Shishi-Keiki)

These devices can serve as target indicators signalling landing spots, navigation markers, disaster spot markers, etc

The following items, which might serve as Markers or Indicators are described in Ref 1: Navy Type 96 Floating Landing Flare (p 241, Fig 395) Ground Signal Flare (p 242, Fig 396)
Type 94 Float Flares (pp 242–43, Figs 397 to 400)
Army Smoke Candles (pp 243–44, Fig 401)
Navy 40-kg Floating Smoke Generator (pp 244–45, Fig 402)
Navy Smoke Float (p 245, Fig 404)
Navy Smoke Flare (pp 245–46, Fig 404)
Type 2 Target Indicator (p 246, Fig 405)
Navy Navigation Markers:
Type 0 Model 1 & Model 2 (p 246, Figs 406 & 407)
Type 2 Model 11 (p 246, Fig 408)
Cardboard Type (p 246, no Fig)
Navy Type 2 Model 1 Torpedo Marker (p 249, Fig 413)

The following items which might serve as Markers or Indicators are described in Ref 2:
Navy Type 94 Float Lights (pp 106–08).
See under FLARES
2-kg and 43-kg Smoke Floats (pp 109–10).
See under Smoke Floats
Navy Sea Markers Type 0, Model 1, Model 2
were light metal cylinders with rounded nose and conical tail, provided with fins. They condt
fine Al dust with 6.5% Zn dust. On impact
with water the plug in the nose was forced into
the body forcing the tail portion to be freed
from the body, thereby scattering Al-Zn pdr,
1 oz of which covered 118 sq ft of water. Used
as distress indicator (pp 110–11, Fig 84)
Navy Sea Marker Type 0 Cardboard Type, on
impact with water broke and scattered Al-Zn
pdr over the surface of water (p 112)
Type 3 No 6 Target Marker Bomb consists of
a steel cylinder with round nose and conical
tail, provided with fins. The filling consisted of
WP-filled steel pellets contd in steel canisters.
The space not occupied by pellets was surrounded
by loose WP. The canisters were covered with
heavy wax coating. The nose-piece contd a
Gaine well surrounded by charge of Type 98
Exp (p 113, Fig 85)
Type 2 2-kg Target Indicator consisted of a
Bakelite cylinder with round nose and tail
provided with fins. The filling consisted of
liquid FM (Titanium Tetrachloride). This
smoke mixture was scattered when the bomb
was released from a plane, hit the target and
broke (pp 113–14, Fig 86)

Matsu. Blasting Gelatin

Mechanical Time Fuze. Takeishiki bakkan

Meiayaku (Army), Sansheki (Navy?) or Mechiru
Nitroamin. Tetral or 2,4,6-Trinitrophenyl-
methylnitramine,

\[
\text{CH}_3
\]
\[
\text{NO}_2; \text{mw} \ 287.16, \text{N} 24.39%;
\]
cream to yellow crs, density 1.73, mp 130°;
Brisance by Sand Test 126% TNT; Impact
Sensitivity with 2kg Wt, BurMinesApp 26cm vs
100 for TNT; Ignition Point 257° in 5 mins;
Detonation Velocity 7850m/s vs 6900 for TNT;
Power by Ballistic Mortar 130% TNT [AMCP
706-177 (1967), p 335]

Used by Japanese in pressed state as a stand-
ard Sub-Booster in Army and Navy Shells and
occasionally as the sole Booster in Navy 25-mm
Shells and as Burster in some Bombs and Shells.
Also used in composite explosive Tanōyaku (qv)
(Ref 1, pp 26, 31 & 33; Ref 5, p 367)

Menkayaku (Guncotton). Comp with Shōkamen
(Nitrocellulose)

Menyaku. Same as Menkayaku

Military (adj). Rikugun no

MINES. Jirai (Land), Kirai (Sea), Suirai
( Amphibian or Water) and Booby Traps
(Yūgeiteki jirai)

Accdg to Ref 1, p 210 & Ref 2, p 203, al-
though the use of Land Mines by the Japanese
Forces was not as extensive as it was in Europe,
they were important defensive weapons in the
Pacific War. Also, because of the Japanese lack
of effective A/Tk Artillery and the inequality of
Armored Forces, mining and similar tactics be-
came a mainstay of defense against mechanized
equipment

Three features of land mining methods were
especially important:
1) The relatively small number of standard, mass-produced mines and
firing devices;
2) The prevalent use of extremely
large expi charges which was wasteful and
3) The emphasis was placed on various types of con-
trolled mines instead of by enemy activated
ones. Firing devices operated by lanyards or
poles were very common.

Japanese mining techniques were characterized by an almost complete lack of uniformity.

In Chapter VI of Ref 1, entitled “Japanese Mines and Booby Traps” are briefly described and rather poorly illustrated various Army and Navy Land Mines (pp. 210 to 228).

More comprehensive descriptions, with good illustrations, are given in Chapter 3 of Ref 2 for the following Land Mines:

Type 93 A/Tk (Antitank) and A/P (Antipersonnel) Land Mine, known as “Tape Measure” Mine, was a light metal cylinder 6¾ inches diam and 1¾ inches high, filled with 2 lbs of cast PA and Booster of pressed PA. It was activated by pressure on Percussion Primer located on top of mine (pp. 202–03, Fig. 152). Described in Ref 7, p. 183 as Land Mine Model 93 (1933).

Anti-Vehicular “Yardstick” Land Mine was an oval tube 3.35” x 1.8” in diam and 36” long, formed by two halves of sheet steel welded together. It contains 8 blocks of cast PA wrapped in paper (wt 6 lbs) and a Fuze which could be activated by strong pressure such as that of a vehicle (pp. 204–06, Fig. 153). Described in Ref 7, p. 187 with Figs.

Type 99 Armor-Penetration Land Mine was a canvas disc 4.75” in diam & 1.5” wide, filled with cast blocks of 50/50–RDX/TNT forming a circle. Four magnets were attached around the circumference, which were designed for attaching the mine to the side of a tank or other vehicle (pp. 206–07, Fig. 154). Described in Ref 7, p. 181 as “Magnetic” Antitank Mine Model 99 (1939).

Antitank “Lunge” Mine (p. 208, Fig. 155) is also described here under “ Hollow Charge Japanese Ammunition” and in the book of Tantum & Hoffschmidt, listed as Ref 7 (p. 184).

Suction-Cup Mine consisted of sheet metal cylinder filled with 4.5 lbs of 53/47–RDX/TNT and provided on top with a 5 ft pole. Two suction cups at the bottom of the mine secured it to target when pressed. Ignition was made by two Friction Pull Igniters which were lashed to the bottom of pole (p. 209, Fig. 156).

Dutch A/Tk–A/P Land Mine, known as “Mushroom Mine” consisted of a pressed steel low cylindrical body and a dome-shaped cover made of the same material. It was filled with TNT (5½ lbs) and contd a fuze. A spring held the cover off the fuze, but a pressure of 50 lbs was sufficient to activate it (pp. 210–11, Fig. 157).

Type 3(A) Antivehicular & Antipersonnel Mine, known as “Flower Pot”, consisted of a terra cotta, low cylinder filled with Type 88 Explosive or with 50/50–Amatol. The fuze located on top of mine could be activated by pressure of ca. 22 lbs (pp. 211–12, Fig. 158). Described in Ref 7, p. 182 as “Navy Type 3 (Flower Pot) Land Mine–Small”.

Type 3(B) A/Vehicular & A/P Land Mine consisted of a wooden box filled with Type 88 Expl (p. 213, Fig. 159).

Beehive A/P Land Mines, recovered on Okinawa, consisted of serrated cast-iron, hemispherical casing filled with ca. 5 lbs TNT. A pressure of one person on a fuze located on the top was sufficient to initiate explosin (p. 214, Fig. 160).

Type 3(C) 5-kg Hemispherical A/Tk Land Mine consisted of light metal casing filled with HE (?) and a fuze mounted opposite the flat bottom. It was a suicide weapon, intended to be placed against an armored vehicle either by hand or by means of a pole attached to the case (p. 215, Fig. 161).

Bangalore Torpedo, described here under DEMOLITION TUBES could be used as a Land Mine (pp. 215–17, Fig. 162).

Pressure and Traction Land Mine consisted of a wooden box, the lid of which was held in place against the internal fingers on the top by stout springs, one in each corner. A wooden block was secured by two bolts to the underside of the lid and served to operate the pull igniter by pressure exerted on the cover. It was filled with 2 lbs of cast PA or TNT (pp. 217–18, Fig. 163).

Friction-Fuzed Land Mine consisted of a wooden box 13” long, 3” wide and 2.37” high contg 5 blocks of cast PA or TNT (3.5 lbs), detonator or igniter. To the igniter was attached a trip wire which extended out thru the end of the box and was secured to a tree or other fixed object (pp. 218–19, Fig. 164).

Improvised Land Mine consisted of a rectangular tin box with a cover fastened by friction tape. Two holes were roughly punctured in the cover thru which a grenade fuze or detonator projected. Contained in the box were: one Type
Hand Grenade and 12 blocks 1\(\frac{1}{2}\)" by \(\frac{3}{8}\)" of 66.6/33.3—RDX/Al compn, each wrapped in waxed paper. It could be used as an A/Tk Mine when fused with an armed grenade or as an A/P Mine or Booby Trap when armed with pull or tension detonator (p 219, Fig 165). Air-Strip Land Mine consisted of 31 100-kg bombs stacked around PA blocks in which electrical detonators were inserted. The ensemble was under a turf-covered piece of sheet iron that would close the circle and fire the charge if the iron were lifted or depressed. A clockwork was also inserted to fire the charge if the iron were not depressed (p 220, Fig 166, upper half).

Improvised Antitank Land Mine consisted of a wooden box serving as a base of Mod 99 Armor-Piercing Mine on top of which was placed 2 lbs prep'd charge of PA. Two Hand Grenades were placed at both sides of AP Mine and the ensemble was covered with a board. Pressure on the board fired fuzes in grenades and this fired by sympathetic detonation AP Mine and prep'd charge of PA (p 220, Fig 166, lower half).

Type JE Antisubmarine Land Mine consisted of a hemispherical steel body 20.24" in diam & 10.62" high contg 46.5 lbs of Type 98 Expl, with PA Booster and Electrical Tetryl Detonator. There were two handles and two acid-lead horns on upper half of the mine. The mine was laid in water, flat-bottom down, and if one of the horns was bent or crushed, an acid vial inside was broken, allowing the acid to drain onto two plates of a small battery which generated sufficient current to fire the detonator. As the wiring was series-parallel, either horn on being touched would act independently to fire the charge (pp 221–22, Fig 167).

Note: This mine is identical with Naval Antisubmarine Mine described on p 185 of the book of Tantum & Hoffschmidt, listed here as Ref 7. It was believed that the mine was designed for anti-invasion purposes to be placed in shallow water against landing craft, vehicles and tanks.

Type JG Antisubmarine Land Mine, which existed in 5 modifications, each being actuated by a single chemical horn screwed into the top. The mines were either bell-shaped or of a truncated cone. Each contd 22 lbs of Type 98 Explosive. The mines were fired when chemical horns were broken (pp 222–23, Fig 168).

No Sea (Amphibian) Mines are described in Ref 2, but the following *Naval Sea Mines* are described in Ref 1, Chapt III, under Japanese Underwater Ordnance (pp 34–53):

**Contact Mines** were generally spherical, with several lead, steel horns screwed into the mine case. Horns were of several types but the most common was a lead chemical horn, such as described above under Type JE Antisub Marine Mine. Some contact mines had a long copper wire antenna extending out of the top or bottom of the case. Should a steel ship touch the antenna while both were immersed in sea water, the mine would explode. Moored contact mines usually had some type of safety mechanism to render them inactive if they bore adrift, as well as devices to make sure the mines were safe while aboard the mine-laying ship, submarine, or plane (p 34).

**Influence Sea Mines**, were of various shapes, but usually they were cylindrical, like a bomb in appearance. They did not require contact with a ship in order to fire, and hence could be laid on the bottom many feet below the ship. Any physical change which took place in the water around the mine to the ship's passage over it could be utilized to actuate the firing mechanism. However, the most common types were magnetic and acoustic. Any mine without horns or antenna was almost certainly either an influence mine, or else a *Controlled Mine*, connected by an electric cable to the shore and fired by sending a current thru that cable (p 35).

Following is the list of Japanese Sea Mines used during WWII:

**Type 88 Model 1 Contact Mine** consisted of two light-metal hemispheres separated by a cylindrical belt and provided by four chemical horns on upper hemisphere. It was filled with 396 lbs of block-fitted Shimose. The depth setting hydrostat and mooring cable were provided (p 36, Fig 1).

**Type 92 Moored Controlled Mine** was ellipsoidal in shape, 55 by 41.5 inches, filled with 1100 lbs of Type 88 Explosive. The combined control and mooring cable entered the mine thru the base plate. An electric detonator and booster assembly were bolted to the chge underneath the top cover plate. A microphone to detect the presence of ships was also provided (p 36, Fig 2).

**Type 92 Model 1 contd** Type 1 Explosive and was without microphone (p 36, no Fig).
Type 93 Model 1 Moored Contact Mine was sphere-shaped 34 inches in diam and contd 220 lbs of Type 98 Explosive. Four chemical horns were located around the upper hemisphere, while the detonator was housed at the base (pp 36–7, Fig 3)

Type 93 Model 4 had six horns and was loaded with Type 1 Explosive (p 38, Fig 4)

Type 94 Model 2 was a ground controlled hemispherical mine with a base diam 28 inches and height 25 inches, contg 190 lbs of Type 88 Explosive. The firing cable entered the Booster and Detonator well thru the side of the case (pp 38–39, Fig 5)

Mark 5 Model 1 Moored Contact Mine had spherical case contg 180 lbs of Shimose. Four chemical horns were fitted to the upper hemisphere (p 39, Fig 5)

Mark 6 Model 1 Moored Contact Mine had spherical case, 41 inches in diam, contg 478 lbs of Shimose. Four chemical horns were set in the upper hemisphere (pp 39–40, Fig 7)

Mark 6 Model 3 contd 440 lbs of Type 88 Expl and was provided with six horns (p 40, no Fig)

Small Model Mine Model 1 was hemispherical in shape, 20 inches in base diam & 10 inches high which contd 45 lbs of Type 98 Expl.

Two chemical horns were located on the upper surface of the case. It was set on the ground in shallow water (pp 40–41, Fig 8)

Small Model Mine Model 2 was truncated cone-shaped case, 14 inches in base diam & 12 inches high, contg 22 lbs of Type 98 Expl. A single chemical horn was placed on top of the mine (p 41, Fig 10)

Type 3 Mark 1 Aircraft Mine Model 1 was cylinder 53 inches long and 24 inches diam, provided with a parachute and four chemical horns. It was filled with 240 lbs of Type 98 Explosive. The detonator and booster were located in a tube running transversely thru the mine case. It was dropped from an aircraft or surface moored (p 42, Figs 11 & 12)

Type 3 Model 2 Aircraft Mine Model 2 was an aircraft-laid drifting mine similar in appearance to fin-tailed bomb, 6 ft long and 14 inches in diam. It was filled with 123 lbs of Type 98 Expl cast in the spherical nose. The detonator and booster were housed in a transverse tube running thru the bomb, while further aft was located the hydrostatic arming switch. The tail section broke free of the mine after laying and floated on the surface. The mine proper was suspended vertically ca 6 ft under the surface of water by a cable attached to the buoyant tail. When the tail was separated from the mine, three sensitive switch horns sprang out around the base of the suspended mine. About 5 lbs pressure on any of the horns could fire the mine (p 43, Figs 14 & 15)

Type 96 Mine was ellipsoid in shape and contd 120 lbs of Type 88 or Type 97 Expl. It was reported to be attached to antisubmarine nets. The firing mechanism of mine was of the tension type which detonated the mine when an enemy's submarine fouled a panel of the net (pp 44–45, Fig 16)

Banana Mine was the moored contact type provided with four chemical horns and loaded with 275 lbs of Type 88 Expl (pp 44–45, Fig 17)

JM Mine was moored, pear-shaped, contact mine which had no horns. The jolt received by the pendulum firing mechanism, when a ship struck the mine, closed an electrical contact and fired the detonator. The mine was filled with ca 110 lbs of a HE (?) (pp 44–45, Fig 18)

Pomegranate Mine was a Vickers antenna mine purchased before WWII. It was spherical, 41 inches in diam, with six chemical horns on the upper hemisphere and two addnl ones welded to the lower hemisphere. An upper antenna and float and/or lower antenna were secured to the mine. No data for HE filler (pp 44–45, Fig 19)

Type 3 Mine was the first Japanese attempt to use influence mines, copied from German S Mine. Their cylindrical Al cases were either 7 or 11 feet long and 21 inches in diam. No data for HE used and the description of the mine given in Ref 1, p 46 is hard to understand

Mark 2 Mod 1 Explosive Hook was actually a device for mine sweeping rather than a mine. It was a cylinder 8 inches in diam and 10 inches long, loaded with 33 lbs max of Type 88 Expl or Shimose. The device was towed underwater and when one of the four projecting arms welded to the body fouled a mine mooring cable, the device was electrically fired from the towing ship (pp 46–47, Fig 21 of Ref 1)

Minefield. Jirai fusetu chitai

Minethrower. Hakugekikō

Mobile Artillery. Kidōshiki kahō

Molotov Cocktail (Tenage kaenbin). See under GRENADERS

Monkan. Friction Primer

Mooring Line. Keiryūsaki

MORTARS (Kyūhō) and SHELLS (Dangan)

The following Mortars are described and illustrated in the book of Tantum & Hoffschmidt, listed as Ref 7:

70-mm Mortar Model 11 (1922) was a rifled bore weapon which fired HE, Smoke or Illuminating Shells, Range 1700 yds (p 159)
81-mm Mortar Model 3 was a smooth-bore weapon, forerunner of Mortar Model 97 (1937). It was manuf'd at Yokosuka Navy Arsenal. No data given for length of barrel and Shells used (p 162)
81-mm Mortar Model 99 (1939) was a short 25.5 inches), smooth-bore, muzzle-loading weapon of high trajectory. Used HE Shells. Range 3200 yds for 7.2-lb Shell (p 163)
81-mm Mortar Model 97 was similar to US 81-mm Mortar M1 with length of tube (including base cup) 49.5 inches. Fired HE Shells (p 165)
90-mm Mortar Model 94 (1934) was smooth-bore, muzzle-loading weapon with length of barrel (including base cup) 11.7 inches. Used HE or Incendiary Shells. Range ca 4050 yds with 11.9-lb Shell (p 167)
90-mm Mortar Model 97 (1937) was a Stokes-Brandt type weapon, very similar to US 81-mm Mortar M1. Length of barrel (interior) 48 inches, range 4050 yds for 11.8-lb HE or Incendiary Shells (p 168)
150-mm Mortar Model 97 (1937) was smooth-bore, muzzle-loading weapon with length of tube (internal) 66 inches. No data for its range and shells used are given (p 169)

There were also two Grenade Dischargers which resembled Mortars in appearance:
50-mm Grenade Discharger Model 10 (1921) consisted of a steel smooth-bore barrel 9.5 inches long attached to a pedestal resting on a base plate. It was muzzle-loaded and was fired by a continuous pull firing mechanism, operated by a lever on the outside of discharger body. Used for firing 91 Hand Grenade, Pyrotechnic Signals and Smoke Shells (p 155)
50-mm Grenade Discharger Model 89 (1929) had steel rifled-bore barrel 10 inches long attached to a pedestal, resting on a base plate. The propellant chge was placed at the bottom of barrel, held at 45 degree angle, and this was followed by projectile inserted thru muzzle (with safety removed). Then the trigger which protruded thru a lengthwise slot in the pedestal was pulled and fired the propnt. It was used for firing Model 89 HE Shell, Model 91 Hand Grenade, Model 95 Smoke Shell and Incendiary Shell. Range 700 yds for shell, 200 for Grenade (p 156 of Ref 7)
32-cm (320-mm) Spigot Mortar. The spigot shown in Fig on p 170 of Ref 7 was a cylinder 31.7 inches long with outside diam 10.1 inches. A cavity was machined in its upper end for the propelling charge. In the bottom of the cavity there was a threaded recess for the igniter assembly. An igniter train hole passed from it to the base of spigot. The steel mounting plate in the shape of a dome was attached to a steel base plate which rested on three layers of heavy wooden beams

Acdg to Ref 3, pp 389-90, the Shell fired by Spigot Mortar was cylindrical in shape (resembling a bomb) with ogival nose and finned tail section hollowed out to inside diam of 10.12 inches and length 14.5 inches so that it could fit over the spigot. The shell contd 103 lbs of Shimose as main chge and PETN as booster. Its Fuze was Type 98. The propellant chge — single-base with BP as Igniter was placed in a brass pot which fitted into the recessed portion of the spigot tube. The chge was contd in a cardboard container having a flash tube passing thru its vertical axis, thru the brass pot and screwed into the spigot. Also there was a drilled and tapped hole near the forward end of the tail section of the shell into which the igniter tube screwed. When in position, the igniter tube was in close proximity to the end of the brass flash tube which led into the propnt chge. Variations in range could be obtb by varying the number of in-
crements selected for the proplnt chge

The following Shells for use in Army and Navy Mortars are described in Ref 3:
Type 89 58-mm HE Mortar Shell was a light steel cylinder with conical nose housing a Fuze. It was filled with 5.4 oz of TNT and it was fired by NC-DPhA flaked proplnt fired from Type 89 Grenade Discharger (pp 272–73, Fig 296)
Type 95 50-mm Smoke Mortar Shell was a forged steel cylinder with conical nose housing a Fuze. It contd 3.7 oz Hexachloroethane Smoke Mixture and was fired by NC-DPhA flaked proplnt. Fired from Type 89 Grenade Discharger (pp 374–75, Fig 297)
Type 89 50-mm Incendiary Mortar Shell was a sheet metal cylinder with hemispherical nose. It contd 10.7 oz of Incendiary Mixture and its base was NC proplnt and two BkPdr delays. Fired from Type 89 Grenade Discharger (pp 375–76, Fig 298)
Type 11-Year 70-mm HE Mortar Shell was a light steel cylinder with conical nose housing a Fuze. Its Main Charge was 15.2 oz TNT with Booster 1.6 oz of PA. Was fired by NC-DPhA flaked proplnt. Fired from Type 11-Year 70-mm Rifled Mortar (p 378, Fig 301)
70-mm HE-AA Barrage Mortar Shell consisted of a light-steel outer shell contg 7 canisters in each of which was an HE cylinder and parachute. A turned steel base was welded to the outer shell, and provision was made for reception of a delay train holder, a shell proplnt container, and an end cover. The delay train led to a BkPdr chge which ejected the canisters from the shell. Total wt of Main chge 12.84g of RDX, with Booster of LA (Lead Azide)

We are not describing the construction of this shell in detail because it is very complicated and requires detailed illustrations. It suffices to say that the Shell was fired by dropping it into a smooth-bore Mortar, 4 feet long, attached to a base plate, a rectangular wooden block and an iron rod which was stuck into the ground to hold the Mortar upright. This weapon, known as 70-mm AA Barrage Mortar, was fired at the time of approach of low-flying enemy aircraft so that it released from each shell 16 parachute-provided cylinders which were suspended in the air forming a kind of barrage. Seven of these cylinders contd HE chge of RDX-LA which was connected to a friction cap attached to parachute cord. Any enemy plane touching a parachute cord of any of the seven HE cylinders would cause an explosion (Ref 3, pp 178–80, Fig 302 and Ref 7, pp 160–61)
Type 97 and Type 100 81-mm HE Mortar Shells were in the shape of streamlined bombs. Its Main Charge was TNT (ca 1.2 lbs), Booster-PA and proplnt which is described here under PROPELLANTS. Was fired from Type 97 or 99 Mortars (pp 380–81, Figs 303 & 304) 81-mm Parachute HE Mortar Shell was cylindrical in shape. Its Main Charge was RDX (one block) and TNT (two blocks) — total wt 4 oz. Its primer, ejection and delay charges were BkPdrs, but no data for its propellant are given. Was fired from 81-mm Smooth Bore Mortars (pp 381–82, Fig 305)
81-mm Parachute HE Smoke Mortar Shell was similar in construction to HE Shell, except that its Main Charge was Tetryl and it contd a smoke pellet (?) (pp 383–84, Fig 306) 81-mm Parachute-Flare Mortar Shell was cylindrical, similar to 81-mm HE Mortar Shell, except that it contd a flare compn for which no data were given (pp 384–85, Fig 307)
Type 94 HE and HE Semisteel Mortar Shells were in the shape of streamlined bombs. Their Main Charge was TNT (2.35 lbs), with pressed PA as Booster. Compn of proplnt not given. Was fired from Type 94 or 97 90-mm Mortars (pp 385–86, Figs 308 & 309)
Type 94 90-mm Incendiary Mortar Shell was similar to Type 94 HE, except that it contd Incendiary filling consisting of WP, CS2 & 40 cylindrical rubber pellets (2.2 lbs). It also contd Burster Chge and Proplnt of unknown compn (pp 386–87, Fig 310)
Type 2 120-mm HE Mortar Shell was streamlined in shape, contg 6.0 lbs of TNT with RDX/Wax Booster. No data for proplnt. Was fired from Type 2 120-mm Smooth Bore Mortar (pp 387–88, Fig 311)
Type 96 150-mm HE Mortar Shell was streamlined in appearance contg 12.9 lbs of TNT as Main Charge with PA as Booster. Compn of proplnt is given under PROPELLANTS. Was fired from Type 96 150-mm Smooth-Bore Mortar (p 388, Fig 312)
Type 97 150-mm HE Mortar Shell was a shorter version of Type 96. It contd 9.98 lbs of TNT
and RDX/Wax Booster. Its proplnt was the same as for Type 96 (pp 388–89, Fig 313) 32-cm Spigot Type Mortar Shell is described under 32-cm (320-mm) Spigot Mortar

**Mountain Artillery. Sampōheii**

**Muenkayaku or Muenyaku.** Smokeless Propellant. See under PROPELLANTS

**Munitions.** Gunjuhin

**Mustard Gas.** Masūtado gasu; Iperitto gasu

**Muzzle.** Hōko (Arty); Jūkō (Small Arms)

**Muzzle-Loader.** Zensōhō

**Muzzle Velocity.** Shosoku

**Naval.** Kaigan no

**Navigation Markers.** See under Markers and Indicators

**Navy.** Kaigun

**Nigotanoyaku-Mk 2.** Pale-yel or cream expl mixts of TNT & RDX (70/30, 60/40 & 50/50). Its 50/50 Compn is called in Ref 2, p 30 “Army Mk 2 Explosive” or Tanōyaku, but acqdg to Ref 8, p 373, Tanōyaku contd TNT, RDX & 7–10% Tetryl. Acqdg to Ref 5, p 368, the 50/50 version was called Chauyaku

Nigotanoyaku expls were used in many types of Army Ordnance (Ref 1, p 27). Acqdg to Ref 5, p 368, they were used in Bombs, Shells, Land Mines and Bangalore Torpedoes. The 53/47–TNT/RDX version was used in Demolition Charges

**Nitroglycerin muenyaku.** Nitroglycerol or Double-Base Propellants. See under PROPELLANTS

**Nitrocotton.** Menkayaku

**Nitroglycerol (Nitroglycerin).** Described in Vol 6 of Encycl, under GLYCEROL TRINITRATE, pp G98ff. It was used in Dynamites (See under Dainamaito) and in Double-Base Propellants (See under PROPELLANTS)

**Nose (of Projectile).** Dantō

**Nose Fuze.** Dantō shinkan

**Obturation.** Kinsoku

**Offensive.** Kosei

**Onayaku.** A yellow, castable expl compn consisting of PA (Picric Acid) 50 & DNN (Dinitronaphthalene) 50%, which melts below 120°. Its Brisance, Power and Rate of Detonation were lower than those for straight PA and it was less sensitive to Impact than PA. Used as Bursting Charge in Artillery Projectiles, such as in 7-cm (75-mm) HE-AA Proj, listed in Ref 3, p 318. There was also, acqdg to documents, mixture of PA 80 & DNN 20% which was identical with Frantsuzskaya Smes' used by the Russians (Ref 1, p 26 & Ref 5, p 368)

**Oshitsuyaku.** Acqdg to Ref 5, pp 367 & 368, it is identical with Koshitsu (qv)

**Oshiyaku.** A yel, press-loaded expl compn consisting of PA 90 & wax 10%. It was much less sensitive to Impact & Friction than straight PA and had lower Brisance, Power and Rate of Detonation. It was used as Bursting Charge in nose of some Armor-Piercing Projs, such as Type 95 7-cm (70-mm) AP Proj listed in Ref 3, p 309 (See also Ref 1, p 26 & Ref 5, p 368)

**Oshokuyaku (Army), Shimose bakuyaku (Navy) or Pikurinsan.** Picric Acid (PA), HOC₆H₂(NO₂)₃; mw 229.11, N 18.34%; yel crysts, density 1.76, mp 122°, Brisance by Sand Test 113% TNT; Explosion Temp 320° (decomp); Power by Ballistic Mortar 112% TNT; Rate of Detonation 7350m/sec for cast (Shimose) vs 6900 for TNT; Impact Sensitivity by Bur Mines App with 2-kg weight 85cm vs 100 for TNT (AMCP 706-177 (1971), p 288). Used by the Japanese Army as a press-loaded Booster Charge in Bombs, Shells, Mines and Standard Demolition Charges. Used
cast, under the name of Shimose, as Bursting Charge in some Bombs, Shells, Torpedoes & Mines. Also used in some expl compns, such as Chōšyaku, Oneyaku & Oshiyaku (Ref 1, pp 25, 26 & 31 and Ref 5, p 369)

Otsu-B, Type A or A (ko) Explosive. A dark green expl compn of TNT 60, HNDPhA (Hexanitrodiphenylamine) 24 & Al pdr 16%. Used by the Navy for loading Torpedo Warheads, Sea Mines and Depth Charges. It was identical with German expl compn Schiesswolle 18 oder TSMVI-101, described in PATR 2510 (1958), p Ger 177-L (See also Ref 1, p 32 & Ref 5, p 361, under A (ko) Explosive)

Paper Balloons. See Balloons, Paper

Parachute Flare (Chōkō shōmeidan). See under FLARES.

Penetration (of Projectile). Shintetsu

Pentolite. See Pentoriru

Pentoriru (Pentolite). A pale yel or cream expl compn of PETN (Pentaerythritol Tetranitrate) 50, coated with TNT 50%, density 1.65, Bri- sanse by Sand Test 125% TNT, Explosion Temp — decomposes at 220° in 5 secs; Power by Ballistic Mortar 126% TNT, Impact Sensitivity by BurMinesApp with 2-kg wt 34 cm vs 100 for TNT; Rate of Detonation for cast 7465m/sec vs 6900 for TNT [AMCP 706-177 (1971), p 272]. Used in Navy 20 & 30-mm Proj (Ref 1, p 33) and in some Shaped Charges (Ref 5, p 369) (See also Ref 3, p 441)

PETN (Pentaerythritol Tetranitrate). See Shoeyaku

Picric Acid (PA) (Pikurinsan). See Oshokuyaku (Army) and Shimose Bakuyaku (Navy)

Pikurinsan Ammonia (Ammonium Picrate) is described as Explosive D in AMCP 706-177 (1971), p 136, (O₂N)₃C₆H₂ONH₄; mw 246.24, N 22.76%; yel crystals, d 1.72, mp=decomp at 265°; Bri- sanse by Sand Test ca 92% TNT; Explosion Temp, decomp in 5 secs at 318°. Impact Sensitivity BurMinesApp, 2-kg Wt >100cm;

Power by Ballistic Mortar 99% TNT; Rate of Detonation 6850m/s vs 6900 for TNT. Used in composite expls: Type 1, Type 1 Mk 5P5, Type 1 Mk 6 and others (Ref 5, p 369)

Point Detonating Fuze. Danōbakkan; Danō shikanen

Powder (Kayaku; Bakuyaku). See PROPELLANTS

Powder Bag. Yakunō

Powder Charge. Sōyaku

Powder Train. Dōkaten

Primer. Bakkan

Primer Charge. Kibakusai

Projectile (Dangan; Shadan). See under AMMUNITION

PROPELLANTS (Hōshayaku).

There were 1) Smoke Propellants (or Non-smokeless Propellants), called Yuenyaku, which were Black Powder called Kokushokuyaku (Ref 1, p 27) and Brown Powder called Kasshokuyaku (Ref 1, p 30). They are described here under K's and 2) Smokeless Propellants of general term Muenkayaku or Muenyaku.

The following Smokeless Propellants were used by the Japanese:

A) Nitrocellulose or Single-Base Propellants (Shōkayaku) contd NC (Nitrocellulose) (ca 12.5% N) 98 & DPhA (Diphenylamine) 2%. Used to propel many projectiles (Ref 1, p 30). A slightly different formulation is given on p 33 of Ref 1: NC 98.0, DPhA 0.8, DNT 0.95 & graphite 0.25%

Single-base propants were subdivided accdg to shape and sizes of grains into Mk 1 (squares of flat strips); Mk2 (squares); Mk3 (strips); Mk 4 (strips) and Mk 5 (strips) (Ref 3, pp 346 & 357). There were also graphited propants made of single-perforated cylinders. They were used in Types 97 & 98 20-mm Ammunition (Ref 3, p 280)

B) Double-Base (NC-NG) Propellants included:

a) NC 67.8, NG 29.3, DPhA 0.7 & NaCl 2.2%.
Used for propelling some Mortar Projectiles (Ref 1, p 30)

b) NC 65.6, NG 29.2, Ethyl Centralite 3.7 & NaCl 1.5%. Used for propelling many Artillery Shells (Ref 1, p 33)
c) NC 60.0, NG 34.5, Ethyl Centralite 3.0 & Diphenylformamide 2.5%. Used to propel Type 1 47-mm HE & AP Shells (Ref 3, pp 298–99)
d) Smokeless Double-Base Propellant B analyzed in US: NC 63.50, NG 27.71, Ethyl Centralite 3.8, Diphenylformamide 3.68, volatiles 1.30, graphite 0.45 and ash 0.34%. Consisted of single perforated cylinders of various lengths for use in 20-cm HE Spin-Stabilized Rockets (Ref 3, p 372)
e) NC & graphite 58.0, NG 7.1, DNT 25.7, DPhA 0.5 & K nitrate 8.7%. Used to propel Type 97 81-mm HE Mortar Shell (Ref 3, p 380), Type 96 & 97 150-mm HE Mortar Shells (Ref 3, pp 388 & 389)
f) Propellant for “Baka” Rocket Bomb: NC 59.9, NG 26.9, MNN (Mononitronaphthalene) 6.1, EtCentralite 2.9 & volatiles 1.3% (Ref 2, p 118)
g) Special DT6 Double-Base Propellant: NC 60.0, NG 29.3, EtCentralite 2.85 & MNN (Mononitronaphthalene) 6.14% consisted of single-perforated grains. Was used to propel 12-cm Incendiary-Shrapnel Rocket (Ref 3, p 514), 20-cm HE Rocket (Ref 3, p 515) and 45-cm HE Rocket (Ref 3, p 516)

Proving Ground. Shiken shagekijō

Proximity Bomb, Type 5 No 25 Mk 33 is described in Ref 2, p 93, Fig 70 on p 92

Pyrotechnic Pistol. Shingō kenjū

Pyrotechnics. Shōgōdan

Pyrotechnic Signal. Kasen shinjō

Quickmatch. Kyūnen kasaku

Raibun (Enka) (Thunder Powder or Smoke and Fire) consisted of K chloride 60 & Arsenic Sulfide 40%. Lt gray to tan powder used in fuze and pyrotechnic primers. Comp with BakuFun (Ref 1, p 25; Ref 5, p 369, where its compn is erroneously given as K chloride 60 & Antimony sulfide 40%)

Raigeki suru. Torpedo

Raikan. Percussion Primer or Cap

Raikō (Army) or Raisansuigin (Navy) “Thunder Mercury”. Mercuric Fulminate, Hg(ONC)₂; mw 284.65, N 9.84%; white to gray pdr, d 4.43, mp–decomp or detonates without melting; Brisance by Sand Test–ca 55% TNT; Expln Temp 210° in 5 sec; Impact Sensitivity, Bur Mines App, 2-kg wt–5cm; Power by Trauzl Test–51% TNT; Rate of Deton 5000 at d 4.0 (AMCP 706-177, listed as Ref 8, p 201). Used straight by the Japanese in Instantaneous Fuzes, in Blasting Caps and in mixture called Bakufun (Ref 1, p 25 & Ref 5, p 369)

Rakkasan. Parachute

Range (of Weapon). Shatei

Recoil. Handō suru or Kōza sura

Relay. Denkeiki

Rempetsuji. Magazine Rifle or Repeating Rifle

Rentaihō. Regimental Cannon

Report (Explosion). BakuSei

ResshaShōhei. Railway Artillery

Ricochet. Chōdan

Rifle. Shūji

Rifle Grenade. Jūyō tekidan

Rikugun. Army, Land Forces

Rin. Phosphorus

ROCKET (or Roketto) (Funshindan) or Rocket Ammunition Raketto Danyaku) can be defined as any self-propelled, unguided missile which is fired from a device called a launcher, as was Ger Faustpatrone, Russian Katiusha or Amer Bazooka. Japanese launchers are briefly described here under ROCKET LAUNCHERS. Each Rocket Missile carried its own propeller
type motor (described here under Rocket Motors) and a warhead consisting of an HE, Incendiary or Chemical Agent (Vol 1 of Encycl, p A384-R) (See also ROCKETS)

ROCKET BOMBS (Roketto bakudan).

The following Navy Rocket Bombs are listed in Ref 2:

Type 5 No 1 Mk 9 Model 1 and Type 3 No 6 Mk 9 Bombs were fin stabilized and closely resembled American Aircraft-launched Rockets. Both carried HE Filler and small charges of Propellant (p 71 of Ref 2; no Fig)

Type 3 No 6 Mk 27 Rocket Bomb Model 1 was fin-stabilized, aircraft-launched, designed for use against formations of large enemy planes. It was cylindrical in shape with conical nose section which cont'd a Fuze, a Burst Charge and an Incendiary Shrapnel, consisting of WP (White Phosphorus) filled steel pellets. Its cylindrical part cont'd Ballistite type Propellant in grains 350mm long and 19.2mm in diam. Its tail section was constricted to form a venturi for the escape of driving gases. This Rocket, along with Type 3 No 1 Mk 28 Model 1, was the only Navy Rocket which was actually under production at the end of WWII (p 84 & Fig 65 on p 85)

Type 3 No 1 Mk 28 Rocket Bomb Model 1 was to a certain degree similar to Type 3 No 6, except that it cont'd HE Filler instead of Incendiary (pp 86–87, Fig 66)

“Baka” Piloted Rocket Bomb was a suicide weapon designed to be controlled by a human pilot. It resembled a plane, was carried beneath the fuselage of a bomber and released near its target. Three Type 4 Mk 1 Rocket Motors provided propulsion after “Baka” was released from the mother plane. The entire HE content of the “Baka” (1135 lbs of Type 91 Expl) (Trinitroanisole) was in the warhead of the nose. “Baka” was 19 ft 10 inches long with wingspread 16 ft 5 inches. Its warhead had Nose and Tail Fuzes (pp 116–17, Fig 88)

Type 4 Mk 1 Launching Rocket (“Baka” Motor) was cylindrical in shape with conical nose and venturi tube tail. It cont'd 97.8 lbs double-base propellant listed under PROPELLANTS, item f. Three of such devices, located in fuselage of “Baka”, served as the main propulsion units (Ref 2, p 118, Fig 89)

ROCKET LAUNCHERS and PROJECTORS (Roketto Hashaki).

Accdg to Refs I & R, Japanese Rocket Launchers were mostly wooden troughs, others were guide rails and steel barrels (tubes). Launchers were provided with some electric ignition device. The most practical launcher was used by the Japanese Army under the name: Type 4 20-cm Rocket Launcher, described in Ref 1, p 204 and shown in Fig 320 on p 206. It consisted of a metal tube on a mounting permitting fine adjustments in elevation and train. The launcher was in three parts which could easily be disassembled for transportation. A rate of fire of from 1 to 2 rounds per minute could be obt'd

Other launchers listed and illustrated in Ref 1 were:

20-cm Rocket Launcher consisting of a wooden trough, 7 feet long, held in an inclined position by a bipod (p 205 & Fig 329 on p 209)

20-cm Rocket Launcher consisted of a steel barrel (tube) mounted on 10-ft wheels (p 205 & Fig 330 on p 209)

45-cm Rocket consisting of a wooden trough mounted on wheels. Was good for one shot only (p 205 & Fig 331 on p 209)

Special Mark 1 Model 21 Launcher for 250-kg Rocket Bomb consisted of a wooden trough 22 feet long (p 25)

12-cm Incendiary Shrapnel Rocket Launcher consisted of a wooden trough installed fore and aft on “Suicide Boats” (qv). Later models were adjustable in elevation (p 25)

In the book of Tantum & Hoffschmidt (Ref 8), the following Rocket Launchers and Rocket Projectiles are described and illustrated: 20-cm Rocket Projectile and Launcher (p 171)

The ground-launched Rocket, 21.05cm in diam, described here under ROCKETS was fired from a trough-shaped launcher ca 7 ft long, which weighed ca 175 lbs. Range of 1970 yds at 50 degree elevation was claimed (See also Ref 1, p 205)

Rocket Launcher and Rocket Motor Model 10 (p 172) was designed to propel the 60-kg aircraft bomb out of an inclined trough. The launcher was constructed of wood and metal with legs made of iron pikes. The launcher channel was a right angle wooden trough, ca 20 ft long with a motor and bomb positioner
made of 1/8-inch pierced sheet metal. The rear of launcher was attached by a pin to a base plate. Elevation was controlled by cables run from the base plate to the legs, and between them.

The Rocket Motor resembled a blunt short-bodied, fin-stabilized, bomb provided with a venturi tube at its rear. The propellant (12.94 lbs) which consisted of three cylindrical sticks tied in a silk bag, was in a cylindrical container located above the venturi tube. The chge was ignited by an Igniter Pad and an Igniter Fuze in the forward part of the motor by means of wires leading to a small Hand Blasting Machine. When fired the motor propelled the Rocket from the launcher and then dropped off. Ranges up to 1300 yds were claimed, but accuracy was very poor.

These devices are described in Ref 2, pp 120–21, Fig 91), under the title “Type 3 and Model 10 Bomb Launching Devices”. The description includes the compn of propint used in Rocket Mortar as: NC 65, NG 30, Et Centralite 3.0 & NaCl 2.0%. It was code-designated by the Japanese as “343 DT₂” and was ignited by an electric squib.

Rocket Motors (Roketto Hasshaki) are devices designed to provide propulsive power (propel or launch) to a Bomb or Rocket Projectile of an inclined trough or barrel called Launchers.

One of such Rocket Motors is described and illustrated in the book of Tantum & Hoffschmidt (Ref 7, p 172) under the title “Rocket Launcher and Rocket Motor Model 10”. It is briefly described here under Rocket Launchers. They are also described in Ref 2, pp 120–1.

Another Rocket Mortar (Type 4 Mk 1) is described here under ROCKET BOMBS as a device used to propel “Baka” Piloted Rocket Bomb (Ref 2, p 118).

There was also “Special Mk 1 Rocket Motor Model 21” which consisted of a cylinder 80.3 inches long & 11.8 inches in diam with rounded nose and a venturi tube attached at the rear. This unit was designed as an integral type bomb launcher for 250-kg Bombs from ground launchers for land bombardment. Its motor weighed 429.2 lbs, while wt of propint was 178.2 lbs, consisting of 20 sticks of “343 Special DT₂”, which contd NC 65, NG 30, Et Centralite 3.0 & NaCl 2.0%. It was ignited by Electric Igniter with BkPdr charge. The unit was launched from a crude wooden trough mounted on a bipod forward and a steel base plate aft (Ref 2, pp 119–20, Fig 90).

ROCKETS (Roketto) or ROCKET PROJECTILES (Roketto Dangan)

Japanese Rockets are subdivided into Army Rockets (Ref 1, p 204 & Ref 3, pp 371–72) and Navy Rockets (Ref 1, pp 204–05 & Ref 3, pp 514–16).

The following Rockets are described with illustrations:

Type 4 20-cm Army Rocket was spin-stabilized, cylindrical in shape, 37 inches long, (including ogive nose) weighing 180 lbs and contg HE chge of TNT and Ballistite as propellant (Ref 1, p 204) 20-cm HE Army Spin-Stabilized Rocket, cylindrical in shape, 38.58 inches long (including ogive nose), weighing 186.3 lbs and contg cast TNT as HE Filler and Smokeless Double-Base Powder B, described here as item d, under Propellants (Ref 3, pp 371–72).

Note: This Rocket seems to be identical with the one described in Ref 1, p 204.

Navy Rockets were HE (two kinds) and Incendiary (one kind). They are listed in Ref 1, pp 204–05, but not described. The following are described in Ref 3:

12-cm Incendiary-Shrapnel Spin-Stabilized Rocket was cylindrical in shape with ogive nose, overall length ca 28.5 inches, weight 51 lbs, which included Fuze, 4.75 oz PA burster chge, 10 lbs 10.75 oz of Incendiary cylinder contg WP pellets, 7 lbs 5 oz Special DT₆ Propellant (listed as item g under PROPELLANTS) and Ignition Mechanism. It was launched from a wooden trough mounted one on each side of the cockpit of a Suicide Boat (qv) (Ref 3, p 514, Fig 423).

20-cm HE Spin-Stabilized Rocket had an artillery-projectile shape body, overall length (w/o Fuze) 40.28 inches, weight ca 200 lbs, which included: Fuze 38.58 lbs cast Type 91 Expl (TNAns), Fuze, Propellant (18.3 lb) Special DT₂ (no data) or Special DT₆ (listed as item g under PROPELLANTS) and Ignition Mechanism. Three signal-round launchers were designed for this Rocket. Two of them were trough-type, while the 3rd was a barrel type launcher. In addn a triple-mount trough was used (Ref 3,
Same Rocket is described in Ref 8, p 171
45-cm HE Spin-Stabilized Rocket was cylindrical in shape with conical nose, length (w/o Fuze) 67.5 inches, weight 1514 lbs, which included 401.0 lbs of Type 98 Expl (PA), 131.5 lbs Special DT6 Propn (listed as item g under PROPELLANTS) and Ignition system. A crude wooden-type launcher was used with this Rocket (Ref 3, p 516, Fig 425)

Roketto. See ROCKETS

Roketto-shiki tekidan. Antitank (A/Tk) Rocket

Ryūdampō. Howitzer

Ryūsandan. Shrapnel Shell

SABOTAGE DEVICES (Bōgyō sōchi; Hasai bōryaku).

Under this term may be included any item intended to produce malicious injury to people, machinery, tools, etc; practiced mostly during the wars.

The following Japanese Sabotage Devices are described in Ref 2, pp 258-63:

Explosive Toothpaste consisted of a toothpaste tube filled with 4.23 oz of a mixture of RDX 80.2% & mineral oil with wax 19.8% and provided with an Igniter (p 258, Fig 195)

Explosive Coal consisted of thin earthenware containers of irregular shape coated with a bituminous type paint. Each piece was filled with RDX and contd an Igniter with Detonator and a small chge of BkPdr. When exposed to fire the heat eventually ignited the BkPdr chge, which in turn set off the Detonator and RDX (p 259, Fig 196)

Explosive Food Cans consisted of regular tin cans with American labels such as “Libby’s Strawberries”. They contd 1.37 lbs of a mixt of RDX 78.3% & mineral oil 21.7% and were provided with an Igniter, Delay and Detonator (pp 260-61, Fig 197)

Metal Incendiary Cylinders were 6½ inches long and 2½ inches in diam filled with Thermite (Incendiary) and provided with two Igniters, Delay and First Fire Charge consisting of KClO₃, Sb₂S₃ & Al pdr (pp 261-62, Fig 198)

Incendiary Brick consists of an incendiary mixt of K chlorate, sulfur, ground coal (or sugar), iron filings and wax compressed in the shape of a regular US brick. It was coated with paint to give it the appearance of a glazed brick. As there was no Igniter, one of the methods used to ignite the brick was to place some K permanganate on top of it and pour conc sulfuric acid over it (p 262, Fig 199)

Incendiary Soap consisted of a mixture of Ba nitrate 30.4, paraffin 19.4, Mg 11.3, Al 11.1, rosín 10.9, ferroso-ferric oxide 9.1, NC 4.4 & gritty siliceous material 2.6%, compressed into a regular soap bar. Its method of ignition by the Japanese is not known.

Following items which are actually Sabotage Devices are described in Ref 1, pp 231-32:

Sucker Traps were devices set in living quarters, abandoned equipment, etc, mainly for the purpose of lowering the enemy’s morale. Besides the various Booby Traps described here under MINES, the following items were used by the Japanese:

Explosive Radio, contd HE charge occupying the space obtnd after removing all batteries except one. When the switch was turned on, the electric circuit closed (p 231, Fig 377)

Explosive Phonograph. An electric contact on the pick-up was so arranged that sufficient movement of the arm to play a record would close the circuit and set off the charge of explosive concealed with battery under floorboard (p 231, Fig 378)

Explosive Telephone. A pull-igniter fitted to an expl chge was placed inside a telephone. The line from the igniter was secured to the crank of the telephone so that an attempt to ring would set off the charge (p 232, Fig 379)

“Tin Can” Booby Trap consisted of a regular American tin can which was filled with expl chge and provided with a friction pull igniter and trip wire (p 232, Fig 380)

Safety Fuze. Dōkusaku

Safety Pin. Anzusen

Sairetsu teryūdan. Fragmentation Grenade

Sairetsu tōka bakudan. Fragmentation Bomb

Sakuma Dainamaito. One of the Japan Dynamites
Sakura No 1 & No 2. Blasting Gelatin

Sakuretsu. Burst, Explosion

Sakuyaku. Bursting Charge

Sampōhei. Mountain Artillery

Sandanjū. Shot-gun

Sanzōki mechi nitroamin (Tetryl). See Meiayaku

Sanshōki toruoru or Type 92. See Chakatsuyaku

Seacoast Artillery. Kaiganho

Sea Markers. See under Smoke Floats and under Markers and Indicators

Seigata (Army), Type 97H (Navy) (German Hexanitrate). A light yel. or orn. castable expl mixture of TNT 60 & HNPPhA (Hexanitrodiphenylamine) 40%. Density 1.65 (cast), Brissance by Cu cellar crush test 96% of PA; Expln Temp 302°; Impact Sensitivity, 14cm max for no explns with 5-kg wt; Power by Ballistic Mortar 92% PA; Rate of Deton 7100m/s vs 6900 for TNT. It was used at the beginning of WWII for loading some Torpedo Warheads, but later replaced by A(ko) (Type A Expl); was also used in some Depth Charges and Navy Rockets (Ref 1, p 32 & Ref 5, p 370)

Self-Propelled Guns. All guns mounted on combat vehicles may be considered Self-Propelled Guns. These include Tankettes (Ref 8, pp 117–19), Tanks (p 120–29) and Armored Cars (pp 132–35). Only one of these is listed as “self-propelled”. It is 150-mm Selbst-Propelled Howitzer, Type 38 Year (1905) which is mounted on medium tank chassis. Its detailed description and illustration are given on p 129 of Ref 8

Semimetal Bomb. Koseisen bakudan

Sensahō. Tank Gun

Sensō. War

Sensuikan. Submarine

Sentai. Groove, Rifling

Sentō. Battle, Combat

Sento bakugekiki. Fighter Bomber

Sentōkan. Battleship

Shadan. Projectile

Shakunetsuzai (Thermite). An incendiary mixture of Al & Fe₃O₄, which can be ignited by Mg. Used in admixture with BlPdr or HE’s for loading some incendiary items of Ammunition, while the majority of such items contd WP (White Phosphorus). Following incendiary items contd Thermite: 1-kg and 5-kg Thermite Incendiary Bombs (Ref 2, p 15)

Type 98 No 7 Mk 6 Model 1 Incendiary Bomb (Ref 2, p 71)

Metal Incendiary Cylinders, used as a Sabotage Device (Ref 2, p 261)

Shakyorii. Range

Shaped Charge. See Hollow Charge [Ta(dan)]

Shell. Hōdan

Shikan. Adapter

Shimose Bakuyaku (Navy) or Ōshokuyaku (Army)

Shingōdan. Pyrotechnics

Shingō kasen. Signal Rocket

Shingō kenjū. Signal Pistol, Very Pistol

Shinkan. Fuze

Shinkan kibakuza. Fuze Detonator

Shin-kiri or Shinkoryokki. Gelatin Dynamite

Shinkō. Cavity (of Projectile)

Shinshoku. Erosion, Corrosion

Shintetsu. Penetration (of Projectile)
Shin-toku shoan. Ammonium Nitrate Dynamite

Shiraume. Gelatin Dynamite

Shoan bakuyaku. An AN (Ammonium Nitrate) explosive compn used in Army Substitute Demolition Charges: AN 79, DNN (Dinitrophenol) 10, sawdust 1 & NaCl 10%. TNT could replace DNN (Ref 1, p 29 & Ref 5, p 371)

Shoanbakuyaku Nos 104 & 201. Ammonium Nitrate Dynamite

Shoanyaku. A series of coal-mining expls manufactured at the Uji Factory (Kyoto Prefecture):

No 1. AN 70, DNN 9, woodmeal 1 & NaCl 20%

No 2. Same as Shoan bakuyaku (See previous item)

No 5. AN 64, TNT 12, woodmeal 3, wheat starch 1 & NaCl 13%

No 7. AN 75, DNN 9, TNT 1.5, woodmeal 1.5 & NaCl 13%

Special: AN 64, DNN 3, TNT 7, GuN (Guanidine Nitrate) 2, Na nitrate 2, woodmeal 2 & NaCl 20%

These expls were reported to have Impact Test Value with 2-kg Weight ca 80cm and Gap Test Values 100 to 150mm. Some of the mixtures were used for military purposes (Ref 5, p 371)

Shōdenyaku. See Anbenyaku

Shoeiyaku. Pentaerythritol Tetranitrate (PETN), C(CH₂ONO₂)₄; mw 316.14, N 17.72%; wh crysts, d 1.77, mp 141°; Brisance by Plate Dent Test 129% TNT; Explosion Temperature 225° (decomp in 5 secs); Impact Sensitivity BurMines-App, 2-kg Wt 17cm (vs 100+ for TNT); Power by Ballistic Mortar Test 145% TNT; Rate of Detonation 8300m/sec (Ref 8, p 276). Pressed PETN was used in Army 7.7 & 12.7-mm Fuzelss Projectiles and 20-mm MG Projs. Also in Boosters. Its mix with TNT is called Pentoriri (qv). PETN with 8.5% wax was used for loading 20-mm Shells. Its mixtures with RDX were used in 7.7 & 12.7-mm Projectiles. PETN was also used in Incendiary Mixtures (Ref 1, p 27 & Ref 5, p 372)

Shōgeki. Impact; Shock

Shōgeki shikan. Impact Fuze

Shōdan. Incendiary Shell

Shōyo bakudan. Incendiary Bomb

Shōyo teki dan. Incendiary Grenade

Shōji. Rifle

Shokaki. Small Arms

Shokamen. Nitrocellulose (NC)

Shokayaku (Nitrocellulose Propellant). See under PROPELLANTS

Shokuhatsu jirai. Contact Mine

Shomeidan. Illuminating Shell; Parachute Flare

Shonayaku. Yel-brn expl mixture of AN 90 & DNN 10% used for loading cast semisteel Shells, Land Mines and Demolition Charges. It was suitable for tunnel work because it did not give toxic fumes (Ref 1, p 27 & Ref 5, p 372)

Shōsan. Nitric Acid

Shosoku. Muzzle Velocity

Shotgun. Sandanji

Shotoyaku (Amatol). A mixture of AN 50 & TNT 50% of brown color, d (cast) 1.59, mp 81°, Brisance by Plate Dent Test 52% TNT; Expln Temp 265° (dec in 5 secs); Impact Sensitivity BurMinesApp 2-kg Wt 95cm; Power by Ballistic Mortar 124% TNT; Rate of Detonation 6430m/sec (Ref 8, p 16). Was used in same* Bombs and in Semisteel (high C cast iron) Shells. Was reported to be found in some Navy Mines (Ref 1, p 27 & Ref 5, p 372)

Shoyaku or Tanayaku (RDX or Cyclonite), O₂N₂N—CH₂—N.NO₂

H₂C—N(NO₂)—CH₂; mw 222.13, N 37.84%
wh crys, d 1.82, mp 204°; Brisance by Plate
Dent Test 135% TNT; Expln Temp 260° (dec
in 5 secs); Impact Sensitivity, BurMinesApp,
2-kg Wt 32cm; Power by Ballistic Mortar 150%
TNT; Rate of Deton 8180m/sec (Ref 8, p 60).
Straight RDX was used as Booster and Sub-
Booster in some Army Fuze, and as Bursting
Cgres in 20-mm HE Shells. When desensitized
with wax, it was used in 37-mm HE Shell
(Ref 1, p 26 & Ref 3, pp 282 & 290)
The following expl comps, contg RDX,
Angayaku, Chauyaku, Haishokuyaku, Koshtsu,
Nigotanoyaku-Mc 2, Tanôyaku, Type 94M
Explosive and Unknown Name Explosives were
used as Bursting Charges of Bombs and Shells
(Ref 5, p 373)

Shrapnel Shell. Ryûsandan

Single-Base Propellants. See under PROPELLANTS

Smoke. Kemurie (en)

Smoke Agent. Hantsuenzai

Smoke Bomb (Hatsuen bakudan). One of such
bombs is described here under “Markers and
Indicators” and also in Ref 2, p 113-R as Type
2 2-kg Target Indicator. A larger Bomb,
weighing 155 lbs, called Type 3, No 6 Target
Marker Bomb is also described here and in
Ref 2, p 113-L

Smoke Candles (Hatsuentô). See under Smoke
Pyrotechnics

Smoke Floats are devices serving as “Sea
Markers”. Following are examples:
2-kg Smoke Float was a bomb-resembling de-
vice, weighing 4.75 lbs, thrown by hand. It
contd an igniter and a smoke-producing compn
(no data) (Ref 2, p 109, Fig 82)
43-kg Smoke Float consisted of long conical
nose and cylindrical tail contg a parachute.
Its nose section contd smoke compn and an
igniter. It was thrown from a plane (Ref 2,
p 110, Fig 83) (See also under Smoke Pyro-
technics)

Smoke Generator (Hatsuenku). See under Smoke
Pyrotechnics

Smoke Grenades. See under Grenades

Smokeless Powder (Moenkayaku). See under
PROPELLANTS

Smokeless Powder B. See item d under PRO-
PELLANTS

Smoke Powder. Yuenyaku (Nonsmokeless
Powder) (Ref 1, p 27)

Smoke-Producing Materials. Following are
examples of materials used by the Japanese
for producing smokes:
a) FS Smoke, chlorosulfonic acid 41, sulfur
trioxide 54 & sulfuric acid 5%. Used in Type
100 50-kg Smoke Bomb (Ref 2, p 21)
b) Mixture of titanium & silicon tetrachloride;
used in Frangible Smoke Grenade (Ref 2, p 239)
c) Mixture of hexachloroethane (HC) 56.2,
Zn 27.6, Zn chloride 2.9 & Zn oxide 13.3%;
used in Smoke Rifle Grenade (Ref 2, p 243)
d) White phosphorus (WP), used in Type 90
75-mm Smoke (WP) Projectile (Ref 3, p 336)
e) WP in Type 13-Year 150-mm Smoke (WP)
Projectile (Ref 3, p 367)
f) Berger Mixtures, probably similar to the ones
described in Vol 2 of Encyc, p 102-L, were used
in Smoke Candles listed in Ref 1, p 255 (Compn
of Japan mixture not given)
g) Hexachloroethane (HC) Mixture used in Type
95 50-mm Smoke Mortar (Ref 3, pp 3 & 14)
h) WP, presumably was used in 140-mm Navy
Smoke Projectile (Ref 3, p 494)
i) FM (Titanium Tetrachloride) used in Navy
Type 2 Target Indicator (Ref 1, p 246)
j) Tannic Acid smoke compn used in Type 94
4-kg Practice Bomb (Ref 2, p 28)

Smoke Projectile or Shell (Hatsuentan). See
under AMMUNITION and in Ref 3, pp 336, 367,
374 & 494

Smoke Projector. Emmaku hōshaki

Smoke Pyrotechnics (seizo-jittsu). The following
devices are described in Ref 1, pp 243–46 and 255
Army Smoke Candles. Exclusive of self-project-
ning and floating smoke candles, all smoke candles
were of the same basic construction, differing
only in size and filling. This type of candle con-
sisted of a cylindrical sheet metal container with a removable metal cover held in place by adhesive tape. The cover protected a match head in the top of the candle and contained a wooden scratch block. To ignite the candle, the scratch block was rubbed against the match head which, after a few seconds delay, ignited the Main Filler. The candle was thrown to, or placed at the spot where smoke was required (Ref 1, p 143). A list of Smoke Candles 1/2-kg, 1-kg and 10-kg is given on p 255. As Main Fillers Berger Mixture, HC (Hexachloroethane) and other smoke-producing mixtures were used.

Army Self-Projecting Candles usually consisted of an outer tube sealed at one end by a wooden block and provided with protective metal covers held on each end with adhesive tape. The smoke mix was held in a snug-fitting inner container. A match head in the block at the base of the candle was ignited by the scratch block contd in the top metal cover. This ignited a delay fuse which set off the propellant charge and shot the inner tube container outward. A delay element was ignited by the expln and in turn set off the smoke mix. A rod encircled the candle and extended along the length of the candle to form a spike which was inserted into the ground to support the candle at the desired angle for firing (Ref 1, p 243). Same smoke mixes are listed in Ref 1, p 255.

Army Type 94 Floating Smoke Candle consisted of a cylindrical metal tube 314 inches long and 3/4 inches diam, equipped with a supporting ring to which an automobile rubber tube was attached. It was inflated to support the candle floating in water. The filling was either Berger Mixture (Type 94A) or HC (Type 94B). A metal top covered the fuze pocket which was closed during shipment. Two types of firing mechanisms were used. One type was the ordinary hand grenade igniter without a blasting cap detonator; this gave a delay of 8 secs. The other type was a friction igniter initiated by a small piece of wood painted with abrasive. This produced a delay of up to 30 secs. The candle burned for ca 3.5 mins, producing a dense white smoke which hovered close to the surface of water (Ref 1, p 243, Fig 401). Navy 40-kg Floating Smoke Generator consisted of a welded steel drum 14 inches high & 12 inches in diam, equipped with carrying handles, a filling hole, a fuze with ignition system, and a spray tube which extended inside to the bottom of the drum. An inflated rubber tube was attached to the lugs around the drum to float it in water. In operation, the safety pin in the firing assembly was removed and the plunger was struck to fire the detonator and ignite the fuse, a wick, and start the combustion of a slow-burning charge. As pressure from the burning built up in the upper part of the drum the FS smoke mix (See item 12 under Smoke-Producing Material) in the lower part was forced out thru the emission tube. Upon contact with air, a dense white smoke was formed (Ref 1, pp 244-45). Navy Smoke Float had an aerial bomb body, ca 18 inches long, with three tail fins. It was filled with yellow or black smoke mixtures, an igniter and a small ejection charge. It was dropped from a height less than 700 meters to serve as an aircraft navigation marker (p 245, Fig 403). Navy Floating Smoke Flare had a long conical nose section and shorter cylindrical tail section with parachute. Overall length 51 inches. The body contd an ejection charge, pull igniter, delay train and a smoke mixture. It was dropped from an aircraft and a parachute slowed the descent of the flare and the delay train permitted the flare to hit the surface of the water before the smoke mix was ignited (pp 245-46, Fig 403).

Navy Type 2 Target Indicator had a bomb-shape thin bakelite body, 17 inches long, 3 inches in diam, filled with FM (Titanium Tetrachloride) smoke mix. The bomb had no fuze because its light body was designed to break (when dropped from a high altitude) on hitting water, thus releasing the smoke mix (Ref 1, p 246 & Fig 404 on p 245).

Smoke Screen. Emmaku

Smoke Signal. Hatsu sen shingō

Smooth-Bore. Kakkō no

Sōkōsha. Armored Vehicle

Spigot Type Mortar. See under MORTARS and in Ref 3, p 389 & Ref 7, p 170.

Stabilizing Fin. Dan-yaku

Stick Charges, 50-mm were projectiles fired from Type 98 Discharger. They consisted of
square cast-iron boxes mounted on wooden poles. Each box contained blocks of PA (Picric Acid) with two Pull Igniters. When the projectile is fired these pull igniters are initiated and in turn set off the bursting charge after a short delay. A Bk Pdr propinth in silk bags, independent of the projectile, is also fired by a pull igniter inserted in the side of the launcher tube. Stick charges may be considered as a special type of army mortar (Ref 3, pp 377–78)

Submarine. Sensuikan

Sucker Traps. See under Sabotage Devices and in Ref 1, pp 231–32

Suicide Boats (Army & Navy). The craft were lightweight plywood, gasoline-driven speedboats 15 to 20 feet long. The Army boat (Ref 1, Fig 39 of p 65) had mounted on its sides two 120-kg depth charges, which released when the boat struck its target. Some boats had a 3rd depth charge mounted over the stern.

The Navy boat, shown in Fig 40 had a 640-lb charge of Type 98 Expl charge built into the nose. The boat was exploded electrically upon contact with the target, or by throwing a switch in the cockpit. A stand-by pull igniter firing device was also fitted into the charge

Suirai hashakan. Torpedo Tube

Suiraitel. Torpedoboot

Sulfuric Acid. Ryūsan

Taihō. A/Tk Gun [Kahō-Gun (general)]

Taijinteki jirai. Antipersonnel Mine

Taikū. Antiaircraft

Tail Fin. Anteiban

Tail Fuze. Danbei shinkan

Taisenshadan. Antitank Bomb

Taisenshatunshindan. Antitank Rocket

Taisensha funshin toshaki. Antitank Rocket Launcher

Taisensha jirai. Antitank Mine

Taka Nos 1, 2 & 3. Gelatin Dynamites

Tama. Bullet

Tanayaku (?). See Shouyaku

Tankei. Ogive

Tanken. Dagger

Tanks (Tanku) and Tankettes are listed here under Combat Vehicles and described, with illustrations, in the book of Tantum & Hoffschmidt, pp 116–29, listed as Ref 7

Tank Gun. Senshabo

Tanōyaku. See under Nigotanoyaku and in Ref 5, p 373, where No 1 consists of RDX 60, TNT 30 & Tetryl 10%, and No 2 consists of RDX 55, TNT 38 & Tetryl 7%. They were used as cast-loaded carton charges for various projectiles

Tekidan. Grenade (qv)

Tekidanki. Rifle Grenade Launcher

Tekidantō. Grenade Discharger

Tekishahō. Howitzer. See under ARTILLERY

Tekkodan. Armor-Piercing Projectile

Tenagedan. Hand Grenade. See under GRENADES

Tenka bakkan. Igniting Primer

Tanka shikan. Igniting Fuze

Tenkayaku. Igniting Charge

Terumitto (Thermite). See Shakunetsuzai

Terumitto shoidan. Thermite Bomb

Teryudan. Tenagedan

Tessen keitei. Trip Wire
Tetryl. See Melayaku

Thermite. Terumitto. See Shakunetsuzai

Thermite Bomb. Termitto shōidan

Thermite Grenade. Termitto tekidan

TNAln (Trinitroanisol). See Type 91 Explosive (Navy)

TNPhnt (Trinitrophenol). See Heineiyaku

TNT (Trinitrotoluene); Sanshōki torūoru or Type 92 Explosive (Navy). See Chakatsuyaku

Tō. Sword

Tōka hakōdan. Armor-Piercing Bomb

Tokai bakudan. Time Bomb

Tokeishiki bakkan. Mechanical Time Fuze

Toku shiraume Nos 1, 2 & 3. Gelatin Dynamites

TORPEDOES (Raigeki suru or Gyora). Accdg to Ref 1, p 54, the Japanese Torpedoes of WWII were of designs developed during 20 years of research. They were unrivaled in speed and range, yet carrying an extra weight of explosives. They had cigar-shaped bodies and usually were steam-driven. Besides the HE filling and engine, they carried machinery necessary to keep the torpedo on a set course and at a set depth to its target. A typical torpedo is shown in Fig 25 on p 54

A table on p 60 of Ref 1 lists torpedoes used before and during WWII, of which several Types are marked “obsolete” or “obsolete”:

Type 44 Mk 2 Mod contd 400 lbs of Type 97 or 98 Expl (Obsolete)

6th Year Type contd 450 lbs of Shimose (obsolete)

8th Year Type Mk 2 Mod 2 contd 850 lbs of Shimose or Type 98 Expl (Obsolete)

Type 89 Mod 1 contd 625 to 660 lbs of Type 94, 97 or Shimose (Obsolete, including its Mod 2)

Type 90 Torpedo contd 880 lbs of Type 94 or Shimose (Obsolete; replaced by Type 93)

Type 93 Model 1 Mod 2 contd 1080 lbs of Type 97 Expl; was 29 ft 6 inches long & 24 inches in diam; used in destroyers and cruisers. It was the standard Navy torpedo and a very superior weapon. There was also a Model 3 carrying 1700 lbs of Type 97 Expl

Type 91 Mods 1, 2, 3, 3 (Special), 4, 6 and 7 were standard aircraft torpedoes of diam 17.7 inches and 17 ft 3 inches to 18 ft 10 inches long which carried 340 to 900 lbs of Shimose, Type 94, 97 or 98 Expl

Type 92 Mod 1, used in submarines, was copied from the Germans and contd 660 lbs of HE (no data). It was electrically driven, while other Japanese torpedoes were steam-driven

Type 94 Mod 1, used in submarines, was 22 ft long & 21 inches in diam. It contd 870 lbs of Type 97 or Shimose

Type 94 Mod 2, used in aircraft, was 17 ft 4 inches long and 17.7 inches in diam. It contd 460 lbs of Shimose

Type 95, apparently the most common submarine weapon, was 23 ft 6 inches long and 21 inches in diam. It contd 880 lbs of Type 98 Expl

Type 97, used in midget subs & PT boats, was 18 ft 5 inches long and 17.7 inches in diam. It contd 790 lbs of Type 97 Expl

Type 2, used in midget subs, was of the same size as Type 97. It contd 774 lbs of Type 97

Type 2 (Special) was an aircraft torpedo, which carried 650 lbs of Type 97

No torpedoes are listed in Refs 2, 3, 5 and 7

Torpedo. Suirairei

Torpedo Tube. Suirai hoshakukan

Torpedo Warhead. Suirai-bakuhatsusen-tōbu

Tracer (Eikōdan) and Tracer Ammunition (Eikōdan-yaku). Many projectiles contg tracers are listed in Ref 3, but no compns of tracers are given

Trajectory. Danô

Trinitroanisol (TNAln). See Type 91 Explosive
Trinitrophenol (TNPht). See Heinelyaku

Trinitrotoluene (TNT). See Chakatsuyaku or Type 92 Explosive (Navy)

"TYPE" Explosives. A series of explosives used by the Japanese Navy, which are listed in Refs 1, 5 and below:

Type A (Explosive) or A (ko). See Otsu-B
Type K (Explosive). A series of expls based on mixtures of inorganic salts, developed during WWI. Their compns are not given but they are probably similar to German Ersatzsprengstoffe (Substitute Explosives), listed in Vol 5 of Encycl, pp E121–E122

Type 1 (Explosive). A greenish-yel expl compn consisting of Ammonium Picrate 81, AI 16, sawdust 2 & heavy oil 1%. Its Brissance by Sand Test was lower than that of TNT; Expln Temp 475°C; Impact Sensitivity, comparable to TNT; Power 138% TNT & Rate of Deton 4300m/sec. It was used as press-loaded chges for Depth Bombs or other Underwater Ordnance (Ref 1, p 32 & Ref 5, p 374)

Type 1, Mk 5, P5 (Explosive). A dark-grn expl compn consisting of Amm Picrate 81, ferrosilicon 16, woodpulp 2 & heavy oil 1%. Its apparent density was 1.16; Brissance by Cu Cylinder Crusher Method 99% of PA; Expln Temp 450°C; Impact Sensitivity 15cm (maximum for no explns with 5-kg wt); Power by Ballistic Mortar 72% PA & Rate of Deton 4100 m/sec. Used press-loaded in some Depth Charges (Ref 5, p 374)

Type 1 Mk 6, P6. A dark-grn expl compn consisting of Amm Picrate 86, ferrosilicon 11, woodpulp 2 & heavy oil 1%. Its apparent d was 1.13; Brissance 95% PA; Expln Temp 450°C; Impact Sensitivity 13cm (max for no explns with 5-kg wt); Rate of Deton 4620m/sec. Used press-loaded in Depth Charges (Ref 5, p 374)

Type 2 (Explosive). See B4 (Incendiary)

Type 4 (Explosive) (Navy). A gray expl compn consisting of Amm Perchlorate 79.2, ferrosilicon 16.4, Al & Fe pdr 1.0, heavy oil 2.5 & unaccounted 0.9%. Its properties are not given, but it was stated in Ref 5 that they were similar to Type 88 Explosive (qv), except that it did not burn as easily. Used in Depth Charges (Ref 1, p 32 & Ref 5, p 375)

Type 4 Mk 1, K1. A lt-grayish expl compn consisting of Amm Perchlorate 80, talc 10, ferrosilicon 8 & chloronaphthalene 2%. Its apparent d was 1.24; Brissance 89% PA; Expln Temp 455°C; Impact Sensitivity 18cm (max for no explns with 5-kg wt); Power by Ballistic Mortar 84% PA & Rate of Deton 3600m/sec. Used press-loaded in Sea Mines and Depth Charges (Ref 5, p 375)

Type 4 Mk 2, K2. A gray expl compn consisting of Amm Perchlorate 89, woodpulp 5 & coal tar 6%. Its apparent d was 1.0; Brissance 70% PA; Expln Temp 470°C; Impact Sensitivity 34cm; Power 82% PA & Rate of Deton 3900m/sec. Used press-loaded in Sea Mines and Depth Charges (Ref 5, p 375)

Type 4 Mk 3, K3. A lt-gray expl compn consisting of Amm Perchlorate 47, Amm Sulfate 32, ferrosilicon 20 and chloronaphthalene 1%. Its apparent d was 1.20; Brissance 83% PA; Expln Temp 470°C; Impact Sensitivity 20cm; Power 81% PA and Rate of Deton 2900m/sec. Used press-loaded in Sea Mines and Depth Charges (Ref 5, p 375)

Type 4 Mk 4, K4. A white expl compn consisting of Na Chlorate with K chlorate 88 & petroleum oil 12%. Its appd d was 1.05; Brissance 77% PA; Expln Temp 580°C; Impact Sensitivity 14cm; Power 81% PA and Rate of Deton, not given. Used press-loaded in Sea Mines and Depth Charges (Ref 5, p 376)

Type 4 Mk 5, K5. A gray expl compn consisting of Amm Perchlorate 55, Amm Nitraxe 29, ferrosilicon 10, woodpulp 5 & heavy oil 1%. Its apparent d was 1.05; Brissance 92% PA; Expln Temp 450°C; Impact Sensitivity 28cm; Power 81% PA; and Rate of Deton, not given. Used in Sea Mines & Depth Charges (Ref 5, p 376)

Type 4 Mk 6, K6. A gray expl compn consisting of Na Chlorate 84, woodpulp 6, & petroleum oil 10%. Its appd d was 1.35; Brissance 78% PA; Impact Sensy 17cm; Power 67% PA and Rate of Deton, not given. Used in sea mines & Depth Charges (Ref 5, p 376)

Type 4 Mk 7, K7. A dark-gray expl compn consisting of Na Chlorate 84, woodpulp 5, charcoal 8 & coal tar 3%. Its appd d was 1.2; Brissance 82% PA; Expln Temp 385°C; Impact Sensy 25cm; Power 63% PA and Rate of Deton, not given. Used in Sea Mines & Depth Charges (Ref 5, p 376)

Type 88 (Explosive) (Navy) or Karitto (Army).
A gray expl compn consisting of Amm Perchlorate 66, silicon carbide 16, woodpulp 12  & petroleum 6%. Its props are not given, except that it was stated in Ref 1, p 31, that it was dangerous to burn and very sensitive to friction. In Ref 5, p 377 it was stated that it was nonhygroscopic, stable in storage and produced on burning very poisonous fumes.

The Navy used this expl compn loosely packed in Mines and Depth Charges, while the Army used it under the name of Karitto in Subtittute Demolition Charges. It was proposed to use it also as a Solid Rocket Propellant.

This expl compn was used under the name of Carlit for industrial purposes and is described in Pamphlet of "The Japan Carlit Co, Ltd" located before WWII in Chiyoda-ku, Tokyo. There are several formulations of Carlit, but all of them contain some Amm Perchlorate, but none 66%. (Pamphlet of Carlit Co and Ref 5, p 377)

Type 88 (Ko) or Haenososan-bakuuyaku, called by "The Japan Carlit Co, Ltd" Carlit Kuro (Black), was a gray expl compn consisting of Amm Perchlorate 75, ferrosilicon 16, woodpulp 6 & heavy oil 3%. Its density was 1.05; Brisance lower than that for TNT; Expln Temp 430°; Impact Sensitivity 17cm (max for no explns with 5-kg wt); Power higher than TNT; and Rate of Deton 4400m/sec. Judged by its high Impact Sensitivity and high Friction Sensitivity, 30 to 40 kg (max pressure betw two rubbing surfaces for no explns), instead of usual 60 kg, it was found unsuitable for loading HE Shells, but was suitable for press-loading Navy Mines.

Was used as an industrial expl by The Japan Carlit Co, Ltd, Tokyo (Ref 5, p 377 and Pamphlet of Carlit Co, Table listing properties)

Type 91 (Explosive) (Navy); Trinitroanisole (TNAs); Methyl Picrate, or 2,4,6-Trinitrophenylmethyl Ether, H₃C.O.C₆H₄(NO₂)₃; mw 243.13, N 17.28%; yel crystals, d 1.7 & 1.6 for cast; mp 68.4° (pure), 65–67° (coml); Brisance by Cu Cylinder Crusher Test 92% PA or 100% TNT; Expln Temp 279°; Friction Sensitivity 60 kg (max pressure betw two rubbing surfaces for no expln); Impact Sensitivity with 5-kg wt 19cm; Power by Ballistic Mortar 101% PA; Rate of Deton 6660m/sec at d 1.59

It was widely used straight for cast loading Bombs and Shells as replacement for Shimose (melting at high temp of 122.5°). Was also used for composite expls: A (ko) or Type A (Explosive), described here as Otsu-B; Type 2 (Explosive), described as B4 (Incendiary); E (Explosive); Type 98 (Explosive), described here as H2 Kongo; Type 94M (Navy) and some other expl comps (Ref 1, p 32, Ref 5, p 377 and Vol 1 of Encycl, pp A450–451, under Anisole)

Type 92 (Explosive). This term is applied by the Navy to straight TNT which was used in Naval 25-mm & 40-mm Shells. TNT was described here as Chakattuyaku.

The term Type 92 is also applied to a compn consisting of TNT 66 & Al pdr 34% which was cast-loaded in Navy 25-mm Shells. This mixture originated in Germany under the name of Tritonal. A similar mixture, but contg TNT 80 & Al 20% was used in the US and described in AMCP 706-177 (1971), p 386 (See Ref 1, p 33 & Ref 5, p 378). Example of use of 66/34 mixture in 25-mm HE Proj is described in Ref 3, p 447

Type 94M (Explosive) (Navy). A cream-yellow expl compn consisting of Trinitroanisole 60 & RDX 40%. Its cast d was 1.64; Brisance by Cu Cylinder Crusher Method 107% PA; Expln Temp 216°; Friction Sensitivity 40–50 kg (max pressure betw two rubbing surfaces for no explns); Impact Sensitivity 13 cm (max for no explns with 5-kg wt); Power by Ballistic Mortar 112% PA; Rate of Deton by Dautriche Method 7700 m/sec. Originated as powerful expl for loading Torpedo Warheads but this use was discontinued due to its sensitivity to Sympathetic Detonation. Later uses included Shaped Charge Grenades and as a Booster Surround (Ref 1, p 32 as Type 94 & Ref 5, p 379 as Type 94M)

Type 97H (Explosive) (Navy) is described here as Seigata (Army)

Type 98 (Explosive) (Army & Navy) is described here as H2 Kongo

Unknown Name Explosives are listed here under X-EXPLOSIVES and also under Japanese Explosives and Related Items Described in Journals & Patents Listed in Chemical Abstracts After WWII
Very Pistol. Veri shingō kenjū

Vomit Gas Projectile was 7cm (75-mm) in diam and contd 0.37 lb of crude diphenylcyanoarsine (See DG in Vol 2 of Encycl, p C167-R) as a vomit warfare agent and 1.02 lbs of TNT desensitized with 30% naphthalene as burster charge (Ref 3, p 341, Fig 268)

War. Sensō

Warhead. Gyorai jitsuyō tōbu

Weapon. Heiki

White Composition consisting of AN, RDX & GuN (Guanidine Nitrate) was used in conjunction with cast TNT in Type 91 10-cm (105-mm) HE Long Pointed Projectile described in Ref 3, p 352, Fig 278

White Phosphorus (WP) (Hakurin). It was used as an Incendiary and in Incendiary Mixtures in Bombs, Grenades and Projectiles, described here under “Incendiaries” and in Refs 2 & 3. WP was also used in Smoke Compositions as described here under Smoke Producing Materials, items d, e, h and in Refs 2 & 3

X-EXPLOSIVES:

X-1. A light gray to tan mixture of KClO₃ & Sb₂S₃ of proportion not given in Ref 1, p 25, but it was the most common mixture for Fuze Primers (See also Ref 5, p 378)
X-2. A chalky-white mixture of PETN 58 & RDX 42% used as filler for Army 7.92-mm and 12.7-mm Ammunition (Ref 1, p 27 & Ref 5, p 379)
X-3. A pressed mixture of RDX 83 & PETN 17% used in 7.7-mm Ammunition (Ref 5, p 379)
X-4. A pressed mixture of RDX 92–95 & wax 8–5% used as a main charge in 20-mm Anti-aircraft Projectiles (Ref 5, p 379) and in Boosters (Ref 3, p 388-R)
X-5. A pressed mixture of RDX 85 & wax 15% used as a Main Charge in 75-mm Armor-Piercing Projectiles (Ref 1, p 27 & Ref 5, p 379)
X-6. A pressed mixture of RDX with Al pdr. Al was evidently added for its incendiary effect (Ref 5, p 379)
X-7. A pressed mixture of PETN & wax. It was used as Main Charge in some Incendiary Shells and as a Booster in 13-mm & 15-mm Shells (Ref 5, p 379)
X-8. A brown expl compr consisting of Amm chlorate 51.5, Ba Nitrste 34.5, Trinitonaphthalene + oil 8.2, woodpulp 5.0 & other ingredients 0.8%. It was used in some Demolition Charges. Dangerous to burn in large quantity (Ref 5, pp 379–80)
X-9. A pressed mixture of RDX & AN (Ammonium Nitrate) used in Type 92 7-cm (70-mm) HE Projectile described in Ref 3, p 304, Fig 236
X-10. A pressed expl compr of RDX 90 & paraffin 10%, coated with graphite and wrapped in Al foil. It was used in Type 1 47-mm and 7-cm (75-mm) Armor-Piercing Projectiles described in Ref 3, pp 299 & 331

Yasenō (Field Gun); Yūgikiteki jirai (Booby Trap). See under MINES

Yuensuku (Black Powder). See Koko Shoku-yaku

Yokosuka Depth Charge. This antiship weapon was not a depth charge in the accepted sense. It was towed under water by escort ships and would explode upon contact with a submerged enemy submarine. The charge was cylindrical in shape, ca 5 ft long & 1 inch in diam, filled with 55 lbs of Type 88 Expl. Vertical and horizontal rudders were fitted on the tail, and an impeller-armed Impact Firing Mechanism was located in the nose (Ref 1, p 65)

References on Japanese Explosives and Related Items Used Before and During WWII
2) Anon, “Japanese Explosive Ordnance” (Bombs, Bomb Fuzes, Land Mines, Grenades, Firing Devices and Sabotage Devices); Dept of the Army and the Air Force, TM 9-1985-4 and TO 398-1A-11, March 1953
4) B.T. Fedoroff et al, “Japanese Ammunition,
Explosives and Weapons”, Picatinny Arsenal Manuscript, Unpublished (1955)
5) O.E. Sheffield, “Handbook of Foreign Explosives”, FSTC 381-5042 (1966), pp 360–387,
US Foreign Science Technology Center, Munitions Building, Washington, DC 20315
6) Anon, “Military Explosives”, Dept of the Army and of the Air Force TM 9-1300-214 and
TO 11A-1-34, Nov 1967, Table 8-2 on pp 8-2 & 8-3 for Explosives and Table 11-7 on p 11-4 for
Propellants
World Combat Series, published in 1968 by WE Inc, Old Greenwich, Connecticut 06870
PATR 1740 (1958), reproduced as Anonymous
AMC (Army Materiel Command) Pamphlet
706-177 (1971)

Additional References on Japanese Explosives
and Related Items
University of Chicago, Chicago, III (1942)
B) M. Wiater, “Examination of Japanese
7.7-mm Ball Ammunition (Model 99)”, Frankford
Arsenal Rept R395 (1943)
C) A. Suzuki, “Crystalline Structure and Explosibility of Furfuril of Mercury”, US Air
Technical Branch Rept 0418 (1944)
D) Anon, “Soldier’s Guide to the Japanese
Army”, Military Intelligence Service, War Dept,
Washington, DC (1944)
E) Anon, “Data on Foreign Explosives”, US
Chemical Warfare Service Field Lab Memo
4-6-2 (1944?) (US Office of Technical Services
PB Rept 11544)
Formate Munition”, US Office of Technical
Services, PB Rept 50950 (1944)
War Dept Tech Manual TM 30-541, US Govt-
PtgOff, Washington, DC (1945)
H) Anon, “Explosives and Demolitions”, US
War Dept Tech Manual FM 5-25, US GovtPtg-
Off, Washington, DC (1945), pp 133–34 (Japanese
Explosive)
US Naval Technical Mission to Japan Dec 1945,
US Office of Tech Services PB Rept 53045
J) Anon, “Allied and Enemy Explosives”,
Aberdeen Proving Ground, Maryland (1946)
K) G.C. Tibbitts et al, “Japanese Powder and
Explosives”, US Ordinance TechIntelligence Rept
6 (1946), US Office of Technical Services PB
Rept 50394
Powders”, Kogyökayakukiyókaishi 20, 275–79
(1958) (US Dept of the Army Translation I-1955,
ID No 2203235 (October 1962), Report No 1
“Measurement of Combustion Temperature of
Delay Powders”; Translation I-1957, ID No
2203237 (Oct 1962), Rept No 3 “On Mechanism
of Combustion of Delay Powders”; Translation
I-1958, ID No 2203238 (Oct 1962), Rept No 6,
“Theory of Burning Rate of Delay Powders”)
M) Anon, “Manufacturing and Construction
(Vehicles, Explosives, Arms and Ammunition)”,
US OrdNtechIntelligence Rept OTHA 5801,
Chapter VI, Section 64, pp 64-1 ff (March 1960)
US OrdNtechIntelligence Agency, Ordn
Liason Group, Durham, North Carolina (1960)
O) M.H.B. Smith & J.E. Smith, “Small Arms
of the World”, Stackpole, Harrisburg, Pennsyl-
vania (1960), pp 498–518
Development Inst, Japan Defense Agency (Sept
Intelligence Rept 2220044563 (1963)]
Q) Heizo Nambo, “Who Invented Explosives”,
Kogyökayakukiyókaishi 28, 322–29 & 403–23
(1967). Engl translation obtainable for $15.00,
surface mail from Japan Publishers Trading Co.
Ltd, PO Box 5030, Tokyo International, Tokyo, Japan
R) K. Sassa & I. Ito, “Studies of Detonation
Pressure by Aquarium Technique”, Kogyökayakukiyókaishi (abbrd as KKK) 32, No 6,
Nov-Dec 1971. Engl translation by Mrs Geti
in 1972 at the Bureau of Mines; available from
US Dept of Commerce, NatlTechInfoService,
Springfield, Va 22151 (Compositions of Japanese
explos Akatsuki, Enoki Dynamite No 2 and
Geogel are given)
S) Picatinny Arsenal Technical Reports (PATR)
and other Reports:
  a) A.B. Schilling, “Grenade, Serrated, Cast
  Iron (With PD Fuze for Japan Type 89
  Grenade Discharger)”, PATR 149 (1943)
b) Ibid., “Grenade, Serrated, Cast Iron (Hand Use or for Japan Type 10 Grenade Discharger)”, PATR 150 (1943)
c) Ibid., “Unfired 20-mm HE Round of Japanese Ammunition”, PATR 1236 (1943)
d) R.M. Dennis, “Drift Signal”, PATR 1203 (1943)
e) W.R. Tomlinson, Jr., “Compilation of Data on the Composition of Foreign Explosives”, PATR 1311 (1943)
f) A.B. Schilling, “Grenade, Hand, Stick Type”, PA Fragmentation Rept 193 (1944)
g) Ibid., “Land Mine and Type 93 Detonator”, PATR 1383 (1944)
h) L.H. Eriksen, “Compilation of Data on the Composition of Foreign Boosters and High Explosives”, PATR 1446 (1944)
i) R.M. Dennis, “Unfired 25-mm HE Shell (Maroon Shell)”, PATR 1350 (1944)
j) Ibid., “Unfired 25-mm HE Shell (Orange Shell, Green Band)”, PATR 1351 (1944)
k) A.B. Schilling, “Unfired AP Rifle Grenade”, PATR 1461 (1945)
l) W.G. Eppler, “Fragmentation of 81-mm HE Mortar Shell, Equipped with PD Fuze”, PA Fragmentation Rept 214 (1945)
m) A.B. Schilling, “Unfired, Serrated Grenade for Hand Use or for Type 10 Grenade Discharger”, PATR 1491 (1945)

Japanese Explosives and Related Items Described in Journals & Patents Listed in Chemical Abstracts After WWII

Following abbreviations are used for the Japanese Journals and Japanese Patents listed in CA:

Japan Kokka — Japanese National Diet
KKK — KōgōKayakuKyōkaishi (Journal of the Industrial Explosives Society of Japan)
KKZ — KōgōKayakuZasshi (Journal of the Chemical Society of Japan. Industrial Chemistry Section)
KKwZ — KōgōKwagakuZasshi (Journal of the Society of Chemical Industry, Japan, abbrd in CA as JourSocChemInd, Japan)
KōjinKwZ — KōjinKwagakuZasshi [Engineers Chemical Journal, Japan, abbrd in CA as Eng.ChemJour(Japan)]
TKShH — TokyoKōgōShikenshoHōkoku (Reports of the Government Chemical Industrial Research Institute, Tokyo)

Following are the titles of Japanese Patents and Papers, arranged in chronological order. A brief description of each subject is given in CA:


3) Y. Suzuki, “Explosive Tube from Liquid Carbon Dioxide”, JP’s 3, 4 & 5 (1950); CA 46, 8376 (1952)

4) Z. Akashi, “Gasless Fuse”, JP’s 3043 & 3045 (1951); CA 47 (1953)


8) T. Hikita, "Progress in the Theory of Detonation", KKK 13, 3-8 & 77-85 (1952) and CA 49, 2736 (1955)
9) T. Kikara & T. Hikita, "Molecular Theory of Detonation", KKK 13, 106-13 (1952) and CA 49, 5841 (1955)
10) T. Okawa, "Studies of the Muzzle Flashes of Explosives by Means of High-Speed Cinematographic Camera", KKK 13, 247-54 (1952) & CA 49, 5841 (1955)
41) Sh. Okubo, K. Shindo & S. Oinuma, "Copper Azide Detonators. I. Preparation of Copper Azide and Its Impact Sensitivity Test", TKShH 52, 311-14 (1957) & CA 52, 8559 (1958)


58) T. Murata & T. Fukuda, “NAP Propellant”, KKK 63, 1900–01 (1960) & CA 57, 1140 (1962) [A new type of solid propelent prop by treating NH₄ClO₄ + polyester resin composite propelant with NG, MeNO₂ or O(CH₂CH₂ONO₂)₂]

59) K. Akimoto et al, “Experimental Studies of Composite Propellants”, KKK 63, 1902–06 (1960) & CA 57, 1140 (1962) [Composite propelent NH₄ClO₄ + epoxy polysulfide resin was found to be superior to propelant NH₄ClO₄ + polyester resin in both its phys & burning props]


70) M. Inagaki, “Rocket Propellants”, JP
4445 (1962) & CA 60, 5269 (1964)
74) K. Tsujihashi, "Coal Mine Explosives", JP 15798 (1962) & CA 60, 5270 (1964)
78) K. Yamazaki et al, "Polyethylene-Ammonium Perchlorate-Base Propellants Having High Specific Impulse and Slow Burning Rate", Tokyo Daigaku Koku Kenkyusho Shuho (Bulletin of the Aeronautical Research Institute, Univ of Tokyo) 3 (6), 381–89 (1963) & CA 60, 359 (1964)
80) T. Watanabe & M. Inagaki, "Rocket Propellants", JP 9594 (1963) & CA 60, 7864 (1964)
82) N. Tomenishi & K. Oishi, "Dynamite Explosives", JP 16148 (1963) & CA 60, 7865 (1964)
84) K. Ito & Y. Inumaru, "Waterproof Commercial Explosives", JP 20394 (1963) & CA 60, 10472 (1964)
87) K. Suzuki, "Properties of Polyether-Epoxy Type Propellants". 5th InternSymp on Space Technology & Science, Tokyo, Japan (2-7 Sept 1963), 61–64. IAA Accession No A65-1429, 3pp (1964) (Engl) & CA 60, 9735 (1965)
87a) M. Uetake & H. Sato, "Ignition of Propellants by Electrostatic Discharge", KKK 24 (2), 67–72 (1963) & CA 60, 11835 (1964)
88) J. Fukuyama & T. Hagi, "Salt Pair Explosive for Coal Mining", KKK 24 (3), 140–47 (1963) & CA 63, 6777 (1965) (Two types NG contg exps with salt pairs NH4Cl/KNO3 and NH4Cl/NaNO3 were investigated. Both types offered greater safety than the usual mining exps. The salt pair NH4Cl/NaNO3 made the expl hygroscopic and gave poorer transmission than NH4Cl/KNO3)

Note: Same paper is listed in CA 60, 11836 (1964) under the title "Ion-Exchange Type Coal Mine Explosives"

90) K. Yamamoto, "Primary Explosives. II. Thermal Decomposition of Diamonitrophenol and Other Primary Explosives", KKK 25 (1), 10–19 (1964) & CA 60, 15673 (1964)
91) E. Yoshikawa, "AN-FO Explosives in Europe", KKK 25 (1), 33–38 (1964) & CA 60, 15674 (1964) (A review)
92) Y. Wakazono & Ch. Sato, "Granular Explosives. I. AN-FO Blasting Agents", KKK 25 (2), 63–69 (1964) & CA 61, 9397 (1964)
93) T. Yoshida et al, "AN-FO Explosives. I", KKK 25 (2), 76–84 (1964) & CA 61, 11838 (1964)
94) Sh. Okubo & Y. Mizushima, "Tests of AN-FO Blasting AGents", KKK 25 (2), 85–90 (1964) & CA 61, 11838 (1964)
96) Ibi, IV. "Initiation of Lead Thiocyanate-Potassium Chlorate", KKK 25 (3), 134–44 (1964) & CA 61, 11839 (1964)
128) H. Kimura & H. Tannada, “Slurry Explo-
161) K. Kishi & T. Yamazaki, "Mixed Solid
188) M. Takeda, "Match Stiker Composition Containing Urea Resin as Adhesive", JapanKokai 23911 (1973) & CA 79, 89 (1973)
195) A. Asano, "Smoke-Controlling Oily Agent


JATO. Abbreviation for Jet Assisted Take Off which refers to the use of an auxiliary rocket motor for added thrust in the take-off of an aircraft. The usual JATO units are solid propellants (Ref 2):

<table>
<thead>
<tr>
<th>Description</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rocket description</td>
<td>12 AS-1000 D-1</td>
</tr>
<tr>
<td>Charge type</td>
<td>End-burning</td>
</tr>
<tr>
<td>Purpose</td>
<td>Assisted take-off of aircraft</td>
</tr>
<tr>
<td>Thrust, lb</td>
<td>1000 (rated)</td>
</tr>
<tr>
<td>Burning time, sec</td>
<td>12 (rated)</td>
</tr>
<tr>
<td>Impulse, lb·sec</td>
<td>16,000 (actual)</td>
</tr>
<tr>
<td>Propellant</td>
<td>Asphalitic composite</td>
</tr>
<tr>
<td>Outside tube diam, in</td>
<td>9.6</td>
</tr>
<tr>
<td>Length, in</td>
<td>36</td>
</tr>
<tr>
<td>Gross weight, lb</td>
<td>205</td>
</tr>
<tr>
<td>Propellant weight, lb</td>
<td>90</td>
</tr>
</tbody>
</table>

However JATO units based on liq propellants have also been used for example the "Tonka" fuels used with white fuming nitric acid (Ref 1). Tonka fuels are mixtures of aniline, monomethylaniline, dimethylaniline, naphtha, triethylamine & iso-hexylamine


JB-2 Bomb. The American copy of the Ger "buzz bomb". The chge consisted of a perforated cylinder, 8.5 inches OD, 2.5 inches ID, 36.5 inches long, and weighed 120 lbs. It developed at 70°F a thrust of 1100 lbs for a period of 1.85 secs. Four of these chges, each encased in a suitable motor, were attached to a launching sled, which ran on a heavy non-portable rail system. By an ingenious development of the US Air Forces, the bomb, sled & rocket motors were converted to what amounted to a free flight rocket. This light, highly portable launching system had a great advantage over the heavy & rigid German launchers

The JB-2 Bomb did not come into combat use, although a requirement for 500 launching units a month existed at the end of WWII. The construction of a production plant with a capacity of 600000 lbs of propht a month (sufficient for 1200 launchings) was nearing completion. Interest in the JB-2 Bomb remained sufficiently high so that the pilot plant was converted to a small production plant for peacetime operation


JB Powders. Smokeless prophts, patented originally in 1888 by Johnson & Borland, and manufd for many years by the EC Powder Co Ltd, Greenhite, England (See EC Powders in Vol 5 of Encycl, p E6-R). The military propht contd NC 50, KNO₃ 40 & roasted starch (partly burned) or lampblack 10%. A sporting propht contd NC 50, KNO₃ 22, Ba(NO₃)₂ 25 & roasted starch or lampblack 3%. These powders are no longer manufactured

Ref: Daniel (1902), pp 386–87
**JCP Powder.** One of the varieties of plastos—menites. It is a sporting smokeless propellant containing NC, DNT & Ba nitrate
*Ref:* Daniel (1902), pp 387 & 634

**Jelly Bag Method of Mixing.** See under Drums, Rotary (Rotating) for Blending (Mixing) of Explosives and Their Components in Vol 5 of Encycl, p D1556-R

**Jelly, Mineral.** Same as vaseline

**Jensen’s Test (For Stability of Explosives).** Test tubes, each containing 0.1g of a NC expl, are suspended in a bath of mineral jelly preheated to 100°. A strip of KI-starch paper is suspended above the sample. Heating of the bath is continued, by raising the temp 5° per min, until the point is reached at which the test paper begins to color. This point is taken as an indication of the stability of the NC propiant. Jensen’s Test is a combination of the deflagration test of Sy and the Abel Heat Test
*Ref:* Reilly (1938), pp 80 & 83

**Jetcord.** An Explosives Technology (Fairfield, Calif) development which is special explos chge similar to the military shaped, or “hollow chge”. It consists of a metal-sheathed, exp Highly filled, linear-shaped chge, having an approx chevron-shaped cross section. Jetcord is available in a size range (1000, 2000 & 4000 grains/ft expl chge) that permits the cutting of a few mils of Al up to 3 inches of steel. Initiating the expl device raised a problem. The use of conventional, highly sensitive electric blasting caps would not be feasible in the vicinity of high-powered elec machinery. An alternative was to use a very insensitive device known as Exploding Bridgewire (EBW) detonators developed during WWII (See in Vol 4 of Encycl, p D807-L and in Vol 6, p E353-R). EBW detonators differ from conventional types in that they require currents of thousands of amps delivered in a few microseconds before they will detonate. Under such elec impulses, the detonator bridgewire literally explodes, and the resulting high temp shock adequately initiates Jetcord

*Typical Jetcord comes in a variety of sizes for special applications. The powerful linear-shaped chge permits demolition contractors to cut supporting steel structures with a few ounces of expl rather than depend on the blast effect of large quantities of expls. Jetcord has been used successfully to demolish buildings & bridges
*Ref:* Industrial Research Magazine (April 1974), pp 67–70

**Jet Formation in Shaped Charges.** See Detonation; BMT (Birkhoff-MacDougal-Pugh-Taylor) Theory of Jet Formation in Shaped Charges in Vol 4 of Encycl, p D226-R; and Detonation, Munroe-Neumann Effect (or Shaped-Charge Effect) and Lined-Cavity Effect in on pp D442-Rff

**JET FUELS.** *Introduction.* These are fuels for air-flow jet propulsion systems (See Jet Propulsion in this Vol). Fuels for rockets are described, in part, under Hypergolic Propellants in this Vol, pp H254ff. Air-flow jet propulsion systems are divided generally into two categories: those needing an external source of ignition and those that are self-igniting or hypergolic

Recent literature on jet fuels is very extensive, eg between 1957 & 1971 there are some seven pages of Refs to jet fuels in CA indexes and these probably do not include all contract research reports and certainly not classified literature.

What follows is a summary of the props & requirements of the most common jet-aircraft fuels (JP series) and a list of recent review articles on these and other jet fuels of special interest

**Jet Fuels (Grades JP-3, JP-4, JP-5 & JP-6).** Specifications for early aircraft jet fuels were based primarily on manuf considerations, since it was believed that aircraft could burn almost anything of the nature of kerosene fuels (Ref 1). Later improvements in aircraft, particularly in high-speed jets, made it necessary to pay more attention to fuel characteristics and less attm to ease of manuf. The most important of these fuel characteristics is fuel stability at high temps. Other problems associated with jet aviation fuels, both for military & civilian use, are minor compared to stability at high temps. Such problems include availability, handling & physical property specifications
The need for thermal stability of jet fuels is based on the following facts. In supersonic flight, the aircraft is subjected to high temperatures. This heat must be removed. Most convenient solution, in many types of jet aircraft, would be to use the fuel as a heat sink for cooling vital aircraft components, such as engine oil.

Theoretically, use of fuel as a coolant is convenient. But, it is well known that heating oils for even short periods of time accelerates gum and sediment formation. (In fact, this is the basis for many stability prediction tests in the petroleum industry.) It was not unexpected then, that when fuels are heated in aircraft fuel systems, temperatures would be high enough to cause some fuel degradation.

Mechanism studies of this disintegration have been resolved into three thoughts:

a) fuel is cracked
b) fuel is oxidized
c) fuel is polymerized

Most workers in this field believe polymerization to be the cause of sediment which plugs filters and deposits lacquers on such vital parts as heat exchanger surfaces.

Although petroleum refiners are not unfamiliar with questions of thermal stability of petroleum products, jet fuel stability requirements (stable in the range 400–500°F) presented a new set of problems. One of the first things to be done was to define limits of acceptable stability. Such limits naturally would depend upon individual engine design and the environment to which the fuel is exposed. Fuels meeting one set of conditions could conceivably fail to meet another set. The solution: devise some sort of laboratory test that would correlate with actual engine performance.

Pioneering efforts in this were carried out by Pratt & Whitney and Du Pont. Working with a scaled down fuel system, they were able to obtain some degree of correlation with engine performance. But, around the same time, others interested in this problem—refiners, engine manufacturers, and air frame designers—began to build test rigs. Soon there were as many devices as there were groups involved in the thermal stability problem. Consequently, a meeting was arranged to suggest a suitable piece of equipment and set up a program for evaluation of jet fuels under defined conditions. The apparatus selected was called the Erdco Jet Fuel Coker. Data correlation was undertaken by the Coordinating Research Council, an organization sustained jointly by the Society of Automotive Engineers, and the American Petroleum Institute.

With this Erdco coking unit, the test fuel under consideration is pumped thru an annular heat exchanger. Hot fuel then passes thru a heated sintered steel filter. Fuel, flow rate, fuel temperature, and filter temperature can be varied over a wide range to control test severity. Insoluble sediment which is formed during the heating operation deposits on the filter. The time required to achieve a given pressure drop across the filter is a measure of fuel stability. Visual observations of the heated tubes are made to estimate heat exchanger fouling tendencies.

More specifically, typical operating conditions are: a flow rate of 4 pounds per hour, a fuel temperature from the heat exchanger of 400°F, a filter temperature of 500°F, and a fuel pressure of 150 pounds per square inch gauge. Fuel stability is then determined by operating until a pressure drop of 25 inches of mercury is obtained across the filter or until 300 minutes have elapsed. No heater tube appearance requirements have been designated.

Some characteristics and requirements of the JP Series of fuels is given below (Ref 1) (more detailed characteristics are presented later):

**JP-3 (Used by Military)**. A mixture of about 70% gasoline and 30% light distillate. Specifications limiting boiling range are a 5 to 7 pound Reid vapor pressure and a maximum 90% evaporated point of 470°F.

**JP-4 (Used by Military and Commercial)**. A mixture of about 65% gasoline and 35% light distillate. Reid vapor pressure specification is 2 to 3 pounds. Product was developed as a replacement for JP-3 because of excessive fuel losses encountered at high altitudes with JP-3.

**JP-4 Referee (Used by Military)**. A special grade of JP-4 having rigidly specified characteristics. Fuel is designed to be typical of lower quality JP-4 fuel which would be available in wartime. Used in developing and testing of military jet engines.

**JP-5 (Military)**. Essentially a specially fraction-
Kerosine (Used by Commercial) has essentially the same requirements as burning kerosine with the exception of freezing point. Commercial airlines usually require freezing point of \(-40^\circ\text{F}\) maximum because of the low temperatures encountered at altitude. Many, but not all, burning kerosines will meet this requirement.

Attempts have been made to increase the stability of the above fuels by:

a) use of processing or treating techniques

b) incorporation of additives

c) segregation of stable stocks

Use of conventional treating processes was the logical starting point for improving thermal stability since refineries already have these facilities. One of the first treating processes considered was sulfur dioxide treatment, although current capacity would possibly be insufficient in an all-out war. In tests carried out at one petroleum laboratory, a JP-5 fuel stability was increased twofold: Untreated fuel had a stability rating of 57 minutes. After sulfur dioxide extraction, the time was increased to 119 minutes. However, for all intents and purposes, this is not considered to offer any benefits since the test defines acceptable stability as 300 minutes.

Another refinery treating procedure is acid treatment. This method is somewhat successful. In further test on a JP-5 fuel at the above mentioned laboratory, the improvement factor was 2.5 or 157 minutes. Actually, effectiveness of treating depends upon the crude oil supply and its appearance after original refining, that is, first separation from the crude. What would be described as mild treatment is good for one refiner, but another, because of crude supply, would need much higher treats, say to the order of 25 to 50 pounds per 42-gallon barrel, before a stable product is obtained. Acid treating is not favorable from the processing side, but it is possible to produce high quality fuels with the process.

A better approach to achieving thermal stability is thru hydrogenation. Hydrogen is now readily available in most refineries because of catalytic reforming developments. Hydrogenation produces fuels of excellent thermal stability. Again citing the laboratory example above, the same JP-5 fuel hydrotreated shows improved stability to 300 plus minutes.

Use of hydrogen is effective in removing trace constituents, often times the bad actors, such as nitrogen and sulfur. Also, hydrogen saturates previously unsaturated olefins or aromatics which may be present. The amount of hydrogenation depends again, upon the source of crude oil. Good crudes may only need mild hydrogenation, other cases, heavy.

Another possible solution to the problem of high temperature stability is the use of additives. Not exactly a stranger to petroleum people (as evidenced by use in gasoline and lubricants) they generally fall into two classes: metallic and non-metallic. The former, for the most part are metal salts of sulfonates or naphthenates, whereas the latter are either amines or amine derivatives (later other organics may prove more effective).

Use of additives in jet fuels, however, must of necessity be approached with caution. As surface active materials, many have a variety of uses and properties. Hence, they must not introduce new problems such as foaming at high altitudes, emulsification, or interference with low temperature flow. These could easily by severe limitations, but additives are under serious consideration throughout the industry.

Additives can be most effective when contaminants causing instability have been minimized by refinery processes such as hydrogenation. In general, however, the prime purpose of an additive is either to dissolve or prevent formation of sludge or else to disperse sediment in fine particles which enables passing thru filters and other vital engine parts. The problem is of prime concern in the case of metallicity type additives which can build up ash deposits, which can seriously affect jet engine operation. An appreciable slug of ash can do extensive damage if suddenly tossed thru a jet engine.

Studies have been made thruout the refining industry in an effort to utilize selected stocks for the production of jet fuels. Basically, this would amount to determining the stability of many stocks, for example, straight run gasolines, distillates, kerosines, alkylate bottoms, and whatever else is available from refinery streams. Those with best heat stability, by laboratory test, could then be blended into jet fuels meeting required
specifications

To meet military supply and demand it often becomes necessary to maintain large supplies of jet fuels at various locations throughout the world. This fuel must be stable so it can be available for immediate use. While fuel can be kept relatively new in some areas by means of stock rotation, this may not be possible in others. This brings up the problem of stability in standing storage.

Jet fuels can be made from a very broad hydrocarbon range, in fact, anywhere from 150°F to 600°F depending upon specifications. Fuels consist of certain quantities of straight run, catalytically cracked, and thermal cracked material. Stability-wise, straight run is most stable; thermal cracked least stable. Hence, most blends end up intermediate.

Standing storage stability can usually be improved by the same general procedures being explored for thermal stability, that is, acid treatment and/or hydrogenation, where improvement is gained by removal of nitrogen and sulfur compounds, olefins and diolefins. Hence, the problem today is actually a corollary of thermal stability.

Another problem encountered with jet fuels is due to contaminants and the consequent plugging of filter systems in the jet aircraft. Handling and distribution systems for jet fuels are usually underground. Fuel is pumped out of the tank by water displacement. Understandably, when fuel is moved, there is some agitation. If an additive is present, emulsification is possible. Also, there may be many contaminants such as rust, dirt, and water. The rate of settling, naturally, is a function of viscosity and density. This is important since jet engines cannot tolerate too much foreign matter.

For the most part, filtration systems can remove the contaminants. However, aside from foreign matter there is the case of insolubles formed from the fuel in the nature of gums. In this respect filterability behavior is apparently related to the character of the deposits formed. The more crystalline the deposit the less tendency to plug filters.

Zabetakis et al (Ref 6) reported that mixes of JP-4 & UDMH (unsym-Dimethyl Hydrazine, See in this Vol, pp H203-R to H204-L) are insensitive to shock.

Recent Review Articles on Jet Fuels.

Hibbard & Olson (Ref 10) discuss turbojet & ramjet fuels and their relation to fuel systems. Carney (Ref 11) reviews the props of “slush” hydrogen and points out its desirability as a space vehicle propellant fuel because of weight savings on long-term storage. “Slush” hydrogen is a fluid mixt contg solid H2 (by wt) 50% in liq H2.

F-O mixts (Flox) with light hydrocarbon (HC) fuels as rocket propellants for upper-stage applications show the unique ability to provide high performance, space storability, hypergolicity, current availability, high d, and capability for both transpiration & regenerative cooling. Methane, (CH4), gives the highest theoretical specific impulse of an HC fuel with Flox, but others show advantages in handling, bulk d, and regenerative & cooling capability. Blending of HC fuels may further improve space storability, fuel d, and fuel cooling ability. Methane is the best HC fuel with Flox in a transpiration-cooled engine or in a regeneratively cooled engine operating at supercritical pressure in the heat exchanger. 1-Butene and a eutectic blend of pentane & iso-pentane are the most promising fuels for regenerative cooled thrust chambers at subcritical pressures. An F/O ratio of ca 82/18 gives a sharp max specific impulse with methane as the fuel. Such high concns give hypergolicity (ignition on contact with fuel) almost like F alone. Data are given on ignition delay times using Flox with several light HC fuels and with H. All give satisfactory short ignition times for ambient sea-level start and during cold-altitude starts (Ref 12).

Lo (Ref 14) predicts that “triergol” or “tribril” propellants will provide max thrust in future rockets & space vehicles. These systems contain a metal (or metal oxide) in addition to H2 as fuel. Examples are: O3/Be/H2, O2/Be/H2 & F2/Li/H2. They give a much greater specific impulse (See Isp in this Vol, p 163-R) than the corresponding “diergol” O3/H2, O2/H2 & F2/H2 by increasing the reaction temp. In practice, 3 tanks are required for the “triergol” system, but one of them can also serve as the combustion chamber.

Zelov (Ref 15) reviews the development of jet fuel technology with emphasis on production methods, composition, energy content, combustion
## Table 1
### Chemical and Physical Requirements and Test Methods

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Distillation:</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Initial boiling point</td>
<td>(1)</td>
<td>(1)</td>
<td>(1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fuel evaporated, 10 percent min. at</td>
<td>(2)</td>
<td>(2)</td>
<td>(2)</td>
<td>400°F F. (204.4°C)</td>
<td>D86</td>
</tr>
<tr>
<td>Fuel evaporated, 20 percent min. at</td>
<td>240°F (116.8°C)</td>
<td>290°F (143°C)</td>
<td>290°F (143°C)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fuel evaporated, 50 percent min. at</td>
<td>250°F (176.7°C)</td>
<td>370°F (187.8°C)</td>
<td>370°F (187.8°C)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fuel evaporated, 90 percent min. at</td>
<td>470°F (243.3°C)</td>
<td>470°F (243.3°C)</td>
<td>470°F (243.3°C)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>End point, max.</td>
<td>(3)</td>
<td>(3)</td>
<td>(3)</td>
<td>550°F (287.8°C)</td>
<td>1001</td>
</tr>
<tr>
<td>Percent evaporated, at 400°F F. (204.4°C)</td>
<td>(4)</td>
<td>(4)</td>
<td>(4)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Residue, vol. percent max.</td>
<td>1%</td>
<td>1%</td>
<td>1%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Distillation loss, vol. percent max.</td>
<td>1%</td>
<td>1%</td>
<td>1%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gravity *API—max. (specific gravity, min.)</td>
<td>0.005</td>
<td>0.005</td>
<td>0.005</td>
<td>0.005 (0.760)</td>
<td>401</td>
</tr>
<tr>
<td>Gravity *API—min. (specific gravity, min.)</td>
<td>0.65</td>
<td>0.65</td>
<td>0.65</td>
<td>0.65 (0.730)</td>
<td>401</td>
</tr>
<tr>
<td>Existent gum, mg/100 ml, max.</td>
<td>7</td>
<td>7</td>
<td>7</td>
<td>7</td>
<td>3302</td>
</tr>
<tr>
<td>Potential gum, 16 hr. aging, mg/100 ml, max.</td>
<td>14</td>
<td>14</td>
<td>14</td>
<td>14</td>
<td>3354</td>
</tr>
<tr>
<td>Sulfur, total, percent wt. max.</td>
<td>0.4</td>
<td>0.4</td>
<td>0.4</td>
<td>0.4</td>
<td>3201</td>
</tr>
<tr>
<td>Mercaptan-sulfur, percent wt. max</td>
<td>0.003</td>
<td>0.003</td>
<td>0.003</td>
<td>0.003 (0.760)</td>
<td>5204</td>
</tr>
<tr>
<td>Reid vapor pressure, 100°F F. psi, min. (gpm cm²/min.)</td>
<td>5.0</td>
<td>5.0</td>
<td>5.0</td>
<td>5.0 (351.6)</td>
<td>1201</td>
</tr>
<tr>
<td>Reid vapor pressure, 100°F F. psi, max. (gpm cm²/min.)</td>
<td>7.0 (462.2)</td>
<td>3.0 (210.9)</td>
<td>3.0 (210.9)</td>
<td>3.0 (210.9)</td>
<td>1201</td>
</tr>
<tr>
<td>Freezing point, °F, max.</td>
<td>-75° (4°C)</td>
<td>-76° (4°C)</td>
<td>-55° (4°C)</td>
<td>4°F</td>
<td>1411</td>
</tr>
<tr>
<td>Thermal value (see 3.2.2) Heat of combustion: (lower or net) BTU/lb min.</td>
<td>18,400</td>
<td>18,400</td>
<td>18,300</td>
<td>18,300</td>
<td>2502</td>
</tr>
<tr>
<td>Viscosity, centistokes at -30°F F. (34.4°C), max.</td>
<td>5.250</td>
<td>5.250</td>
<td>4.500</td>
<td>4.500</td>
<td>3601 and 401</td>
</tr>
<tr>
<td>Aromatics, vol. percent max.</td>
<td>25.0</td>
<td>25.0</td>
<td>25.0</td>
<td>25.0</td>
<td>3703</td>
</tr>
<tr>
<td>Olefin, vol. percent max.</td>
<td>6.0</td>
<td>5.0</td>
<td>5.0</td>
<td>5.0</td>
<td>3703</td>
</tr>
<tr>
<td>Smoke point, mm, min.</td>
<td>(5)</td>
<td>(5)</td>
<td>(5)</td>
<td>(5)</td>
<td>2107</td>
</tr>
<tr>
<td>Explosiveness, percent max.</td>
<td>10.0</td>
<td>10.0</td>
<td>10.0</td>
<td>10.0</td>
<td>1151</td>
</tr>
<tr>
<td>Smoke volatility index, min. (see 3.2.3)</td>
<td>52.0</td>
<td>52.0</td>
<td>52.0</td>
<td>52.0</td>
<td>12322</td>
</tr>
<tr>
<td>Copper strip corrosion, ASTM classification, max.</td>
<td>No. 1</td>
<td>No. 1</td>
<td>No. 1</td>
<td>No. 1</td>
<td>D1319</td>
</tr>
<tr>
<td>Water reaction, vol. change, ml, max. (see 3.2.1)</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>4251</td>
</tr>
<tr>
<td>Flash point, min.</td>
<td>(6)</td>
<td>(6)</td>
<td>(6)</td>
<td>140°F F. (60.0°C)</td>
<td>1102</td>
</tr>
<tr>
<td>Thermal stability</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>D93</td>
</tr>
<tr>
<td>Change in pressure drop in 5 hr, in. Hg</td>
<td>(7)</td>
<td>(7)</td>
<td>(7)</td>
<td>3464</td>
<td>3464</td>
</tr>
<tr>
<td>Preheater deposit</td>
<td>(8)</td>
<td>(8)</td>
<td>(8)</td>
<td>3464</td>
<td>3464</td>
</tr>
</tbody>
</table>

1 To be reported — not limited
2 Use steam jet method of D381 after oxidation
3 The mercaptan–sulfur determination may be waived at the option of the Inspector, if the fuel is considered “Doctor sweet” when tested in accordance with Method 5203 of Federal Test Method Standard No. 791
4 May be reported as Bromine No when specified by the armed service for whom the material is approved
5 To be performed in accordance with paragraph titled “Tests at 212°F for Aircraft Engine Fuels” of ASTM D130-56. Earlier ASTM methods are not applicable since they require a dry sample
6 See 4.5.1 for exception to Method 3251
7 See 4.5.2 for test conditions under Method 3464
characteristics, scale formation & corrosion tendencies, volatility, thermal & oxidation stability, low temp props, and expln & fire hazards

Giacco (Ref 16) reviews solid propulents for use in rockets

Koecker (Ref 17) reviews specification requirements of fuels for industrial gas turbines, conventional jet turbines (kerosene & gas oils), supersonic jet engines (JP-5 & JP-6), and future hypersonic jets (endothemic fuels)

Back (Ref 18) discusses propulnt toxicology, including toxicological effects on blood pressure, heart rate, and effects on autonomic & central nervous systems caused by N₂H₄ & derivs, penta- & decaboranes, F compds and Be compds

The handling of hazardous fuels & oxidizers is reviewed by Cloyd & Murphy (Ref 13). Data are presented on some highly reactive materials that have been studied in the search for fuels & oxidizers for space work, including liq H, pentaborane, F, ClF₃, O₃, N₂O₄ & N₂H₄ and its derivs. Both the hazards that result in these materials and the procedures by which they were handled & stored safely are described. Refs are given to work done by NASA & other investigators

Jet Fuel Patents

The patent literature on jet fuels is very extensive. Some examples of patents granted on jet fuels are as follows: Fox & Britton (Ref 2) claim the incorporation of viscosity-index improvers in jet fuels to improve engine start-up & combustion efficiency over a wide range of temps. Materials claimed to be effective are ethyl glycol, Acryloid & Sanotex

Maisner (Ref 3) claims that incorporating nitroaraffin gel improves jet fuels

Burhardt (Ref 4) proposes propulents, for reaction engines, with compns which contain finely divided metals such as Zn, Cd, Mn, Fe, Pb & W

Thomas (Ref 5) claims systems useful for propulsion of rocket shells, JATO & airborne vehicles, consisting of a mix of finely divided MnO₂ with fuels which are nitro compds, such as NGu, Amm Picrate, etc. The addn of Al can modify burning characteristics. Binders are chlorinated polyphenyls or urea resins. The ingredients are milled on rolls just hot enough to soften the resin and then the mixt is pelletted

Beatty & Gluckstein (Ref 8) propose a mixt of Me₃Al 5 – 40 & Et₃Al 95 – 60% as a jet fuel igniter in jet aircraft engines

Becker (Ref 9) claims that wetting the combustor surface of a jet engine with di-methyl silicone or Si(OEt)₄ prevents decompn of Be₂O₃ on the engine walls, from Be contg fuels, such as a 4/1 mixt of trimethoxyboroxime-acetone

Because of the importance of these fuels, the Military Specifications for Jet Fuels, JP-3, JP-4 & JP-5 (Ref 4) are listed:

1. SCOPE

1.1 Scope. This specification covers fuel for aircraft turbine engines, ramjet engines, and rocket engines

1.2 Classification. Jet fuels shall be of the following grades, as specified:

<table>
<thead>
<tr>
<th>Grade</th>
<th>NATO symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>JP-3</td>
<td>None</td>
<td>High vapor pressure type</td>
</tr>
<tr>
<td>JP-4</td>
<td>F-40</td>
<td>Low vapor pressure type (NATO description: wide-cut, gasoline type)</td>
</tr>
<tr>
<td>JP-5</td>
<td>F-42</td>
<td>High flash point, kerosene type</td>
</tr>
</tbody>
</table>

3. REQUIREMENTS

3.1 Materials. The fuel shall consist completely of hydrocarbon compounds, except as otherwise specified herein

3.2 Chemical and physical requirements. The chemical and physical requirements of the fuel shall conform to those listed in Table 1, when tested in accordance with the applicable tests. Requirements contained herein are absolute and are not subject to correction for tolerance of test methods. However, if multiple determinations are made, average results shall be used

3.2.1 Water reaction. The fuel shall separate sharply from the water layer with only a few small bubbles around the periphery of the interface and no shreds of lace & film at the interface. Neither layer shall have changed in volume by more than 1 milliliter

3.2.2 Heating value. The heat of combustion determination may be waived at the option of the Inspector if the aniline-gravity product of the fuel is not less than the numerical values specified in Table 1. The aniline-gravity product
is defined as the product of the gravity of the fuel in degrees API and the aniline point of the fuel in degrees Fahrenheit. If the aniline-gravity product of the fuel is less than the value specified, the fuel shall be accepted or rejected on the basis of the heat of combustion requirement

3.2.3 Smoke volatility index. The smoke volatility index shall be computed from the following equation:

\[ \text{S.V.I.} = \text{S.P.} + [0.42 \times \text{volume percent boiling under } 400^\circ F (204.4^\circ C)] \]

where: S.V.I. = Smoke volatility index
S.P. = Smoke point in millimeters as determined by ASTM Method D1322

3.3 Additives. The additives listed herein may be used singly or in combination in amounts not to exceed those specified.

3.3.1 Antioxidants. The following active inhibitors may be added separately or in combination to the fuel in total concentration not in excess of 8.4 pounds of inhibitor (not including weight of solvent) per 1000 barrels of fuel (9.1gm/100 US gal, 24mg/liter or 109 mg/imp gal) in order to prevent the formation of gum:

a) 2,6-ditertiary butyl-4-methyl phenol
b) N,N'-Disecodary butyl paraphenylenediamine
c) 2,4-dimethyl-6-tertiary-butylphenol
d) 2,6-ditertiary-butylphenol

3.3.2 Metal deactivator. A metal deactivator, N,N'-dialicyclidene-1, 2-propane-diamine, or N,N'-dialicyclidene 1,2-ethylene diamine may be added in an amount not to exceed 2 pounds of active ingredient per 1000 barrels of fuel (2.2gm/100 US gal, 5.8mg/liter or 26mg/imp gal)

3.3.3 Corrosion inhibitor. An approved corrosion inhibitor shall be added. The corrosion inhibitor furnished under this specification shall be product which has been approved under Specification MIL-I-25017. The amount added shall be listed in QPL-25017 (latest revision). The contractor shall maintain and, upon request, shall make available to the Government, evidence that all inhibitor products used are equal in every respect to the product qualified under Specification MIL-I-25017

3.4 Workmanship. The fuel shall be free from undissolved water, sediment, and suspended matter. No substances of known dangerous toxicity under usual conditions of handling and use shall be added.

4. QUALITY ASSURANCE PROVISIONS

4.1 General. All the tests required herein for the testing of fuel are classified as acceptance tests, for which necessary sampling techniques and methods of testing are specified in this section.

4.2 Inspection. Unless otherwise specified by the procuring activity, inspection shall be in accordance with Federal Test Method Standard No 791, Method 9601

4.3 Sampling. Sampling shall be in accordance with ASTM Method D270 titled Sampling of Petroleum and Petroleum Products.

4.3.1 When required, a 10-gallon sample, taken in accordance with ASTM Method D270, shall be forwarded to the laboratory designated by the procuring activity for testing as specified herein.

4.4 Examination of product. Each container of fuel may be examined to determine conformance with this specification.

4.5 Test methods. Tests as specified in 3.2 to determine conformance to chemical and physical requirements shall be conducted in accordance with Federal Test Method Standard No 791 or ASTM standards, using applicable methods as listed in Table I, except for the following.

4.5.1 Water reaction. The water reaction test shall be conducted in accordance with Method 3251 of Federal Test Method Standard No 791, except that the water used for the test shall be a buffer solution consisting of 1.15g K₂HPO₄ (anhydrous, reagent grade) plus 0.46g KH₂PO₄ (anhydrous, reagent grade) per 100ml of distilled water. This solution shall have a pH of 7.

4.5.2 Thermal stability. A manual, semi-automatic or automatic CFR fuel coker shall be used. The equipment and operating procedures shall be in strict accordance with Method 3464 of Federal Test Method Standard No 791. No deviation from this method is allowed.

4.5.2.1 Test conditions. The test conditions, as applicable, shall be in accordance with Table II.
Table II. Thermal Stability Test Conditions

<table>
<thead>
<tr>
<th></th>
<th>Grade JP-4</th>
<th>Grade JP-5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Preheater temperature (° F.)</td>
<td>300</td>
<td>400</td>
</tr>
<tr>
<td>Filter temperature (° F.)</td>
<td>400</td>
<td>500</td>
</tr>
<tr>
<td>Fuel flow (lb/hr)</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>Test time (minutes)</td>
<td>300</td>
<td>300</td>
</tr>
</tbody>
</table>

4.5.2.2 Reported data. The following data shall be reported:

a) Differential pressure in inches of mercury at 300 minutes or time to a differential pressure of 25 inches of mercury, whichever comes first

b) Preheater deposits at the end of the test

4.6 Rejection and retest. Material not conforming to the requirements of this specification shall be rejected. Rejected material shall not be resubmitted without furnishing full particulars concerning previous rejection and measures taken to overcome defects

6. NOTES

6.1 Intended use. The fuel covered by this specification is intended for use in aircraft turbine engines, ramjet engines, and rocket engines

6.3 Precaution for mixing inhibitors. To prevent any possible reaction between the concentrated forms of different corrosion inhibitors, the fuel supplier is cautioned not to commingle corrosion inhibitors prior to their addition to the fuels

written by J. ROTH

Refs: 1) Staff, C&EN 33, 4502 (24 Oct 1955)
2) H.M. Fox & S.C. Britton, USP 2712726
   (1955) & CA 49, 14298 (1956) 3) H. Maisner,
   USP 2712989 (1955) & Ordn 40, 476 (1955)
4) W. Burkhardt, GerP 1021238 (1957) & CA
   54, 10328 (1960) 5) C.A. Thomas, USP
   2857258 (1958) & CA 53, 2580 (1959)
6) M.J. Zabetaakis et al, US BurMines RI 5635
   (1960) & CA 54, 21763 (1960) 7) J. Kofen-
   bach & T. Wasserbach, USP 2938779 (1960)
   & CA 54, 17851 (1960) 8) H.A. Beauty &
   M.E. Glueckstein, USP 2935839 (1960) & CA
   54, 17889 (1960) 9) E. Becker, USP 2986876

Hibbard & W.I. Olson, CombustPropellenti-
Nuovi, AttiConvMilan 1963, 217 (Eng) & CA
61, 15904 (1961) 11) R.R. Carney, Adv-
CryogenEng 9, 529 (1964) & CA 61, 13115
(1964) 12) A.I. Masters, JSpacecraftRockets
3 (6), 905 (1966) & CA 65, 10415 (1966)
13) D.R. Cloyd & W.J. Murphy, NASA Rept
SP-5032 (1965), AEC Access No 1727 &
CA 65, 10415 (1966) 14) R. Lo, ChemEng-
Tech 39 (15), 923 (1967) & CA 67, 92451
(1967) 15) V.N. Zrelov, IzogiNaukiTehnol-
OrgVeshchestv 1966, 5 (Prib 1968) & CA 72,
4854 (1970) 16) M. Giacco, QuadMercurial
7 (1), 63 (1968) (Ital) & CA 71, 83156 (1969)
17) M.H. Koecker, BrennstChem 50 (4), 105

Jet Perforators & Jet Tappers. Two commercial
applications of the Shaped Charge Effect,
marketed by the duPont Co, are described in
the Blasters Handbook (Ref 1). The principle
upon which they are based is as follows:

“...The jet principle is simply the focusing of a
greater than normal amount of the heat and
energy of an explosion against a very small
area. This is accomplished by using a high
density, high strength explosive formed around
a conical liner of metal or glass. When the explo-

Jet Perforators are explosive shaped
cases and explosive charge contained in plastic
or metal bodies of different sizes depending upon
the specific well to be perforated.
When detonated, duPont Jet Perforators emit a high velocity jet of tiny copper fragments which penetrate the oil-well casing and formation leaving a deep, large diameter hole through which the oil can flow into the well.

Perforators for both "carrier" and "through-tubing" type oil well completions are available. In "carrier" type guns ranging from 2 5/8 to 5 inches OD, the Jet Perforators are threaded on a strand of "Primacord" and loaded into the gun, a steel tube which is sealed at both ends. The gun serves to control the standoff distance and prevents the well fluid from interfering with the jet formation. For through-tubing completion a smaller version of the carrier-type perforator can be supplied.

duPont pioneered the development of shaped charges for oil well use and these perforators and auxiliary devices have been used dependably for several years in the oil well industry.
duPont Jet Tappers are designed for tapping open hearth steel furnaces. They consist of a two ounce charge of a very powerful explosive in a plastic case with a copper cone. The charge is contained in a bullet-shaped insulating jacket. When the Jet Tapper is assembled for use, a special high temperature Jet Tapper Blasting Cap is threaded through a hollow fiberboard pole eight feet in length, the cap is inserted in the charge, and the insulating jacket is slipped onto the end of the loading pole.

The explosive used in the Jet Tapper is relatively insensitive to impact, friction, heat, and other causes of accidental detonation as compared to most commercial explosives. If heated to a high enough temperature it will burn without detonating. In one test a case of Jet Tappers was completely burned in a bonfire of kerosene-soaked wood without detonating. In another test, six Jet Tappers were laid side by side between steel plates and a 150 pound weight was dropped nine feet onto the upper plate. The charges were completely crushed but there was no detonation.

The special Jet Tapper Blasting Caps have been designed to withstand prolonged exposure to high temperatures. Experience has shown that they will stay in a hot tap hole for from three to eight minutes before detonating when they are used as part of the Jet Tapper assembly.

The only phase of open hearth furnace tapping, as it is now practiced, that is replaced by the use of the Jet Tapper is the lancing operation or the burning through of the tap hole facing with an oxygen lance. The tap hole must be dug out to the facing as is the present practice. The Jet Tapper is then slid into the tap hole until the nose of the insulating jacket touches the facing, as shown in the diagram. The person placing the Jet Tapper then retires to the firing point and the Tapper is detonated.

Field experience indicates that over-all a 90 to 95 percent average of successful taps is being obtained using duPont Jet Tappers, and that the unsuccessful taps are very easy to lance. The unsuccessful taps can usually be traced to: (1)
a poorly dug out tap hole, (2) steel which has leaked through the facing, or (3) excessively high furnace bottoms

The use of Jet Tappers has proved to have the following advantages over the conventional method of tapping:
(1) No one is at the runner when the metal flow starts
(2) The danger of burns from faulty connections on oxygen lance is eliminated
(3) Heats can be tapped at exactly the desired time, avoiding off-specification steel resulting from loss of volatile additives during delayed or slow taps
(4) Taps start out at full flow rate, thus reducing the over-all tapping time and ladle skull
(5) Tap hole maintenance is reduced and perfect alignment of the tap hole is insured
(6) The need for bars to knock down ridges in front of the tap hole is practically eliminated

Because of these major advantages, it is expected that Jet Tappers will eventually replace the oxygen lance method of tapping entirely. In fact they have already been adopted for more than two-thirds of all the open hearth furnaces in this country.

Cook (Ref 2) states that further commercial use of the Jet Effect is hindered by the high cost of well-made lined-cavity charges

Refs: 1) Blasters Hdb (1958), pp 91-96  2) Cook (1958) p 259

JET PROPULSION
Introduction. "In the absence of all contact with an external solid, a prime mover placed in a fluid of finite or even zero density can propel itself by ejecting a fluid or solid mass toward the rear. By convention we say that this mover is propelled by jet propulsion, although the propulsive thrust really results from the effects of pressure and friction exerted on the wall of the hollow interior of the mover by the solids or fluids moving in the interior toward the exhaust nozzle."

"In all jet propulsion engines, just as for all propulsion of thermal origin, the source of available energy is an exothermic chemical transformation of solids, liquids, or gases, carried on board and called propellants. For brevity, we include all these propellants under the single term fuel."

"For propulsion in the atmosphere, where ambient air is available, this air may be inducted by the jet engine in order to participate essentially in the chemical transformation of the fuel: the engine is then properly called an air flow jet engine in contrast to the rocket, which does not use any air and is the only jet engine which can be used for propulsion in vacuum."

These quotations are from Sec B of a classic book on Jet Propulsion entitled Jet Propulsion Engines (Ref 1). The contents of this excellent treatise are summarized in its preface:

"This volume considers those principles and problems encountered in combining components to form a complete engine. It relies heavily upon the other volumes which deal with basic principles or principles and problems related to components of an engine."

Section A gives a concise history of the development of rockets and air flow jet engines. Section B gives definitions of thrust and various efficiencies and derives relationships for the performance of the different jet propulsion systems. Section C gives the performance analysis of turbojets based on the internal solution of matching the compressor, combustor, turbine, and nozzle. It includes a discussion of off-design performance and describes the problems of control and testing which are unique to a complete unit. Section D treats the turboprop in a somewhat similar manner. It gives the logic for interest in a turboprop and discusses the additional complications. Section E is devoted to the ramjet, its performance, controls, and methods of testing. Section F discusses the wave engines in general, and in particular the pulse jet and the compex. Section G treats the liquid rocket engine, from the consideration of appropriate fuels (both monopropellant and bipropellant) to the designing and testing of the motor. Section H gives a similar treatment for solid propelled rockets, with special stress on the stability and characteristics of burning. The possibility of a variety of hybrid engines, part rocket, part turbine, or more generally part jet and part rotating machinery, is introduced in Sections I and J which treat two such cases—the ram-rocket and the jet rotor.
Each section derives the possible performance and outlines the possible use of these engines. Section K deals with the problems in making a nuclear jet power plant suitable for aircraft. It gives the theory related to the shielding, heat transfer, and the production and control of a small lightweight reactor. The final section does not quite give a peek into the future, but it gives a systematic procedure for exploring the many possibilities of the types of jet engines.

In what follows we will briefly describe the various types of jet propulsion systems and give a list of references to the voluminous literature that has appeared on this subject since the publication of the above book (1959). The reader is also referred to the following Encyclopedia articles: Combustion; Hypergolic Propellants; Ignition; Propellants and Rockets.

In general, in all jet propulsion engines, the driving energy source is the exothermic chemical transformation of solids, liquids or gases (propellants) carried on board the engine.

For propulsion in the atmosphere, where ambient air is available, this air may be induced by the jet engine in order to participate essentially in the chemical transformation of the fuel: the engine is then properly called an air flow jet engine (in Fig 1 the Turbojet, Ramjet & Pulse Jet are examples of air flow jet engines) in contrast to the rocket, which does not use any air and is the only jet engine which can be used for propulsion in vacuum.

A flow of ambient air may also be induced exclusively for augmenting the thrust by increasing the mass flow of the discharged gases: thus we can conceive of an air flow jet engine with two flows, an example of which is a turbojet with a ducted fan.

Whereas the internal flow of a rocket is usually continuous, and even quasi-steady, that of an air flow jet engine can be continuous or discontinuous, i.e. intermittent. The machines which work on the internal flow of a jet engine and have a continuous flow require purely rotating machines, i.e. turbomachines; the engine is called a turboprop.

In the turboprop, the exothermic reaction not only drives the propeller but the exhaust of the reaction products through the nozzle contributes to the total thrust.

If compression and expansion of the internal flow is not produced by rotating or reciprocating machinery but by oscillatory motion of the flow, i.e. wave motion, the jet engine is called a pulse jet.

If, in a continuous flow jet, only the compression resulting from ram effect in the inlet diffuser is utilized, the compression and expansion machinery is eliminated, and the turbojet becomes a ramjet.

Types of Jet Propulsion Engines. The basic jet propulsion engines are illustrated schematically in Fig 1:

![Diagram of Jet Propulsion Engines](image_url)

Fig 1. Types of Jet Propulsion Engines

In these diagrams, D is the intake duct, C is the compressor, B is the burner, T is the turbine and N is the exhaust nozzle. The mode of operation of these engines will be described in the next section.
Modes of Operation of Various Jet Engines.

1) Rockets. As clearly stated a rocket is differentiated from a fluid flow engine in that a rocket generates its thrust entirely from reacting material carried within it, while a fluid flow system takes in material (air) from the outside. The subject of rockets will be described in more detail under Rockets and some material pertaining to rockets has already been described under Hypergolic Propellants and under Ignition. Here we just outline the type of rockets in use or under development:

   1) Solid propellant rocket motor
      a) Constant volume combustion
      b) Constant pressure combustion
      (i) Restricted burning
      (ii) Unrestricted burning
   2) Liquid propellant rocket engine
      a) Gas pressure feed system
      b) Turbopump feed system
   3) Gas or vapor propellant rocket engine
   4) Ionic rockets

Turboprop & Turbojet. The basic principles involved in generating the potential power of a turbojet and a turboprop are identical. They each have a compressor, a burner, and a turbine which runs the compressor. This common part of each produces energy in the form of a hot gas at a high pressure. The only difference in the engines is the manner in which the available energy is converted to useful thrust. In a turbojet the energy is converted to thrust by expanding the gases through a nozzle to form a jet at high velocity. By directing this jet backward the reaction gives a forward component of thrust. In a turboprop, part of the potential energy is removed by additional turbine stages that drive a propeller. Any residual energy is then used to produce a jet stream similar to that of a turbojet, but because of the lower energy level the jet velocity is much lower (See Fig 2 below)

For convenience of thought in comparing the two engines, the part that is common to the turbojet and the turboprop is usually referred to as the hot gas generator, or power generator, and the means for converting the energy to thrust power is called the propulsive system. The hot gas generator is analogous to the steam boiler on a steam engine, in that each produces available energy stored in a gaseous medium at a high temperature and pressure.

![Fig 2. Comparison of Turbojet & Turboprop Engines](image)

Note: The power generator system for the turboprop is essentially the same as in the turbojet.

The advantages of a turboprop are in its greater efficiency, i.e. more of the gas generator energy is converted into useful work than in the turbojet. Its main disadvantages lie in the heavier and more cumbersome machinery that it requires.

Ramjet. We quote from Sect E of Ref 1:

"The ramjet engine, in common with all jet engines, produces propulsive power by increasing the momentum of the working fluid through some form of heat release so that the momentum of the exiting jet exceeds the momentum of the entering air stream. In contrast to other air-breathing jet engines the working cycle is accomplished without mechanical compression of the working fluid and without intermittent semiclosed combustion cycles;"
consequently the ramjet is mechanically the least complicated air-breathing engine yet devised.

The basic elements of the engine can be represented schematically as in Fig 3. The charge air is inducted to the combustion or heat release chamber through a diffuser, so-called because the entering air stream is generally decelerated from the forward speed of the engine to a lower combustion entrance velocity. In engines employing liquid or gaseous fuels for the energy addition to the charge air, the fuel is injected through orifices or nozzles at some point in the diffuser process. The location of the point of fuel injection depends upon the desired degree of atomization and vaporization of the fuel and the desired mixture pattern of the fuel and charge air at the entrance to the combustion chamber.

At the end of the diffuser passage is located a flame holder, a turbulence-generating device offering shelter to the subsequent combustion flame. The fuel-air mixture entering the combustion chamber is ignited in the sheltered regions of the flame holder by some independent ignition source. In most engines this ignitor is only required to function at the onset of combustion, the combustion process being self-propagating for continuous operation.

The combustion of the fuel-air mixture is ideally completed in the combustion chamber, normally a constant area passage. (In ramjets utilizing an external heat source such as a nuclear reactor, the fuel-injection, flame-holding combustion process is replaced by a heat exchanger from which the uncontaminated charge air is discharged at an elevated temperature.)

The products of combustion leaving the combustion chamber are, in general, at a higher pressure than the free stream air surrounding the engine. The exhaust nozzle is used to expand the flow and thereby to convert the pressure into velocity. The flow velocities after combustion are subsonic; hence an initial contraction of the flow area is necessary. This contraction must be sufficient to either reduce the internal pressure to the ambient value or to accelerate the internal flow to sonic velocity. At low design flight speeds, ambient pressure is achieved with unchoked flow and the nozzle terminates with the convergence. At high flight speeds the internal flow chokes at pressures above ambient and for maximum thrust realization a diverging passage is added downstream of the throat in which the flow continues to expand to supersonic velocity until ambient pressure is attained.*

The fundamental limitation of the ramjet is its requirement of sufficiently rapid motion for the engine to become operative. "The engine will not yield an internal thrust force until the flight speed is high enough to produce a diffuser pressure rise that is sufficiently great to exceed the pressure losses in the flame holder, combustion chamber, and exhaust nozzle. An effective propulsive thrust force will not be obtained until the flight speed is enough greater than this minimum value for the internal thrust to compensate the drag directly chargeable to the engine installation. A yet higher speed must be obtained before the propulsive thrust is great enough to overcome the drag of the vehicle in which the engine is installed and thus to qualify the engine as a useful propulsive device."

The minimum flight speed at which the ramjet can qualify as a propulsion system for aircraft depends upon the degree to which the internal pressure losses and the installation drag are minimized and upon the drag of the propelled airframe. In general it has been found that the minimum speed application of the ramjet is not less than about 400 mph. (This speed is the relative speed of the engine and not necessarily of the airframe. Thus, ramjet engines, tip-mounted on the rotors of helicopters, have proved to be adequate propulsive devices at zero forward airplane speed.) The efficiency of the engine is very low at these minimum speeds, however, and applications of
interest at subsonic speeds are confined to installations in which the attractions of light weight, mechanical simplicity, and low cost outweigh considerations of fuel consumption.

As the engine flight speed is increased the thrust increases as a function of the square of the flight speed in the same order as the airframe drag. Thus a ramjet engine, properly designed, is capable of producing the very high thrusts necessary for supersonic propulsion. The engine efficiency, and hence the specific fuel consumption, ideally improve as flight speed increases so that the principal range of application of the ramjet as a competitive power plant appears to lie in the propulsion of supersonic vehicles.

The inability of the engine to produce a useful propulsive force at zero or low flight speeds necessitates the use of an auxiliary power plant to initially accelerate the vehicle to the required take-over speed of the ramjet. Such auxiliary power is also required for controlled landing of the vehicle. As a consequence the ramjet engine is not well-suited for conventional aircraft applications (with the exception of the helicopter) and the principal application of the engine appears to be further restricted to missiles or other similar vehicles of a one-flight expendable nature.

**Pulse Jet.** We quote from Sect F, Chapter 4 of Ref 1:

"In its standard form the pulse jet consists of a shaped tube fitted with flow-check valves at the front end (Fig 4). The air flowing into the engine through the valves is mixed with a continuously sprayed fuel. The mixture is then fired. As a result of the pressure rise which accompanies the explosion, the inlet valves close and the exhaust gases are forced out through the tailpipe. Expansion waves which are generated in the discharge reduce the pressure behind the check valves until they open again to admit a fresh charge of air, and the cycle is repeated. A spark is needed only for starting; once a regular cycle is established, each fresh charge of mixture is ignited by the hot gases from the preceding explosion and operation proceeds without further use of the spark plug.

Despite the mechanical simplicity of the pulse jet, a great deal of information is still lacking about such important details as its operating cycle as the flow behavior through the intake valves and at the discharge end, the timing and mechanism of ignition, and, above all, the process of combustion. This lack of basic information has necessitated the use of quite arbitrary assumptions in all theoretical investigations of the pulse jet cycle."

---

![Fig 4](image-url)
**Comparison of Performance Parameters of Jet Engines.** The excellent diagram below is taken from pp 96 & 97 of Ref 1

<table>
<thead>
<tr>
<th>Propulsion Method</th>
<th>Picture</th>
<th>Mass of air or gas handled m&lt;sup&gt;3&lt;/sup&gt;</th>
<th>Thrust F proportional to mass handled and change in momentum</th>
<th>Efficiency η of converting fuel energy to mechanical energy</th>
<th>Propulsive efficiency</th>
<th>Overall efficiency</th>
<th>Relative frontal area (drag)</th>
<th>Relative weight of fuel for a given duration</th>
<th>Probable range of maximum flight speeds</th>
</tr>
</thead>
</table>
| Rocket            | ![Rocket](image) | 1.0                                   | 1.0                                                         | 0.1                                                        | 0.25                 | 0.13              | 0.07                        |                                | 1. Above 600 mph  
|                   |         | 1.0                                   | 1.0                                                         | 0.1                                                        | 0.25                 | 0.13              | 0.07                        |                                | 2. For aid in take-off  
|                   |         |                                       |                                                             |                                                            |                      |                   |                             |                                | 3. For flying bombs |
| Ramjet            | ![Ramjet](image) | 1.0                                   | 1.0                                                         | 0.1                                                        | 0.25                 | 0.13              | 0.05                        |                                | 1. Above 500 mph  
|                   |         |                                       |                                                             |                                                            |                      |                   |                             |                                | 2. For flying bombs |
| Pulse jet         | ![Pulse jet](image) | 1.0                                   | 1.0                                                         | 0.1                                                        | 0.25                 | 0.13              | 0.04                        |                                | 1. 300 to 600 mph  
|                   |         |                                       |                                                             |                                                            |                      |                   |                             |                                | 2. For flying bombs |
| Turbojet          | ![Turbojet](image) | 1.0                                   | 1.0                                                         | 0.1                                                        | 0.25                 | 0.13              | 0.10                        |                                | 400 to 1500 mph |
| Turboprop         | ![Turboprop](image) | 1.0                                   | 1.0                                                         | 0.1                                                        | 0.25                 | 0.13              | 0.10                        |                                | 300 to 600 mph |
| Supercharged reciprocating engine with geared propeller and exhaust jet | ![Supercharged reciprocating engine](image) | 1.0                                   | 1.0                                                         | 0.1                                                        | 0.25                 | 0.13              | 0.10                        |                                | 150 to 450 mph |

Note that overall efficiency of all the above engines is primarily controlled by the cycle fuel efficiency & the propulsive efficiency.

Written by J. Roth

7) M. Barrere, “Research in the field of chemical propulsion,” NASA Accession No
N65-21452, Rept No CPIA-67, 4-6 (1964).

Jet Propulsion, History of. Early uses of the principles of jet propulsion and their evolution into modern jet aircraft are summarized in the tabulation below (Ref 2)

c 100 BC Hero built a steam jet engine, called an aeolipile, in Alexandria, Egypt
1232 Chinese used rockets to frighten enemy soldiers
1500 Leonardo da Vinci proposed using upward movement of hot gas in fireplace chimney to turn a spit for cooking meat
1629 Italian engineer Giovanni Branca built a crude steam turbine which drove a machine
1678 Ferdinand Verbiest, a Jesuit in China, built a model carriage, using a jet of steam for power
1687 Sir Isaac Newton stated the law of action and reaction
1791 John Barber, an Englishman, patented a gas turbine that was an ancestor of the turboset
1908 Rene Lorin, a French engineer, made detailed drawings of a proposed ramjet engine
1926 A. A. Griffith in Great Britain described theories of using gas turbines in aircraft
1930 Frank Whittle of Great Britain patented design for a jet-aircraft engine
1939 Heinkel Company in Germany built and flew the first jet-engine airplane
1941 First successful turboset airplane was flown in Great Britain. Engine designed by Whittle
1944 First jet-propelled fighters used in WWII by Germans
1947 First supersonic (faster than speed of sound) rocket-powered airplane was flown in the US by Charles Yeager
1952 First scheduled airline flights by jet transports begun by Great Britain
1955 First jet-engined convertiplane that takes off and lands straight up and down was tested in the US
1957 High-energy fuels that increase jet ranges almost 50% were put into commercial production
1958 First commercial jet passenger service between New York and London
1960 Turbofan engines using fans for added thrust in place of afterburners came into commercial use

More detailed, fairly recent historical data on Air Flow Jet Engines are given in Ref 1, pp 29-53 (including 201 refs). The subjects covered in this excellent review are: Piston
Engine Jet Development; Turbojet Development; Ramjet Development; Development of Intermittent Jets; Other Forms of Air Flow Jet

A.D. Baxter, the author of this chapt of Ref 1 summarizes his review as follows:

"In conclusion, it may be observed that in the ten years following the 1939-1945 war, air flow jet engines supplanted piston engines in almost all high performance military aircraft, they made large inroads on the civil aviation field including the first regular turbojet passenger service in 1951, they made supersonic flight a commonplace, and finally they doubled the world air speed record"


Jet Propulsion Unit “Decipede”. A jet propulsion unit, which develops 1000 pounds thrust for 43 secs, was developed at the request of the Engineer Board Field Station. The unit is the motor power of an amphibious “Snake” employed in Marine & Army land operations. It burns ten 8½-inch diameter grains of CP 401 propellant simultaneously in a single motor tube. The propellant gases come out of five pairs of nozzles spaced along the side of the motor tube. This unit, informally known as the “Decipede” was developed to the point of satisfactory operation

Ref: Central Res Lab, Monsanto Chemical Co, “The ‘Decipede’, a 1000 Pound Thrust, 43 Second Burning Time, Jet Propulsion Unit”, OSRD 5704 (Nov 1945)

Jet Pumps are devices for transferring liquids from one point to another, using the fluid pressure as an operating medium. They also may be used for mixing liquids. Two types of jet pumps are of interest to the chemical engineer. One of them is the ejector (siphon, exhauster or eductor), which is designed for use in operations where the head pumped is low and is less than the head of the fluid used for pumping by pressure. Another device is the injector. It is operated by steam and used for boiler feed or similar services, in which the fluid being pumped is discharged into a space under the same pressure as that of the steam that is used to operate the injector

Ref: Perry (1950), 1439; (1963), pp 5-17 & 6-13

Jewel Powder. One of the early American smokeless propells of the Ballistite type (See Vol 2 of Encycl, p B8-L) proposed by Monroe

Ref: Daniel (1902), pp 387 & 447

Jewler Explosives. See Ievler Explosives in this Vol, p 14-R

Johnite Explosives. See Jahnite Powders in this Vol

Johnson & Alexander Priming Composition. These inventors were issued British patents in 1856-57 for a priming compn containing amorphous P 8.3 & lead nitrate, [Pb(NO3)2], 91.7%

Ref: Daniel (1902), pp 8 & 388

Jolt & Jumble Tests. These tests have been used for many years to establish the safety and general ruggedness of fuze & booster designs under the application of repeated shocks in several directions. Both tests were designed originally as a simulation of shocks received during transportation of US Army caissons over rough terrain. In present applications the tests are intended to accurately represent actual conditions which may be encountered in transportation, handling or use of fuzes. They are rather a deliberate exaggeration of severe conditions to which an item might be exposed during transportation or use. As a result of the long use of these tests, much information has been accumulated regarding the behavior of a wide variety of fuzes in the tests. Although it is not a requirement that the fuzes be operable afterward, some fuze designers do require that their fuzes remain operable. In such cases, operability is judged by examination only, although firings may be
conducted in addn where considered appropriate by the designer.

**Jolt Test.** It consists of assembling the test item into the end of a pivoted arm which under cam action falls by gravity thru 4 inches on a heavy bed plate, giving a considerable jolt. Each item is tested for 1750 jolts, at a std speed of 35 jars per min, in each of three positions: vertically up, vertically down & horizontal. All fuze expl elements shall be present in the fuze during the test. The jolt testing machine is shown on US Ordnance Corps Drawing 81-3-30. It takes 50 mins to test a sample in each of the three positions, or ca 2½ hrs for a complete test.

The criteria by which samples are judged to have withstood this test are: 1) no element shall explode and 2) no parts shall be broken, deformed, be displaced, come apart, or arm in such a manner as to make the assembly unsafe to handle, or dangerous to use. Breakdown & inspection, together with engineering judgment, are usually the basis for the decision.

**Jumble Test.** It consists of placing the test item in a std steel fixture which completely encloses the item, then putting the fixture inside a rectangular hardwood maple box, which is rotated about its diagonal corners. The fixture containing the item rolls inside the box, receiving bumps at random. The box rotates at 35 rpm and the test consists of 3600 revolutions. It takes about 2½ hrs. The jumble test machine is shown on US Ordnance Corps Drawing 81-3-35. The criteria for passing the test are the same as given under Jolt Test.

**Note:** Particular care should be exercised in handling test items after the jolt & jumble tests; a safety shield should be used. The articles may become armed or the expl chge may have shifted out. In these cases, a slight jar may initiate a defective fuze. The jolt test is always run first. Total time for both tests is ca 5½ hrs.


---

**Jones Detonator Loader.** An automatic loading machine developed in 1943 by R.A. Jones of Ohio in cooperation with engineers at Picatinny Arsenal. This machine proved to be a great success during WWII and saved about 175000 man hours per year.

Ref: E. Cox, ArmyOrdn 29, No 152, 282–84 (1945)

**Jones Blasting Explosives.** Jones patented the following compn: chlorates such as NaClO₃ are used with finely divided carbonaceous substances & the o-isomer of nitrotoluene, which keeps the mixt plastic. The chlorate may be mixed with a powdered carbonaceous absorbent material such as sawdust & a liq mixt of o-nitrotoluene & melted TNT.

Ref: L.T.W. Jones, USP 1820567 & 1820568 (1931) & CA 25, 5771 (1931)

**Jones Dynamite.** An Australian Dynamite consisting of NG 30–35% absorbed by a mixt of Ca sulfate & kieselguhr 70–65%.

Ref: Daniel (1902), p 388

**Jones Smokeless Powder.** One of the first progressive-burning smokeless proplnts patented in 1897 in Austria. It was prep’d by coating grains of proplnt with a thin layer of resin, fatty acids or carnauba wax. About ½ part of coating per 100 parts smokeless proplnt was used. This method retarded combustion, especially at the start, and allowed attainment of a fairly progressive increase in the rate of burning without increasing the pressure in the gun barrel.

Ref: Daniel (1902), p 388

**Jordan Pulping Machine.** It is used in preparing cotton for nitration. The machine operates on the centrifugal principle, by means of which the entering cotton is thrown from the large end of the heater to the small tapered end, and thence to a storage tank above the machine. The more efficient way to operate these units is to have them adjusted in such a manner that
the cotton is given successive cuts. In other words, the first treatment in the Jordan machine will give only a fairly coarse cut, the next a little finer, and so on, until the cotton has finally been pulped to the necessary degree of fineness.

Jouguet, (Jacques-Charles) Émile (1871–1943). French physicist, general inspector of mines and professor of mechanics École des Mines, École Polytechnique, member French Academy of Science (1930). He was the author of “Mécanique des Explosifs” (1917) and conducted research on wave diffusion, movement of fluids, explosives; and fundamental work on the hydrodynamic theory of detonation. His name is associated with that of Chapman in the famous Chapman-Jouguet condition. In their honor parameters of a steady detonation wave are usually designated by the subscript CJ.


Jouguet’s Rule. See in Vol 4 of Encycl, p D607-L

JP, JPH and JPN Propellants. See under Ballistite in Vol 2 of Encycl, p B9-R

Judson Dynamite or Powders. American expls patented by E. Judson in 1876 and manufd in Drakesville, NJ. These expls were, in fact, a combination of Black Powder & NG and were much more powerful than straight BlkPdr, although some of them contained only a small amt of NG. For example, one compn contained: NG 5, NaNO₃ 64, sulfur 16 & bituminous coal (cannel coal) 15%. Judson’s expls were also called RRP (meaning Railroad Powders). Judson’s patent may have been pre-dated by Nobel’s patent (Ref 1) for a mixt of NG & Black Powder.


Juinite. See Ethylenebisurethane and Derivatives in Vol 6 of Encycl, p E234-L

Jump Firing Test. In tabulating the elevations & azimuths required to obtain a given range and deflection of a gun, account is taken of the fact that usually the direction of the target to the trajectory at the origin differs from that of the axis of the bore of the gun before it is fired. This difference in direction is called jump. The jump may be in any direction, but as a matter of convenience it is described by two coordinates, vertical & horizontal.


Jumping Detonation. See Detonation, Jumping in Vol 4 of Encycl, p D421-L

Junk Test. See Bergmann-Junk Test in Vol 2 of Encycl, p B102-R

Juno. A space research vehicle which is a four-stage version of the Juniper C (qv). The first stage burns a high energy hydrazine fuel, called Hydyne (developed by the North American Aviation Co), and liq oxygen. The 2nd, 3rd & 4th stage thrusts are solid propellants. Characteristics of Juno 1 are length 68.6", diam 70", wt loaded 64000 lbs, useful pay load 10–15 lbs, altitude 2000 miles, and velocity 19000 miles/hr. The rocket was used to launch several of the Explorer satellites.


Juno 2. A space research vehicle having 4 stages of thrust. The first stage burns a hydrocarbon, RP-1, and liq oxygen. The other three
stages burn solid propellants. Length of the rocket is 76', diam 105", wt loaded 121000 lbs, and pay load 13–80 lbs. The vehicle was used to orbit Explorer 7

Juniper C. A space research vehicle having 3 stages of thrust. The first stage burns either liq oxygen & ethyl alcohol or liq oxygen & Hydyne (See under Juno 1, above). The second & third stages are solid propellants. The rocket is 66.6' in length, diam 70", wt loaded 64200 lbs, pay load 300 lbs, range 1600 miles, altitude 400 miles, and velocity 14700 miles/hr

Jupiter Dynamite or Powder. One of the older American type No 2 Dynamites, similar in compn to Neptune or Vulcan powders
Ref: Daniel (1902), p 389

Justice Powder. Justice patented in 1888 an expl mixt consisting of a nitrate & a chlorate, such as KClO₄, consolidated by means of molten paraffin or naphthalene

Jute, C₁₂H₁₈O₉; mw 306.26, O 47.02%; fibers of East Indian, South American & Chinese plants, such as Corchorus, Olitorius, Capsularis & Abyltron Avicenna. It is used extensively in the manuf of gunny sacks, bags & twine (Ref 6). On nitration it yields:
Nitrojutes, C₁₂H₁₅O₆(NO₃)₃, N 9.52% to C₁₂H₁₄O₅(NO₃)₄, N 11.52%; expl compds prepd in 1889 by Cross & Bevan (Ref 1) by nitrating jute with mixed acid contg about equal vols of concd nitric & sulfuric acids. The authors do not state the ratio of acid to jute. Temp of nitration was 18°, time 30 mins, N content 10.5% which corresponds to a compn between the above formulas
Mühlhaeuser (Ref 2) nitrated jute previously purified by boiling in a 1% Na₂CO₃ soln followed by a water rinse. By using 1p dry jute & 15 parts of acid, 2/1 concd H₂SO₄/concd HNO₃, a product with 12.26% N and 132% yield was obtd. The props of Nitrojute are similar to those of nitrated lignocelluloses (See Lignin Nitrate in this Vol)

Since the cost of jute is higher than that of cotton or wood pulp, and since Nitrojutes are no better than ordinary NC, there is no advantage in manuf Nitrojutes. They were never used in expls, except for experimental purposes
K (Pulver). See under Erosion of Gun Barrels in Vol 5, p E116-L

K1 and K2 (of Muraour). For a proplnt whose burning rate, V, may be expressed by V=a+bp, where a and b are constants and p is the ambient pressure, Muraour defines a constant, K1, which is the total area under an ideal pressure-time curve, and not the real pressure-time curve. The ideal combustion rate V1 is given by V1=bp. At low ambient press, the real \( \int p\,dt \), based on V, is always less than the “ideal” \( \int p\,dt \) which is obtained from V1. Thus K1= \( \int p\,dt \) (based on V1) is always greater than the real \( \int p\,dt \) (based on V). The constant K2 is obtained by dividing K1 by the proplnt thickness, and is inversely proportional to the rate of diminution of proplnt thickness for a proplnt ignited on two opposite faces.


K-1 Explosive (Russ K-1 Splav, K-1 Fusion, Mixture K-1). Mixture consisting of TNT 70 and DNB 30%, used during WWII for filling some land mines made of cast iron. Although this mixt was less brisant than TNT, it was nevertheless too brisant for a container made of cast iron material, and sometimes broke it into fragments too small to be effective against personnel. For this reason, the brisance of K-1 was reduced by the insertion in its mass of long blocks of a less brisant expl, such as Schneiderite. Because of the toxicity of K-1, it was preferable to use K-2, described below.

Refts: 1) Shilling (1946), 240 2) PATR 2145 (1955), Russ 10

K-2 Explosive (Russ K-2 Splav, K-2 Fusion, Mixture K-2). Mixture containing TNT 80 and DNN 20%, used for loading 82mm Land Mines made from cast iron. It was less toxic than K-1, described above.

Refts: 1) Shilling (1946), 240-41 2) PATR 2145 (1955), Russ 10

Kadinite. A Dynamite containing NG 26, Na nitrate 56, sulfur 10, carbon 4 & ligneous materials 4%. Daniel (Ref 1) uses the name Kadmit for the same compn

Refts: 1) Daniel (1902), 390 2) M. Guia, Trattato 6 (1) (1959), 388

Kailpinites. Expls resembling Cheddites (See Vol 2 of Encycl, p C155-L) developed in Fr based on the study of the Belg Yongkites. Kailpinite O No 12 contains Amm perchlorate 38, Na nitrate 28 & TNT 34%; Kailpinite O No 13 consists of Amm perchlorate 38, Na nitrate 31 & TNT 28%. They are used for demolition and mining purposes.


Kalialmatrit No 55. See under Almatrites in Vol 1, p A140-L

Kallenites. Dynamites patented in Australia in 1899 by Callaghan & Fraser, recommended for underground work because they did not produce poisonous gases. They contained NG, absorbed by finely ground eucalyptus leaves and tree bark. Some varieties contained K nitrate and NC.

See Fraser & Callaghan in Vol 5, p F186-L

Reft: Daniel (1902), 390

Kamikaze Bomb. See under Baka Bomb in Vol 2, p B4-R

Kanite. A US AN expl containing a substance akin to oxidized (blown) oil.

Reft: W.H. Blumenstein, Interstate Corn Com. Commission Opinion No 1557 (1 June 1911) & CA 5, 2949 (1911)

Kanone. Ger or Swiss for Cannon

Kapsiul. Russ for blasting cap, detonator or igniter.

Reft: B.T. Fedoroff et al, PATR 2145 (1955), p Russ 28
**Kapsiul' Detonator.** Russ blasting cap containing the following expl comps (Ref 1)

<table>
<thead>
<tr>
<th>Composition (grams)</th>
<th>Type</th>
<th>MF</th>
<th>Stab Mixt</th>
<th>Lead Stypnate</th>
<th>LA</th>
<th>Base Charge</th>
<th>Use</th>
</tr>
</thead>
<tbody>
<tr>
<td>No 8 &amp; No 8M</td>
<td>0.5</td>
<td></td>
<td></td>
<td></td>
<td>1.02</td>
<td>1.00 PETN or RDX</td>
<td>Blasting</td>
</tr>
<tr>
<td>No 8A</td>
<td></td>
<td>0.10</td>
<td>0.20</td>
<td></td>
<td>0.11</td>
<td>Tetryl</td>
<td>same Blasting</td>
</tr>
<tr>
<td>TAT-1 &amp; TAT-2</td>
<td></td>
<td>0.06</td>
<td>0.21</td>
<td></td>
<td>0.11</td>
<td>Tetryl</td>
<td>Fuze</td>
</tr>
<tr>
<td>M-1</td>
<td></td>
<td>0.20</td>
<td>0.10</td>
<td></td>
<td></td>
<td>Tetryl</td>
<td>Fuze</td>
</tr>
</tbody>
</table>

**Kapsiul' Vosplamenitel'.** Igniter cap, usually referred to as *slib*. If followed by “udarnykh sostavov”, it refers to percussion igniters. The formulation used in rifle (“vintovochnyi”) percussion caps is MF 16.7, K chlorate 55.5 & Sb trisulfide 27.8%; for pistol (“revolv'ernyi”) percussion caps, MF 25.0, K chlorate 37.5 & Sb trisulfide 37.5%; and for mine throwers (“minomietnyi”), MF 35.0, K chlorate 40.0 & Sb trisulfide 25.0% (Ref 2)

Igniter caps containing non-corrosive (“nekorrodiruyou-shchikh”) percussion comps: MF 67.8, Ba nitrate 29.6 & Sb trisulfide 2.6% (for rifles); Czech “Oksid”: Lead Stypnate 45.0, Tetraene 5.0, Ba nitrate 30.0 & Sb trisulfide 20.0% (for pistols) (Ref 3)

**Kapsiul' Vosplamenitel', Trubochnye.** Tubular igniter caps containing the following percussion mixts: a) KTM cap — MF 25.0, K chlorate 37.5 & Sb trisulfide 37.5%; and b) RGM cap — MF 50.0, K chlorate 25.0 & Sb trisulfide 25.0% (Ref 4)


**Kardox Cartridge.** See under Cardox in Vol 2, p C67-R

**Karioo.** See under Carlit in Vol 2, p C68-R and under Japanese Explosives in this Vol

**Kariitto.** See under Carlit in Vol 2, p C68-R


**Karman, Theodor von** (1881–1963). Hungarian scientist who worked in Germany & the US on aeronautical problems, on rockets, and on combustion phenomena

Ref: Anon, Explosivist 1963, 134

**KA-Salz.** One of several Ger names for RDX. See under Cyclotrimethylenetritramine, Cyclonite or RDX in Vol 3, pp C611-L to C626-L

**Kast Brisance Meter.** See Brisance in Vol 1, p IX; Compression Tests in Vol 1, p X; also see under Brisance or Shattering Effect in Vol 2, pp B265-L to B297-L, and Brisance Test Methods in Vol 2, pp B299-L to B300-R

**Kast, H** (1869–1927). A Ger scientist who specialized in expls, and was for many years associated with the Chemisch-Technische Reichsaustalt. He designed an apparatus for the detn of brisance of expls (brisance meter), and developed a formula for theoretical calculations of brisance. Kast also developed an impact machine (Kast Stauchapparat) that was
for many years the “standard” impact test for expls. He was the author of several books and articles on expls
Ref: F. Lenze, SS 22, 305 (1927) & CA 22, 1238 (1927)

Katiusha (Katyusha or Kostikov’s Gun). Russ 132mm rocket launcher M-13. It consisted of a truck-mounted system for firing 16 rockets from eight rails. The M-13 weighed 7.1 tons and fired 94-lb rockets to a max range of 9846 yards
Ref: 1) Anon, FieldArtil (Nov 1943), 816
2) G. Underhill, InfantryJ (May 1945), 44–49

KDNBF. Acronym for the K salt of Dinitrobenzofuroxan. See Vol 2, p B68-R

Keeping Test. One of the tests used in England to determine the exudation of Dynamites; other methods include press and centrifugal tests. In this procedure, a previously weighed cartridge of Dynamite is stored in a vertical position for 6 days at 40°, and then reweighed. If leakage exceeds 5%, the expl is considered unsatisfactory. With straight Dynamites, this test correlates well with the pressure test; for other Dynamites, agreement is not as good
Ref: Marshall 2 (1917), 421

Keil Explosive. A mixt of nitrodetroglucone (predp from starch) with K nitrate or chloride & vegetable fibers
Ref: Daniel (1902), 391

Kekulé Oil. A liq expl (d=1.47g/cc) prep’d in 1869 by passing ethylene gas into mixed nitric-sulfuric acid. The oil consisted of a mixt (about 50/50) of Nitroglycerol, CH₃(ONO₂)₂, CH₂(ONO₂), and beta-Nitroethyl Nitrate, CH₂(ONO₂)CH₂NO₂
Refs: 1) Naoum (1928), 220–21 2) Davis (1943), 228

Kelbar Powder Company Explosive. The Kelbar Powder Co of Avondale, Pa developed an expl contg Amm Nitrate 93.37, resin 6.57 & moisture 0.05%. It had satisfactory brisance and stability was comparable to TNT & 80/20 Amatol, but was sensitive to bullet impact and was rendered very insensitive by compression. Its high mp (about 160°) made loading of shells or bombs by casting or extrusion impracticable, therefore the expl was thought to offer no particular promise
Ref: J.D. Hopper, “Study Ammonium Nitrate Explosive Received from Kelbar Powder Company”, PATR 1009 (Oct 1939)

Kelbetz Explosives. Expls patented in England in 1896 by an Austrian, Kelbetz. They consisted of AN mixed with oxalates of aromatic amines, such as aniline and toluidine, and sometimes also contained small quantities of charcoal. Other Kelbetz expls contained about 95% AN, fatty acids such as stearic, palmitic, oleic, etc (with or without their metallic soaps, such as Ca), and small quantities of charcoal
Ref: Daniel (1902), 391

Kellow and Short. Patented in 1862 in England an expl mixt consisting of K nitrate, Na nitrate, K chloride, sulfur, barkmeal or sawdust
Ref: Daniel (1902), 391

Kelly, Bell and Kirk Explosive. Australian Dynamite patented in 1899, containing NG, K nitrate, cork and calcined eucalyptus leaves
Ref: Daniel (1902), 391

Kent, Robert H (1886–1956). American ballisticsian at Aberdeen Proving Ground, Maryland from 1922. His achievements are described in a series of articles in Ordnance 40, 769–81 (1956)

Kent Powder. A Brit coal mine expl no longer on the “permitted” list. It contained NG 24,
K nitrate 32.5%, wood meal 33.5% & Amm oxalate 10%. It had a charge limit of 32 oz, and power by ballistic pendulum of 2.01".
Ref: Marshall, Dict (1920), 53

Kentite. A Brit "permitted" coal mine expl of the Favier type. It contained AN 34, K nitrate 34, TNT 15 & Amm chloride 17%. It had a charge limit of 18 oz; power by ballistic pendulum 2.64", compared with 3.27" for Brit standard 60% Gelfignite (See Vol 5 of Encyc, pp G57-Rff)

Kerosene (Kerosine, Coal Oil, Astral Oil). A mixt of petroleum hydrocarbons, chiefly of the methane series having from 10 to 16 carbon atoms per molecule. It constitutes the fifth fraction in the distn of petroleum, after the petroleum ethers and before the oils. Pale yell or w-white, mobile liq; characteristic, not altogether disagreeable odor; bp 175–325°, flash pt 150–185°F, d about 0.809/cc. Insol in w, misc with other petroleum solvents. Besides uses as a fuel, illuminant and cleansing agent, it is used as a rocket and jet engine fuel (Refs 1, 3 & 4)

An underground storage tank, partially filled with kerosene, exploded, killing 29 people. The expln occurred after CO₂ was pumped into the tank to test a fire-extinguishing installation (Ref 2)

Kerosene Nitrate. Fiala claims the nitration of kerosene by adding 3–5% of concd nitric acid per liter of kerosene with vigorous stirring, and allowing it to stand for several days
Ref: F. Fiala, USP 2511433 (1950) & CA 44, 8103 (1950)

Ketten Explosives (Kessensprengstoffe in Ger). Several expl mixts were proposed by W. Kessen of WASA-G (Wastfälsich-Anhaltsische Sprengstoff Aktiengesellschaft). One of such compns, patented in 1938 (Ref 1), consisted of BlkPdr used for blasting mixed with NG and/or nitroglycerin and a large amt of Na bicarbonate. It was intended for use in gaseous coal mines.

Another patent issued to Kessen (Ref 2) dealt with the manuf of moist AN expls contg large amounts of carbonaceous materials
Refs: 1) W. Kessen & WASA-G, BritP 493984 (1938) & CA 33, 2719 (1939) 2) Ibid, GerP 679511 (1939) & CA 33, 9647 (1939) 3) PATR 2510 (1958), Ger 101-L

Ketocyclopentane. See Cyclopentanone in Vol 3, pp C603-R to C603-L

Ketohexamethylene. See Cyclohexanone in Vol 3, p C597-L

Ketone Peroxides. Bjorklund and Hatcher (Ref 1) prepd several expl compds by treating ketones with hydrogen peroxide–nitric acid mixts. The probable compns of these compounds are as follows:
1) (C₃H₆O₂)₃; white solid; mp 95–97°; was obtained by adding acetone to a mixt of H₂O₂ and 70% HNO₃; it exploded violently on impact, sudden heating or treatment with concd H₂SO₄
2) (C₃H₆O₂)₂; white solid; mp 131–132°; was obtained by adding a mixt of H₂O₂–70% HNO₃ to acetone; it is a violent explosive
3) (C₅H₈O₂)₃; white solid; mp (decomp 166–167° and explodes violently at higher temps); was prepd from cyclopentanone, H₂O₂ and HNO₃
4) (C₅H₈O₂)₂; white solid; mp 126–27°; was prepd from cyclohexanone, H₂O₂ and HNO₃; it is a violent explosive
5) Explosive oils of unknown compn obtained by treating methylethylketone, ethylethylketone and 3-methylocyclohexanone with a H₂O₂–HNO₃ mixt
2-Ketotrimethyleneimine. See 2-Azetidinone in Vol 1, p A519-L

Ketoximes and Derivatives. The Dow Corning Corp reported an exp1 hazard inherent to ketoximes and many of their derivatives. In manuf ketoximosilanes,

\[ RSi\left(O-N=C<_{\text{Et}}^\text{Me}\right)_{3} \]

two potentially devastating explosions were experienced. An analysis of the incidents strongly suggests that acidic conditions were inadvertently generated which markedly lowered the threshold temps required for expl decompn of these materials. It is suspected that conditions were unintentionally produced under which the highly exothermic Beckman rearrangement preceded, and that the resulting high temps resulted in the explosively rapid formation of huge volumes of gaseous degradation products.

Subsequent expts led to the following findings: 1) Methylthylketoxime can be distilled at atmos press (152°F) only if highly purified; 2) The presence of impurities, especially acidic impurities, drastically lowers the temp at which degradation occurs; the hydrochloride salt of methylthylketoxime undergoes violent degradation when heated to 50–70°F; 3) Although the derived ketoximosilanes can be distd without difficulty at reduced press, here again the presence of acidic impurities drastically lowers their thermal stability. FeCl₃ at 10ppm did not trigger the degradation even at 250°F, but 500ppm yielded an expln at 150°F, and 2% lowers the onset of violent degradation to 50°F. As with the ketoxime itself, the presence of HCl salts can cause expl degradation at relatively low temps.


Keystonite. See under Alkalies, Action on Aromatic Nitrocompounds in Vol 1, p A126-R

The wrapped charges are dried at 60–70°F and then dipped in paraffin. They were used as bursting charges in mines.

Ref: 1) Anon, PB Rept No 925 (1945), 48 2) PATR 2510 (1958), Ger 101-L

Kickless or Recoilless Weapons. Weapons where the recoil effect is reduced and counteracted by vents in the breech which deflect up to 80% of the expansion gases to the rear. Recoilless weapons can therefore use artillery-sized ammo without generating artillery-sized recoil, or any recoil at all. Thus, the heavy montings and recoil-absorbing devices needed with artillery weapons are not required; with internal stress and muzzle energies low, the barrel can be much thinner and lighter than that of a full-recoil gun firing the same weight of shell. Naturally, the effective range is also much smaller than with full-recoil weapons.

For calibers of about 100mm, a recoilless rifle (RR) will have a max range of 1000–2000 yards, as opposed to a full-recoil gun range of about 25000 yards. But the ratio of weapon weights is also of the order of 1 to 20 even counting the wheeled mounts used with some RR’s, and cost ratios are also in this range. Recoilless weapons are used whenever the range is secondary to the size of shell delivered, and where high velocities are not required. RR’s are primarily intended for antitank use where their “hollow charge” (HEAT) shells penetrate armor without needing the kinetic energy required by conventional solid shot fired from full-recoil artillery.

The US 106mm RR and two earlier US models which can be fired from the shoulder, the 75 and 57mm, are widely used, as is the Russ B-11 107mm RR and the older B-10 82mm RR. Another widely used type is the one-man 84mm Carl Gustav, manuf in Sweden and used by, among others, the Brit army.

The largest recoilless weapon, the Brit Wombat 120mm, is not widely used. These are all properly called recoilless rifles since their barrels are rifled as in the case of conventional artillery. Some light recoilless weapons have smooth barrels and project a HEAT charge attached to a reduced caliber “stick” which fits into the barrel. The West Ger Panzerfaust and the Russ
Kieselguhr or Guhr (Diatomite, Tripoli Powder). Soft bulky solid material, 88% silica, composed of skeletons of small prehistoric aquatic plants related to algae (diatoms). It has the property of absorbing up to three times its weight of NG and other liquids. It was formerly used extensively in the manuf of Guhr Dynamites, invented by Nobel in 1866 (See under Dynamite in Vol 5, pp D1595-L to D1596-R). Present-day straight Dynamites have various proportions of active constituents substituted for kieselguhr with resulting higher performance (Ref 3).

Kieselguhr has also been used as a coating agent for porous prilled AN in Nitro-carbonitrate (NCN) blasting agents, where it aids in the absorption of fuel oil (Ref 3).


Kinenite. AN expl containing DNB, somewhat similar in compn to Bellite (Vol 2, p B32-R), Roburite, Sekurite & Tonite
Ref: Colver (1918), 141

KINETICS IN EXPLOSION PHENOMENA

Introduction. In a very broad sense, kinetics (often called chemical kinetics) may be defined as the study of the rate of change of chemical reactions under non-equilibrium conditions. This incomplete definition will be augmented below, but first we must consider why kinetics are such an important facet in the practical and theoretical understanding of explosion phenomena.

There is general agreement that the energy sources for detonations, deflagrations, thermal explosions and various transition phenomena are exothermic chemical reactions. Thus, it is immediately clear that the rate of energy supply, ie, the rate of the chemical reactions, will have a most important bearing on the initiation, growth and propagation of the above processes. Obviously, the influence of kinetics on explosion phenomena is an immensely broad subject. Consequently, we must arbitrarily limit

RPG both consist of a small caliber tube (about 40mm) within which the charge and “stick” are set. When the weapon is fired, the “stick” and the much larger HEAT charge affixed to its end are projected out. These are short-range, one-man weapons fired from the shoulder (Ref 1).

A novel approach to reducing recoil in conventional artillery is the US XM-204 Soft Recoil Gun, still in an early stage of development. The barrel and recoiling parts are held in a firing position against a spring; when released they move forward. At their maximum forward velocity, the charge is fired. Much of the recoil energy is expended in checking the forward momentum of the barrel, the remainder is absorbed conventionally and “recocks” the spring used for the first part of the sequence. Recoil forces transmitted to the carriage are reported to have been halved (Ref 2).

Note: See also under Cannon in Vol 2, p C28-R

Kier or Keir. A large vat in which cotton, textile goods etc are boiled, bleached, etc

Kiering. Digestion of raw cotton, or other materials, in kettles called Kiers. For instance, raw cotton may be purified by digesting it in a kier with a solution of caustic soda
Ref: Marshall 3 (1932), p 29

Kiernan and Bowen Explosive. An expl mixt developed by Kiernan & Bowen of New Orleans, La, contg Na chlorate 75, turpentine 10, bone meal 10 & dried blood 5%. It was concluded that this expl had no value from a military standpoint because of its low brisance, high degree of sensitivity to impact, and the ease with which it lost turpentine, which resulted in decreased sensitivity to initiating agents
ourselves to a description of only a relatively few of its most important aspects, namely, detonations and thermal explosions, to the extent that the latter are required for a description of detonation effects. The kinetics of individual explosives, if known, will be described under the individual explosives, and, as already stated, we will be primarily concerned with the rate of energy supply, i.e., the rate of the thermal decomposition of explosives and the resulting exothermic reactions. The kinetics involved in the synthesis of explosive compounds, e.g., nitration kinetics, will be described elsewhere in this Encycl

_A Cursory Review of Chemical Kinetics._ In what follows we will employ certain concepts and terms widely used in the extensive literature on chemical kinetics. These concepts and terms will now be reviewed briefly. For details, the reader is directed to the general references listed at the end of this section

For a generalized chemical reaction whose stoichiometric (overall) equation is

\[ aA + bB = mM + nN, \]

the reaction rate \( R \) is defined by

\[
R = -\frac{1}{a} \frac{dA}{dt} = -\frac{1}{b} \frac{dB}{dt} = \frac{1}{m} \frac{dM}{dt} = \frac{1}{n} \frac{dN}{dt} \tag{1}
\]

where \( a, b, \text{ etc} \) are the number of moles of chemical species \( A, B, \text{ etc} \), and \( \frac{dA}{dt}, \frac{dB}{dt} \) etc., are the rate of change of species \( A, B, \text{ etc} \). If experiments show that the reaction rate is independent of reactant \( A \), the reaction is called zero-order in \( A \); if the rate depends on the instantaneous value of \( A \), the reaction is termed first-order in \( A \); if the rate depends on the instantaneous value of \( A^2 \), the reaction is called second-order in \( A \), and so on. In most explosion phenomena we will deal only with zero-order and first-order reactions. The rate equations for zero-order and first-order reactions are, respectively

\[
\frac{dA}{dt} = k_0 \quad \text{or} \quad A = A_0 - k_0 t = 1 - f \tag{2}
\]

and

\[
\frac{dA}{dt} = k_1 A \quad \text{or} \quad A = A_0 e^{-k_1 t} \tag{3}
\]

where \( k_0 \) and \( k_1 \) are the zero-order and first-order rate constants, respectively; \( A_0 \) is the concentration of species \( A \) at zero time; \( f \) is \( A/A_0 \), the fraction of \( A_0 \) reacted in time \( t \). The rate constants \( k_0 \) and \( k_1 \) have the units of moles/liter-sec and sec\(^{-1} \), respectively, if the concentrations are expressed in moles/liter. Obviously, the above definitions could have been made in terms of \( B \) or \( M \) or \( N \), except that in accordance with Eq 1, the minus signs of the differential equations for \( M \) and \( N \) (Eqs 2 & 3) must be replaced by plus signs.

An excellent review of higher order reactions, opposing reactions, consecutive reactions, etc, is given in Chaps II & III of Ref. 10 and in Ref 11

Rate constants are “constant” only at a fixed temp and vary strongly as the temp of the system changes. The following empirical dependence of the rate constant \( k \) on the absolute temp \( T \) is known as the Arrhenius equation

\[
\frac{dlnk}{dT} = \frac{E}{RT^2} \quad \text{or} \quad k = Z e^{-E/RT} \tag{4}
\]

where the Arrhenius parameters \( E \) and \( Z \) are called activation energy and frequency factor or pre-exponential factor, respectively. The usual dimensions of \( E \) are cal/mole. The dimensions of \( Z \) are the same as those of \( k \); thus for a zero-order reaction \( Z \) is in moles/liter-sec, and for a first-order reaction \( Z \) is in sec\(^{-1} \). The term \( E/RT \) is dimensionless. Based on kinetic theory, \( Z \) is interpreted (Refs 1, 2, 6, 10 & 11) as the number of collisions per unit volume per second of molecules \( A + A \) or \( A + B \), and the term \( e^{-E/RT} \) is a measure of the fraction of colliding molecules that result in reaction. Thus, if \( E=0 \), then \( e^{-E/RT}=1 \), and all colliding molecules react; conversely, if \( E/RT >> 1 \), almost none of the colliding molecules react.

An alternate approach, the so-called transition state or activated complex theory, is based on quantum mechanics and thermodynamics.

It postulates that the necessary condition for reaction is the crossing of a potential energy barrier. The initial configuration (reactants) passes over by continuous changes of coordinates to the final configuration (products), and there is always an intermediate configuration (the transition state or activated complex) which is critical for the process. Once this critical configuration is attained there is a high
probability that the reaction will go to completion (Refs 5, 6, 10 & 11). In terms of this theory \( E = \Delta H^* + RT \), where \( \Delta H^* \) is the difference in enthalpies between the activated complex and the initial state, and

\[
Z = \frac{k_B T}{h} e^{\Delta S^*/R}
\]

where \( k_B \) and \( h \) are the Boltzmann and Planck constants, respectively, and \( \Delta S^* \) is the entropy change in passing from the initial state to the activated complex.

Both the collision and activated complex theories predict a mild dependence of \( Z \) on \( T \), and the latter also predicts a mild dependence of \( E \) on \( T \). In practice, over the limited temp ranges of the usual expnl conditions, these mild dependencies are rarely observed. Both theories also predict that “normal” values of \( Z \) should be \( 10^{13} \) to \( 10^{14} \) sec\(^{-1}\) for unimolecular processes. This agrees with many expnl observations. In many cases, however, because of steric effects, \( Z \) can be much smaller than “normal”. Benson (Ref 12) presents evidence that \( Z \) for certain unimolecular gas reactions producing two free radicals, or for reactions involving the opening of a “small” carbon ring, is larger than “normal” and is of the order \( 10^{16} \) sec\(^{-1}\).

All of the above discussion is strictly applicable only to homogeneous gas phase reactions. Usually the above considerations do apply reasonably well to non-polar liquids and non-polar solutions, although “normal” \( Z \) values may be an order of magnitude less than for gas reactions. Reactions in solids are often much more complex, since they are usually heterogeneous, involve catalytic effects, reactions at preferential sites (dislocations, etc), and nucleation phenomena. These complicated processes are quite beyond the scope of the present article. For some description of these phenomena, and further references, the reader should consult Refs 9, 10 & 11.

Another important aspect that we have so far ignored is that reactions almost never actually proceed as represented by a stoichiometric equation (eg, Eq 1). Usually a stoichiometric equation is the algebraic sum of a number of steps called elementary reactions. Frequently one of these elementary reactions is much slower than the others, and thus controls the overall reaction rate. For obvious reasons such a reaction is called rate-controlling, and the overall rate constant is essentially that of this rate-controlling step.

For complex reactions, some of whose elementary steps are of comparable rates, but others are slower, the so-called steady state hypothesis (Refs 4, 6, 10 & 11) can occasionally lead to a simple theoretical description (mechanism) of the complex reaction. A famous example of this is the thermal decomp of \( N_2O_5 \), where the observed kinetics for this reaction are accurately first-order, even though the reaction is complex (Ref 10).

For overall Arrhenius parameters (based on measurements of the overall rate constant as a function of temp), there are no “normal” values if the reaction involves several elementary steps of comparable rates. Indiscriminate use of such parameters in assessing detonation phenomena (eg, hot spots) can lead to gross errors. The reader is reminded that much of the existing kinetic data for explosives is based on the measurement of the overall pressure changes in the system under study. Unless the detailed reaction sequence, usually called the reaction mechanism, is known, Arrhenius parameters based on pressure change measurement can be most unreliable.


**Application of Kinetics to Explosion Phenomena.**

Brief mention was made earlier of the various explosion phenomena that are strongly dependent on the kinetics of the exothermic chemical processes involved in explosions and detonations. This matter will now be considered in more detail as follows:

1) Steady Detonation, 2) Initiation Processes, 3) Stability and Storage, and 4) The “Ideal” Explosive

In what follows, the experimental Arrhenius parameters E and Z may be for the overall exothermic process and may not always refer to the E and Z of a simple elementary reaction in that process

**Steady Detonation.** The thermo-hydrodynamic theory of detonation is very successful in describing ideal detonation parameters (CJ states) without any reference to chemical kinetics. This is so because reaction rates at CJ conditions are extremely fast — at least for military explosives. Based primarily on the dependence of detonation velocity on explosive charge diameter, Jones, Eyring and others (Refs 2, 3, 10 & 11) have estimated that reaction times for ideal detonations of military explosives are the order of 0.1 to 1 µsec. Thus for ideal detonation conditions, f, the fraction of explosive reacted, must approach unity in times that are appreciably less than 0.1 to 1 µsec. In Table I we show that this is indeed the case. The calculations of f were made under the assumption of isothermal first-order decomposition at $T_{CJ}$ (conservatively estimated in the tabulation), and the kinetic parameters of Table III of a following section

---

**Table I**

<table>
<thead>
<tr>
<th>Explosive</th>
<th>Arrhenius Parameters*</th>
<th>$\tau$ (nanosec)</th>
<th>$T_{CJ}$ (°K)</th>
<th>Fraction Reacted (f)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TNT</td>
<td>Zinn &amp; Rogers</td>
<td>10</td>
<td>2250</td>
<td>1</td>
</tr>
<tr>
<td>TNT</td>
<td>Zinn &amp; Rogers</td>
<td>1</td>
<td>2250</td>
<td>0.70</td>
</tr>
<tr>
<td>NMe</td>
<td>Makovsky &amp; Gruenwald</td>
<td>1</td>
<td>3000</td>
<td>1</td>
</tr>
<tr>
<td>NMe</td>
<td>Makovsky &amp; Gruenwald</td>
<td>0.1</td>
<td>3000</td>
<td>0.74</td>
</tr>
<tr>
<td>PETN</td>
<td>Robertson</td>
<td>0.1</td>
<td>2550</td>
<td>1</td>
</tr>
<tr>
<td>PETN</td>
<td>Roth</td>
<td>1</td>
<td>2550</td>
<td>1</td>
</tr>
<tr>
<td>PETN</td>
<td>Roth</td>
<td>0.1</td>
<td>2550</td>
<td>0.99</td>
</tr>
</tbody>
</table>

*See Table III

**Initiation Processes.** For an adiabatic thermal explosion, the delay $\tau$ is given by (See Vol 4, p D620 and this Vol, under Hot Spots, p H170ff)

$$\tau = \frac{CRT^2}{QZE} \exp \left( \frac{E}{RT} \right)$$  \hspace{1cm} (5)

where Q is the heat of explosion and c is the specific heat. The controlling variable in this equation is the activation energy E obtained from kinetic studies. This is shown in Table II

---

**Table II**

<table>
<thead>
<tr>
<th>Arhenius Parameters*</th>
<th>$T$ (°K)</th>
<th>$\tau$ (sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Robertson</td>
<td>2550</td>
<td>$8.8 \times 10^{-18}$</td>
</tr>
<tr>
<td>Andrew &amp; Kaydimov</td>
<td>2550</td>
<td>$4.7 \times 10^{-14}$</td>
</tr>
<tr>
<td>Roth</td>
<td>2550</td>
<td>$1.7 \times 10^{-12}$</td>
</tr>
<tr>
<td>Robertson</td>
<td>1000</td>
<td>$2.7 \times 10^{-12}$</td>
</tr>
<tr>
<td>Andrew &amp; Kaydimov</td>
<td>1000</td>
<td>$8.6 \times 10^{-10}$</td>
</tr>
<tr>
<td>Roth</td>
<td>1000</td>
<td>$5 \times 10^{-9}$</td>
</tr>
<tr>
<td>Robertson</td>
<td>750</td>
<td>$4 \times 10^{-9}$</td>
</tr>
<tr>
<td>Andrew &amp; Kaydimov</td>
<td>750</td>
<td>$3.3 \times 10^{-7}$</td>
</tr>
<tr>
<td>Roth</td>
<td>750</td>
<td>$5.1 \times 10^{-7}$</td>
</tr>
<tr>
<td>Robertson</td>
<td>600</td>
<td>$7.8 \times 10^{-6}$</td>
</tr>
<tr>
<td>Andrew &amp; Kaydimov</td>
<td>600</td>
<td>$1.5 \times 10^{-4}$</td>
</tr>
<tr>
<td>Roth</td>
<td>600</td>
<td>$7.5 \times 10^{-5}$</td>
</tr>
<tr>
<td>Robertson</td>
<td>500</td>
<td>$1.3 \times 10^{-2}$</td>
</tr>
<tr>
<td>Andrew &amp; Kaydimov</td>
<td>500</td>
<td>$1.1 \times 10^{-1}$</td>
</tr>
<tr>
<td>Roth</td>
<td>500</td>
<td>$1.0 \times 10^{-2}$</td>
</tr>
</tbody>
</table>

*See Table III
Note that variation in Arrhenius parameters strongly influences \( t \) at high temperatures, but regardless of the source of the kinetic data, the conclusion that chemical reaction is exceedingly fast at CJ temperatures is still valid. At lower temperatures (the region where initiation processes presumably occur) considerable compensating effects are evident and the computed explosion times are much less dependent on data source than they are at high temperatures.

Campbell et al (Ref 7) have successfully used Eq 5 to interpret their observations on the shock initiation of homogeneous explosives such as NM, liq TNT & single crystal PETN.

Heterogeneous initiation phenomena, generally interpreted in terms of hot-spot mechanisms, are also strongly influenced by the kinetics of exothermic decomp of the explosive being initiated. The general relations between hot spot parameters and decomp kinetics parameters are given in Eqs 1 thru 5 of the article on Hot Spots in this Vol, pp H170ff. Of special interest is a recent interpretation of the initiation of explosives by impact presented by Afanasev & Bobolev (Ref 12), described under Impact, Initiation of Explosion by in this Vol, pp 135ff. In Ref 12, the relation between the variables controlling impact initiation and the kinetic parameters of the explosive impacted are given by their Eqs 11 thru 14. A rather different treatment of impact initiation, based on adiabatic compression of small occluded gas bubbles in explosives, and the dependence of these processes on chemical kinetics are presented by Bowden and Yoffe (Refs 5 & 6).

The large uncertainty in the quantitative interpretation of heterogeneous initiation phenomena results not only from the uncertainty of the kinetic parameters used (See section on Kinetic Data), but also from the large uncertainty of the temps of the initiation sites (hot spots).

Storage and Stability. It is obvious that practical explosives must possess long storage life. Kinetic data are most useful in evaluating the storage characteristics of an explosive, but they must be the right kind of data. Most explosives are solids under normal storage conditions. Thus, kinetic data obtained for a particular explosive in its vapor or liquid state and then used to estimate its storage life, could give very misleading results. Furthermore, most explosives decompose autocatalytically, i.e., they produce decomposition products that accelerate further decomposition. Also, impurities not removed during manufacture can greatly shorten storage life. All of these facts must be taken into account.

As an example, let us consider the storage characteristics of liquid NG at 21°C (294°K). Unfortunately, the Arrhenius parameters for NG given in the literature (Table III) vary widely. An average of the Serbinov-Roth parameters (Tables III & IV) gives an adiabatic explosion time of 32 years at 21°, while the Robertson parameters (Table III) give a time of 2500 years! Since conditions will not be adiabatic over long periods of time, the calculations show that pure NG has good storage life — certainly longer than 32 years. However, Gorbunov & Svetlov (Ref 8) have shown that induction periods at 40°, to the onset of rapid gas production, are approximately in the ratio 7:3.5:1 for pure NG, NG with 0.16% w, and NG with 0.17% nitric acid. Thus, the storage life of NG containing minor amounts of nitric acid can be expected to be at least seven-fold less than that of pure NG. Nitric acid in NG can be an impurity not totally removed during manuf, and it can also be a decomp product (3NO₂ + H₂O = 2HNO₃ + NO) The “Ideal” Explosive. From a kinetics point of view the “ideal” explosive is readily identifiable, but to find such an explosive is quite another matter. To possess good storage life and be impervious to moderate energy stimuli, the ideal explosive should have a large E, small Z, be readily purifiable, non-hygroscopic, and produce no decomp products that accelerate further decomp. It will explode only upon being subjected to large, externally applied stimuli, such as the nearby detonation of another explosive. At present, theory offers little guidance as to what type of compounds have a large E for decomp, yet produce appreciable heat once they start to decom (large Q). For a low value of Z, the activated complex theory indicates that \( A \times e^{\frac{Q}{RT}} \) (See Cursory Review of Chemical Kinetics section) should be negative. In terms of molecular structure,
this suggests that the activated complex has a very rigid molecular structure, while the un-decomposed explosive has a very "floppy" molecular structure.

**Kinetic Data.** In a previous section we made extensive use of the Arrhenius parameters $E$ and $Z$, and to a lesser extent of $Q$, the net heat of decomposition reactions. A compilation of this data is given in Table III. Unfortunately, many of these data are wildly discordant, eg, for PETN, $E$ and $\log Z$ range from...

<table>
<thead>
<tr>
<th>Explosive</th>
<th>Temp Range (°C)</th>
<th>$Q$ (cal/g)</th>
<th>$E$ (kcal/mole)</th>
<th>log $Z$ (sec$^{-1}$)</th>
<th>Experimental Method</th>
<th>Refs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amm Nitrate</td>
<td>217–267</td>
<td>–</td>
<td>38.3</td>
<td>12.3</td>
<td>Weight loss</td>
<td>Cook &amp; Abegg (7)</td>
</tr>
<tr>
<td>Amm Nitrate</td>
<td>243–361</td>
<td>–</td>
<td>40.5</td>
<td>13.8</td>
<td>Pressure</td>
<td>Robertson (2)</td>
</tr>
<tr>
<td>Diaminotinitro-</td>
<td></td>
<td>–</td>
<td>46.3</td>
<td>15.1</td>
<td>DSC (a)</td>
<td>Rogers (18)</td>
</tr>
<tr>
<td>benzene (DATB)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DINA</td>
<td>150–170</td>
<td>680–720</td>
<td>45.0</td>
<td>18.6</td>
<td>Weight loss;</td>
<td>Dubovitsky et al (10)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Pressure &amp; Heat Evol</td>
<td></td>
</tr>
<tr>
<td>DINA</td>
<td>130–175</td>
<td>–</td>
<td>35.5</td>
<td>13.4</td>
<td>Adiabatic furnace</td>
<td>Gross &amp; Amster (14)</td>
</tr>
<tr>
<td>EDNA</td>
<td>184–254</td>
<td>–</td>
<td>30.5</td>
<td>12.8</td>
<td>Pressure</td>
<td>Robertson (2)</td>
</tr>
<tr>
<td>EDNA</td>
<td>144–167</td>
<td>–</td>
<td>30.8</td>
<td>11.1</td>
<td>Weight loss</td>
<td>Cook &amp; Abegg (7)</td>
</tr>
<tr>
<td>EGDN</td>
<td>95–105</td>
<td>–</td>
<td>39.0</td>
<td>15.9</td>
<td>Pressure</td>
<td>Phillips (1)</td>
</tr>
<tr>
<td>EGDN (vap)</td>
<td>140–170</td>
<td>–</td>
<td>35.7</td>
<td>14.3</td>
<td>Pressure</td>
<td>Andreev &amp; Belyaev (9)</td>
</tr>
<tr>
<td>HMX</td>
<td>270–295</td>
<td>–</td>
<td>52.7</td>
<td>19.7</td>
<td>Pressure</td>
<td>Robertson (5)</td>
</tr>
<tr>
<td>HMX</td>
<td></td>
<td>560</td>
<td>57.3</td>
<td>18.8</td>
<td>DSC (a)</td>
<td>Rogers (17)</td>
</tr>
<tr>
<td>(Ref 17)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hydrazine Nitrate</td>
<td>189–200</td>
<td>–</td>
<td>38.1</td>
<td>12.2</td>
<td>Weight loss</td>
<td>Cook &amp; Abegg (7)</td>
</tr>
<tr>
<td>Hexanitrostilbene</td>
<td>(HNS)</td>
<td>–</td>
<td>30.3</td>
<td>9.2</td>
<td>DSC (a)</td>
<td>Rogers (18)</td>
</tr>
<tr>
<td>NG</td>
<td>75–105</td>
<td>–</td>
<td>40.3</td>
<td>17.1</td>
<td>Pressure</td>
<td>Phillips (1)</td>
</tr>
<tr>
<td>NG</td>
<td>115–282</td>
<td>~35</td>
<td>~35</td>
<td>~14</td>
<td>Thermal explosion</td>
<td>Serbinov (8)</td>
</tr>
<tr>
<td>NG (vap)</td>
<td>150–160</td>
<td>–</td>
<td>36</td>
<td>15.5</td>
<td>Pressure</td>
<td>Andreev &amp; Belyaev (9)</td>
</tr>
<tr>
<td>PETN</td>
<td>160–225</td>
<td>–</td>
<td>47.0</td>
<td>19.8</td>
<td>Pressure</td>
<td>Robertson (2)</td>
</tr>
<tr>
<td>PETN</td>
<td>145–171</td>
<td>–</td>
<td>39.0</td>
<td>15.6</td>
<td>Pressure</td>
<td>Andreev &amp; Kaydimov (13)</td>
</tr>
<tr>
<td>PETN</td>
<td>177–222</td>
<td>–</td>
<td>46.5</td>
<td>19.4</td>
<td>DSC (a)</td>
<td>Rogers (18)</td>
</tr>
<tr>
<td>PETN</td>
<td>149–201</td>
<td>–</td>
<td>31.8</td>
<td>13.4</td>
<td>Chemical analysis</td>
<td>Roth (6)</td>
</tr>
<tr>
<td>RDX</td>
<td>213–299</td>
<td>–</td>
<td>47.5</td>
<td>18.5</td>
<td>Pressure</td>
<td>Robertson (4)</td>
</tr>
<tr>
<td>RDX</td>
<td>?</td>
<td>43.1</td>
<td>16.4</td>
<td>DSC (a)</td>
<td>Rogers (18)</td>
<td></td>
</tr>
<tr>
<td>RDX</td>
<td>?</td>
<td>615</td>
<td>45.2</td>
<td>–</td>
<td>DSC (a)</td>
<td>Hall (16)</td>
</tr>
<tr>
<td>Tetryl</td>
<td>211–260</td>
<td>–</td>
<td>38.4</td>
<td>15.4</td>
<td>Pressure</td>
<td>Rideal &amp; Robertson (3)</td>
</tr>
<tr>
<td>Tetryl</td>
<td>140–165</td>
<td>–</td>
<td>40.0</td>
<td>16.0</td>
<td>Pressure</td>
<td>Dubovitsky et al (12)</td>
</tr>
<tr>
<td>Tetryl</td>
<td>145–155</td>
<td>340</td>
<td>36.0</td>
<td>15.6</td>
<td>Heat Evoln</td>
<td>Dubovitsky et al (11)</td>
</tr>
<tr>
<td>Tetryl</td>
<td>?</td>
<td>330</td>
<td>–</td>
<td>–</td>
<td>DSC (a)</td>
<td>Hall (16)</td>
</tr>
<tr>
<td>Tetryl</td>
<td>132–164</td>
<td>–</td>
<td>34.9</td>
<td>12.9</td>
<td>Weight loss</td>
<td>Cook &amp; Abegg (7)</td>
</tr>
<tr>
<td>TNT</td>
<td>257–310</td>
<td>–</td>
<td>34.4</td>
<td>11.4</td>
<td>Pressure</td>
<td>Robertson (4)</td>
</tr>
<tr>
<td>TNT</td>
<td>237–277</td>
<td>–</td>
<td>43.4</td>
<td>12.2</td>
<td>Weight loss</td>
<td>Cook &amp; Abegg (7)</td>
</tr>
<tr>
<td>TNT</td>
<td>288 (b)</td>
<td>–</td>
<td>41.1</td>
<td>13.2</td>
<td>Thermal explosion</td>
<td>Zinn &amp; Rogers (15)</td>
</tr>
<tr>
<td>TNT</td>
<td>220–260</td>
<td>300 (c)</td>
<td>37.0</td>
<td>11.2</td>
<td>Adiabatic furnace</td>
<td>Gross &amp; Amster (14)</td>
</tr>
</tbody>
</table>

a) Differential Scanning Calorimeter
b) Min temp at which explosions were observed in 1-mm thick samples
c) Assumed
a high of 47.0 kcal/mole and $10^{19.8}$ sec$^{-1}$ to a low of 31.8 kcal/mole and $10^{13.4}$ sec$^{-1}$, respectively. Moreover, the temp ranges over which the kinetic parameters were measured are relatively low. Thus, there is no assurance that the parameters hold at considerably higher temps, eg, detonation temps.

Some insight into what are the “best” parameters, and how valid they are at higher temps, may be gained by examining the results of the so-called Wenograd test. In this test liq or molten expl is loaded into a hypodermic needle which is then heated rapidly to a controlled temp by condenser discharge, and the explosion time of the test explosive at that temp is measured. Roth (Ref 9) obtained “Wenograd” kinetic parameters for NG, EGDN, TNT and Petrin. He analyzed the results in terms of Eq (5) and the following Eq based on Frank-Kamenetskii (Ref 1) and Chambre (Ref 4)

$$
\frac{E}{T_{\text{min}}} = 4.57 \log \left( \frac{\rho Q - \varepsilon}{\delta \lambda R} \frac{r^2}{T_{\text{min}}^2} \right) \quad (6)
$$

where: $\delta$ is a geometric factor = 2 for cylinders
$\lambda$ is the heat conductivity of the expl
$r$ is the radius of the expl cylinder
$T_{\text{min}}$ is the lowest temp at which explosions are recorded.

The results in Table IV show agreement with the lower values of $E$ and $Z$ of Table III, except for TNT, for which “Wenograd” data are not reliable.

**Written by J. ROTH**

**Refs:**
1) L. Phillips, Nature 160, 753 (1947)  
2) A.J.B. Robertson, JSocChemInd(London) 61, 221 (1948)  
4) A.J.B. Robertson, TransFaradSoc 44, 977 (1948)  
5) Ibid, 45, 85 (1949)  
6) J. Roth, Addendum to Bulletin Sixth Army-Navy Solid Propellant Group Meeting (1950), p 41  
7) M.A. Cook & M.T. Abegg, IEC 48, 1090 (1956)  
8) A.I. Serbinov, ZhFizKh 33, 2641 (1959)  
10) F.I. Dubovitsky, TransIzvAkadNauk(Chem) No 6, 1126 (1960)  
11) Ibid, 1763 (1960)  
14) D. Gross & A.B. Amster, 8th SympCombustn (1962), 729  
16) P.G. Hall, TransFaradSoc 67 (2), 556 (1971)  
17) R.N. Rogers, Thermochimica Acta 187, 1 (1972)  
18) R.N. Rogers, Private Communication (1974)

**Table IV**

**Arrhenius Parameters Based on the Wenograd Test**

(0.018 cm inner diameter tubes)

<table>
<thead>
<tr>
<th>Explosive</th>
<th>Temp Range $\Delta$K</th>
<th>E (kcal/mole)</th>
<th>Z (sec$^{-1}$)</th>
<th>Min Tube Temp $^\circ$K</th>
<th>Min HE Temp $^\circ$K</th>
<th>Z** (sec$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NG</td>
<td>555–650</td>
<td>35</td>
<td>$2.3 \times 10^{14}$</td>
<td>555</td>
<td>515</td>
<td>$\sim 2 \times 10^{15}$</td>
</tr>
<tr>
<td>EGDN</td>
<td>570–645</td>
<td>35</td>
<td>$2.8 \times 10^{14}$</td>
<td>570</td>
<td>530</td>
<td>$\sim 9 \times 10^{14}$</td>
</tr>
<tr>
<td>TNT</td>
<td></td>
<td>--</td>
<td>--</td>
<td>700</td>
<td>650</td>
<td>$\sim 4 \times 10^{14}$ (a)</td>
</tr>
<tr>
<td>Petrin</td>
<td>550–630</td>
<td>28</td>
<td>$12 \times 10^{11}$</td>
<td>540</td>
<td>510</td>
<td>$\sim 4 \times 10^{12}$</td>
</tr>
<tr>
<td>(PE-trinitrate)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*From slope of explosion time vs $1/0.93T_{\text{tube}}$ data and Eq (5), with $Q \equiv 500$ cal/g for NG, EGDN & Petrin and $Q \equiv 300$ cal/g for TNT.

**From Eq (6) using min HE Temp and “direct” $E$.

(a) Assumed $E = 41.1$ kcal/mole
Kirsanov Explosive. A Sprengel-type expl prepd by mixing just before use, a mixt of K chlorate 80 and Mn dioxide 20%, with a mixt of turpentine 90 and phenol 10%. The solid oxidizing mixt acts as an absorbent for the liq fuel

Refs: 1) Davis (1943), 355 2) A. Pérez Ara (1945), 231

KI-Starch Test. See Abel’s Test in Vol 1, p A2-L

Kittitas. Conditioning agents commonly used for reducing the tendency of AN to cake are: Kaolin or other forms of clay, and various types of Kieselguhr (qv), such as Kittitas, Celite & Dicalite

Refs: 1) Anon, IEC 36, 1088 (1944)
2) Not found in CA

Kiwit. A Ger chlorate expl introduced during WWI. It contains not more than 77% Na or K chlorate, carbon carriers such as paraffin, naphthalene, vaseline, meal or oil, and not more than 15% DNT. It may contain DNT, Dinitronaphthalene, NaCl and not more than 4% Guncotton

Ref: Marshall, Dict (1920), 54

Kjeldahl, J. (1849–1900). Danish analytical chemist, inventor of the method for determining nitrogen in organic compounds, which is still used extensively (See below)

Refs: 1) Hackh’s (1944), 470 2) S. Veibelm, JChemEduc 25, 459 (1949) & CA 43, 776 (1949)

Kjeldahl Method. A method of detg the N in an organic compd by digesting the substance with concd sulfuric acid in the presence of catalysts such as selenium, Devarda’s alloy, etc. This treatment transforms the N into Amm sulfate. By adding an excess of caustic and distg the liberated ammonia into a measured quantity of standard sulfuric acid (which is later titrated), it is possible to determine the amount of N in the substance

In using Devarda’s alloy (Cu/Al/Zn–50/45/5) for N content analysis, it is advisable not to
heat the flask until the evolution of hydrogen and other gases has ceased. This minimizes the risk of initiating a hydrogen-air expln. As an added precaution, the entire app should be placed behind a safety shield (Ref 6)


Klaaffke's Explosive. A blasting powder composed of K nitrate 73.9, C 13.4 & cellulose 12.7% Ref: A. Klaaffke, FrP 396496 (1908) & CA 4, 2733 (1910)

Klepsydra. See Clepsydra under Chronographs in Vol 3, p C308-L

KMA Block. A Ger substitute expl. See under Ersatzsprengstoffe in Vol 5, p E122, Table E15

KMP (Powder). See under Plastomenite

Knallgelert. A Russ pre-WWI expl used for rock blasting contained NG 88.6, pyroxylin 6.7 & DNT 4.7% Ref: Anon, SS 12, 409 (1917)

Knallgas (Ger). An expl mixt of H₂ and O₂, or air. The effect of initial mixt density on the initiation of detonation in knallgas and the detonation properties of “heavy” knallgas (2D₂ + O₂) have been studied

Initial mixt composition ranges of knallgas and w vapor in a detonation tube were varied by controlling temps at 100, 200 & 300°, for knallgas densities of 0.64–1.8, 8.5–16 & 46–62g/liter, respectively. Ignition was by hot wire. The threshold composition decreased as initial mixt density increased. Threshold compositions were 62, 44 & ~36% knallgas at 100, 200 & 300°, or initial mixt densities of 1.22, 13 & 63g/liter, respectively. The % knallgas reacted at final press and temp was a function of initial concn and rose abruptly at the threshold composition. The partial molal density of knallgas, which is proportional to the potential heat release per unit, increased with temp (Ref 1)

Calculated detonation properties of knallgas and heavy knallgas were compared from 1 to 15 atm using an 18ft, 1" diameter detonation tube. The detonation velocity was determined by using ionization probes to measure the time for a wave to traverse 8.984ft. Pressures were measured with an oscillogram instrumented transducer. Knallgas mixts were prepared by premixing H₂ or D₂ and O₂ and storing underground for 2–5 days. Theoretical calculations were based on Chapman-Jouguet detonation theory with assumption of equilibrium. Stable detonation velocities in heavy knallgas at 25° agreed with those predicted by Chapman-Jouguet detonation theory from 1 to 15 atm (Ref 2)


Knallquecksilber (Ger). See under Mercuric Fulminate or Mercury Fulminate in Vol 6, pp F217-L to F223-L

Knallsilber (Ger). See under Silver Fulminate in Vol 6, pp F223-R to F224-R

Knallzündschnur (Ger & Swiss). Detonating Fuse or Primacord. See under Detonating Cords or Detonating Fuses in Vol 3, pp D103-R to D107-L

Knecht Compound. Knecht (Refs 1 & 2) treated cellulose with 65% nitric acid (d 1.14 g/cc) and obtained an addition product. After removing the acid retained mechanically he established that the addition compd contained nitric acid corresponding to 7.7% N. This is approximately equivalent to one NO₂ group for
every anhydroglucose unit. Drowning the addition product in water caused nitric acid to split off, and Knecht thought that it was an exceptionally labile nitrate. Häusserman (Ref 3) found it to be a rather unstable addition product. Champetier & Marton (Ref 8) examined the ultra-violet reflection spectrum, and found that it contains a band at 270–305µ which is not present in the spectrum of NC. They considered that this supports the view that the Knecht compound is an individual substance differing from NC.

Cellulose regenerated from the addition compound may demonstrate a certain degree of nitration (0.5–2.2% N). The properties of the compound are similar to those of hydrocellulose, e.g., glittering fibers, increased hygroscopicity and higher reactivity.

X-ray investigations by Hess & Katz (Ref 4) suggest that the addition product is probably compatible with the formula C₆H₁₀O₅.HNO₃.

The X-ray diagram of the compound is a characteristic one differing from that of cellulose. They concluded that it was produced by the action of 86% nitric acid.

According to the expts of Andress (Ref 5) the composition of the Knecht compound after being kept under reduced press for some time should be denoted by the formula 2C₆H₁₀O₅.HNO₃. He demonstrated that the compound produces a characteristic X-ray fiber diagram.

Later, Wilson (Ref 7) established that its composition after drying under a high vacuum is 2C₆H₁₀O₅.HNO₃.H₂O. The product of absorption of nitric acid vapors by cellulose is, according to Wilson, not a Knecht compound, since it did not give the characteristic X-ray diagram though its chemical composition is approx that of the Knecht compound.

Troegus (Ref 6) collected data concerning the concn of nitric acid to produce the Knecht compound, and concluded that it is formed when a mixt of almost equivalent quantities of the hydrates HNO₃.H₂O and HNO₃.3H₂O act on cellulose.


Knetmaschine (Ger for Kneading Machine).

An apparatus for mixing solid ingredients in the presence of liquids. Several types were used in Ger for mixing propulstn & exprts, such as the Columnar type (Saulenknetmaschine) (Ref 2, pp 105–6; Ref 3, p 237); Werner-Pfleiderer Misch- und Knetmaschine (Ref 1, p 75; Ref 3, p 227; Ref 4, pp 585–87), and others. In France, Chaudel-Page kneaders are in general use (Ref 4, pp 588–9).


Kochsalzprengstoffe (Kitchen Salt Explosives). Ger substitute expl mixts containing a large amount of Na chloride, used during WWII. See under Ersatzprengstoffe (Substitute Explosives) in Vol 5, pp E121-L to E122-R

Koffler Micro Hot State Apparatus. Electrically heated apparatus which might be considered a modification and improvement of the Fisher-Johns apparatus. It permits microscopic studies of fusion, sublimation, crystallization and explosion phenomena.


Kohlen-Albit (Ger). One of the Kohlensprengstoffe (coal mining explosives). Its compn was similar to Gesteins-Koronit. It was used as a
permissible expl in coal mines for a short time, but because of its large flame on expln was replaced by AN expls

Refs: 1) Naoum, NG (1928), p 428
2) Clift & Fedoroff 2 (1943), p 401

Kohlenprengstoffe (Coal Explosives). A group of Ger expls permitted for use in coal mines:

Kohlen-Carbonit. NG 25, K nitrate 34, Ba nitrate 1, flour 38.5, spent tan meal 1 & soda ash 0.5%. Qe 506 kcal/kg, Temp of Expl 1561°, Vel of Deton 3160m/sec, d 1.16g/cc, Trauzl test value 235cc, Charge limit 900g, equivalent to 501g of Brit standard Gellignite


Kohlen-Koronit III. NG 4, K chlorate 68, Na chloride 14, paraffin 8, MNN 5 & wood meal 1%. OB -2.6%, Trauzl test value, 195cc

Refs: 1) Naoum, Expl (1927), 147 2) PATR 2510 (1958), Ger 101-R

Kohlen-Salit. NG (gelatinized with NC) 12.5, vegetable meal 2.5, aromatic nitrocompounds 7.0, AN 41.0 & alkali chlorides 37.0%. OB -2.6%, Trauzl test value 260cc

Refs: 1) Naoum, NG (1928), 411 2) Fedoroff & Clift 4 (1946), 49 3) PATR 2510 (1958), Ger 101-R

Kohlen-Silesia 4A. K chlorate 80, resin 16 & Nitroresin 4%. Silesia No 4 contd K chlorate 80 & resin 20%

Refs: 1) Barnett (1919), 111 2) Fedoroff & Clift 4 (1946), 50

Kohlen-Westfalt I. NG 4.0, AN 83.0, K nitrate 7.0, Ba nitrate 2.0, meal 2.0 & TNT 2.0%. OB +16.4%, Trauzl test value 230cc

Refs: 1) Naoum, NG (1928), 435 2) PATR 2510 (1958), p Ger 102-L

Kohlen-Westfalt IV. NG 3.2, AN 73.0, K nitrate 2.8, alkali chloride 15.0, meal 1.0 & DNT 5.0%. OB +8.8%, Trauzl test value 200cc

Refs: 1) Naoum, NG (1928), 435 2) PATR 2510 (1958), p Ger 102-L

Kohler Powder. K chlorate 70, sulfur 20.0 & charcoal 10%. Very sensitive and dangerous mixt, similar to Berthelot's Powder (See Vol 2, p B106-L

Ref: Perez Ara (1945), p 206

Kolax. A Ger expl of the Carbonite type cntg NG 25, K nitrate 26, Ba nitrate 5, woodmeal 34 & starch 10%. Super-Kolax was a modification of the above used in England: NG 28.5, Collodion Cotton 1.0, K nitrate 16.5, Ba nitrate 5.0, Amm oxalate 9.5, woodmeal 30.5 & starch 9.0%. Both are no longer on the permitted list


Kolfit (Ger). An expl mixt patented by H. Kolf in 1890 for use as a smokeless propi, but also suitable as a blasting expl. It was prep'd by nitration of residues from flour mills, starch and sugar plants, distilleries, breweries, etc. Oil cakes, moss, and pine needles were also nitrated. The resulting nitrated substances were treated under 5 atm press with sulfides or polysulfides, and then mixed with saltpeter previously saturated with Nitrobenzene


Kolf's Blasting Powder. Ger mining expl patented in 1892. A mixt of equal parts of NC, Nitroglycerin & Nitrostarch 50, Nitrosugar 38, NG 8, saltpeter 2 & aniline 2%

Refs: 1) Daniel (1902), 394 2) Fedoroff & Clift 4 (1946), p 50

Kolf's Powder. A propi (greenish-brown leaflets) used at the end of the 19th century in England. It consisted of equal parts of NC, Nitrochris & Nitrolignin, gelatinized by a volatile solvent, and mixed with about 0.5% sulfur and 2% K nitrate

Ref: Daniel (1902), p 394
Kollodiumwolle. Ger & Swiss for Collodion Cotton

Kölner Dynamit Fabrik. A method, patented in 1885, of impregnating Dynamite cartridges with such materials as Al oleate in order to make them more flexible
Ref: Daniel (1902), 395

Kölner-Rottweiler Sicherheits-Sprengpulver.
One of the older Ger safety expls consisting of: Amm Nitratre 93.0, vegetable oil 4.9, sulfur 1.2 & Ba nitrate 0.9%.
Ref: Daniel (1902), p 395

Kolowratnik's Explosives. Chlorate expls contg PA 40, K chlorate 10–20, Na nitrate 25–35, Na carbonate 5 & woodflour 10%
Refs: 1) R. Kolowratnik, BritP 26884 (1912) & CA 8, 1672 (1914) 2) Colver (1918), 325

König Explosives. J.B. König patented in 1890 a method of prepn of expls by nitration of high bp hydrocarbons derived from the distn of coal, bituminous shales or residues of petroleum refineries, paraffins and ozokerite
Ref: Daniel (1902), 395

Konovaloff Reaction. A color test for primary or secondary nitro-compd. Samples are treated with Na hydroxide soln, the salt formed is extrd with w, and ether is added to this extr. Upon dropwise addition of ferric chloride, a red to reddish-brown color develops in the ether

Kontinuierliche Verfahren. Continuous methods, such as those of Schmid, Meissner & Biauzzi, for the prepn and purification of expls and proplnts, have been used extensively in Ger. Some of these methods will be described under Nitratron in Vol 8 of Encycl


Koronit. One of the Ger expls of the chlorate class, developed during WWI in order to conserve NG. It contains K nitrate and/or Na nitrate 70–80, Nitroderivatives of toluene and/or diphenylamine and/or naphthalene 12–20, vegetable meal 1–5, solid hydrocarbons and/or oils and fats 3–5 & NG (not gelatinized) 2–6%
Refs: 1) Marshall 3 (1932), p 112 2) Davis (1943), 361

Koronit or Favorit is, according to Marshall (Ref 1), a Ger chlorate blasting expl. Its nonpermissible modification, Gesteins-Koronit T1, (See Vol 6, p G73-L) contains Na chlorate 72, DNT & TNT 20, NG 3–4, vegetable meal 1–2 & paraffin 3–4% (Ref 2). Bebie (Ref 3) and Warren et al (Ref 4) list this expl under the name Coronit (See Vol 3, p C543-R). The permissible modification, Kohlen-Koronit (qy), contains according to Ref 1, not more than 68 parts of K or Na chlorate, not more than 12 parts of aromatic hydrocarbons or nitrocompds (but no trinitrocompds), not more than 4 parts Blasting Gelatin, not more than 4 parts of powdered coal, with the remainder consisting of Na chloride, paraffin, etc

These expls were eventually replaced by Perkornits, which are similar to Koronits except that they contain perchlorates rather than chlorates

Koronit V. A Ger permissible Dynamite contg NG 4, K chlorate 65, Na chloride 14, naphthalene 10, MNN 5 & woodmeal 2% (Ref 1). According to Marshall (Ref 2), the name Koronit was given in 1931 to Chloratit 1 (See Vol 2, p C209-L), which contains K and/or
Na chlorate 70–80, Nitroderivatives of toluene and/or naphthalene and/or diphenylamine 12–
20, vegetable meal 1–5, solid hydrocarbons and/or oils and fats 3–5 & NG (not gelatinized)
2–6%
Refs: 1) Naoum, Expls (1927), 147
2) Marshall 3 (1932), 112
3) PATR 2510 (1958), p Ger 102-R

Kostevich, Michael M. (1877–1957). A former
Colonel of the Imperial Russian Guard Artillery
and head of the Russian chemical warfare
section during WWI. After the revolution he
worked in Engl, Fr and Belg. He built and
directed a chemical laboratory for the Czech
Skoda works. From 1930–1936 he taught
chemistry at the Technical Institute in Paris.
From 1937 he was chief of the research
department in the explosives plant at Villa
Maria, Argentina. Author of numerous publica-
tions on ordnance and expls including books
entitled, “Trinitrobenzol and Trinitrotoluol”
and “Methods for Removing Explosives from
Artillery Shells by Burning”
Ref: A.A. Schrimpff, SS 32, 319 (1937)

Kraft Dynamite. Patented by C.G. Bjorkmann
at the end of the 19th century. It contained
NG 55.36, K chlorate 16.96, K nitrate 15.18
& pulverized cork 12.50%
Ref: Daniel (1902), 396

Kraft Paper. A strong, relatively insensitive
paper made primarily from pine by digestion
with a mixt of caustic soda, Na carbonate,
Na sulfate & Na sulfide (Ref 3). It is used
extensively in Dynamite cartridges (Ref 2)
and in pyrotechnic flare and signal cases (Ref 1)
Refs: 1) S.M. Kaye et al, “New Flare Case
Materials”, PATR 3215 (1965) 2) Blasters’
Hdb (1969), 30–33
3) CondChemDict (1971), 499-L

Kraftzahl (KZ or Strength Number). It has
been recognized for some time that enlarge-
ments in the lead block cavity, obtained in
the Trauzl test (See Vol 1, pp XXV–XXVI),
are not exactly proportional to the power of
the expls. For this reason, the Trauzl test
has been the subject of numerous inves-
tigations, principally by Kast, Selle, Neubner
(Refs 1, 2 & 3), and at the US BurMines
(Ref 5)

In the course of these investigations, Neubner
(Ref 3) found that better results are obtained
if, instead of firing a standard 10g charge, the
weights are varied until expansions between
250 and 325cc are obtained. The weight
(w) corresponding to a 300cc expansion is
calculated by interpolation. By dividing 300
by w and multiplying by 10, a value is obtained
which is more representative than the old
Trauzl value. This new value is called the
Kraftzahl or KZ of the expl. The following
table compares these values with older Trauzl
test values determined by various investigators
(Kast, Stettbacher, Zschokke, Will & Brunswig):

<table>
<thead>
<tr>
<th>Explosive</th>
<th>Old Value</th>
<th>KZ Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitromannite</td>
<td>650</td>
<td>675</td>
</tr>
<tr>
<td>Blasting Gelatin</td>
<td>520–610</td>
<td>554–575</td>
</tr>
<tr>
<td>NG</td>
<td>515–563</td>
<td>540–545</td>
</tr>
<tr>
<td>Nitropentaerythrite</td>
<td>460</td>
<td>500</td>
</tr>
<tr>
<td>Gelatin Dynamite (60%)</td>
<td>410</td>
<td>465–472</td>
</tr>
<tr>
<td>Tetranitoanisol</td>
<td>390</td>
<td>440</td>
</tr>
<tr>
<td>Guhr Dynamite</td>
<td>350</td>
<td>419</td>
</tr>
<tr>
<td>NC (13% N)</td>
<td>325–420</td>
<td>400–426</td>
</tr>
<tr>
<td>Tetryl</td>
<td>340–350</td>
<td>405</td>
</tr>
<tr>
<td>TNB</td>
<td>330</td>
<td>386</td>
</tr>
<tr>
<td>Hexanitrodiphenylsulfide</td>
<td>325</td>
<td>380</td>
</tr>
<tr>
<td>Hexanitrodiphenylamine</td>
<td>320</td>
<td>376</td>
</tr>
<tr>
<td>Picric Acid</td>
<td>300–365</td>
<td>376–385</td>
</tr>
<tr>
<td>TNT</td>
<td>285–300</td>
<td>350–353</td>
</tr>
<tr>
<td>Trinitroresol</td>
<td>275</td>
<td>336</td>
</tr>
<tr>
<td>Dinitrobenzene</td>
<td>250</td>
<td>311</td>
</tr>
<tr>
<td>Single-base Proplnt</td>
<td>150</td>
<td>226</td>
</tr>
<tr>
<td>(13% NC)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Double-base Proplnt</td>
<td>150</td>
<td>226</td>
</tr>
<tr>
<td>(40% NG)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lead Azide</td>
<td>115</td>
<td>165</td>
</tr>
<tr>
<td>Mercury Fulminate</td>
<td>110–150</td>
<td>226</td>
</tr>
<tr>
<td>Black Powder</td>
<td>30–112</td>
<td>95</td>
</tr>
</tbody>
</table>

Refs: 1) H. Kast, SS 15, 171 (1920)
2) H. Kast & H. Selle, Glickauf, 1927, 900
3) R. Neubner, SS 15, 3, 53–4, 82, 126,
162, 194 (1928) 4) Marshall 3 (1932),
143 5) N.A. Tolch & G.St.J. Ferrott,
“The Trauzl Block Strength Test of Dynamites”, US BurMines RT 3099 (1930)
6) Stettbacher (1948), 113
Kratites. Expls containing Amm Perchlorate, NG and NC in various proportions, based on requirements for strength, brittleness, sensitivity, etc
Refs: 1) Daniel (1902), 396 2) Pérez Ara (1945), 219

Kreulen’s Aluminum Block. See Aluminum Block (of Kreulen) in Vol 1, p A145-L

K-Salz (Ger). See under Cyclonite in Vol 3, p C611-L

KSG Powder. An older Brit 33-grain sporting smokeless powder which is no longer used
Ref: Marshall 1 (1917), 327

KS Powder. An older Brit 42-grain sporting smokeless powder which is no longer used
Ref: Marshall 1 (1917), 326

K₃-Stoff (Ger). A highly dispersed silica, prep'd by a special process, used during WW11 in some “Tetan” expls
Ref: G. Römer, “Reports on Explosives (Germany)”, PBL Rept No 85160 (1946), p 3

Ksalt (Russ, Xylite in Fr, Ksalit in Polish, TNX). A term given to commercial Trinitroxylen (TNX) which consists chiefly of the trinitro derivatives of meta-, with some ortho- and para-xylenes. It is a greyish-yellow substance prep'd by the nitration of commercial xylene by mixed nitric-sulfuric acid. It is insol in w. Ksalt is slightly more sensitive to initiation, and about as sensitive to impact and bullet test as TNT. It was used in composite expls as a partial substitute for TNT
Refs: 1) PATR 2145 (1955), p Rus 10-L 2) Gorst (1957), 100 3) Urbanski 1 (1964), 396

Kubin Explosives. A safety expl, patented in 1893, containing AN 75-95 and Nitraniline 25-5%. It could also contain up to 20% of added Amm oxalate, chloride or sulfate. Other nitrated amines, such as those of toluene, xylene or naphthalene, could be used in lieu of aniline
Ref: Daniel (1902), 397

Kubin & Sierch. Patented in England in 1884 a safety expl consisting of a regular Dynamite to which was added 20-50% of Amm chloride or sulfate. This expl can be considered as an early Wetterdynamite
Ref: Daniel (1902), 398

Kumulatwnyi Zariad (Shaped Charge). Mixts containing Gekosgen (RDX) or TEN (PETN) with TNT or other expls or phlegmatizers were used in Russ shaped charges
Ref: Gorst (1957), 95, 98 & 100

Küp Powder. A mixt containing Ba nitrate 80, sulfur 20%, plus charcoal
Ref: Daniel (1902), 398

Kurzschlusszünder. Ger short-circuit primers or igniters, such as the Schaffler, Reinecke systems as well as the Endrahtzünder (single wire igniters) are described in the Ref
Ref: Beyling-Drekovf (1936), pp 216-22

Kynarkite. An early permitted expl containing NG 25, woodmeal 35, Ba nitrate 3, K nitrate 28, Amm oxalate 5 & w 3.7%. BalPend value for 4oz sample, 2.21” swing compared to 3.27” for Brit 60% Gelignite. Charge limit in gallery test 20oz
Refs: 1) Barnett (1919), 135 2) Marshall, Dict (1920), 55-6

Kynite. A Brit safety expl invented by A.T. Cocking of Kynoch, Ltd: NG 25-27, Ba & K nitrates 30-36, woodmeal & starch 40-43 & Na carbonate (added) 0.5%. To this could be added 0.5 to 10% Amm oxalate
Refs: 1) Daniel (1902), 398 2) Marshall, Dict (1920), 56
Kynite, Condensed. Brit safety expl manufd by Kynoch, Ltd and packaged in a non-waterproof parchment paper wrapper: NG 24–26, starch 32.5–35.0, woodmeal 2.5–3.5, Ba nitrate 31.5–34.5 & Ca carbonate 0 to 0.5% 

Ref: Daniel (1902), 340

Kynoch Limited. A Brit expls company with plants at Kynochtown (Essex) and Ferrybank Arklow (Ireland). This company prospered greatly during the South African War of 1901, when holders of its preferred stock received dividends equal to 7 times the price of the stock. At that time, the company manufd Dynamite, Gelignite, NC, Kynite, Cordite & Smokeless Pdr, as well as various munitions Ref: Daniel (1902), 309

Kynoch Smokeless Powder. One of the older Brit 30-grain sporting powders. It contains NC 52.1, DNT 19.5, K nitrate 1.4, Ba nitrate 22.2, woodmeal 2.7, ash 0.9 & volatiles 1.2%. Qc 807cal/g; gas vol 726 l/kg at STP

KZ. See Kraftzahl in this Vol
L-Alloy. A Rus fuseable expl consisting of 95% TNT and 5% Trinitroxylenol, mp 74°. It has an
explosion power similar to that of TNT, although it differs from the latter in detonating more readily
due to the fact that during the cooling process
very fine TNT crystals are formed
Refs: 1) Ya.I. Leitman (according to A.G.
Gorst), “Porokha i Vzryvchatyiye Veshchestva”,
Oboronizd, Moskva (1949) 2) Urbanski 3
(1967), 248–49

Labels for Shipping Explosives and Other Dang-
erous Materials. The US Interstate Commerce
Commission Regulations specify that the fol-
lowing labels be provided when expls or other
dangerous materials are shipped or transported:
Flammable liquids: diamond-shaped label,
bright red in color, with each side measuring
4". Printing must be in black letters inside of a
black line border, measuring 3.5" on each side
Flammable solids and oxidizing materials:
same shape, size and lettering as above, but
the color shall be bright yellow
Corrosive liquids, alkalies or acids: same shape,
size and lettering as above, but the label shall be
white in color
Compressed gas: same shape, size and lettering
as above, but the color of the label shall be
red for flammable gases and bright green for
nonflammable gases
Poison (gas, liquid or solid) and tear gases:
same shape and size of label as above, but with
a white label with letters in red
Explosives, propellant powders and special
fireworks: rectangular white labels with red
letters reading Explosives A or Explosives B
Radioactive materials (Groups I or II): diamond-
shaped label, white in color, with each side
measuring 4". Printing is in red letters inside
a red-line border. Group I materials are those
which emit any gamma radiation, either alone
or with electrically charged particles or cor-
puscles (alpha, beta, etc). Group II materials
are those which emit neutrons and either or
both of the types of radiation characteristics
of Group I materials
Radioactive materials (Group III): same as
above except that the lettering and border line
are in blue. Group III materials are those which
emit electrically charged corpuscular rays only
(alpha, beta, etc), or any other that is so shielded
that the gamma radiation at the surface of the
package does not exceed 10 milliroentgens per
24 hours at any time during transport
For more detailed information, consult the
ICC Regulations (Ref 1) or Ref 2
Refs: 1) Agent T.C. Georges Tariff No 19,
published by Interstate Commerce Commission
Regulations for Transportation of Explosives
and other Dangerous Articles by Land and
Water in Rail Freight Service and by Motor
Vehicle (Highway) and Water, including Spe-
cifications for Shipping Containers; issued 5
Aug 1966; available from T.C. George, 63
Vesey Street, New York, NY 10007 2) Sax
(1968), 308–62

Labile Nitrocellulose (Knecht Compound).
Knecht found that when cotton is immersed
in 60% nitric acid, it forms a “labile” nitrate
from which the nitric acid can be removed by
washing with w. Using an acid with a d of
1.415, he obtained a cellulose with 35.8% abso-
rated acid, corresponding to C₆H₇NO₃HNO₃
Refs: 1) E. Knecht, Ber 37, 549 (1904)
2) Marshall 1 (1917), 151
NOTE: See Cellulose Nitrates in Vol 2 of
Encycl, pp C100-Lff

Lactic Acid Nitrates (Nitrolactic Acid),
CH₃CH(ONO₂)COOH; mw 135.09, N 10.37%;
light yellowish oil misc with w, alc, eth & benz;
insol in lignin. May be prepd from lactic acid
using Zn acetate in a mixt of fuming nitric
and concd sulfuric acids. Decomp at RT, too
unstable for use
Refs: 1) Beil 3, 279 2) A.H. Blatt & F.C.
Whitmore, “A Literature Survey of Explosives”,
OSRD 1085, 56 (1942)

LACTOSE AND DERIVATIVES

Lactose (Milk Sugar), C₁₂H₂₂O₁₁,H₂O; mw
360.31, monoclinic spheroidal crystals (from w),
mp 201–202° (becomes anhyd at 120°), bp
(dec), d 1.53g/cc at 20°. Present in the milk
of mammals, and is produced commercially
from whey, a by-product of the cheese industry.
It is hydrolyzed by dil mineral acids to glucose and galactose. Termansen (Ref 2) found that a homogeneous mix of lactose with 12.5% PETN is not expl, however, lactose contg 12.5% Hexanitromannitol has expl props. Lactose yields expls on nitration (See below).


**Lactose Hexanitrate**, $C_{12}H_{16}O_{5}(ONO_2)_6$; a white amorphous material with mp about 70°C, was isolated from the alc mother liquors in the crsyt of the octanitrate (Ref 6). It is a less powerful expl than the octa-compd.

Sjöberg, in 1888, patented expls contg mixts of nitrotactose, nitromolasses and Na nitrate, as well as admixts of the above with Anm nitrate, naphthalene and paraffin. These expls were unduly heat sensitive (Ref 2). Crater patented the use of nitrotactose with materials such as MF and K chloride, or LA or DDNP in blasting cap charges (Ref 4), as well as their utilization together with oxidizers such as Anm nitrate in blasting expls (Ref 5).


**Lactose Octanitrate** (Nitrotactose), $C_{12}H_{14}O_{3}(ONO_2)_6$; mw 702.31, N 15.96%, OB to CO$_2$ -9.1%, white monoclinic needles (from methanol), mp 145–46°C (dec), bp (ignites at 246–48°C), d 1.684g/cc; insol in w, sol in acet, methanol and AcOH; sol in cold alc, sol in hot; reduces Fehling’s soln on warming.

May be prep’d by stirrning lactose with mixed nitric-sulfuric acids until the mass forms a solid agglomerate [1p lactose in 15p acid, consisting of 1 vol nitric (d 1.5) and 2 vols concd sulfuric]. The mixt is transferred to ice w and pulverized under w. The product may be purified by crystall from alc (Ref 1).

Nitrotactose is comparable in brisance (sand crush test) to Tetryl, but has a lower vel of deton (4225m/sec as detd in 1.25" x 8" cartridges). Its sensitivity to impact is comparable to PETN, and it is fairly insensitive to friction and insensitive to static electricity. The thermal stability of Nitrotactose is satisfactory; 65.5°C KI test – 28 minutes; loss on heating for 8 days at 50°C – 0.7%; at 75°C for 24 hours – 1%; 54 hours – 23%; at 100°C for 10 hours – 20% loss (Ref 7).

**Lafaye Explosives**. Lafaye patented in Fr in 1888, a method of prep望着 NC from pure wood cellulose, the resulting product being cheaper than NC obtained from cotton

Ref: Daniel (1902), 400

**Laflin and Rand Powder Co.** One of the first expl manufg companies in the USA. Its first BlkPdr plant was erected near Orange, NJ in 1808 (Ref 1). Another plant was built at Orange Mills, near Newburgh, NY, and a plant at Pompton Lakes, NJ, manuf proptnt pdr for small arms. It consisted of Guncotton 67.25%, NG 30 and metallic salts 2.75%, gela-tinized with acetone. A dense shotgun pdr of similar compn was also made (Ref 2). The Pompton Lakes, NJ plant was purchased by duPont and expls including initiators, such as MF, LA, blasting caps, etc were manufd there


**Lake’s Explosives**. Mining expls patented in Engl in 1894. Typical formulations included:

1) Na nitrate 76%, nitronaphthalene 8% & nitrophenol 16%
2) Na nitrate 58%, nitronaphthalene 8% & nitrophenol 34%
3) Na nitrate 74%, nitrophenol 10%, sulfur 8%, charcoal 4% & rosin 4%

Ref: Daniel (1902), 401

**Lambotte Explosives**. Complicated mixts of nitroglycerine, sawdust, Pb oxide, Ba sulfate, etc

Ref: Daniel (1902), 401
Laminac 4116. A binder and adhesive most often specified for use in pyrotechnic flares and signals. It is a proprietary product manufactured by the American Cyanamid Co, and belongs to the class of thermosetting alkyl-styrene resins more commonly called unsaturated polyesters (with added styrene). Since the liquid resin is relatively low in viscosity, it can be added to solid powders without any volatile solvent. For activation, 1–2% of methyl ethyl ketone peroxyde (tradename Luperol DDM) is mixed with the resin. This “catalyst”, actually a curing agent that reacts chemically, converts the resin first into the gel state, then in strongly exothermic reaction into a hard solid. The reaction is accelerated by the promoting agent Co napthenate (tradename Nuodex). Laminac is usually admixed as 4–9% of the formula wt

Laminated Powders. Progressive burning propellants originated by the U.S. during WWI and used in large caliber guns. Of these powders was prep'd as follows: A quantity of Ballistite, prep'd by mixing 50p of NG with 50p of NC (12% N), was rolled between heated rollers until a sheet of desired thickness was obtained. Another portion, consisting of 50p of DNT and 50p of NC (12% N), was rolled to form a sheet thinner than the previous one. A strip of the first mix was sandwiched between two strips of the second mix, and the ensemble combined into a laminated product by pressing between warm rollers. In the resulting “laminated” product, the outer layers burned relatively slowly with a temp of about 1500°, while the inner slab burned rapidly with a temp of about 3000°. Similar propellants were prep'd during WWII
Ref: Davis (1943), 318

Lamm Explosives. Lamm patented, in 1888 in Fr, the use of waxes, such as carnauba, as coating agents for hygroscopic ingredients of expls. He also proposed several expl mixts contg DNB in combination with K or Amm nitrates
Ref: Daniel (1902), 401

LaMotte's Explosive. DNN 7p and TNT 5p were fused together and mixed with 88p of Amm nitrate. After the mix cooled, 2p of pyrites were added
Ref: E. LaMotte, USP 911019 (1909) & CA 3, 1088 (1909); BritP 7921 (1908) & CA 3, 2382 (1909)

Lance Missile (XMGM-52A). US surface-to-surface inmissile weighing approx 2814 lbs with a 30 mile range. It is launched from a converted M113 armored personnel carrier chassis, and is intended as a divisional support weapon. As such it will replace the Honest John and Sergeant rockets. It is powered by a storable liq fuel rocket motor and guided by a simple inertial system. It has a small nuclear warhead, but other types are also available. An extended range Lance is under development with a 50–60 mile range. The Sea Lance XMGM-52B is a marine conversion for firing from a gyroscopically stabilized launcher

Land. One of the raised ridges in the bore of a rifled gun barrel.
Note: The caliber of small-arms ammo, except for shotgun ammo, is expressed as the diameter
of the bore of the weapon in inches. Thus, caliber .30 ammo is intended for a weapon having a bore of 0.30 inch diameter across the lands and, although the outside diameter of the bullet is a few thousandths of an inch greater than the bore diameter, it is customary to speak of this ammo as caliber .30. The same is applied to the bore of the gun
Refs: 1) Ohart (1946), 64 & 98 2) Ord-TechTerm (1962), 172-R

Lanthanum Triazide. See in Vol 1 of Encycl, p A554-R

Laser, Application to Explosives and Weapons Technology. A device used for the amplification or generation of coherent light waves, the term laser being an acronym for light amplification by stimulated emission of radiation. Unlike the waves emitted by an ordinary light source, such as an electric lamp, the laser produces a light beam that does not diffuse. Moreover, a laser organizes the energy waves emitted by a stimulated atom so that they travel in the same direction, at the same frequency, and perfectly in step with the stimulating radiation. This property is known as coherence. A laser thus produces a very narrow band of frequencies similar to that of a radio oscillator, but in the infrared and visible-light portions of the spectrum (Ref 15)

The highly parallel, intense beam of coherent radiation emitted by the laser has found a multitude of important applications in the military area. These range from anti-missile applications, communications, surveillance, range finding (ground and airborne), tracking (ground, sea, air), weather reconnaissance, metal working, nondestructive measurement, defensive and offensive weaponry, to field surgery (Refs 1, 16, 27 & 29)

Picatinny Arsenal adapted the laser for remote, wireless initiation of thermal batteries and other devices contg expls, propmts and pyrotechnics in expls to demonstrate the feasibility of using a thin beam of infrared radiation in lieu of a method requiring wires. The wires, which can act as an antenna to the detriment of the device contg them, were removed and replaced by a thin transparent window to provide an optical path between the laser beam and the reactive material to be initiated. Reportedly offering a savings in the cost of thermal batteries and squibs, the method also contributes to increased safety in environments which could cause undesired ignition. The laser beam, programmed to deliver a series of pulses as required, can be directed thru the air on a line-of-sight path or thru a maze of fiber optics in any direction dictated by the geometry of the equip-

Landauer Explosives. Fr expls, patented in 1891, in which chlorates and perchlorates are desensitized by coating them with fatty materials, tar or nitrated hydrocarbons. Following are some expl mixts proposed by Landauer: 1) K chlorate 48.8, tar 24.4, DNN 24.4 & sulfur 2.4%; 2) K chlorate 20, tar 20, NC 40 & coconut oil 20%; 3) K chlorate 22.2, tar 55.6 & NG 22.2%; 4) K perchlorate 3, tar 1, DDN 10, NC6, NG 75 & sawdust 5%; 5) K perchlorate 2.0, tar 18.4, DNN 4.0, K nitrate 42.9 & Amm nitrate 32.7%
Ref: Daniel (1902), 402–403

Land Mines. See Mines (Military) in Vol 8, of Encycl

Landshof Dynamite. Dynamite contg NG 75, kieselguhr 20 & Amm urate 5%
Refs: 1) Daniel (1902), 403 2) Fedoroff & Clift 4, 51 (1946)

Landshof Powder. BlkPdr type compn contg K nitrate 73, Amm urate 9, charcoal 9 & sulfur 9
Ref: Daniel (1902), 403

Landskrona Powder. Swiss propmt, invented by E. Schenker before 1890 and manufd in Landskrona, Sweden. It was made by gela-

atinizing pure NC with amyl acetate, and was used by the Swiss and Danish armed forces
Ref: L. Gody, “Traite des Matières Explosives”, Namur (1902), 641
ment used. Upon striking the expl material, the pulse excites electronic and vibrational energy levels, causing it to ignite (Ref 3).

A laser-initiated pyrotechnic system (Zr-Amn perchlorate) for spacecraft application has been shown to be superior to conventional electroexplosion initiation (Ref 22). Primary high expls such as LA and LSt can be initiated directly. Q-switched-mode lasers can directly deton PETN, RDX, and Tetryl in a properly designed device and can be useful where microsecond simultaneity is important (Ref 6).

A "hardened" laser optic system that resists X-rays was developed by Space Ordnance Systems, Inc, to assist missiles in reaching their target. The system can withstand very high temps and the tremendous bursts of X-rays from high altitude thermonuclear explosions which may be used to destroy incoming ballistic missile warheads. The X-rays would be released by defensive missiles fired in the general vicinity of incoming weapons. Called LEED, an acronym for laser energized explosive device, the system was developed to end the danger of premature expln or failure of prophts pyrotechnics in the presence of radiofrequency, electrostatic energy, X-rays, or dense electron fields. If, while still in their high-energy state, the X-rays released from a nuclear expl impact upon a warhead, they are quickly stopped and their electromagnetic energy is converted into heat energy. The resulting surge of energy might shatter the outer casing of the warhead and create a shock wave that could damage the interior mechanisms. LEED, constructed of low "Z" materials, utilizes a laser energy pulse transmitted along nonmetallic, fiber optic conductors instead of conventional connecting metallic electrical cables, bridgewires, and spark gaps such as those used in interior warhead mechanisms (Ref 2).

The US Army (Ref 29) successfully tested a laser-guided artillery shell that could revolutionize the role of artillery on the battlefield. With the new shell, whose course can be changed in flight to bring it down on a target with high accuracy, artillery would have, for the first time, a capability of attacking moving targets, such as tanks.

The guided projectile homes in on a target in the following manner. A forward artillery observer, either on the ground or in an aircraft, trains a laser beam on a target. A laser-seeker in the nose of the artillery shell, which is fired in the general vicinity of the target, picks up the laser spot on the target. On the basis of information received from the laser-seeker, a guidance system sends out course directions to fins that deploy after the shell leaves the gun barrel. The fins then change the course of the shell so it will hit the target.

On the basis of test firings, accuracies on the order of 1–3 ft at any range can be obtained with this shell, compared with a probable error of 45–60 ft with conventional 155mm ammo. The shell can be used in the 155mm howitzer with no modifications to the artillery piece.

Another area of laser use applied to expl materials involves its employment to excite Raman spectra for studies of crystal structure, lattice dynamics, phase transitions and vibrational mode frequencies. Compds studied include TiN3 (Refs 10, 17 & 23), NaN3 (Ref 18), KN3 and RbN3 (Ref 4), NH4N3 (Ref 7), BaN3 (Refs 5, 8 & 24), LA (Ref 9), HMX (Ref 25), RDX (Ref 11) and Amm perchlorate (Ref 26).

In the course of an investigation to provide bright light sources of min volume and weight, a search was conducted for new sources of chemically generated light. Chemical systems were investigated in a flash photolysis laser apparatus in order to discover new output wave lengths and new molecules which produce lasing (Ref 12).

The principle of using the products and energy from an expl detonation to produce a population inversion suitable for lasing has been studied. The design of a 175mm disposable laser detonation cartridge (LDC) was explored, and it was predicted that the energy stored in 30 grams of expl can be converted into a 60 kilojoule laser pulse with a one millisecond pulse duration. A program is outlined to construct a prototype LDC and to measure its performance (Ref 19).

Lasers have also been employed to study fast chemical reactions, those taking place in the time domain of a few billions of a second. Capellos et al at PicArsn, have employed Q-switched Ruby and neodymium lasers, which provide an intense pulse of monochromatic
light of 347 and 265nm, respectively, with time durations of the order of 12 nanoseconds. The ultra-violet light induces electromagnetic excitation in molecules absorbing the laser energy, and by subsequently applying the principle of absorption spectroscopy, kinetic and spectroscopic information relating to the electronically excited states of various energetic molecules have been derived. The systems studied to-date include s-TNB, s-TNT, triphenylamine, and mono- as well as di-nitronaphthalenes (Ref 13, 14, 20, 21 & 28)

Written by S. M. KAYE


Launchers. See Vol 2 of Encyc, p C28-R (Rocket Launcher) and Vol 6 (1974), p G139-R (Grenade Launcher) & G185 (Guided Missile Launcher)

Launoy Powder. A mixt of Na nitrate, sawdust, nitrated bran and sulfur
Ref: Daniel (1902), 423

Lava Fire Bomb. USA incendiary bomb developed during WWII, designated as M74. It weighs 10 lbs and consists of a hexagonal pipe 19" long, inside of which is a plastic cup filled with "synthetic lava" consisting of GOOP (Mg pdr coated with asphalt particles), white P and other ingredients. When clusters of these bombs are impacted on targets, their burning contents flow much like volcanic lava
Ref: Anon, ArmyOrdn 29, 208 (1945)

Lavoisier, A.L. (1743-1794). Fr Chemist, regarded as the father of modern chemistry by virtue of his study of combsn and the role of oxygen. He formulated the theory of the conservation of matter, and laid the basis for chemical nomenclature. From 1775 to 1791 he was in charge of exp1 manufg in Fr ("Regisseur des Poudres"). He was unjustly accused and executed during the Fr revolution
Ref: Hackh's (1944), 481-R; (1969), 381-R
**LAW.** Acronym for **light assault weapon.** The USA M72 66mm anti-tank rocket-launcher weighs 4.5 lbs with one unguided rocket, and has a range of 500–600 yds. The disposable launcher-rocket package is 25° long and extends to 34° for firing.


---

**Lead (Plumbum).** Pb, at wt 207.19, bluish-white, silvery, gray metal, mp 327.4°, bp 1740°, d 11.34 at 20°/4°, d at mp 10.65. Highly lustrous when freshly cut, tarnishes upon exposure to air; very soft and malleable, easily melted, cast, rolled and extruded. Occurs chiefly as sulfide in galena (86.8% Pb), also as carbonate in cerusite (77.5% Pb) and as sulfate in anglesite (68.3% Pb). May be prepd by roasting and reducing Pb ores. Sol in hot concd nitric acid, in boiling concd HCl or sulfuric acid, in acetic acid. Attacked by pure w, weak organic acids in the presence of O. Resistant to tap w, HF, brine & solvents (Ref 2).

Its general uses as a metal are innumerable (Ref 2). In the manuf of expts it is used, in either pure or alloy form, in many types of equipment such as nitrators, drying pans, bearing metal, pipes, tanks, etc. In ammo, it is used as the core of bullets, shrapnel balls and shot, and for the prep of oxides and salts used in expl and pyrotechnic mixts.

One of the most recent uses for Pb is the addition, in the form of foil or coil of wire, to proplnt pdr as a decocker agent for the bore of the gun. The Pb is placed between the proplnt charge and the projectile.

Pb is highly toxic by inhalation of dust or fumes. Tolerance=0.2mg/cc in air. A cumulative poison, FDA regulations require zero Pb content in foods (Refs 3 & 4).


---

**Lead Acetate.** See in Vol 1 of Encycl, p A29–I

**Lead Aceto-Bromate.** See in Vol 1 of Encycl, p A29–L

**Lead Aceto-Chlorate.** See in Vol 1 of Encycl, p A29-L

**Lead Aceto-Perchlorate.** See in Vol 1 of Encycl, p A29-L

**Lead Aceto-Sodium Perchlorate.** See in Vol 1 of Encycl, p A29-L

**Lead Acetylide.** See in Vol 1 of Encycl, p A76-R

**Lead-Antimony (For Use in Ammunition).** Material used by the USA Armed Forces for the manuf of bullet cores, base fillers and point fillers. Three grades are covered in specification MIL-L-13283B (MR) (19 Aug 1970) with the following compn requirements:

<table>
<thead>
<tr>
<th>Grade 1</th>
<th>Grade 2</th>
<th>Grade 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lead + antimony, %, min</td>
<td>99.2</td>
<td>99.2</td>
</tr>
<tr>
<td>Antimony, %</td>
<td>1.0–2.5</td>
<td>9.0–10.5</td>
</tr>
<tr>
<td>Copper, %, max</td>
<td>0.10</td>
<td>0.10</td>
</tr>
</tbody>
</table>

The material shall be clean and of uniform compn, and shall be free from segregations, dross, oxides, blow holes, hard spots, foreign material and other injurious defects. The material shall be furnished in commercial cylindrical ingots of specified dimensions.


---

**Lead Azide.** See in Encycl, Vol 1, A545–A587

*Addl Refs:* 1) B. Reitzner, “Influence of Silver Coatings on Ignition Behavior of Colloidal Lead Azide”, PATR FRL-TR2 (1960), PB148915 2) B. Reitzner, “Influence of Water on Thermal Decomposition of Alpha...
Lead Azide*, PATR FRL-TR5 (1960), PB-148916  
[Decomp of alpha LA]  
5) H.W. Voight Jr & F.H. Schmitt, PATM 1673 (1965) (Low-impulse LA film to be used in applications calling for light-initiated plane wave explosives)  
6) Spec, "Lead Azide (Special Purpose, For Use in Ammunition)", MIL-L-14758 (10 May 1968)  
9) G. Cohn, Ed, Expls&Pyrots 3 (3) (1970)  
(Scanning electron microscopic examination of pure, dextrinated and RD 1333 LA)  
13) Director, Explosives Research and Development Establishment, Waltham Abbey, Essex, England [Abstract reported by G. Cohn, Ed, Expls&Pyrots 7 (4) (1974); New Lead Azide comps RD 1343 & RD 1352]

**Lead Azidodithiocarbonate.** See in Vol 1 of Encycl, p A637-L

**Lead Bichromate.** See Lead Dichromate in Vol 3 of Encycl, C284-R

**Lead Block Compression Test.** See under Compression (or Crusher) Tests for Determination of Brisance in Vol 3 of Encycl, C492-L to C493-L

**Lead Block Expansion Test.** See under Trauzl Test in Vol 1 of Encycl, XXV–XXVI

**Lead Bromate (Monohydrate).** Pb(BrO3)_2·H_2O; mw 481.06, colorless crystals, mp (decomp at 180°), d 5.53g/cc; sl sol in cold w, moderately in hot w. *Poisonous!* Pure Pb bromate is not dangerous, but when prepd by the action of Pb acetate on an alkali bromate, the unstable *diazeto-diplumbo-bromate* is always present. This complex explodes violently on heating, striking or rubbing

a) E. Güzel & E. Marcus, ZAngChem 38, 929 (1925)  
2) I. Victor, ZAngChem 40, 841 (1927) & CA 21, 3324 (1927)  
3) Anon, Chem&Ind 46, 690 (1927) & CA 21, 3170 (1927)  
4) Merck (1968), 612-R  
5) Sax (1968), 864-L  
6) CondChemDict (1971), 508-R

**Lead Carbonate.** See in Encycl 2 (1962), C59-L

**Lead Carbonate, Basic.** See in Encycl 2 (1962), C59-L

**Lead Chlorate (Normal).** See in Encycl 2 (1962), C188-R

**Lead Chlorate (Basic).** See in Encycl 2 (1962), C189-L

**Lead Chlorite.** See in Encycl 3 (1966), C245-R

**Lead Chromate.** See in Encycl 3 (1966), C280-L

**Lead Dichromate.** See in Encycl 3 (1966), C284-R

**Lead 2,4-Dinitroresorcinare (LDNR).** See in Encycl 5 (1972), D1274-R

**Lead 4,6-Dinitroresorcinare, Basic.** See in Encycl 5 (1972), D1275-R
LEAD HYDROXIDE-2,4,6-TRINITRORESORCINATE. See in Encycl 5 (1972), D 1277-L

LEAD IMIDE. See in Encycl 1 (1960), A 169-L

LEAD MONONITRORESORCINATE. See in Encycl 5 (1972), D 1271-L

LEAD NITRATE. See under Nitrates in Vol 8 of Encycl

LEAD NITROAMINOGUANIDINE. See in Vol 1 of Encycl, p A 212-R

LEAD OXIDES

LEAD OXIDE, YELLOW, OR LITHARGE (LEAD MONOXIDE, LEAD PROTOXIDE OR PLUMBOUS OXIDE). PbO, mw 223.21, yellow to yellowish-red, heavy, odorless prdr or minute, cryst scales; mp 888 °C, bp (vol at red heat); d 9,53g/cc; insol in w, alc; sol in acet ac, dil nitric acid, in warm solns of fixed alkalihydroxides. It may be prepd in the lab by heating Pb nitrate, carbonate or hydroxide; commercially, it is made by heating Pb to a temp considerably above its mp and continually skimming off the litharge produced. It is used in some primer compns

Litharge is toxic as a dust. Wear dust mask, and wash thoroughly before eating or smoking.

Keep away from feed or food products


LEAD OXIDE, BROWN OR LEAD DIOXIDE (LEAD SUPEROXIDE OR PEROXIDE). PbO₂, mw 239.21, dark brown tetragonal crystls, mp (dec to PbO and O₂ at 290 °C), d 9.38g/cc. May be prepd from Pb acetate and Ca hypochlorite. Insol in w and alc; sol in acet ac and hot alk

It is an extremely strong oxidizing agent, so care must be taken in mixing or storing with comb materials. When PbO₂ is gently rubbed with sulfur or red phosphorus, the mass ignites. It is used with amorph P as an ignition surface for matches, and in pyrotechnic applications (Ref 5). Anderson, in 1908, proposed its use as an additive to expls to increase their power; for example 1) PA 40 & PbO₂ 69%; 2) PA 20, TNT 10, Guncotton 15, NG 10 & PbO₂ 45%. Colver (Ref 1a) had considerable doubt as to the safety of mixts of this type. McNutt (Ref 2) proposed using PbO₂ in primer mixts, for example, PbO₂ 25, Pb dinitrophenolazide 15, Ba nitrate 30, Sb trisulfide 18, Ca silicide 6 & TNR 6%

The USA military specification (Ref 6) details the following requirements for Type 1 (low alkalinity), Type II (high alkalinity), Class 1 (subsize) and class 2 (100 mesh, nominal) PbO₂

<table>
<thead>
<tr>
<th>Composition</th>
<th>Type I</th>
<th>Type II</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lead dioxide, min, %</td>
<td>95.0</td>
<td>95.0</td>
</tr>
<tr>
<td>Water-soluble salts, max, %</td>
<td>0.05</td>
<td>0.05</td>
</tr>
<tr>
<td>Acidity</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Alkalinity as Na₂CO₃, max, %</td>
<td>0.01</td>
<td>0.10</td>
</tr>
<tr>
<td>Ammonium salts</td>
<td>None</td>
<td>None</td>
</tr>
</tbody>
</table>

Particle Size Distribution

<table>
<thead>
<tr>
<th>Percent Maximum</th>
<th>Class 1</th>
<th>Class 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Retained on a No 100 (149 micron) sieve</td>
<td>–</td>
<td>0.1</td>
</tr>
<tr>
<td>Retained on a No 140 (105 micron) sieve</td>
<td>–</td>
<td>10.0</td>
</tr>
<tr>
<td>Retained on a No 200 (74 micron) sieve</td>
<td>0.0</td>
<td>–</td>
</tr>
<tr>
<td>Retained on a No 325 (44 micron) sieve</td>
<td>5.0</td>
<td>–</td>
</tr>
<tr>
<td>Larger than 20 microns</td>
<td>20.0</td>
<td>–</td>
</tr>
<tr>
<td>Smaller than 5 microns</td>
<td>15.0</td>
<td>–</td>
</tr>
</tbody>
</table>

2) J.D. McNutt, USP 1906394 (1933), USP 1930653 (1933), USP 2009556 (1935)
8) Cond-ChemDict (1971), 509-L

LEAD OXIDE OR LEAD TETROXIDE (Minium, Red Lead, Mineral Orange, Mineral Red, Plumbo, Puce, Orthoplumbate, Paris or Saturn Red).
Pb₃O₄, mw 685.63, bright red scales or amorph pdr, mp (dec betw 500–300°), d 8.32 to 9.16 g/cc; insol in w and alc, sol in acet ac and hot HCl. May be prep'd by carefully heating un-fused litharge at 470–800° for several hours in a current of air. The hot pdr acquires a deeper tint, becoming violet and then black. On cooling, the color changes to red. Used as an oxidizer in pyrotechnics and as an ingredient in some expls, eg, in the Austrian expl “Lederite” to the extent of 20%.

The USA military specification (Ref 3) contains the following chemical requirements:

<table>
<thead>
<tr>
<th>Percentage by Weight</th>
<th>Minimum</th>
<th>Maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>Assay as Pb₃O₄</td>
<td>98.0</td>
<td>—</td>
</tr>
<tr>
<td>Insoluble matter in nitric acid</td>
<td>—</td>
<td>0.10</td>
</tr>
<tr>
<td>Water-soluble substances</td>
<td>—</td>
<td>0.05</td>
</tr>
<tr>
<td>Carbon compounds (as C)</td>
<td>—</td>
<td>0.010</td>
</tr>
<tr>
<td>Manganese (Mn)</td>
<td>—</td>
<td>0.0005</td>
</tr>
</tbody>
</table>

Lead oxide is highly toxic as a dust. Use with adequate ventilation, and keep away from food and food products.


Lead Perchlorate. See under Perchlorates

Lead Picrate. See under Picrates

Lead Plate Test for Detonators. See under Plate Tests for Explosives

Lead Salts of Nitrocompounds. Hopper (Ref), at PicArsn, prep'd and characterized Pb dinitroresorcinlate, Pb dinitrophenolate, and Pb dinitrophthalate as possible substitutes for MF in priming compn. Only the dinitroresorcinlate salt was found suitable. He also prep'd and studied the expl props of Pb Trinitro-N-methyl-nitroamine Resorcinate & Phenolate, and Pb Trinitrobenzoate for possible use as deton. agents. All were unsatisfactory.

Ref: J.D. Hopper, “Study of Lead Salts of Nitrocompounds as Substitutes for Mercury Fulminate”, PATR 480 (1934)

Lead-Shot Metal. An alloy of Pb and As. The As content may range from 0.3 to 0.8% and may be added either in the form of white As or arsenical dross. The As imparts a greater fluidity to the metal and increases its tendency to assume a spherical shape in passing thru the air when dropped from the top of a Pb shot tower into cold w. About 0.025% Na sulfide is added to the w in order to prevent oxidation of the shot.

Ref: CondChemDict (1950), 393-R; not found in later editions

Lead-Sodium Thiosulfate. (Lead-sodium Hyposulfite, Sodium-lead Thiosulfate). Na₄Pb(S₂O₃)₃, mw 635.59, white, small, heavy crysts, mp (dec at 30–40°); sparingly sol in w; readily sol in Na acetate or Na thiosulfate solns. May be prep'd (Ref 1) by mixing solns of Na thiosulfate and Pb acetate, followed by alc addition. On concg the soln, the liq separates into two layers. The bottom layer is removed and treated with more alc, whereupon the mix solidifies to a white, amorph, gelatinous mass of variable compn. On din, the amorph mass turns cryst, which on drying, corresponds to the above formula. It is used in the prepn of matches. Poisonous.


Lead Stearate. See under Stearates

Lead Stypnate. (Lead 2,4,6-Trinitroresorcinate). See Encycl 5, D1277-Lff
Lead Styphnate. Basic. (Lead Hydroxide Styphnate, Lead Hydroxide-2,4,6-Trinitroresorcinate). See Encycl 5, D1277-L

Lead Sulfocyanate. See Lead Thiocyanate

Lead Tetraethyl (Tetraethyl lead, Lead Tetraethide, Tetraethyl Plumbane). Pb(C₂H₅)₄, mw 323.45, colorless liq, fr p -136°, bp about 200°, also stated as 227.7° with decomp, d 1.653g/cc at 20°; insol in w; sl sol in alc, sol in benz and eth. May be prep'd by the action of Pb chloride on Zn ethyl or on a Grignard reag. Used extensively as an anti-knock addition to gasoline, and has been proposed by Fr investigators as a flash reducer in propnts (Ref 2)

Lead Thiocyanate (Lead Sulfocyanate, Lead Rhodanide). Pb(SCN)₂, mw 323.38, white, odorless pdr, mp (dec 190–95°), d 3.82g/cc; sol in about 200p cold, 50p boiling w; sol in alkali hydroxide and thiocyanate solns. May be prep'd by the action of a sol Pb salt (acetate or nitrate) on K thiocyanate (Ref 1). It has found extensive use in stab priming and ignition mixts. Examples are: 1) priming mixt: Pb(SCN)₂ 25, Sb₂S₃ 17, LA 5 & KClO₃ 53%; 2) igniting mixt: Pb(SCN)₂ 45 & KClO₃ 55% (Ref 2)

The requirements of the USA armed forces are covered by a military specification (Ref 3) which contains the following criteria: (1) form – discrete crystals, (2) color – white or yellow, (3) granulation – 100% shall pass thru US standard sieve No 140 (105 microns), and 75% thru sieve No 325 (44 microns), (4) Pb as Pb(SCN)₂, 99.3% min, (5) thiocyanate as Pb(SCN)₂, 99.3% min, (6) chloride, as PbCl₂, 0.20% max, (7) Na, as thiocyanate, 0.20% max, (8) moisture, 0.20% max, and (9) insoluble matter, 0.20% max

Lead Trinitoresorcinates. See under Trinitro-derivatives of Dihydroxybenzene in Vol 5 of Encycl, pp D1277-Lif


Lechoth. A mixt of the diglycerides of stearic, palmitic and oleic acids, linked to the choline ester of phosphoric acid. Yellowish-white, waxy mass, obtained either from eggolk or soybeans. Insol in w; sol in alc, chlf and eth. Lechoth was used in Composition C (RDX 88.3, nonexp oily plasticizer 11.1 & lechoth 0.6%) to help prevent the formation of large crystals of RDX which would increase the sensitivity of the compn (Refs 1 & 4)

Lecithin. A mixt of the diglycerides of stearic, palmitic and oleic acids, linked to the choline ester of phosphoric acid. Yellowish-white, waxy mass, obtained either from eggolk or soybeans. Insol in w; sol in alc, chlf and eth. Lechoth was used in Composition C (RDX 88.3, nonexp oily plasticizer 11.1 & lechoth 0.6%) to help prevent the formation of large crystals of RDX which would increase the sensitivity of the compn (Refs 1 & 4)

Lechoth used by the USA armed forces is covered by a military specification (Ref 3) contg the following requirements: (1) moisture, 1.0% max, (2) benz insol matter, 0.1% max, (3) acid no, 24 max, (4) acetone insoluble matter, 68% min, and (5) Lecithin, 19.0% min

Lécorché-Jovinet Sensitivity Test. Test for the stability of propnts contg NG. Any nitrous acid present is absorbed by diethyldiphenylurea to form ethyldiphenylnitrosamine, which can be detected by testing with α-naphthylamine-HCl

LEDCC. See Low Energy Detonating Cord

Ledere. An older Austrian mining expl contg K nitrate 45, red lead (Pb₃O₄) 20, leather scrap 18, sulfur 15 & PA 2%
Ref: Daniel (1902), 404
LEED. Acronym for Laser Energized Explosive Device. See under Laser

Lennard-Jones, Devonshire Equation of State. Derived for gases at high d, in terms of interatomic forces, using statistical mechanics. The atom in a dense gas was considered similar to that in a liq or crys, subject to multiple collisions at all times (Ref 1). This method was later extended to liqs (Ref 2) and solids (Reps 3 & 4), and was used by Murgai and others for the calcn of the expl properties of TNT and PETN (Reps 5 & 6).


Note: See Encycl 4, pp D287-L to D288-L

Le Marechal Powders. Prep'd by mixing finely pulverized K, Na or Amm chloride (about 84%) with molten stearic or palmitic acid (about 16%). After cooling, the mass was pulverized and mixed with a small quantity of finely powdered charcoal, which served to increase the flammability of the product. The resulting expd was loaded into cartridges by extrusion.

Ref: Daniel (1902), 404

Lenite. A mixt of PA and collodion cotton

Ref: Daniel (1902), 404

Leonard's Smokeless Powders. Several varieties of propltn manufd at the end of the 19th century by the Leonard Smokeless Powder Co of Manchester, NJ, using as one of the ingredients, urea dissolved in acet. Typical comps contd: 1) NG 31.6, Guncotton 52.6, lycopodium 10.5 & urea 5.3%; 2) NG 70.1, Guncotton 23.3, lycopodium 4.7 & urea 1.9%. For use in cannon, these powds were mixed with about 3.5% cottonseed oil as a waterproofing agent.

Reps: 1) Daniel (1902), 405 2) VanGelder & Schlatter (1927), 859

Leonit (Leonite). Ger permisssible Dynamite contg NG 4, K perchlorate 35, AN 10, Na nitrate 3, crude TNT 11, woodmeal 7 & alkali chloride 30%

Reps: 1) M. Giua, "Dizionario di Chimica", Torino (1951), 166 2) B.T. Fedoroff et al, "Dictionary of Explosives, Ammunition and Weapons" (German Section), PATR 2510 (1958), 107-R

Lesmok Powder. One of the older American sporting powds manufd by DuPont

Ref: Marshall 1 (1917), 330

Le Sueur's Explosive. Prep'd by mixing 12.5p of molten paraffin wax with 11p sulfur and 54p Na nitrate. After cooling, the mixt is granulated thru a screen and blended with 22.5p of pulverized K chloride.

Ref: E.A. Le Sueur, USP 923435 (1909) & CA 3, 2227 (1909)

Lauschel Explosives. Ger exps prepd by impregnating moss with glucose, starch, sugar or glycerin solns, drying the mixt and nitrating the resulting product

Ref: Daniel (1902), 405

Levoglucosan Trinitrate. See under Fructosan Trinitrates, Vol 6 of Encycl, F208-L

Levulose Trinitrate. See under Fructose, Vol 6 of Encycl, F208-L

Addnl Refs: 1) Beil 1, (460) 2) H.A. Lewis, USP 1947530 (1934) & CA 28, 2538 (1934) 3) Ibid, CanP 340567 (1934) & CA 28, 4234 (1934)
Lewin Explosives. Patented in Fr in 1887, consisting of nitrated residues of cane sugar plants, either alone or mixed with NG, NC, Na nitrate, rye flour, paraffin, tar, etc. They were also known as "Sandhouiltes"
Ref: Daniel (1902), 406

Lewisite (L). Dichloro (2-chloro-vinyl) arsine.

\[ \text{CICH-CHCl}_2 \]; mw 207.35, dk-grn oily liq, f.p. -18\(^{\circ}\), bp 190\(^{\circ}\), liq d 1.89 at 20\(^{\circ}\), vap d 7.2 (compared to air), decomp temp (above 100\(^{\circ}\)). Vap press 0.087mm Hg at 0\(^{\circ}\), 0.394mm at 20\(^{\circ}\), 32.5o nm at 100\(^{\circ}\). Volatility 967mg/m\(^3\) at 0\(^{\circ}\), 2300mg/m\(^3\) at 20\(^{\circ}\), 8890mg/m\(^3\) at 30\(^{\circ}\). Odor, usually geranium-like, very little odor when pure. Flash p. none; rapidly hydrolyzed in liq or vap state to HCl and chlorovinylarsenic oxide. The later is a nonvol blister-forming solid not readily washed away. L is stable in steel or glass containers
In use, a moderately delayed-action casualty gas. A "blister gas", toxic lung irritant and systemic poison. It produces immediate and strong stinging sensation of the skin
Note: See Encycl 2 (1962), C168-R

Lezinsky's Explosive. A mixt of whole wheat flour 27p with resin 53p is treated with 20p nitric acid and, without washing, combined with 100 to 300p of K chloride
Ref: G. Lezinsky, USP 909915 (1909) & CA 3, 1088 (1909)

L.F. Dynamites. Abbr for Low-Freezing Dynamites. See Vol 5 of Encycl, D1584-L & D1588-R to D1593-L

L.G. Powder. Brit abbr for Large Grain BlkPdr formerly used in cannons
Ref: Daniel (1902), 406

Liardet Explosives. Liardet of Australia patented, between 1889 and 1894, several expls named "Nico Powder" and Acme Powder
(Encycl 1 (1960), A93-R)
Ref: Daniel (1902), 5 & 406

Lichenin (Moss Starch). C\(_6\)H\(_{10}\)O\(_5\); mw 162.14, mp 10\(^{\circ}\). White gelatinous mass, resembling starch in props. SL sol in cold w; sol in boiling w giving a colloidal soln and sol in HCl. On nitrination it yields an exp1, Lichenin Pentanitrate (qv)

Lichenin Pentanitrate. C\(_{12}\)H\(_{14}\)O\(_3\)(ONO\(_2\))\(_5\); mw 549.28, white solid, N 12.75% (theory), 12.4% (found). May be prepd by method described by Reilly (Ref 2) as follows: At RT with stirring, gradually add 1p of finely powd lichenin to 50p of a mixt consisting of 3p by wt of H\(_2\)SO\(_4\) (d 1.84) and 1p of HNO\(_3\) (d 1.50). Stop the agitation, allow the nitrate to settle, and decant the supernatant liq. Pour the ppt and remaining spent acid into ice w. Filter and purify the pentanitrate, first by boiling in sl acidic w, then in sl alkaline w, and finally with neutral distd w. Dry at a low temp
Lichenin Pentanitrate is insol in w, eth and benz; readily sol in acet, et acet and amyl acet; sol in eth-alc mixts. It is an expl compd, decomps at high temps, giving off oxides of nitrogen
Refs: 1) Bell, not found 2) Reilly (1938), 42

Liebert. Patented in 1889 in Fr, a method of nitrating glycerin with mixed nitric-sulfuric acids in the presence of Fe sulfate or AN
Ref: Daniel (1902), 407

Liebert Dynamites. Low-freezing dynamites prepd from NG mixed with 3-5% iso-amyl alc nitrate. It was claimed that these were more powerful than those prepd from NG alone
Ref: Daniel (1902), 407
Life of Guns. See under Erosion of Gun Barrels in Encycl 5 (1972), E112-R to E120-R

Ligdyn. A South African Dynamite contg NG 40, Na nitrate 45, woodmeal 13 & wheat flour 2% (Ref 2). An expn of 4210 lbs of Ligdyn in 1913 was reported caused by friction in a packing machine (Ref 1)
Refs: 1) A.B. Denne, JSCI 32, 627 (1913) & CA 7, 3229 (1913) 2) Marshall 1, 362 (1917)

Light, Effect on Explosives. Many expls undergo slight changes in compn when subjected to direct light, especially sunlight or ultraviolet radiation. Mitra & Shrinivasan (Ref 5) studied the effect of sunlight on thin layers of TNT and Tetryl. In the case of TNT, they noted a lowering of its mp, a reduction in sensitivity, and a color change from light buff to yellow-orange to dark brown with continued exposure. Tetryl exhibited a mp decrease with no change in sensitivity. Nitrous fumes were evolved and exposed surfaces became brown, but much more slowly than for TNT. Krauz and Turek (Ref 1) noted a TNT mp decrease from 81.4° to 73.5° after 4 months direct sunlight exposure, and extracted Trinitrobenzoic acid and Trinitrophenol as breakdown products

Urbanski (Ref 6) followed the photochem decompn of NG, NC (11.9% N) and Hexogen on exposure to UV radiation by the liberation of iodine from K iodide soln, and derived rate equations. NG continued to decomp even after the UV source was removed, becoming increasingly acidic, and was judged less stable than NC. The NC rate of decompn remained constant with time. NC evolved oxides of N₂, became acidic, and was finally judged useless as an expl material. Hexogen, although changing color from yellow to brown, produced no volatile products capable of oxidizing K iodide

Münzinger (Ref 2) exposed pure NC to sunlight and noted only slight yellowing after several months. However, NC contg gelatinizers such as castor oil or tricresyl phosphate was affected in a matter of days, yellowing and becoming less sol in et acet or acet

Diazodinitrophenol (DADNPh) was found to be stable for long periods of time in diffused light. Samples standing in the laboratory, protected from direct sunlight, showed no signs of discoloration after 6 months. In direct sunlight, however, DADNPh darkened rapidly eventually acquiring a dark brown color. To detn the effect of exposure on its expl strength, DADNPh samples subjected to from 0 to 20 hours of sunlight were fired in a Sand-Test bomb. The results showed that DADNPh undergoes no loss of strength on one hour exposure, with no marked decrease even after 3 to 5 hours, despite a marked darkening in color (Ref 3)

Eggert (Ref 8) and McAuslan (Ref 7) reported on the initiation of explosions by light. Evidence was presented for photochemical and/or photothermic effects in various nitrides, azides, acetylides and perchlorates

Light Emission (Luminosity Effects) from Detonations and Explosions. See "Detonation (and Explosion), Luminosity (Luminescence) Produced on" in Encycl 4, (1969), D425-L to D434-L, and "Detonation (and Explosion); Spectra and Spectrographic Measurements in", D548-R to D549-L

Light, Production of. See under Pyrotechnics

Lightning Protection. It is policy in the USA to install lightning protection on blgs and structures used for manufg, processing, handling or storing expls, amm, expl ingredients, and other hazardous materials, particularly where
operations cannot be shut down during electrical storms and personnel evacuated.

Approved lightning protection systems are the integrally mounted system, the separately mounted shielding system (mast type), and the separately mounted shielding system (overhead ground wire). Details of all of these systems are described in the Ref.

The purpose of these installations is to provide a metal path of low resistance for the discharge of electrical currents from the air to the ground without damage to the structure or contents. These systems also serve to prevent the charging of structural metal components as a result of induction when lightning strikes nearby.

Ref: Anon, "Safety, Safety Manual", AMCR 385-100 (April 1970), Chapt 8, 8-1 to 8-36

Lignin. A polymer found in wood (25–30%). The structure of the lignin monomer is still not completely known. Its similarity to coniferyl alcohol, noted more than 75 years ago (Ref 1), is confirmed by the fact that it can be oxidized to vanillin and hydrogenated to compounds of the cyclohexylpropyl type. Lignin is removed from wood by both the sulfate and soda paper pulp processes, and limited amounts have been recovered from these sources and other wood waste. It has been used as a component of Dynamites, and has been nitrated.


Lignin Nitrates (Nitrolignin). A general term employed to designate nitrated products containing lignin, such as wood, straws, jute, esparto grass, flax and hemp fibers (Ref 1). The earliest experiments devoted to the nitration of lignin, previously isolated from wood pulp by means of HCl, were reported by Hägglund (Ref 3). Lignin mixed with fuming nitric acid was heated on a water bath until the lignin was completely dissolved, whereupon the nitration product was ppt by dil the soln with water.

Fischer and Schrader (Ref 4) nitrated lignin by treating it with 31.5% nitric acid at RT, completing the operation with moderate heating. Only 4.3% N was found in the prod, to which the formula C_{39}H_{39}O_{21}N_{3} (OCH_{3})_{3} was assigned on the basis of analysis. At the same time, König (Ref 5) nitrated lignin at low temp to obtain a prod contg 3% N.

Powell and Whittaker (Ref 6) nitrated lignin with mixed nitric-sulfuric acids at 5°C, and isolated the nitration prod by pptn with ice and water. It was a red powd with the empirical compn C_{42}H_{39}O_{13}(NO_{2})_{3}.

Kürschner et al (Refs 7, 8 & 11) prep nitrolignin by direct nitration of wood pulp. A soln of nitric acid (d 1.40) in alc was used at the temp of boiling alc. Under these conditions cellulose remained unchanged. Lignin, on the other hand, was nitratet and dissolved in the alc soln. Nitrolignin was then separated from the soln with water, in the form of an amorphous yellow powd. The yield of nitrolignin was 15–17% of the wt of wood pulp. According to the type of wood used to prepare the lignin, the N content of the fully nitrated prod varied from 3.0 to 4.5%. On the basis of his own experiments, Kürschner suggested the following functional compn for nitrolignin:

C_{18}H_{31}O_{2}(COOH)(OH)_{2}(OCH_{3})(NO_{2})

In order to separate lignin from carbohydrates, stronger reagents were used in the experiments of Friese et al (Refs 9 & 12) for the nitration of wood pulp, namely normal mixed acids. In this way nitrolignin with a higher N content was obtained. When nitric-sulfuric acids (1:2 ratio) were used, a prod contg 8.45% N was obtained with a yield of 55%.

Although nitrolignin was used in early propellants, mining exps, and sporting powds (Refs 1 & 2), Urbanfiski (Ref 14) states that it is unsuitable for manufg nitro compds likely to be of practical value as exps.

Lignin Dynamites. Mixts prepd by impregnating sawdust with NG were known by this name. They sometimes contd metallic nitrates.

Ref: Daniel (1902), 408.
Lignite (Brown Coal). A brownish-black coal in which the alteration of vegetable matter has proceeded further than in peat, but not as far as in sub-bituminous coal. There is no sharp distinction between these three materials, but in general, lignite is denser, darker in color, and contains more C than peat. Lignite consists of w 9–12, vol comb matter 31–38, fixed C 27–43, and ash 5–20% (Ref 3).

Lignite in powd form has been used as a component of blasting expls, for example, Explosive of Kolowratnik (1912) contains PA 45, Na nitrate 40, sawdust 9 & lignite 6% (Ref 1). Marcusson (Ref 2) nitrated lignite with fuming nitric (d 1.52) and mixed nitric-sulfuric acids. The prod was sol in acet, and consisted of various nitrocompds 

**Refs:** 1) Colver (1918), 325 2) J. Marcusson, ZAngChem 34, 521–2 (1921) & CA 16, 1496 (1922) 3) CondChemDict (1950), 398; (1971), 516-R

Limpet Charge. HE grenade or mine which uses a magnet for adherence to the metal sides of tanks or ships 


Lindeman Explosive. BritP of 1899 for an expl consisting of NG, K chlorate and MNB or DNB 

**Ref:** Daniel (1902), 408

Lindner Explosive. FrP of 1895 for an expl contg AN 93.2%, naphthalene 5.5 and K chlorate 1.3% 

**Ref:** Daniel (1902), 409

Linear Shaped Explosive Charges. Picatinny Arsenal developed a linear shaped demolition charge (Ref 1) capable of felling trees up to 40” in diam. The charge is 7” long, 2.4” high and 4.7” wide, and contains 1.5 lbs of Comp B. It is more efficient than other shaped charges, such as conical shaped and...
platter charges, because of its exceptional side-cutting ability. The claim is made that this charge uses 50-70\% less expl than more conventional bulk and ring tree-cutting expl devices.

Moses (Ref 3) describes linear expls consisting of tube-like containers filled with burning or detong expls of the following types:
1) Safety fuze used for mining and quarrying consists of BkPdr in cotton and polyethylene covers, wax dipped; 2) Detonating cord used for general blasting and demolition consists of PETN or RDX in a waterproof textile sheath; 3) Mild deton fuze is smaller than type (2); it is used to transfer deton, and has a metal sheath; 4) Confined deton fuze is similar to type (3) except that the metal sheath is wrapped in fiberglass and polyethylene to contain the fragments; 5) Flexible linear shaped charge is a detong cord with a V-shaped cross-section used to cut metal.

Linear shaped charges as well as expl cords have been developed for use in commercial cutting operations. One special application is a cast iron pipe cutter called “Jetcutter”. It makes a straight and clean cut in cast iron pipe from 6 to 12\" in diameter by means of a reusable fixture and an expendable shaped charge insert. A forced entry tool, dubbed “Jet-Axe” cuts thru 5\" of roofing material and thru fire doors and similar structures. Rapid egress systems are being developed for aircraft and other passenger carrying vehicles (Ref 3).


Lined Cavities of Explosives (Lined Cavity Effect). See under Detonation, Munroe-Neumann Effect (or Shaped Charge Effect) And Lined-Cavity Effect In, Encycl 4 (1969), D444-R to D450-R

Line Wave Generator. Devices for controlling the sequence of arrival of detonation waves at various points in an expl charge. Those of the manifold type are made by loading expls into channels machined, molded, or cast into metal or other inert components and by constructing arrays of detong cord. These arrays are limited to relatively large systems by the spacing needed to prevent initiation or damage due to radial blast effects of adjacent cords (See Fig).

Line Wave Generator of the Manifold Type

The advent of mild detonating cord (MDC) has opened new possibilities in manifold type wave shaping devices.

Another line wave generator of the manifold type consists of perforated sheet expl (See Fig).

Line Wave Generator of Sheet Explosive

A duPont version (Ref 1) is in the form of perforated equilateral triangles measuring 10.9\" on a side and are prep'd from “Detsheet” flexible expl. The sheet expl is perforated in such a manner that a deton initiated at any one of the apexes will proceed as a straight-line deton zone to the opposite edge.

In addition to such generators, warped surfaces may be used to produce line waves of any desired curvature. The circular front generated by the point initiated deton of a plane charge may also be modified by warping the plane and by transmitting to other expl surfaces. An example, illustrated below, is the generation of a straight-line wave by means of warped sheet expl.
Line Wave Generator of Warped Sheet of Explosive


Linen Fiber (Flax, Byssus). The bast fiber of the flax plant, an annual herb about 2 ft high. It is grown in Europe and Egypt for the fiber and in the USA, USSR & Argentina for the seeds (linseed oil). The nitrated unbleached prod was proposed by Trench (Ref 1, p 773) for use in expl compns

Nitrated Linen Fiber (Nitroflax, in Fr Nitrolen). Product resembling cotton NC and prepd by nitratrung unbleached flax. When nitrated under identical conditions, flax gives a more viscous prod than cotton. In order to lower the viscosity of nitroflax, the temp of nitration is raised. Nitroflax has been used in some celluloids (Ref 1, p 120), and was proposed (Ref 1, p 773) as a basic ingredient in commer-cial expls, other components being collodion cotton, glycerin, ozokerite, resin, etc

Refs: 1) Daniel (1902), 120 (under Celluloid) & 773 (under Trench) 2) Hackh’s (1944), 345-R

Linolein (Glyceryl linoleate). A glyceride of linoleic acid. It is one of the constituents of linseed oil which induces drying. Yel oil; may be prepd by treating K linolate with trichlorhydrin at 160° in a stream of H₂. V sol in eth, benz and chlf; less sol in alc, methanol, ligroin & C₅

The nitrated compd was proposed by Reid and Earle (Ref 2) as a constituent of expl mixts, eg, 1p nitrolinolein and 9p NC Linolein Nitrate (Nitrolinolein). A prod obtd

by nitrination of linolein (See above) and used in some expl mixts


Linseed Oil (Boiled Oil, Bung Oil or Bung Hole Oil). The oil obtained by hot pressing the seeds of the flax plant. Consists of glycerides of linolenic, linoleic, oleic, stearic, palmitic and myristic acids. Yellowish liq, peculiar odor, bland taste

“Boiled linseed oil” is a misnomer, since the oil does not boil. Small amts of driers (oxides of Mn, Pb or Co, or their naphthenates, resinates or linoleates) are added to hot linseed oil to accelerate drying. The “boiled oil” becomes thicker and darker. It is combustible, with a flash p of 403°F and an autoignition temp of 650°F (Ref 4)

For many years, boiled linseed oil has been used to coat Mg powd to protect it from corroson when used in pyrotechnics. Mg powd and oil are mixed together and allowed to stand in a warm place in shallow trays for about 48 hours, before the other chemicals are added. Lately there has been a tendency to replace linseed oil with polyesters, or to use no coating at all, but there can be no doubt that linseed oil renders good protection. Stores made with Mg coated with linseed oil are good for several years, which is not the case with uncoated Mg (Refs 2 & 5). Linseed oil has also been used for coating K chlorate used in expls (Ref 1)

The requirements of the USA armed forces for boiled linseed oil are covered by Federal Spec TT-L-190C (Ref 3) and are as follows: (1) Set-to-touch (hours), 16 max; (2) Loss on heating at 105°-120° (by wt), 0.3 max; (3) Acid number, 7.5 max; (4) Saponification number, 189-195; (5) Unsaponifiable matter
better sepn of the linters and to have less short fibers adhering to the seed, deSegundo devised a special machine. He concluded that it is very difficult to completely remove all the fuzz without employing chemical methods.

Note: See also Cotton Linters in Vol 3 of Encyclopedia of Materials, p 547.


Liquefaction Test. A test used to deter the exudation of Dynamites. In Eng. the test was conducted as follows: A cylinder of approx equal length and diameter was cut from a cartridge of Gelatin Dynamite to be tested. With paper removed, the base of the cylinder was placed on a sheet of pasteboard and held fast by a pin driven vertically thru its center. After conditioning at 29–32°F (85–90°F) for 144 hrs, the height of the cylinder was measured. If it did not shrink more than 25% of its original height, the Dynamite passed the test. Another condition of acceptability was that the upper cut surface retain its flatness and sharpness of its edges.

In Germany, the test was conducted by heating an entire cartridge for 5 days at 30°C (86°F). The requirement was that no NG should exude and, after cooling, that the cartridge dimensions remain unchanged.

Refs: 1) Marshall 1 (1917), 163; 2) Naoum, NG (1928), 314

LIQUID AIR AND LIQUID OXYGEN EXPLOSIVES

Liquid Air and Liquid Oxygen Explosives originally consisted of porous combustible materials impregnated with liquid air. Soon after liq oxygen became commercially available it began to replace liq air in these explosives. Consequently this article is devoted almost entirely to Liquid Oxygen Explosives commonly called LOX. It should be noted that LOX are not to be confused with Fuel-Air Explosives (See FAE in Vol 6, p F3). For LOX the oxidizer is liq and the fuels are either solids or liquids, while in FAE the ox-

Linters. Fleecy short fibers (1/8" to 1/4" in length), consisting chiefly of cellulose, which adhere to cotton seed after it has been passed once thru a cotton-gin. These are usually removed from the seed by a second and third ginning to yield first and second cut linters. The first cut fibers are the longer and are used mainly for padding, upholstery, mattresses, etc. Second cut linters are suitable for the prep of various NC's after special treatment consisting of boiling in caustic soda, followed by bleaching.

Purified linters are used extensively in the USA for the prep of NC, but longer fibers are preferred in Eng. During WWI, Engl also used material called “divers” (Ref 1, 3, p 30), which was staple cotton in the form of unspun strips – an intermediate stage in the manuf of yarn. Although more expensive than cotton waste, it yielded purer and more stable Cordite.

In Fr, three kinds of cotton material were used in NC manuf: (1) Waste No 1, which consisted of bleached waste from spinning and weaving mills using long-staple cotton of Egyptian and American origin; (2) Waste No 2, consisting of unbleached spinning-mill waste and (3) Linters, similar to American linters, which must be free of cotton waste. According to deSegundo (Ref 2), some short fiber cotton called “fuzz” still remains on the seed after removing the long fibers and linters by ginning and delinting machines. In order to effect
L. 20

dizer is atmospheric oxygen and the fuels are usually gaseous or liquid droplets at the time of explosion. General references on liq air and liq oxygen expls are Refs 2, 3, 5, 7, 8, 14, 16, 19 & 52.

In what follows we will examine:
1) The history of LOX
2) Typical LOX compositions
3) Uses
4) Detonation and sensitivity characteristics
5) Recent Patents

1) History. LOX, or more precisely Liquid Air Explosives, were invented by Linde in 1895, who called these expls Oxyliquits (Ref 1). They were made by impregnating porous combustible solids with liquid air shortly before firing the charge. Usually the combustibles, in some type of combustible cartridge, were soaked in liq air just before loading into the bore hole. Liq air expls were used extensively in 1899 in the driving of the Simplon tunnel between Italy and Switzerland.

Liq nitrogen has a much lower heat of vaporization than liq oxygen, and thus evaporates more readily. Because of this, liq air (a mixt of nitrogen and oxygen) becomes progressively richer in oxygen as it is warmed or even in storage. This makes it almost impossible to control the oxygen content of a liq air expl charge, even if it is fired promptly after preparation. Variable oxygen content can result in poor performance, or even non-performance, and in uncontrolled explosive fumes. Thus, the natural tendency to replace liq air with liq oxygen in these expls began as soon as liq oxygen became commercially available. This occurred some years prior to WWI.

During WWI the Germans used LOX (and also liq air explosives) extensively in coal, iron and potassium mines, in tunneling and in demolition work. In 1922 LOX were used in Mexican silver mining and a few years later they were introduced into the copper mines of Peru and Chile.

In the USA, LOX have been used primarily in the strip mining of coal. For example, in 1950, 99.5% of all the LOX used was in coal mining (Ref 18). Most of this must have been for moving overburden, since LOX are non-permissible (not allowed by law in gassy mines), and their fume characteristics make them unsuitable in many underground mines even if they are non-gassy. The following tabulation (from Ref 18) shows that LOX consumption in the 1950's amounted to some two to three percent of the total expls used.

Table 1

<table>
<thead>
<tr>
<th>Year</th>
<th>Black</th>
<th>Blasting Powder</th>
<th>Permissibles</th>
<th>Other than Permissibles</th>
<th>LOX</th>
</tr>
</thead>
<tbody>
<tr>
<td>1935-1939</td>
<td>65.4</td>
<td>45.7</td>
<td>253.6</td>
<td>Not available</td>
<td></td>
</tr>
<tr>
<td>1949</td>
<td>20.1</td>
<td>91.6</td>
<td>505.6</td>
<td>13.9</td>
<td></td>
</tr>
<tr>
<td>1950</td>
<td>20.7</td>
<td>109.4</td>
<td>576.0</td>
<td>13.8</td>
<td></td>
</tr>
<tr>
<td>1951</td>
<td>14.0</td>
<td>108.3</td>
<td>611.2</td>
<td>20.3</td>
<td></td>
</tr>
<tr>
<td>1952</td>
<td>10.6</td>
<td>95.5</td>
<td>636.7</td>
<td>21.9</td>
<td></td>
</tr>
<tr>
<td>1953</td>
<td>9.5</td>
<td>89.9</td>
<td>669.0</td>
<td>22.5</td>
<td></td>
</tr>
<tr>
<td>1954</td>
<td>10.3</td>
<td>75.9</td>
<td>615.8</td>
<td>17.7</td>
<td></td>
</tr>
<tr>
<td>1955</td>
<td>6.6</td>
<td>93.7</td>
<td>687.2</td>
<td>19.3</td>
<td></td>
</tr>
<tr>
<td>1956</td>
<td>5.6</td>
<td>97.7</td>
<td>(814.4)</td>
<td>Not available</td>
<td></td>
</tr>
</tbody>
</table>

*Data kindly supplied by Robert W. Van Dolah, Chief, Division of Explosives Technology, Bureau of Mines, Pittsburgh, Pa

Since then the use of LOX has declined greatly, as ANFO and Slurry Explosives began to replace them. An indication of the decline in the use of LOX is given in the patent literature. For example, a 1936 review (Ref 8) lists 64 German patents on LOX. Undoubtedly by 1936 there were also many patents issued in the USA, UK and France. In the period of 1936-1960 a considerable number of patents on LOX is listed in CA. Some of these are abstracted in Section 5 below. Since 1960, however, the number of LOX patents has declined drastically. Almost none is listed in the most recent Quintennial Index of CA.

2) Typical LOX Compositions

Both Liq Air and Liq Oxygen expls contain porous combustible material as fuel. These fuels are generally contained in paper or cloth cartridges. Occasionally liq fuels such as petroleum are mixed with the porous solids; eg, some of Linde's early compositions (Ref 1) contained kieselguhr mixed with petroleum. Carbonized cork was also used in early Liq Air Expls, although charcoal was the original absorbent. More recently, as liq air was replaced
### Table 2

<table>
<thead>
<tr>
<th>Initial Composition (a)</th>
<th>Initial Sp Gr, g/cc</th>
<th>Relative Strength (b)</th>
<th>Detonation Rate (m/sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>38/225 LampBlack/LO</td>
<td>0.23 (c)</td>
<td>0.95</td>
<td>4200</td>
</tr>
<tr>
<td>57/230 LampBlack/LO</td>
<td>0.33 (c)</td>
<td>1.14</td>
<td>5000</td>
</tr>
<tr>
<td>65/225 Gas Black/LO</td>
<td>0.33 (c)</td>
<td>1.16</td>
<td>5000</td>
</tr>
<tr>
<td>49/12/215 Gas Black/FeSi/LO</td>
<td>—</td>
<td>0.94</td>
<td>—</td>
</tr>
<tr>
<td>36/28/193 Woodpulp/Kieselguhr/LO</td>
<td>1.07</td>
<td>0.92</td>
<td>4180</td>
</tr>
<tr>
<td>49/12/216 Woodpulp/Lampblack/LO</td>
<td>0.76</td>
<td>0.80</td>
<td>3350</td>
</tr>
<tr>
<td>58/7.3/167 Woodpulp/Kerosene/LO</td>
<td>0.93</td>
<td>0.95</td>
<td>4660</td>
</tr>
<tr>
<td>64/26/182 Woodpulp/Kerosene/LO</td>
<td>1.09</td>
<td>0.80</td>
<td>4080</td>
</tr>
<tr>
<td>33/49/218 Fuel Oil/MgCO₃/LO</td>
<td>—</td>
<td>0.99</td>
<td>4000</td>
</tr>
<tr>
<td>47/210 Carbene /LO (d)</td>
<td>0.24 (c)</td>
<td>1.13</td>
<td>5200</td>
</tr>
</tbody>
</table>

(a) Before any appreciable evaporation of liq oxygen (LO)
(b) Relative to 40% Dynamite on a volume basis
(c) "Unsoaked" fuel; "soaked" sp gr not given
(d) Carbene is polymerized acetylene
(e) In an iron tube; presumably all other detonation rates are for unconfined cartridges

by liq oxygen, lampblack became the absorbent most commonly used in LOX (Ref 12). The fuel content of LOX compositions should be capable of absorbing 5 to 6 times their weight of liq oxygen. Many other fuels such as soot, turf, corkmeal, powdered anthracite, woodmeal, carbene (polymerized acetylene), calcium hydride, and spongy aluminum have been tried. Physical and chemical properties of many LOX fuels are given by Howell et al (Ref 3) and O'Neil & Van Fleet (Ref 5a).

Several of the potentially useful LOX compositions with some of their physical and detonation characteristics are listed in Table 2 (taken from Ref 6). Some recent LOX compositions are given in Section 5.

3) Uses. In section 1, we showed that most of the LOX in the USA are used in the strip mining of coal. In Europe LOX were also extensively used in open pit mining, tunneling and construction. Indeed, in the first half of this century, LOX were used in most explos applications, although not extensively, where fumes were not a problem. O'Neil & Van Fleet (Ref 5a) consider LOX economical and safe (See Sect 4 on LOX safety). La Magna (Ref 9) prefers LOX to Dynamites. In the last decade LOX have been almost entirely replaced by ANFO or Slurry Explosives.

In actual practice LOX were always prepared near the explosion site. Usually a paper cartridge containing the absorbent fuel was "soaked" in liq oxygen. The soaked cartridges were quickly placed in the borehole and fired promptly. Extensive tests (Ref 3) showed that firing had to occur within 5 to 15 minutes after soaking, otherwise enough liq oxygen evapd to affect LOX performance or even cause misfires. The LOX charges were generally fired with blasting caps. Under favorable conditions, LOX charges can be initiated by flame, but this type of initiation is uncertain, and the performance of LOX thus initiated tends to decrease. As discussed in Section 4, the sensitivity of LOX to flame can be a safety hazard.

In some operations, a cartridge packed with absorbent fuel was inserted in the borehole and filled with liq oxygen thru a tube reaching to the bottom of the cartridge. A vent for evap'd oxygen had to be provided.

Yet another method was to make cartridges with two compartments, one for absorbent and the other for liq oxygen. After insertion in the borehole, the partition was ruptured (either by pressure from the oxygen or by mechanical
means from outside) to mix the liq oxygen with absorbent. To increase the allowable time between liq oxygen impregnation and firing (from 10 minutes to 16–22 minutes), Wakabayashi (Ref 21) suggests precooling the borehole by pouring small amounts of liq oxygen into it.

4) Detonation and Sensitivity Characteristics

a) Detonation Characteristics. LOX is commercial explosives and as such are used primarily in breaking and moving rock and overburden. From a practical point of view, it is important to have a measure of the effectiveness of LOX blast in fracturing and moving the "burden". No single universally accepted measure of blast effectiveness exists today, and certainly none existed in the 1920–1940 period when most of the expl studies of LOX were carried out. Practical experience suggests that the effectiveness of an explosive for fracturing "burden" is related to its brisance. Brisance is a measure of the shattering power of an explosive and is closely related to the detonation press (commonly called P_{CJ} or Chapman-Jouguet pressure) of the explosive (See Brisance in Vol 2, p B265–300). The effectiveness of an explosive for moving "burden" is related to its strength or power (See Vol 4, p D730-L). It is customary to rate explosive strength on a relative basis, i.e., as a percentage of the strength of some standard explosive — usually TNT or some standard Dynamite. This rating is based on comparison tests, the most common of which is the Ballistic Mortar Test (See Vol 2, p B6-R).

Unlike the relation between brisance and P_{CJ}, expl strength is not readily related to some detonation characteristic of the explosive. Attempts to relate strength to detonation energy are not wholly successful. Relative strength, based on ballistic mortar tests, correlates rather well with computed nRT, where n & T are the computed moles of gas and detonation temp of the explosive, and R is the gas constant. Although n & T can differ appreciably with the equation of state used in the computation, it appears that ratios of nRT (at least for similar explosives) do not suffer from this drawback.

In the early expls literature (and much of the LOX work is in the "early" literature) there is a great deal of confusion between brisance and strength.

Now, with the above caveat, we can examine what is known of LOX detonation characteristics.

Perrott (Ref 4) examined the effect of packing density on the relative strength of six liq oxygen-lamp black comps, and measured their detonation rates, D. Relative strengths on a weight basis increased as packing density increased except that the relative strength at the highest packing density tested (0.46g/cc "unsoaked") was low. On a volume basis the compn at the lowest packing density (0.19g/cc "unsoaked") had the highest relative strength. Detonation rate increased from 4500 to 6000m/sec, although it is not clear whether this increase, as expected, occurred as packing density was increased. Perrott also found that substituting Al for some of the absorbent did not increase D but made the compn easier to detonate.

In a subsequent study (Ref 5) Perrott found that 1½ inch diameter cartridges of LOX, containing lamp black, gave optimum blast results at an "unsoaked" packing density of 0.30g/cc. This takes into account not only the strength of the LOX but also their effective "life", i.e., the maximum allowable time between "soaking" and firing. For example, 5 minutes after soaking, LOX cartridges were found to have a relative strength (on a volume basis) 115% that of a standard Dynamite, but after 25 minutes their strength was only 65% of the Dynamite. Perrott suggests using as large a cartridge diameter as practicable to reduce oxygen evapn. He also made further measurements of detonation rate and found that it is controlled by the finest particle size component of the absorbent. Clark & La Motta (Ref 7) also found that D increases as absorbent particle size decreases. For gas-black LOX, D varied from 4000 to 6200m/sec, while for lamp black LOX D varied from 4200 to 5000 m/sec. Perrott states that LOX will burn without detonation when unconfined but will detonate erratically when ignited under confinement.

Okada (Ref 15) states that the oxygen to carbon ratio for LOX of maximum brisance is 2.6. This is essentially the theoretical ratio to convert all the carbon to CO2. This writer believes that Okada really meant maximum strength rather than maximum brisance, as the latter depends not only on compn but also on pack-
Okada's conclusions are supported by Streng & Kirshenbaum (Ref 20) who found that a stoichiometric mixture (33 mole% CH₄) of liq methane and liq oxygen had a higher brittleness (in this case they really measured brittleness) and a higher detonation rate than other mixtures containing from 6 to 80 mole % CH₄. They also determined the expl limits and detonation rates of these mixtures and examined the sensitivity of the stoichiometric mixt to impact, shock waves, and flame & sparks. Their results are summarized below:

<table>
<thead>
<tr>
<th>Mole % CH₄</th>
<th>Table 3</th>
<th>Density</th>
<th>Detonation Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>6</td>
<td>1.05</td>
<td>Failed</td>
<td></td>
</tr>
<tr>
<td>20</td>
<td>0.98</td>
<td>3325</td>
<td></td>
</tr>
<tr>
<td>33</td>
<td>0.88</td>
<td>5110, 5130</td>
<td></td>
</tr>
<tr>
<td>40</td>
<td>0.83</td>
<td>5110</td>
<td></td>
</tr>
<tr>
<td>50</td>
<td>0.76</td>
<td>4620, 4610</td>
<td></td>
</tr>
<tr>
<td>67</td>
<td>0.65</td>
<td>Detonated</td>
<td></td>
</tr>
<tr>
<td>80</td>
<td>0.50</td>
<td>Failed</td>
<td></td>
</tr>
</tbody>
</table>

Although the above are not practical LOX compositions, these results are of considerable interest. They show that the detonation limits of LOX comps are quite wide, and that D (at least for liquid methane-liquid oxygen mixtures) is not strongly affected by considerable changes in compn or even density.

Cook (Ref 22) studied a chemically similar mixt consisting of 78/22 LOX/kerosene (stoichiometric to CO₂ & H₂O), but the kerosene, ie, the fuel, in his case was frozen, so that he dealt with a slurry rather than a solution. His results are as follows:

<table>
<thead>
<tr>
<th>Density</th>
<th>D</th>
<th>P_CJ</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.04g/cc</td>
<td>2240m/sec</td>
<td>10.4kbar</td>
</tr>
</tbody>
</table>

Note that D for the slurry is much lower than for the liq CH₄–liq oxygen solns even though the apparent slurry density is higher than those in Table 3. Possibly this is due to incomplete reaction, even though Cook emphasizes that he used mechanical stirring. Cook's D is also very low compared to the D's in Table 2, for practical LOX comps containing solid absorbent fuels.

Cook (Ref 18a) presents theoretical calculations of the detonation parameters of several carbon black–liq oxygen comps. He gives no comparisons with experimental results. In this writer's opinion, Cook's values (Table 12.20 of Ref 18a) of T_CJ are too high and his P_CJ are too low; the products probably contain more free carbon than shown. However, there is little doubt that most LOX detonations are "hot", i.e., they will readily ignite firedamp (Refs 2, 3, 12 & 14). Also the computed detonation velocities (estimated from Cook's P_CJ) show a much greater variation with compn and density than the experimental data of Table 3.

b) Sensitivity Characteristics. Early investigators (Refs 5a & 9) tended to overestimate the safety of LOX because they were non-explosive before mixing, and because they became non-explosive, even after a misfire, as the oxygen evapd. An interesting comment, by Wakabayashi (Ref 21), significant in these days of terrorist bombings, is that LOX are burglar-proof. Nevertheless, modern consensus is that LOX are more dangerous to handle than conventional Dynamites (Refs 7, 12, 14, 17 & 21).

One of the most serious faults of liq oxygen explosives is the ease with which they inflame and the rapidity with which they burn, amounting practically, in the majority of cases, to their exploding from fire. Denues (Refs 10 & 11) has found that treatment of the granular carbonaceous absorbent with an aqueous solution of phosphoric acid results in an explosive which is nonflammable by cigarettes, matches, and other igniting agents. Mono- and diammonium phosphate, ammonium chloride, and phosphoric acid were found to be suitable for fireproofing the canvas wrappers. Liq oxygen expls made up from the fireproofed absorbent are still capable of being detonated by a blasting cap. Their strength, velocity of detonation, and length of life after impregnation are slightly, but not significantly, less than those of expls made from ordinary non-fireproofed absorbents containing the same amount of moisture.

Streng & Kirshenbaum (Ref 20) found that a stoichiometric mixture of liq methane and liq oxygen will explode from the flame of a safety fuse.

Some LOX comps are liable to self ignite (Ref 12). Cook (Ref 18a) makes the interesting suggestion that many LOX comps do not
self-ignite only because they are so cold

Clark & La Motta (Ref 7) showed that LOX made with gas black or lamp black are more sensitive to impact than the standard Bureau of Mines 40% straight Dynamite. Impact sensitivity increased when small amounts of iron oxides, aluminum dust or ferro-silicon were added to the LOX. Impact sensitivity also increased as absorbent particle size was reduced. As the oxygen evaporated, impact sensitivity, as expected, decreased.

In tests of materials saturated with liq oxygen and subjected to 71–75 lb drop weight tests, the following were found acceptable (one detonation/40 impacts or none/20): fluorocarbon oils & greases, graphite, halogenated biphenyl & molybdenum disulfide lubricants, polyethylene & pure poly fluorocarbons. The following exploded: synthetic elastomers & Thiokols, cellulose-based papers, silicone-based oils & greases, thermoplastics (except pure Teflon), thermostetting plastics, petroleum-based oils and greases (Ref 17).

LOX are sensitive to sympathetic detonation, i.e., detonation initiated by a nearby charge separated from the LOX by an air gap (Refs 7 & 20).

The pseudo-LOX of liq methane–liq oxygen are exploded by bullet impact (Ref 20).

LOX compositions are sensitive to friction (Refs 12 & 14) and to static discharge. Asonov (Ref 13) attributes the premature explosions of liq oxygen on adsorbents (fuels) to small dust particles. As the charge is dropped into the borehole, the small particles become detached from the adsorbent. These particles are carried upward by the oxygen vapors and become electrostatically charged by friction.

Potentials of the order of 15000–20000 volts were measured on these particles. A discharge of such potentials suffices to initiate the LOX. Dust can be minimized by moistening the adsorbent with 20–25% of water or by providing special capsules for the expl. Good results were obtained by using briquetted fibrous vegetable matter, peat, straw, wood pulp, and the like as adsorbents. Such briquets did not produce dust when dropped from a height of 25m. The brisance of the LOX made of such briquets was only 3–4% less than that of ordinary LOX.

Streng & Kirchenbaum (Ref 20) exploded stoichiometric liq methane–liq oxygen by the discharge of a 0.1 microfarad condenser, charged to 1500V, across 1–3 mm air gaps.

5) List of Recent Patents on LOX


Liq oxygen is used with an intimate mixture of a main mass of a solid combustible material such as lamp black together with starch in a proportion materially to reduce the sensitivity of the explosive (eg, 10–30% of 300 mesh starch). H. Sauvage, USP 2076279 (1937) & CA 32, 3697 (1937).

Granular carbonaceous material such as "Bugbird carbon" is impregnated with a solution of a fire-proofing material such as Amm phosphate or Ca chloride and the treated material is used as a carrier for liq oxygen. G. B. Holderer, USP 2119050 (1938) & CA 32, 5630 (1938).

A carbonaceous absorbent is used together with at least 5% of phosphoric acid and at least 2% water, based on the weight of the dry carbonaceous material, serving to avoid undue flammability. A.R.T. Denues, USP 2297538 (1943) & CA 37, 1606 (1943).

The prepn of a liq oxygen explosive, especially useful in mining operations, is described. It is safe to handle and is characterized by high brisance and power. Activated or absorbant C (150–250 mesh), free of hydrophobic and of low-temperature volatile materials or impurities having high ignition points, is treated with phosphoric acid or a salt of phosphoric acid in a weight ratio of 2.5 to 18% of C, and sufficient water to form a slurry. The slurry is dried with agitation at 250–450°F, and the resulting product is then placed in fireproofed canvas bags and dipped in liq oxygen. The amount of oxygen taken up by each part of the treated C is 8.4 parts. This product can be ignited but not detonated by burning paper, and is not affected by rifle fire of 0.30 caliber soft nose bullets. It can be detonated with a Primacord detonator. L.P. Barlow, USP 2723188 (1955) & CA 50, 4510 (1956).

H. Sauvage, BritP 791930 (1958) & CA 52,
17716 (1958) claims preparation of combustible material (starch) for LOX in the form of compressed pellets using a press of at least 15kg/cm². The disruptive power of these exps compares very well with permanent expls.

F.L. Shea, Jr., USP 2872305 (1959) & CA 53, 7597 (1959) claims preparation of LOX by grinding a suitable bituminous material to a preferred particle size of 65–95% thru 200 mesh, and flash-calcing at >1150°F, preferably 1350–1650°F, with an oxidizing gas containing sufficient oxygen to oxidize all but about 10% of the volatile matter in the feed, which volatile matter should be ~15–20% of the feed. The product thus obtained is then immersed in liq oxygen to form the expl.

H. Sauvage, FrP 1112288 (1956) & CA 53, 8630 (1959) claims preparation of LOX for open-air shooting by the same processes as usual, but adjusting the sacking of the carbon black in such a way that CO is formed in the explosion. The brisance of these exps is claimed to be about 50% higher than that of PA.

J.P. Perdrizet, FrP 1114985 (1956) & CA 53, 15570 (1959) describes LOX which are insensitive to shock and have a prolonged life. They contain as a fuel a dry starch paste obtained from potatoes or manioc, which is free of fatty matter and has a grain size between 12 and 64 mesh/cm. In cartridges containing this type of filling and used in the vertical position, the very finely pulverized starch is divided into several compartments by horizontal and vertical divisions of impregnated paper or tissue, impermeable to liq oxygen, to diminish the height of the liq oxygen and thereby its flowing out. The filling is compressed in a higher or lower degree to give cakes of variable density, preferably having an apparent density of 0.45–0.65g/cc.

S.W. Martin, USP 2812246 (1957) & CA 52, 3346 (1958) claims prepns of impact-resistant LOX from an adsorbent carbon obtained by flash-oxidation-calcination of a finely divided swelling bituminous coal (with a volatile content of >12%) in a stream of oxygen-containing gas at 800° or more. An expl prep from this carbon was insensitive to impacts of >1200ft lb. It failed to detonate when burned in semi-confinement with a 0.25 inch orifice. Its vel of deton was 14050ft/sec.

J.P. Perdrizet, FrP 1132039 (1957) & CA 53, 18487 (1959) claims highly brisant, small-diameter, low-cost, shock-resistant LOX cartridges are made with fuels obtained from pulp residue from the extraction of starch from potatoes, and sugar from beets.

F.W. Brown, USP 2879149 (1959) & CA 53, 11840 (1959) claims prepns of a low-cost expl having high brisance and gas-forming ability but much lower fire and detonation sensitivity than conventional liq oxygen-carbon black expls by mixing 10–80% (preferably 20%) lampblack or carbon black with 90–20% AN and then saturating with liq oxygen. Such an expl has a rate of deton of 6000m/sec. Resistance to fire is substantially increased by incorporation of up to ~10% water based on the solids.

Andrew Hyslop, Jr., USP 2886424 (1959) & CA 53, 15570 (1959) describes processes for making mixts of liq oxygen and finely divided fibrous or spherical hydrocarbons (I) for use as expls. These mixts are less hazardous than similar prior-art mixts. The mixing chamber is cylindrical, open at the top, with a funnel-shaped bottom. Two nozzles, jets of atomized oxygen are made to intersect near the middle of the chamber. A downward directed spray of atomized liq (I) is congealed as spheres or strands on mixing with the oxygen. The mix of oxygen and congealed (I) is collected in a container below the congelation chamber. Hydraulic, diesel, lubricating oils, kerosene and gasoline may be used. The particles of (I) are preferably <0.01 inch in diameter. Liq nitrogen may be used to make the divided (I) for later mixing with liq oxygen. Water but not carbon tetrachloride, may be atomized into the chamber with 4 parts oil to produce less-sensitive expls.

C. McKinley, USP 2939778 (1960) & CA 54, 17888 (1960) claims prepns of a liq expl which is relatively safe to prepare, store and use under controlled conditions, by dissolving in liq oxygen a fuel, preferably methane or its admixture with small amounts of heavier, normally gaseous hydrocarbons; natural gas is an excellent fuel. Stoichiometric proportions are used for maximum disruptive effect.

Written by J. ROTH
LIQUID EXPLOSIVES

A liquid explosive, as the term implies, is an explosive substance that is liquid at ambient temp. Liquid Explosives (LE) can be pure compounds or mixtures of miscible compounds, including aqueous solutions. Explosives containing a mixture of liquid and solid phases, eg, the so-called Slurry Explosives, as well as Liquid Oxygen Explosives (LOX), are described in separate articles.

In the present article we will review the history, uses and characteristics of LE under the following headings:

1) History
2) Typical LE
3) Specialized Uses of LE
4) Detonation Phenomena in LE
5) Initiation Phenomena in LE
6) Brief Abstracts of Recent Publications on LE which were not specifically drawn upon in the discussion of the items above. Greater detail on specific LE will be found in Encyclo articles on the individual explosive, eg, under Ethylene Glycol Dinitrate (EGDN) (Vol 6, pp E259ff), Glycerol Trinitrate (NG) (Vol 6, pp G98ff), etc.

1) History. The first LE of practical importance was NG. It was discovered by an Italian chemist, Ascanio Sobrero, in 1846. The “taming” of NG by Alfred Nobel is described in the article on Dynamite in Vol 5, p D1586-R. Even today NG and its close chemical relative EGDN are the best known sensitizers for conventional Dynamites. In recent years, however, Slurry Explosives, whose liquid phase is primarily aqueous Ammonium Nitrate (AN), and which contain no NG or EGDN, have been steadily replacing Dynamites (Ref 32).

Late in the nineteenth century, Hermann Sprengel patented a series of simple oxidation-reduction mixtures for use in commercial blasting. These so-called Sprengel explosives typically consisted of concd nitric acid, or liquid NO₂, mixed with oil fuels such as nitrobenzene, carbon disulfide, petroleum, etc. They were intended to be mixed immediately before use. Because of handling difficulties Sprengel expls never became very popular (Refs 4 & 6).

Mixes of liquid NO₂ with liquid fuels are also known as Pancastites. They are very sen-
sitive to shock and must be handled most carefully after mixing (Ref 4). In the 1950's liquid $\text{N}_2\text{O}_4$/kerosene mixts were briefly used in strip mine blasting of overburden. Handling difficulties again led to the abandonment of these expls.

Use of LE in minefield clearance during WWII is described in section 3.

In recent years liq Nitromethane, NM, although rarely used as a practical expl, has been studied from the point of view of initiation and detonation theory probably more than any other expl, liq, solid or gas. Some of these studies are discussed in sections 4 & 5.

2) Typical Liquid Explosives. Probably the most important class of LE are the liq nitrate esters. The most important members of this family are NG (See Vol 6, p G98-R), EGDN (See Vol 6, p E259-R) & DEGDN (See Vol 5, p D1232-L). The mono esters, MeONO$_2$ and EtONO$_2$ (See Vol 6, p E143-R), are also expl but are relatively unimportant in both expl practice and theory.

Another class of LE are the Nitromethanes. As already mentioned, NM is probably the most thoroughly studied expl. Tetranitromethane, TNM, is expl by itself, and forms very powerful and very sensitive mixts with liq fuels (See below).

In recent years, many Fluoro derivatives have been studied as potential high energy constituents of proplnts. Many of these compounds are liquids and some are highly explosive. The Bs(2-Fluoro-2,2-Dinitroethyl) Formaldehyde, FEFO has been studied by the military and the AEC (Ref 30). Some theoretical studies of Bis-Difluoramo Alkanes have been reported (Refs 26 & 26c).

LE mixts abound although few are used in practical applications. Some of these are listed below:

Acenina is an equimolar mixture of HNO$_3$/H$_2$O/CH$_2$CN. It was used in a theoretical study by Davis et al (Ref 14).

Anilites are mixts of liq N$_2$O$_4$ and benzene. They are not used at present (See Vol 1, p A443-R).

Dithakite is 63/24/14 HNO$_3$/Nitrobenzene/H$_2$O. Cook used it in his investigation of "detonation plasma" (Refs 6, pp 190–94 & Ref 7).

LX-01, also called NTN, an all-purpose AEC LE, consists of 51.7/33.2/15.1 NM/TNM/1-Nitropropone (Ref 30, pp 18–25)

MEN-II, also called RX-01-AC, consists of 72.2/23.4/4.4 NM/MeOH/Ethylendiamine (Ref 30, pp 3–4). A similar mixt was used in WWII for minefield clearance (Ref 5).

Monomethylamine Nitrate solutions containing 10 to 15% water are used as sensitzers in Slurry Explosives. Very recently a tank car filled with such a so1n blew up in a railway yard in Wenatchee, Washington.

N$_2$O$_4$/fuel, usually in stoichiometric proportions, has been used in commercial blasting and in Anilites (Ref 2 & Vol 1, p A443-R).

PLX, a 95/5 mixt of NM and ethylenediamine, was used in clearing minefields (Ref 27).

TNM/fuel mixtures, such as stoichiometric mixts of TNM and hexane, benzene or nitrobenzene, have been used in specialized applications. They are extremely sensitive since several accidents have been reported (Ref 3). This great sensitivity may be associated with their very small critical diameter (See Vol 4, p D653-L). For example, the critical diameter of 87.5/12.5 TNM/NBz is <0.1mm as compared to ~1mm for NG (Ref 8a).

3) Specialized Uses of Liquid Explosives.

Liq NG, EGDN, or their mixts are sensitzers for Ammonium Nitrate, which is the main expl ingredient of modern Dynamites. Why they are so effective is still somewhat of a mystery. Possibly it is because both these liquids, and their mixts, are easy to initiate, and because, once initiated, they will propagate in small (1mm diameter) columns.

Both NG and EGDN are gelled by Nitrocellulose (NC) and form the matrix of water-resistant Gelatin Dynamites. The NG–NC gels are also the main constituents of double-base smokeless powders and of triple-base powders (which also contain Nitroglycerin). Nitroglycerin is also a constituent of many modern high-energy composite propellants.

The use of a conc aqueous soln of Monomethylamine Nitrate as a sensitizer for Slurry Explosives was mentioned in the preceding section.
The above constitute the main uses of LE, amounting to millions of pounds yearly. However, none of these uses is for the LE by itself, ie, solely as a liq. Neat NG has been used in "shooting" oil wells during the secondary recovery of oil. When oil production from a primary well began to lag, four secondary holes were drilled around the primary well so that all the holes, primary and secondary, were in the form of a "five spot". The secondary holes were then loaded with neat NG and shot. Water was then pumped into the secondary holes which forced oil into the primary hole by displacement. It was claimed that neat NG produced better rock fracturing and fissuring than Dynamites, and thus facilitated oil displacement by the water that penetrated into the cracked strata (Ref 1)

LE may be used for cutting breaks in a forest fire. This is done by stretching plastic tubing along the path where the break is desired, filling it with LE and detonating it. A path free of underbrush and foliage is thus cleared with very little danger of ignition of flammables along the path (Refs 21 & 23)

Minefield clearance by LE was studied during WWII. A system developed in these studies consisted of a number of rubber hoses, placed in parallel on a suspected minefield. They were then loaded with a liquid mixture of NM and ethylenediamine and detonated. A variation of this scheme was to gel the above mix (Santocel ARD proved to be a usable gelling agent), spread the gel over the suspected minefield and detonate it (Refs 5, 5a, 5b & 5c)

Conversely, a recent development is claimed to have produced a "liquid" land mine. Liquid Astrolite, a proprietary expl compon of the Explosive Corp of America, Issaquah, Washington (See Sect 6), is spread over the ground where it remains "active" for up to 4 days. This "Landmine" is not detectable by standard mine detectors. It is fired by a standard blasting cap (Refs 24 & 25)

An interesting use of LE (usually NM) is in the Explosive Gas Gun (See Vol 6, p E419-R)

4) Detonation Phenomena in Liquid Explosives. Detonation parameters of LE are similar to those of solid expls of comparable density and energy content. This is shown in Table 1 which is to be compared with Table 3-5 of Ref 29

Nevertheless, LE detonations exhibit certain phenomena not usually found in comparable solid expls. Many of the unique characteristics of LE detonations are ascribable to their homogeneity, whereas most solid expls are heterogeneous. These effects, peculiar to LE, will now be described in some detail

A) Similarities in Detonation in Liquid and Gaseous Explosives

Above we have stressed that detonation parameters of LE and solid explosives are comparable. Nevertheless, LE often exhibit detonation phenomena that resemble gaseous detonations more than detonations of solid expls.

A simplistic view, but one that has some merit, rationalizes the "dual" nature of LE as follows: the resemblance to solid expls is due to the similar densities of most LE and solid explosives; the similarities with gaseous expls are due to the homogeneity of LE and gaseous expls (solid expls except for perfect single crystals are heterogeneous)

An excellent review (with 27 references) of the similarities of LE and gaseous expls is given by Dremin & Rozanov (Ref 17a). They point out the following main similarities:

a) Detonation velocity changes insignificantly with charge diameter until a critical diameter is reached below which propagation of detonation ceases abruptly

b) The critical diameter depends strongly on the energy content of the expl. Thus dilution of the expl with inert additives lowers the critical diameter sharply

c) The detonation is attenuated upon passage from a narrow container into a large volume, eg, in passing from a cylinder into an expanding cone

Dremin & Rozanov suggest that these similarities can be rationalized on the basis of a similar reaction mechanism for both LE and
### Table 1

Experimental Detonation Properties of Some Liquid Explosives (a)

<table>
<thead>
<tr>
<th>LE</th>
<th>Density g/cc</th>
<th>Detonation Velocity D (m/sec)</th>
<th>Chapman-Jouguet Pressure $P_{CJ}$ (kb)</th>
<th>Heat of Detonation $\Delta H_{det}$ (b) Kcal/g</th>
<th>Chapman-Jouguet Temperature of Detonation $T_{CJ}$ (°K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LX-01</td>
<td>1.24</td>
<td>6840 (30, pp 18–25)</td>
<td>156 (30, pp 18–25)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>NG</td>
<td>1.60</td>
<td>7700 (30, pp 18–45)</td>
<td>253 (30, pp 18–45)</td>
<td>1.48 (d)</td>
<td>4000 (9)</td>
</tr>
<tr>
<td>NM</td>
<td>1.16</td>
<td>6370 (14)</td>
<td>141 (10)</td>
<td>1.23 (30, pp 18–47)</td>
<td>3700 (9)</td>
</tr>
<tr>
<td>55.5/44.5 NM/TNM</td>
<td>1.31</td>
<td>6880 (12)</td>
<td>156 (12)</td>
<td>-</td>
<td>3750 (12)</td>
</tr>
<tr>
<td>TNM</td>
<td>1.64</td>
<td>6360 (12)</td>
<td>159 (12)</td>
<td>-</td>
<td>4650 (g) (10)</td>
</tr>
<tr>
<td>TNT (liq)</td>
<td>1.45 (e)</td>
<td>6590 (e) (14)</td>
<td>182 (e) (14)</td>
<td>1.11 (f)</td>
<td>3030 (12)</td>
</tr>
</tbody>
</table>

(a) Bracketted numbers are Refs
(b) Highly confined samples in a calorimeter; H₂O (liq)
(c) 51.7/33.2/15.1 NM/TNM/1-Nitropropane
(d) Computed; should be reliable for this nearly oxygen-balanced explosive
(e) At 93°C
(f) For solid TNT with 0.02 kcal/g correction for heat of fusion
(g) For a 50/50 NM/TNM mixture

Gaseous expls. We will examine this suggestion in a later section.

B) Experimental Detonation Temperatures

The last column of Table 1 lists some experimental detonation temperatures ($T_{CJ}$) obtained by optical methods. Although there is considerable disagreement between measurements made by different investigators, these $T_{CJ}$ values are probably the best that are now available. Detonation temperature is a very important parameter in detonation theory, inasmuch as it provides: 1) the best test for the validity of an equation of state of the detonation products (See Vol 4, pp D268–298) and 2) insight into the chemical reaction rates in the detonation process.

Most of the many equations of state (See Vol 4, pp D268ff) proposed for detonation products are capable of providing computed detonation velocities, $D$, in good accord with experiment. Indeed, many authors use measured $D$'s as a normalization parameter or adjustment factor for their equation of state. Agreement between computed and measured detonation pressure is a better test of the validity of an equation of state, but this test is still far less sensitive than that provided by agreement between computed and measured $T_{CJ}$.

Usually detonation temps are high enough to make reaction rates (extrapolated from data obtained at much lower temps) at $T_{CJ}$ very fast – possibly so fast as to be no longer rate controlling (See Kinetics in this Vol). Thus, from the point of view of chemical kinetics, temps in the pre-detonation shock are of greater significance than $T_{CJ}$ (See Sect 5).

There are conceptual difficulties in the optional methods of measuring $T_{CJ}$ for LE. However, these difficulties are much less severe than for solid heterogeneous expls for which
“Detonation Light” may have little relationship to the detonation process (Ref 16). The main problems in T_cj measurements for LE arise from the fact that: 1) the detonation zone is “screened” by a highly absorbent shock zone ahead of it; 2) the radiation received by the detector comes primarily from the foremost front layers of the detonation zone and may not necessarily correspond to T_cj which is an equilibrium temp and 3) the radiation may not be blackbody or graybody. Nevertheless, these “detonation temperatures” for LE are the best we have at present. Incidentally, the values listed in Table 1 appear to agree better with the BKW equation of state (Ref 12) than with some other equations of state.

C) Low Velocity Detonation (LVD). LVD is a phenomenon commonly encountered with LE. LVD is a reaction wave phenomenon that propagates in LE at a constant velocity just slightly greater than the speed of sound in the unreacted liquid. Thus a typical LVD propagates at around 2000 m/sec whereas a C-J or High Velocity Detonation (HVD) in LE propagates at 6000 to 8000 m/sec (See Table 1). Both LVD and HVD can occur with the same LE depending on confinement, charge diameter and conditions of initiation. LVD can turn into HVD but not conversely. A stable LVD requires that the speed of sound in the container wall exceeds the LVD velocity.

The commonly accepted model for LVD is the so-called cavitation model. It is believed that precursor shocks, propagating thru the charge container, produce cavitation zones in the liq ahead of the LVD. The vapor bubbles (cavities) then act as sites for ignition and growth of chemical reaction to support the LVD wave (Ref 15).

Chalken (Ref 28) has shown that the cavitation model is consistent with the classical C-J picture of detonation, provided that additional constraints, due to the several rate processes that take place in LVD, are imposed on the classical treatment. In essence, he has shown that not all Hugoniot adiabatic states can be accessible from a given initial state, if the precursor shock acts to couple rate processes ahead of the reaction shock front with rate processes behind this front. The solution of the Rankine-Hugoniot (classical detonation) equations for a steady LVD then becomes a minimization problem where the LVD velocity is the minimum velocity consistent with the constraints imposed by the coupled rate processes.

D) Inhomogeneity of the Detonation Wave Front in LE. The studies of Shchelkin (Ref 8), White (Ref 11) and others have demonstrated that the detonation in most gaseous explosives is unstable and the detonation wave front has a turbulent structure. The investigations of Dremin and co-workers (Ref 13a, 17a, 17b, 20 & 22), Mallory (Ref 19), Urtiew & Kusubov (Ref 26b), Watson (Ref 26d), Persson (Ref 26c), Seely et al (Ref 26) & Cook and co-workers (Ref 32) have shown similar effects in the detonation wave fronts of LE. However, inhomogeneities in the wave front were observed only for rather “insensitive” expls, particularly in expls diluted with inert additives. This is understandable if, as suggested by Shchelkin (Ref 8), the size of the inhomogeneities is determined by the reaction time of the expl under classical (C-J) conditions. For “sensitive” LE (eg, NG), these reaction times are short and inhomogeneities are too small to be resolved by instrumentation now available. The problem of wave front instability in all LE is still unresolved. There may be certain LE that behave classically, ie, their detonation fronts are stable, uniform and one dimensional (Ref 18).

The detonation process, at least in “insensitive” LE, can be visualized as follows (Ref 17a): Microinhomogeneities in the LE (eg, fluctuation in density or composition) result in non-uniform reaction rates in the shocked LE. Because reaction rates are so strongly dependent on temp, these perturbations do not attenuate and eventually reach the shock front of the detonation wave and “bend” it, thus creating oblique shocks at its leading edge. Oblique or transverse shocks lead to the formation of triple shock configurations, and to an increase in the temp and pressure behind the oblique shocks over that obtainable behind smooth shocks. This, in turn, makes for more favorable conditions for the initiation of further reaction. At triple wave intersections, conditions for reaction are particularly favorable. Thus the wave front becomes a complex three-dimensional cellular structure, quite different from the classical one-dimensional uniform wave front.
E) Failure Diameter Theory. Closely associated with the inhomogeneity of the detonation front in LE is the abrupt change from stable constant velocity detonation to detonation failure as the diameter of the LE is decreased. Typically the detonation velocity of many LE will change very slowly as charge diameter is decreased, and at a further slight decrease in diameter to \( d_f \), the failure diameter, detonation failure occurs abruptly, even though the detonation velocity has dropped only about 1% below that at infinite diameter. This phenomenon is inexplicable in terms of the classical theories of the diameter effect on detonation velocity (See Vol 4, pp D641–43). Dremin and co-workers (Refs 13a, 17b & 22) have developed a theory of failure diameter for LE which have non-uniform detonation wave fronts. In its simplest terms this theory states that detonation failure occurs when chemical reactions cease because oblique (transverse) wave interactions disappear. As discussed in the preceding section, most of the chemical reaction for these LE occurs at sites of transverse wave interaction, but it is entirely possible that some interaction sites will not produce chemical reaction. Such no-reaction regions propagate as dark waves. Obviously if the dark waves cover the entire cross-section of the charge, transverse wave interactions cease, and, unless the LE is “sensitive” enough to sustain detonation in a “smooth” shock, failure occurs. However, if the dark waves cover only part of the charge cross-section, detonation proceeds without change in velocity. The mathematical consequences of this theory are given in Ref 17b and further expanded by Enig & Petrone (Ref 26a). Although these quantitative treatments confirm the theory, they are not very useful for \textit{a priori} computations of \( d_f \) because they involve parameters that are not usually available for new LE or LE mixtures.

That \( d_f \) for “insensitive” LE is associated with dark waves is shown by several experimental studies (Refs 26, 26b & 26d) in addition to the Dremin refs cited above.

5) Initiation of Detonation in LE. The initiation of detonation in LE occurs either heterogeneously or homogeneously. Impact initiation of powdery solid expls is a prime example of heterogeneous initiation. It is described in detail in the article on Impact, Initiation of Explosion By (pp 13S-Rff). Another example of heterogeneous initiation is the shock initiation of LE containing purposely introduced inhomogeneities such as air bubbles, Al or glass beads, etc. Detonations initiated by such hydrodynamic hot spots are described in the article on Hot Spots (pp H170-Lff). The uniform bulk heating of LE is an obvious example of homogeneous initiation, but it is little different from the bulk heating of solid expls which has been described under ignition (pp 111ff). Several related articles are in Vol 4 such as Decomposition, Thermal; Deflagration; Detonation, Thermal Theories & Thermochemistry and in Vol 2 under Burning and Burning Characteristics. In what follows we will examine in detail the shock initiation of homogeneous (no bubbles etc) LE, since this is an initiation process unique to LE, although it can also occur in perfect single crystals.

The classical paper on the homogeneous initiation of LE is that of Campbell et al (Ref 11a). Their views have since received considerable international support (Refs 17, 20, 26e & 31), but have been criticized by Cook (Ref 32, pp 3–4). In this writer’s opinion there is considerable confusion and possibly a misreading of the original paper in Cook’s criticisms, and they will be omitted, although some of Cook’s ideas on the initiation of LE will be presented later.
tonation travels at a velocity \( D^* > D_0 \) in the shock-compressed LE until it catches up with the original shock. At this point \((d, t_1 \text{ in the figure})\) there is an overdrive in the detonation of the virgin LE, but this decays fairly rapidly to a steady detonation propagating at a velocity \( D_0 \). The overdrive occurs because the particle velocity associated with \( D^* \) is higher than that of a normal detonation. Note that the "hyper-velocity" detonation \( D^* \) travels a shorter distance than \( d \) because the initial shock has imparted a particle velocity \( U_p \) to the interface which has consequently moved away from \( x = 0 \) over a distance \( U_p t^* \).

In the original Campbell et al paper, NM, molten TNT, molten DINA & Dithekite were the LE examined. Smear cameras, framing cameras & ionization pin pairs were the diagnostic tools. Later studies, by Travis (Ref 15a) using electrical transducers, Dremin (Ref 17), using electromagnetic measurements of particle velocity, and Persson & Sjölin (Ref 26c), using photomultipliers to record detonation luminosity, are in excellent accord with the Campbell et al findings. Subsequent studies by Dremin (Ref 20) and Persson (Ref 26c) showed that NG and EGDN also follow this initiation mechanism. Berke et al (Ref 26e) also found that several liq fluoramino alkanes behave analogously to NM, NG etc. Most of the original and later studies were made with NM as the principal LE. Consequently all our subsequent discussion refers to NM.

Table 2 gives average values of the NM parameters measured by Campbell et al (Ref 11a).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shock Velocity ( U_S ) ((\text{mm/μsec}))</td>
<td>4.5</td>
</tr>
<tr>
<td>Particle Velocity ( U_p ) ((\text{mm/μsec}))</td>
<td>1.7</td>
</tr>
<tr>
<td>Input Pressure ((\text{kbar}))</td>
<td>86</td>
</tr>
<tr>
<td>Detonation Velocity in Compressed NM, ( D^* ) ((\text{mm/μsec}))</td>
<td>10.4</td>
</tr>
<tr>
<td>Detonation Velocity Steady State, ( D_0 ) ((\text{mm/μsec}))</td>
<td>6.26</td>
</tr>
<tr>
<td>Induction Time, ( t^* ) ((\text{μsec}))</td>
<td>( \approx 2 )</td>
</tr>
</tbody>
</table>

The induction time \( t^* \) is of particular interest, since it can be compared to the induction time computed for an adiabatic thermal explosion (See Ref 6, pp 173–74 or Eq 6 of Article on Hot Spots, p H172-R) to provide a check on the correctness of the supposition that the input shock generates a thermal explosion (at the shock entry face). Unfortunately, an exact quantitative treatment of the induction times of shock-generated thermal explosions suffers from: a) uncertainty of the shock-generated temperature in the LE and b) uncertainty in the Arrhenius kinetic parameters (activation energy and pre-exponential factor) (See Kinetics in this Vol).

Campbell et al (Ref 11a) claim to have obtained reasonable agreement between observed and computed induction times. They computed their shock-generated temps on the basis of constant specific heat, i.e., no increase in \( C_V \) with an increase in temp. It is to be expected that \( C_V \) for liquids increases with increasing temp and this expectation has been confirmed for NM (Ref 31). Thus the Campbell et al temperatures of about 1170K for the conditions shown in Table 2 are too high. Based on recent studies (Ref 31) these temps should be about 1050K. This means that a lower activation energy than the used by Campbell et al is necessary to get agreement between observed and computed induction times for NM shock-heated to 1050K. Although the above discussion suggests that quantitative agreement between experiment and theory is still to be achieved, the qualitative model for the shock initiation of LE appears to be on solid ground.

Cook and his co-workers have championed a different model for the shock initiation of LE (Refs 6, pp 173–94 & 32, pp 292–96). In essence, this model states that initiation occurs when a shocked region of LE becomes "thermally superconductive" (as a result of rising temp due to partial decomp of the shocked LE) and a "heat pulse" flashes across the shocked LE and catches up with the original shock front. As described in the article on Heatpulse (p H59-L), alternate explanations are possible for some of the observations that Cook considers to be the main experimental support for his "heat pulse". Similarly, Dremin et al (Ref 17) have suggested an alternate ex-
6) Brief Abstracts of Recent Publications on LE:

a) Effect of Pressure on the Burning of LE. An abrupt change in the burning of some LE occurs as pressure on the LE is increased. Two alternate explanations for this effect are: 1) gas phase reactions are moved closer to the liq surface by increasing pressure, thus transferring more heat to the LE and 2) increase in the burning rate generates turbulence in the combustion front which then increases the heat exchange between combustion gases and the LE. Experiments with EGDN and EGDN gelled with a small amount of NC favor explanation (2)


b) Initiation of LE by Gas Phase Detonation. Detonation of stoichiometric methane-oxygen gas mixes was used to initiate liquid TNM/benzene mixtures. For 1.5/1 (by volume) TNM/benzene, initiation occurred only at initial gas pressures above 2 atm. Initiation delay decreased with increasing initial gas pressure. For 4/1 (by volume) TNM/benzene (stoichiometric) the critical initial gas pressure was 0.66 atm. The influence of initial pressure is explained in terms of more efficient heat transfer from the gas detonation to the LE, but there are compensating effects and the surface temp of the LE varies only mildly with initial pressure

Ref: V.E. Gordeev et al, FizGorenlya i Vzryva 1 (2), 12 (1965) & CA 64, 1894 (1966)

c) A Generalized Shock Hugoniot for Organic Liquids. The shock compressibility (Hugoniot) (See pp H179-Lff) of organic liquids, including several LE, is expressed in the following general form:

\[ U = 1.2C_0 + 1.7\mu \]

where U is shock velocity, \( C_0 \) is ambient sound velocity in the liq and \( \mu \) is particle velocity

Ref: I.M. Voskoboinikov et al, FizGorenlya i Vzryva 3 (4), 585 (1967) & CA 69, 37522 (1968)

d) Initiation of LE by High Intensity Light. Stoichiometric mixes of TNM/benzene (See abstract b) were initiated by the radiation of an argon flash bomb. Chemical decompn of NG was observed under these conditions but no detonation. The possibility of light, produced by the shock compression of air pockets in porous exps, contributing to the initiation and propagation of detonation in porous exps is discussed


e) Deflagration to Detonation Transition in LE. High-speed photography was used to study the growth of explosion in various liq exps. The liquids investigated included NG, NM/HNO3 mixts, H2O2/Ethanol mixts and DEGN. In one experimental arrangement, a thin film of expn was confined between transparent plates. Deflagration at the center of the film was initiated by the rapid discharge of a condenser across a spark gap. The study allowed the growth of the burning and the transfer to detonation to be photographed in detail


f) LE consisting of hydrazine, Hydrazine Nitrate, water (with & without added ammonia or AN) & a thickening agent is claimed. This appears to be one of the “Astrolite” exps

Ref: R.M. Bridgeforth et al, USP 3523047 & CA 74, 14701 (1971)

g) Calculated Shock Temperatures for LE. A modification of Walsh-Christian model for metals was used in calculating shock temps in NM, liq TNT and four liquid Bis(Difluoromino) Alkanes. In this modification the heat capacity increases with temp rather than staying constant as in the Walsh-Christian model


Written by J. ROTH


LIQUID PROPELLANTS

Definitions & Overview. Liquid Propellants (LP) are liq substances that can be transformed into gases (usually hot) which act as driving jets in a propulsion system. Propulsion systems can be divided into rockets and jet engines. The forward thrust of both rockets and jet engines is generated by the rearward ejection of a fluid jet thru nozzles mounted on the rocket or jet engine (Ref 12, pp 439–41). In a rocket this fluid jet originates entirely from containers (tanks) within the rocket. Jet engines, on the other hand, ingest the surrounding medium (air or water) to produce their exhaust jet. Thus rockets can function in a vacuum (space) whereas turbojets, ramjets etc., cannot. Since Jets are described in this Vol (See under Jet Fuels, pp J68-R ff) this article deals primarily with rockets and liquid rocket propellants.

It is usually understood that a rocket motor is that part of the propulsion system in which the propellants are transformed into the exhaust jet, while a rocket engine is the entire system, ie, the rocket motor, the propellant containers, pumps, etc. In conventional solid propellant rockets, the motor and the engine happen to be the same piece of apparatus, but this is not so in liq propellant rockets.

Liquid Propellants, sometimes called Propergols if used for rocket propulsion (Ref 23), are divided into two classes: monopropellants and bipropellants. A monopropellant is a single substance or a homogeneous compatible mix that can be caused to react in the rocket motor (combustion changer) to generate the gases that form the exhaust jet. A bipropellant consists of two liq reactants — usually a fuel and an oxidizer. The gas-forming process in a monopropellant is initiated by heat/or catalysts.
These agents can also initiate the gas formation of bipropellants after the components are mixed. Certain bipropellants can, however, self-ignite merely upon mixing. These are called Hypergolic Propellants or Hypergols (See Hypergolic Propellants in this Vol, pp H254-L to H259-R).

Levy (quoted in Ref 23) also suggested the following additional designations: CATergols are LP that are decomposed by the action of catalysis; Monergols are homogeneous liquid bipropellants; Lithergols are heterogenous bipropellants, eg, solid C & liquid oxygen.

LP are often classified into storable and cryogenic liquids. The former, as implied, are stable liquids under ambient operating conditions. The latter require a liquefying plant as they are gases under ambient conditions.

A universally accepted measure of the effectiveness of LP is the specific impulse, usually designated by $I_{sp}$. It is defined by:

$$I_{sp} = \frac{c}{g}$$

where

- $V_e$ = exhaust jet velocity
- $p_e$ = static pressure at the exhaust
- $p_a$ = ambient pressure
- $A_e$ = area of exhaust nozzle
- $m$ = rate of exhaust mass flow
- $g$ = acceleration of gravity

The units of $I_{sp}$ are (lb·sec)/lb or, as commonly used, sec (by cancelling the lbs in the numerator and denominator). For further details on the measurement & computation of $I_{sp}$ see Ref 12, pp 440–4 & 464–71.


General references on the subject of LP are: Refs 7, 9–15, 17, 18, 21, 23 & 27–29.

In the present article we will review:

1) LP History

2) Representative LP Compositions

3) Representative LP Applications

4) Performance Characteristics of LP

5) Sensitivity and Hazards of LP

1) History. An excellent overview of the history of liquid propellants, LP, and propulsion systems using LP is given by Malina (Ref 12, pp 10–23) and Truax (Ref 12, pp 23–25). In what follows we draw heavily on this overview.

The history of LP is conveniently divided into three periods: Pre-WWII, WWII & Post-WWII.

Pre-WWII. The use of LP for space travel appears to have been first suggested by Ziolkowski in 1898, but not published until 1903 (Ref 1). He also suggested the "piggy-back" launch, ie, a rocket launched from another rocket, although this principle had been used much earlier in fireworks displays.

The subject of LP remained quiescent early in the 20th century and thru WWI. In 1928, Esnault-Pelterie (Ref 2) lectured on LP combustion problems. Between 1930 & 1933 the Verein für Raumschiffahrt (Society for Space Travel) experimented with LP engines using combinations of liq oxygen and gasoline or aq alcohol (Ref 12, pp 10–23). This society was the precursor of the group that developed the V-2 rocket during WWII.

The American Rocket Society began experimenting with LP in 1932, using primarily liq oxygen-gasoline systems. Their work was later summarized by Wyld (Ref 6). In 1941 several members of this society formed Reaction Motors, Inc to produce the liq oxygen-alcohol engine for the Bell X-1 supersonic aircraft. In 1936 Goddard summarized his studies of LP (Ref 3). His studies dealt more with combustion chamber design and functioning than with specific LP.

The Guggenheim Aeronautical Laboratory was started in the same year at the California Institute of Technology for the purpose of studying design fundamentals of solid and liq propulsion systems. A year later, Malina (Ref 4) published a paper on the characteristics of LP motors.

These developments up to 1940 are well summarized by Malina (Ref 12, pp 10–23) and we quote:

"The literature published up to 1940 shows that the basic problems of designing successful liquid propellant engines were correctly stated,
at least in major outline. The properties of a number of propellants were fairly well known, especially those combinations using liquid oxygen as the oxidizer. In connection with the rocket motor, the thermodynamics of the combustion process and of the flow of the gaseous products thru the exhaust nozzle to produce thrust, were understood; and a beginning had been made in solving the cooling problem, and the preliminary stages of development of suitable systems of propellant supply had also been reached. It may be noted that the greatest advance to be recorded in this article was the mechanical design of the complete engine, although a number of basic discoveries were made, especially in the domain of propellants, and the theory for guiding design was greatly expanded.”

WWII. From 1932 onward, the study of LP and LP engines proceeded under a cloak of secrecy in Germany. Much of this effort was concentrated at an Army establishment at Peenemunde. This work culminated in the development of the V-2 rocket (originally called the A-4) in 1942. The LP of the V-2 consisted of liq oxygen mixed with 75/25 ethanol/water. Mixing pumps were driven by an impulse turbine powered by steam from a hydrogen peroxide generator. This rocket motor delivered a 55,000 lb thrust, at sea level, for 65 seconds to give an average specific impulse of 223 lb-sec/lb. The original V-2 traveled a distance of 120 miles.

Similar LP systems were also developed and used at Peenemunde for jet-assisted take-off for aeroplanes (See JATO in this Vol, p.67-L).

The Germans also developed LP systems based on hydrogen peroxide both by itself and mixed with fuels. The specific impulse of decomposing 80% hydrogen peroxide (“cold” engine) was a rather low 155 lb-sec/lb. This was raised to 210 lb-sec/lb by burning hydrogen peroxide with a fuel mixture of 30/57/13 hydrazine hydrate/methanol/water. Mixtures of hydrogen peroxide and hydrazine hydrate are hypergolic (Self-igniting — See Hypergolic Propellants (pp H254-L to H259-R) and Hydrazine, Its Salts and Derivatives, pp H190-L to H206-R, in this Vol. These LP were used mostly in turbo pumps for Jet aircraft.

Work on LP engines using nitric acid as the oxidizer was carried out at the Bayerische Motoren Werke (BMW). By 1945 some 6000 fuel-nitric acid mixtures had been tested. Some of these were hypergolic.

During WWII LP research in the USA was concentrated at the Jet Propulsion Laboratory of the California Institute of Technology, Aerojet Engineering Corp, Reaction Motors, Inc, and the Navy Bureau of Aeronautics. These efforts are summarized in books by Zucrow (Ref 7) and Sutton (Ref 9).

One of the main LP developments of the Jet Propulsion Lab efforts, later commercialized by Aerojet, was the nitric acid-aniline JATO engine. Monopropellant engines were also investigated.

Liquid oxygen-alcohol propulants were studied at Reaction Motors, and resulted in the 1500 N4C engine, which powered the experimental X-1 supersonic airplane (Ref 5).

The Bureau of Aeronautics investigated nitric acid-aniline propulants for JATO's for large flying boats such as the PBY-2 (Ref 8). They also developed a prototype engine for the Gorgon, a radio-controlled winged missile. The propellant for Gorgon was a combination of mixed nitric-sulfuric acids and monooctylamine.

Post-WWII. After WWII most LP R&D shifted from Germany to the USA and the USSR. Technical progress since 1945 in LP for rockets can be characterized by a host of minor improvements rather than major advances, and some spectacular applications of LP in rockets for military use and for space exploration, eg, in ICBM's (Intercontinental Ballistic Missiles) and in USA & USSR Moon, Mars & Venus “shots”. Thus the major developments since WWII have been primarily in the application of existing, though improved, propulsion systems to such highly publicized rockets as Sputnik, Vanguard, Soyuz, Saturn, Vostok, Apollo, Explorer, etc.

Efforts in LP per se have been largely restricted to improving performance by increasing the combustion pressure, and the developing and testing of “exotic” mono and bipropellant, such as the NF compounds and Fluorine-NH₃ and Fluorine-H₂ bipropellants (See Sect 2). So far there is no mention in the open literature of the practical use of “exotic” propellants (Ref 30, p 312).

2) Representative LP Compositions. Compared to the vast number of possible or even
<table>
<thead>
<tr>
<th>Constituents</th>
<th>Oxidant: Combustible Ratio</th>
<th>Specific Gravity, g/cc</th>
<th>Specific Impulse, sec</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>100% Nitric acid</td>
<td>Turpentine</td>
<td>4.4</td>
<td>-</td>
<td>221</td>
</tr>
<tr>
<td>Fuming Nitric acid (FNA)</td>
<td>Ethanol</td>
<td>2.5</td>
<td>-</td>
<td>219</td>
</tr>
<tr>
<td>FNA</td>
<td>Aniline</td>
<td>3.0</td>
<td>-</td>
<td>221</td>
</tr>
<tr>
<td>FNA</td>
<td>Ammonia</td>
<td>2.2</td>
<td>-</td>
<td>225</td>
</tr>
<tr>
<td>FNA</td>
<td>JP-4</td>
<td>-</td>
<td>-</td>
<td>225</td>
</tr>
<tr>
<td>99% Hydrogen peroxide</td>
<td>Ethanol</td>
<td>4.0</td>
<td>-</td>
<td>230</td>
</tr>
<tr>
<td>99% Hydrogen peroxide</td>
<td>JP-4</td>
<td>6.5</td>
<td>-</td>
<td>233</td>
</tr>
<tr>
<td>99% Hydrogen peroxide</td>
<td>Hydrazine</td>
<td>-</td>
<td>-</td>
<td>245</td>
</tr>
<tr>
<td>Liquid oxygen (LOX)</td>
<td>Ethanol</td>
<td>1.5</td>
<td>0.97</td>
<td>242</td>
</tr>
<tr>
<td>LOX</td>
<td>JP-4</td>
<td>2.2</td>
<td>1.02</td>
<td>248</td>
</tr>
<tr>
<td>LOX</td>
<td>Turpentine</td>
<td>2.2</td>
<td>-</td>
<td>249</td>
</tr>
<tr>
<td>LOX</td>
<td>Ammonia</td>
<td>1.3</td>
<td>-</td>
<td>250</td>
</tr>
<tr>
<td>Hydrogen peroxide</td>
<td>Nitromethane</td>
<td>-</td>
<td>-</td>
<td>227</td>
</tr>
<tr>
<td>N₂O₄</td>
<td>Hydrazine</td>
<td>-</td>
<td>-</td>
<td>249</td>
</tr>
<tr>
<td>N₂O₄</td>
<td>Hydrogen</td>
<td>11.5</td>
<td>0.565</td>
<td>279</td>
</tr>
<tr>
<td>FNA</td>
<td>Hydrogen</td>
<td>12.6</td>
<td>0.60</td>
<td>298</td>
</tr>
<tr>
<td>LOX</td>
<td>Hydrogen</td>
<td>2.9</td>
<td>0.23</td>
<td>345</td>
</tr>
<tr>
<td>70% LOX</td>
<td></td>
<td></td>
<td></td>
<td>So-called</td>
</tr>
<tr>
<td>30% Ozone</td>
<td></td>
<td></td>
<td></td>
<td>“Zip-fuels”</td>
</tr>
<tr>
<td>100% Ozone</td>
<td></td>
<td></td>
<td></td>
<td>(high energy fuels</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>in the USA)</td>
</tr>
<tr>
<td>100% Ozone</td>
<td>Ammonia</td>
<td>1.13</td>
<td>-</td>
<td>267</td>
</tr>
<tr>
<td>100% Ozone</td>
<td>Hydrazine</td>
<td>0.63</td>
<td>-</td>
<td>277</td>
</tr>
<tr>
<td>Fluorine</td>
<td>JP-4</td>
<td>2.6</td>
<td>-</td>
<td>265</td>
</tr>
<tr>
<td>Fluorine</td>
<td>Ammonia</td>
<td>2.6</td>
<td>-</td>
<td>288</td>
</tr>
<tr>
<td>Fluorine</td>
<td>Diborane</td>
<td>5.0</td>
<td>-</td>
<td>291</td>
</tr>
<tr>
<td>Fluorine</td>
<td>(B₂H₆)</td>
<td>2.37</td>
<td>-</td>
<td>296</td>
</tr>
<tr>
<td>Fluorine</td>
<td>Methanol</td>
<td>1.98</td>
<td>-</td>
<td>298</td>
</tr>
<tr>
<td>Fluorine</td>
<td>Hydrazine</td>
<td>4.5</td>
<td>-</td>
<td>352</td>
</tr>
<tr>
<td>Fluorine</td>
<td>Hydrogen</td>
<td>9.4</td>
<td>0.46</td>
<td>371</td>
</tr>
<tr>
<td>100% Ozone</td>
<td>Hydrogen</td>
<td>3.2</td>
<td>-</td>
<td>369</td>
</tr>
<tr>
<td>100% Ozone</td>
<td>Hydrogen</td>
<td>2.65</td>
<td>0.23</td>
<td>373</td>
</tr>
</tbody>
</table>

practical biproplnt compositions, very few monopropuls have been studied and even fewer are used. The following monopropuls have been investigated:

Hydrogen Peroxide, Hydrazine, Nitromethane, sym-Dinitromethane and Methyl Nitrate in ethanol (Refs 10 & 23). Of these only Hydrazine and Hydrogen Peroxide have been used and are being used in practical propulsion systems

As stated above, many oxidizer-fuel combinations are usable as biproplnts. However, the number of usable oxidizers is fairly limited. Storable oxidizers include: nitric acid (with and without added NO₂), Hydrogen Peroxide, perchloric acid, Tetranitromethane, N₂O₄, & ClF₃. Of these only nitric acid and Hydrogen Peroxide are of practical use. Perchloric acid or Tetranitromethane are too explosive when mixed with fuels, and N₂O₄ or ClF₃ are not really storable, as they boil at 22° and 12° respectively (Refs 10 & 23)

The most commonly used cryogenic oxidizer is liquid oxygen (LO). Other cryogenic oxidizers
include: \( \text{N}_2\text{O}_4 \), ozone, fluorine, chlorine-trifluoride (\( \text{ClF}_3 \)), nitrogen tetrafluoride (\( \text{NF}_4 \)), bromine pentafluoride (\( \text{BrF}_5 \)), perchloryl fluoride (\( \text{ClO}_2\text{F} \)), fluorine monoxide (\( \text{F}_2\text{O} \)), and fluorine dioxide (\( \text{F}_2\text{O}_2 \)) (Refs 10, 23 & 28)

Vast numbers of fuels may be combined with the above oxidizers to form bipropellant systems. Many hypergolic bipropellant systems are listed in this Vol. pp H254-L to H259-1, under Hypergolic Propellants. The following tabulation (Table 72, p 317 of Ref 23) lists several “common” bipropellants.

Other, generally more “exotic” bipropellants are given in Table 10.2, p. 165 of Ref 28 which is shown as Table 2 below:

Table 2

<table>
<thead>
<tr>
<th>Fuel</th>
<th>Oxidant</th>
<th>Oxidant/fuel ratio</th>
<th>( I_5 ) (sec)</th>
<th>( \Delta H_f^{298,0} ) (kcal/mole)</th>
<th>( T_C ) (°K)</th>
<th>( T_E ) (°K)</th>
<th>( M_E ) (&gt; 5% of total)</th>
<th>Products at ( T_E )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{H}_2 )</td>
<td>( \text{O}_2 )</td>
<td>4:00</td>
<td>391</td>
<td>0</td>
<td>2980</td>
<td>1350</td>
<td>10:08</td>
<td>( \text{H}_2\text{O}, \text{H}_2 )</td>
</tr>
<tr>
<td>( \text{B}_2\text{H}_6 )</td>
<td>( \text{O}_2 )</td>
<td>2:00</td>
<td>344</td>
<td>7:5</td>
<td>3846</td>
<td>2592</td>
<td>20:32</td>
<td>( \text{H}_2, \text{HBO}_2, \text{B}_2\text{O}_3 )</td>
</tr>
<tr>
<td>( \text{C}_2\text{H}_6 )</td>
<td>( \text{O}_2 )</td>
<td>1:80</td>
<td>327</td>
<td>54:2</td>
<td>4172</td>
<td>2600</td>
<td>24:07</td>
<td>( \text{CO}, \text{H}_2\text{O}, \text{H}_2, \text{CO}_2 )</td>
</tr>
<tr>
<td>( \text{N}_2\text{H}_4 )</td>
<td>( \text{O}_2 )</td>
<td>0:90</td>
<td>313</td>
<td>22:8</td>
<td>3410</td>
<td>1928</td>
<td>20:29</td>
<td>( \text{H}_2\text{O}, \text{N}_2, \text{H}_2 )</td>
</tr>
<tr>
<td>Kerosene</td>
<td>( \text{O}_2 )</td>
<td>2:60</td>
<td>301</td>
<td>-5:9</td>
<td>3623</td>
<td>2228</td>
<td>25:29</td>
<td>( \text{H}_2\text{O}, \text{CO}, \text{CO}_2, \text{H}_2 )</td>
</tr>
<tr>
<td>( \text{H}_2 )</td>
<td>( \text{F}_2 )</td>
<td>9:00</td>
<td>410</td>
<td>0</td>
<td>4117</td>
<td>2018</td>
<td>13:64</td>
<td>( \text{HF}, \text{H}_2 )</td>
</tr>
<tr>
<td>( \text{B}_2\text{H}_6 )</td>
<td>( \text{F}_2 )</td>
<td>5:20</td>
<td>371</td>
<td>7:5</td>
<td>4934</td>
<td>3130</td>
<td>22:52</td>
<td>( \text{HF}, \text{BF}, \text{H}_2, \text{H} )</td>
</tr>
<tr>
<td>( \text{C}_2\text{H}_2 )</td>
<td>( \text{F}_2 )</td>
<td>1:50</td>
<td>306</td>
<td>54:2</td>
<td>4109</td>
<td>3294</td>
<td>29:96</td>
<td>( \text{HF}, \text{C}(s) )</td>
</tr>
<tr>
<td>( \text{N}_2\text{H}_4 )</td>
<td>( \text{F}_2 )</td>
<td>2:30</td>
<td>363</td>
<td>22:8</td>
<td>4687</td>
<td>2702</td>
<td>21:33</td>
<td>( \text{HF}, \text{N}_2, \text{H}_2 )</td>
</tr>
<tr>
<td>Kerosene</td>
<td>( \text{F}_2 )</td>
<td>2:40</td>
<td>317</td>
<td>-5:9</td>
<td>3917</td>
<td>2748</td>
<td>25:26</td>
<td>( \text{HF}, \text{C}(s) )</td>
</tr>
<tr>
<td>( \text{H}_2 )</td>
<td>( \text{F}_2 )</td>
<td>9:00</td>
<td>410</td>
<td>0</td>
<td>4117</td>
<td>2018</td>
<td>13:64</td>
<td>( \text{HF}, \text{H}_2 )</td>
</tr>
<tr>
<td>( \text{H}_2 )</td>
<td>( \text{O}_2 )</td>
<td>3:50</td>
<td>424</td>
<td>34:1</td>
<td>3123</td>
<td>1426</td>
<td>9:07</td>
<td>( \text{H}_2, \text{H}_2\text{O} )</td>
</tr>
<tr>
<td>( \text{H}_2 )</td>
<td>( \text{F}_2\text{O} )</td>
<td>5:90</td>
<td>410</td>
<td>8</td>
<td>3589</td>
<td>1662</td>
<td>11:36</td>
<td>( \text{H}_2, \text{HF}, \text{H}_2\text{O} )</td>
</tr>
<tr>
<td>( \text{H}_2 )</td>
<td>( \text{NF}_3 )</td>
<td>13:30</td>
<td>350</td>
<td>-29:7</td>
<td>3868</td>
<td>1682</td>
<td>16:43</td>
<td>( \text{HF}, \text{H}_2, \text{N}_2 )</td>
</tr>
<tr>
<td>( \text{H}_2 )</td>
<td>( \text{ClF}_3 )</td>
<td>11:50</td>
<td>318</td>
<td>-39</td>
<td>3390</td>
<td>1356</td>
<td>16:78</td>
<td>( \text{HF}, \text{H}_2, \text{HCl} )</td>
</tr>
<tr>
<td>( \text{H}_2 )</td>
<td>( \text{F}_2\text{O}_2 )</td>
<td>5:00</td>
<td>407</td>
<td>4:7</td>
<td>3362</td>
<td>1504</td>
<td>10:57</td>
<td>( \text{H}_2, \text{HF}, \text{H}_2\text{O} )</td>
</tr>
</tbody>
</table>

Penner (Ref 10) has listed eleven attributes that make an ideal liq bipropellant system. We quote some of them below:

"a) Small negative or preferably positive standard heats of formation of the reactants;

b) The reaction products should have low molecular weights and large negative heats of formation. If conditions (1) and (2) are met, then the reaction products will consist of low molecular weight compounds at high temps, thus assuring good performance;

c) The propellants should have large densities in order to minimize the dead weight of storage tanks;

d) The oxidizers and reducing agents are best handled as liquids. Hence it is desirable to obtain propellants which are normally liq in the operating temp range of service units (i.e., from about -40 to +60°). For substances such as liquid oxygen and hydrogen special cooling units must be provided;

e) Since it may be necessary to store the propellants for long periods of time before use, good propellants should have high storage stability, i.e., they must not decompose or change chemically in any way during storage so that their use as a propellant is impaired;

f) For large-scale use it is, of course, imperative that propellants which are readily available and preferably also of low cost are employed. In practice this last requirement is
essential since experience has shown that rare and expensive chemicals which are needed in large quantities usually become cheap and readily available in the course of time;

g) The bipropellant mixt in a liq-fuel rocket should be spontaneously combustible with minimum time lag; and

h) The reaction products should not be excessively corrosive or form solid deposits thereby leading either to increased or decreased nozzle throat diameters

3) Representative LP Applications. Before proceeding with a description of the specific uses of LP, it is instructive to compare them with solid propellants. The choice between LP and a solid propellant for a given application depends on several factors such as: specific impulse, safety, storability, rocket engine design & special requirements

LP generally have the following advantages over solid propellants:
a) Higher Specific Impulse; b) Rocket engines with LP can be precisely calibrated prior to launching; and c) Rocket engines with LP can be throttled and even restarted after complete throttling

LP generally have the following disadvantages compared to solid propellants:
a) More complex and less rugged rocket engines; b) Poorer stability; and c) Poorer reliability

The most spectacular, though hardly the most elegant, use of LP was in the first stages of early ICBM and space exploration rockets. These used clusters of LP engines to achieve lift-off via "brute force". For example, the original Sputnik, Vostok, Atlas, Titan, Vanguard, Thor, Apollo & Saturn used clusters of engines powered with liquid oxygen/kerosene mixtures for the lift-off stage (Refs 30a & 32). More specifically, Saturn I had a cluster of eight liquid oxygen (LO)/kerosene H-1 rockets. Later Satums and Titans used LP of higher ISP, eg: Saturn-IVB used LO/liq Hydrogen; Post 1959 Titans used N₂O₄/Hydrazine derivatives; Jupiter-C used LO/Hydrazine derivatives; and the French Diamant I space booster used nitric acid/turpentine (Ref 32)

More sophisticated uses of LP, which take advantage of the LP engine's capability of being throttled and restarted, are in thrusters.

Thrusters are rockets for maneuvering and controlling the attitude of space vehicles. The usual monopropellant for thrusters is catalytically decomposed Hydrazine. The usual catalysts are iridium, rhodium or ruthenium and their mixts. For a review of Hydrazine thrusters see Refs 33, 34 & 35. Russi (Ref 34) emphasizes that, in spite of many studies and the general acceptance and apparent success of hydrazine thrusters, new rocket motor design is still largely empirical. A bipropellant consisting of Hydrazine mixed with Hydrazine Nitrate has also been tried in thrusters but is no longer popular.

An informative compilation of the uses of LP is given in Table 3 from Ref 30a

The two interesting applications of LP are in gas generation and in specialized ammunition.

The use of a monopropellant gas generator offers significant advantages over conventional solid propellants because of its capability for variable demand operation, high performance, solids-free exhaust and constancy of flow over a wide temp range. The monopropellant gas generator eliminates the need for filters and design compromises such as oversized propellant grains to account for the variation in burn rate over the temp range of −65 to +165°F (Ref 24)

Ammunition containing bulk-loaded monopropellants have the characteristics of low flame temp, high energy, reduced smoke and flash, and reduced fouling and longer barrel life. There is one disadvantage, however, that has prevented exploitation of these good attributes. Liq monopropellants do not burn stably under the conditions existing in bulk-loaded cartridges. High and erratic pressures occur, accompanied by high frequency, high amplitude pressure excursions. It was shown that bulk loaded monopropellants, eg, alkyl nitrate mixtures, encapsulated to form small spheres, produced propellants of very repeatable and uniform initial surface area similar to conventional smokeless ball powder. Proper ignition and the resulting combustion involving the liquid droplets prevents agglomeration into a large unstable bulk (Ref 25)
### Table 3

Current Liquid Bipropellant Applications

<table>
<thead>
<tr>
<th>Propellant System, Common Designation</th>
<th>Thrust Level, lbs.</th>
<th>Application Area</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>LO$_2$/RP-1 (a)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Thor</td>
<td>170,000</td>
<td>Space booster, IRBM (Delta 1st stage)</td>
</tr>
<tr>
<td>H-1</td>
<td>200,000</td>
<td>Upgraded Saturn I</td>
</tr>
<tr>
<td>Blue Streak</td>
<td>300,000</td>
<td>Space Booster</td>
</tr>
<tr>
<td>Titan I</td>
<td>300,000 (1st stage) 80,000 (2nd stage)</td>
<td>ICBM</td>
</tr>
<tr>
<td>Atlas</td>
<td>370,000–390,000</td>
<td>Space booster, ICBM</td>
</tr>
<tr>
<td>F-1</td>
<td>1,500,000</td>
<td>Saturn V, S-IC</td>
</tr>
<tr>
<td><strong>LO$_2$/H$_2$</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Centaur</td>
<td>15,000</td>
<td>Upgraded Saturn I SIII (2 engines)</td>
</tr>
<tr>
<td>J-2</td>
<td>200,000</td>
<td>Saturn V, SII &amp; SIII</td>
</tr>
<tr>
<td><strong>LO$_2$/NH$_3$</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>X-15</td>
<td>15,000–58,000</td>
<td>Experimental rocket plane</td>
</tr>
<tr>
<td><strong>LO$_2$/C$_2$H$_5$OH</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Redstone</td>
<td>75,000</td>
<td>SRBM</td>
</tr>
<tr>
<td><strong>H$_2$O$_2$/JP-4 (1), JP-5 (a)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AR-2, 3</td>
<td>3300–6600</td>
<td>Auxiliary rocket engine for aircraft</td>
</tr>
<tr>
<td>Warrior</td>
<td>3500–10,200</td>
<td>Auxiliary rocket engine for aircraft</td>
</tr>
<tr>
<td><strong>IRFNA (b)/UDMH (c)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AJ10-118</td>
<td>7500</td>
<td>Space booster (Delta 2nd stage)</td>
</tr>
<tr>
<td>Lance</td>
<td></td>
<td>Surface-to-surface missile</td>
</tr>
<tr>
<td>Agena</td>
<td>16,000</td>
<td>Space booster (2nd stage)</td>
</tr>
<tr>
<td><strong>IRFNA (b)/MAF-1 (d)</strong></td>
<td>Bullpup</td>
<td>Air-to-surface missile</td>
</tr>
<tr>
<td><strong>IRFNA (b)/MAF-3 (e)</strong></td>
<td>TD-174</td>
<td>Air-to-air missile</td>
</tr>
<tr>
<td><strong>IRFNA (b)/MAF-4 (f)</strong></td>
<td>P4</td>
<td>Target drone</td>
</tr>
<tr>
<td></td>
<td>550 (booster)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>106 (sustainer)</td>
<td></td>
</tr>
<tr>
<td><strong>IRFNA (b)/JP-4 (a)</strong></td>
<td>Aerobee 100</td>
<td>Sounding rocket</td>
</tr>
<tr>
<td></td>
<td>sustainer</td>
<td></td>
</tr>
<tr>
<td><strong>IRFNA (b)/aniline-furfuryl alcohol</strong></td>
<td>Aerobee 150 and 150A sustainer</td>
<td>Sounding rocket</td>
</tr>
<tr>
<td><strong>IWFNA (g)/turpentine</strong></td>
<td>Emeaude</td>
<td>Space booster</td>
</tr>
<tr>
<td><strong>N$_2$O$_4$/N$_2$H$_4$—UDMH (c) (50-50)</strong></td>
<td>Apollo service module RCS</td>
<td>Reaction control</td>
</tr>
<tr>
<td></td>
<td>Ullage rocket</td>
<td>Saturn V, S-IVB Ullage Control</td>
</tr>
</tbody>
</table>

(continued)
Table 3 (continuation)

<table>
<thead>
<tr>
<th>Propellant System, Common Designation</th>
<th>Engine System</th>
<th>Thrust Level, lbs</th>
<th>Application Area</th>
</tr>
</thead>
<tbody>
<tr>
<td>N\textsubscript{2}O\textsubscript{4}/N\textsubscript{2}H\textsubscript{4}—UDMH (c) (50-50)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>AJ10-131</td>
<td>2200</td>
<td>General purpose space engine</td>
</tr>
<tr>
<td></td>
<td>F750 L2.2K</td>
<td>2200</td>
<td>Multiple restart space engine</td>
</tr>
<tr>
<td></td>
<td>Lunar module ascent engine</td>
<td>3500</td>
<td>Lunar module liftoff (moon)</td>
</tr>
<tr>
<td></td>
<td>F720 L8.0K</td>
<td>8000</td>
<td>Multiple restart space engine</td>
</tr>
<tr>
<td></td>
<td>Transtage</td>
<td>8000</td>
<td>Upper stage propulsion</td>
</tr>
<tr>
<td>N\textsubscript{2}O\textsubscript{4}/N\textsubscript{2}H\textsubscript{4}—UDMH (c) (50-50)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Lunar module descent engine</td>
<td>1050—10,500</td>
<td>Lunar landing engine</td>
</tr>
<tr>
<td></td>
<td>Apollo service module</td>
<td>21,900</td>
<td>Space propulsion</td>
</tr>
<tr>
<td></td>
<td>YLR113-AJ-1</td>
<td>50,000—150,000</td>
<td>Rocket sled</td>
</tr>
<tr>
<td></td>
<td>Titan II</td>
<td>430,000 (1st stage) 100,000 (2nd stage)</td>
<td>Space booster, ICBM</td>
</tr>
<tr>
<td>N\textsubscript{2}O\textsubscript{4}/MMH (h)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Advanced Syncom RCS</td>
<td>5</td>
<td>Reaction control</td>
</tr>
<tr>
<td></td>
<td>Gemini RCS</td>
<td>25 (two 8-engine sets)</td>
<td>Reaction control</td>
</tr>
<tr>
<td></td>
<td>Transtage ACS</td>
<td>25 (4 engines) 45 (4 engines)</td>
<td>Attitude control</td>
</tr>
<tr>
<td></td>
<td>Apollo command module</td>
<td>93 (two 6-engine sets)</td>
<td>Attitude control</td>
</tr>
<tr>
<td>N\textsubscript{2}O\textsubscript{4}/MMH (h)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Gemini OAMS</td>
<td>25 (8 engines) 85 (2 engines) 100 (6 engines)</td>
<td>Orbital change</td>
</tr>
<tr>
<td></td>
<td>Radiomic</td>
<td>85—100</td>
<td>General purpose attitude control</td>
</tr>
</tbody>
</table>

a) RP-1, JP-4 & JP-5: Kerosene Fractions  
b) IRFNA: Inhibited Red Fuming Nitric Acid  
c) UDMH: Unsym Dimethyl Hydrazine  
d) MAF-1: 50.5% DETA, 40.5% UDMH, 9% CH\textsubscript{3}CN  
e) MAF-3: 80% DETA, 20% UDMH  
f) MAF-4: 40% DETA, 60% UDMH  
g) IW/NA: Inhibited White Fuming Nitric Acid  
h) MMH: Monomethyl Hydrazine
4) Performance Characteristics of LP. Specific impulse, \( I_{sp} \), probably the most accepted measure of LP performance, was defined in Sect 1 and tabulated in Tables 1 & 2. The thrust, \( F \), of a rocket is given by

\[
F = \pi r c
\]

where \( r \) and \( c \) were also defined earlier. Typical thrust levels for “practical” LP systems are given in Table 3. For an ideal rocket other performance parameters are: \( T_C \) = adiabatic combustion temperature; \( T_E \) = exhaust gas temperature; \( c^* \) = characteristic velocity; \( C_F \) = thrust coefficient; \( M_E \) = mean molecular weight of exhaust gases; \( \bar{\gamma} \) = average ratio of specific heats of exhaust gases.

According to Summerfield (Ref 12, p 442):

“The ideal rocket motor analysis rests on the following simplifications: (a) the propellant gas obeys the perfect gas law; (b) its specific heat is constant, independent of temp.; (c) the flow is parallel to the axis of the motor and uniform in every plane normal to the axis, thus constituting a one-dimensional problem; (d) there is no frictional dissipation in the chamber or nozzle; (e) there is no heat transfer to the motor walls; (f) the flow velocity in the chamber before the nozzle entrance is zero; (g) combustion or heat addition is completed in the chamber at constant pressure and does not occur in the nozzle; and (h) the process is steady in time.”

With these simplifications typical computed values of LP rocket performance (according to Ref 12, pp 453–64) are given in Table 4.

Additional computational results are given in Table 2 which also shows major exhaust gas products. Parametric charts of rocket performance for specific missions are given by Jortner (Ref 17, p 471). Further data on rocket performance from the point of view of weight and volume limited systems are presented by Mellish & Gibb (Ref 17, p 447).

At the high combustion and exhaust temps shown in Tables 2 & 4, many of the combustion products will dissociate (e.g., \( CO_2 = CO + O_2; H_2 = 2H \), etc.). The effects of product gas dissociation constitute the greatest source of inaccuracy in the above ideal rocket analysis.

Additional departures from ideal behavior result from:

a) Conical divergence of the exhaust jet;

b) Surface friction and flow disturbances in the exhaust nozzle;

c) Constriction of the exit area due to boundary layer build up;

d) Jet detachment;

e) Heat loss from the hot gas to the cold motor walls;

f) Suspended liquid or solid particles in the exhaust jet; and

g) Pressure drop in the combustion chamber due to heat release.

They are discussed in detail in Ref 12, pp 464–71.

A detailed description of LP Rocket Systems is beyond the scope of this article. An excellent account of many of the facets of LP rockets, such as combustor design, cooling, pressurization prior to combustion, and pump systems, is given in Ref 12, pp 475–516.

---

Table 4

<table>
<thead>
<tr>
<th>Performance parameter</th>
<th>Ordinary range</th>
<th>High range</th>
</tr>
</thead>
<tbody>
<tr>
<td>( T_C )</td>
<td>2000–3000 K</td>
<td>3000–5000 K</td>
</tr>
<tr>
<td>( c^* )</td>
<td>4000–5500 ft/sec</td>
<td>5000–8000 ft/sec</td>
</tr>
<tr>
<td>( C_F )</td>
<td>1.3–1.5</td>
<td>1.5–1.6</td>
</tr>
<tr>
<td>( I_{sp} )</td>
<td>200–270 lb-sec/lb</td>
<td>270–400 lb-sec/lb</td>
</tr>
<tr>
<td>( M_E )</td>
<td>20–25</td>
<td>8–20</td>
</tr>
<tr>
<td>( \bar{\gamma} )</td>
<td>1.15–1.25</td>
<td>1.15–1.20</td>
</tr>
</tbody>
</table>
5) **Sensitivity and Hazards of LP.** Some monopropellants and some well-mixed bipropellants exhibit detonation characteristics typical of Liquid Explosives (See Sects 4 & 5 of article on Liquid Explosives in this Vol). However, bipropellants usually do not sustain complete detonation, i.e., a rather small portion of the bipropellant undergoes something akin to detonation and the remainder deflagrates (Ref 26). Of course even this partial detonation can be very dangerous and destructive. LP are also subject to another phenomenon which is potentially destructive (at least to the rocket), namely combustion instability (Refs 16 & 22).

The two-stage explosion process of most bipropellants, i.e., "detonation" followed by deflagration, manifests itself in the explosion fire-ball and the impulse delivered by such explosions. Both fireball and impulse (at least close to the explosion) for typical bipropellants are larger than for the same weight of TNT, but shock pressure is considerably less. Thus the main hazard of a distant bipropellant explosion is its thermal effects. Nearby, this explosion can also be destructive to organisms and structures that are affected by impulse and large low-velocity fragments. It has been suggested that the nature of a bipropellant explosion is primarily the result of "poor" mixing (on a molecular scale) of the bipropellant fuel and oxidizer, and that turbulence created by the "detonation" disrupts further mixing and extinguishes the "detonation". In liq explosives, by way of contrast, the "oxidizer" and "fuel" are generally contained within the same molecule and there is no mechanism for extinguishing detonation (Ref 26).

Combustion instability in LP results from pressure oscillations during combustion. If these oscillations are of low frequency, e.g., as a consequence of fluctuations in the propellant flow rate, the resulting instability is called chugging. At high oscillation frequencies, leading to what is sometimes referred to as screaming, longitudinal waves are set up in the combustion chamber as well as cylindrical wave motion. Above a critical amplitude, these cylindrical motions can increase enormously and damage the chamber.

Levine (Ref 22, p 1083) has studied high frequency combustion instability of LP via high speed photography and pressure-time measurements. Crocco has examined these phenomena theoretically (Ref 22, p 1101). Bernard & Dufour (Ref 16, p 1074) make the interesting suggestion that instability in amine/nitric acid bipropellants is associated with the production of intermediates, such as amine-nitrates, which decompose explosively on rapid temp and pressure rises.

The compatibility of LP with materials that they may contact can create hazard problems. Hollister (Ref 19) reviews the compatibility of several LP systems in contact with elastomeric materials.

Farber (Ref 31) points out that LP rocket destruct systems (specifically Saturn V) can initiate dangerous secondary explosions in the discharged fuel/oxidant mixture.

Specific tests of the detonation sensitivity of LP are not very different from those used with liq explosives, but the interpretation of LP sensitivity tests is subject to large uncertainties. This is so, because practical considerations dictate that relatively small amounts of LP be tested, but for many LP systems the increase in sensitivity with increasing test sample amount is not known.

Tannenbaum & Beardell (Ref 30, p 344) suggest the following "routine" tests:

a) A modified Trauzl lead block test (See Vol 1 under Physical Tests) with a 0.5 "to 2g LP sample and a No 8 blasting cap initiator;

b) Bureau of Mines drop-weight test;

c) Adiabatic compression test in which a piston rapidly compresses a gas bubble above the test liq;

d) Self-heating test in which the sample is heated in a bomb and the sample and heating bath temps are monitored;

e) Card-gap test (See Vol 1 under Physical Tests to test shock-sensitivity); and

f) Critical diameter established by measuring detonation velocity in columns of decreasing diameter.

Glatts (Ref 20) tested four monopropellants for sensitivity to fire and bullet impact. Hydrazine, Hydrazine Nitrate solution, PrNO₃, and ethylene oxide were all tested in 1-gallon aluminum containers (half of them filled to 25% and half to 95% capacity) by being subjected to oil and wood bonfires, and 20-mm incendiary, 20-mm HE incendiary, and 0.30-caliber rifle fire. Aviation gasoline was tested in the same manner. In the bonfire tests, all four monopropellants generally gave equivalent explosion intensities regardless of the type of bonfire or the amount of liq in the test container. Occasionally, however, with the first 3 monopropellants, anomalous re-
results occurred: there was either no explosion or an explosion of above average intensity. Ethylene oxide more nearly approximated the behavior of aviation gasoline than did the other monopropellants. In the rifle-fire tests, PrNO₃ was the most bullet-sensitive. It was exploded by all three types of ammunition. Hydrazine and Hydrazine Nitrate solution were exploded by the HE ammunition but not by incendiary and HE incendiary ammunition.

Written by J. ROTH


Liquid TNT. See under Drip Oil in Encycl 5, (1972), D1546-R to D1547-L.

Litharge. See Lead Monoxide in this Vol.

Lithium-Aluminum Hydride. See Aluminum-Lithium Hydride in Encycl 1, (1960), A154-R to A155-L, and under Hydrides in this Vol.
Lithium Nitrate. LiNO₃, mw 68.95, colorless delq granules, mp 261°, d 2.38g/cc; sol in w and alc. Used in pyrotechnics as an oxidizer to color burning comps red (Ref 2), and in rocket proplnts (Ref 1a).

Refs: 1) Gmelin, Syst Nr 20 (1927), 98ff
1a) D.R. Stern, USP 2949006 (1960) & CA 55, 4964 (1961)
3) CondChemDict (1971), 525-L

Lithium Perchlorate. LiClO₄, mw 106.40, colorless delq crys, mp 236°, d 2.429g/cc; sol in w and alc. Decomp starts at about 400° and becomes rapid at 430°, yielding LiCl and O₂. It has more available O₂ on a vol basis than liq O₂, and is thus used in “oxygen candles” (Ref 2, p 237). It is used as an oxidizer in solid rocket proplnts and in pyrotechnics to color burning comps red (Refs 1 & 2).

2) Eltern (1968), 124, 237, 271 & 337
3) CondChemDict (1971), 525-L

Lithoclastites. Dynamites patented in Fr by Roca in 1884 contd NG mixed with hydrocarbons and other substances

Ref: Daniel (1902), 690

Lithofracteur. Brit mining expl manufd between 1882 and 1899. Its approx compn was NG 54, Ba nitrate 15, kieselguhr 17, wood flour 2, bran 1, sulfur 4, Mn dioxide 2 & soda ash 2% (Refs 1 & 3). A Ger version, called Lithofracteur Krebs, contd NG 52, pulverized coal 12, sulfur 2, Na nitrate 4 & kieselguhr 30% (Ref 4). Gody (Ref 2) lists a Lithofracteur Dynamite which contd saltpeter, sulfur, sawdust and bran. He also lists a Lithofracteur of Newton, known as Poudre Newton, which consisted of Ba nitrate 77, K nitrate 2 & charcoal 21%

Refs: 1) Daniel (1902), 409
2) Gody (1907), 173
3) CondChemDict (1942), 290
4) Pérez Ara (1945), 331

Lithorite. A Belg mining expl used at the end of the 19th century. It consisted of K nitrate 50.0, Na nitrate 16.0, sulfur 16.0, nitrated sawdust 8.0, Amm picrate 3.5, K ferrocyanide 3.0 & charcoal 3.5%

Ref: Daniel (1902), 410

Little David. An experimental rifled mortar (914mm mortar T1) of 36” diameter developed in the USA during WWII. This mortar and its special ammo were secretly developed for reducing Ger fortifications; however, this proved to be unnecessary. The mortar weighed 172900 lbs; its proj weighed 3650 lbs of which 1589 lbs was HE. It was fired by a max propelling charge of 218 lbs of proplnt, and had a range of about 9000 yards


Livens Projector. Invented during WWI by Capt J. Livens of the Brit Army. It was a crude form of a trench mortar, consisting of a smooth bore tube (8” ID, 37.5” long) which was used to project Livens Drums by means of a proplnt charge ignited by an electric primer. The drums were ellipsoidal-like containers (7-5/8” diam, 20” long) filled with either gaseous agents or incendiary materials, and contained a burster charge. The filled drum weight was about 61 lbs, and its range was between 910 and 1450 yards.

From 400 to 800 projectors were covertly installed in a specially prep’d trench, whose front wall was cut at a 45° angle. Their only mounting was a simple metal base placed to prevent the tube from being driven into the ground on discharge. The projectors were held in place by earth filling. They were electrically connected in sets of 25, and on firing of 800 projectors simultaneously, a charge of 11 tons of phosgene gas would be deposited over 50 to 100 square yard area without preliminary warning to the enemy.

2) A.B. Ray, IEC 13, 716-17 (1921)
3) C. Wachtel, “Chemical Warfare”, Chemical
LOADING AND FABRICATION OF EXPLOSIVES

Process Selection.

Most solid high expls are manufd by processes that yield granular material. Their bulk densities are generally somewhat less than 1 g/cu cm. They are used in military applications as solids of well defined configurations, usually at densities between 1.5 and 1.7 g/cu cm.

The two principal loading techniques are casting and pressing. All expls in common military use can be pressed, however, those that are castable are usually cast because of the greater convenience and flexibility of this process. As a rule of thumb, main bursting charges of large cal munitions are cast while small explosive components (initiators to boosters) are pressed.

More pounds of military expl are cast than are loaded by all other processes. Essentially, the casting of an expl involves only melting it and pouring it into a charge case or mold. In practice, like most fundamentally simple processes, the procedures necessary to cast charges of the quality needed for acceptable performance and safety can become quite elaborate. A suitable pour viscosity is of over-riding importance.

The most common procedure for pressing powd expls is that of pouring the powd into a mold and pressing it with a ram that fits snugly. The pressure most frequently specified for charges used in military items is 10000 psi. Charges may be pressed directly into their containers or pressed into molds and ejected as pellets. Where they are pressed into containers of lengths greater than the diameter, the explosive is usually loaded in increments.

After pressing or casting, it is sometimes necessary to machine explosives, either to provide a smooth surface or a fuze cavity at the filling hole, or to produce complex contours required for some specialized purposes. In some cases, mating contours of two charges are cemented together. Cavities are also formed using a special tool on final pressing.

Of increasing importance are the plastic bonded expls (PBX). These are exactly what the name implies, and like plastics can be obtained in many different forms. Hence, PBX's are available for casting, pressing, or extruding. They vary from rigid to rubbery consistencies depending on the type of plastic used as the binders — thermoplastic or thermosetting — and the degree of polymerization permitted. High mechanical strength and high thermal stability are possible (Ref 10).

Other considerations for process selection include fabrication facilities and suitability of the explosive for its intended application (Ref 13).

Casting.

Projectile Preparation

As part of the manufg process, the interior wall of the proj is sprayed with paint or varnish, primarily to prevent rusting of the proj in storage. The requirements of the coating are that it be compatible with the expl, adhere well to the proj wall, and offer a good bonding surface for the expl. The latter requirement is necessary to prevent rotation of the charge relative to the spinning proj. The finished coating at the base of the proj should be thin enough to assure thorough drying and be sufficiently smooth to eliminate irregularities that could otherwise form air pockets.

The molten expl is usually poured thru a funnel Former. This tool is specially designed to furnish the desired surface contour upon removal and to hold a sufficient reservoir of molten expl to replenish the shrinking, cooling mass beneath it. A thin film of silicone grease is applied sometimes to the former to aid in its release when the expl has solidified.

Effect of Casting Procedure On Charge Characteristics.

1) Porosity and Cavitation.

The porosity of an expl charge is usually introduced by two principal causes, entrained air bubbles and dissolved gases, and shrinkage that occurs as the charge solidifies and cools. The higher the temp of casting and the more fluid the melt, the larger is the fraction of the entrained air that forms into bubbles and floats out of the charge. On the other hand, these conditions maximize cavitation due to shrinkage. The most serious effect of shrinkage is that known to metal founders as "piping".
The casting solidifies from the outside and consequent shrinkage is that of an isolated mass at the center where no additional material is available to fill the volume left by the shrinkage. The result is a single large void at the center of the casting.

In a cast charge (unlike in a pressed charge), both density and pore or cavity size are determined by the casting procedure. Both of these factors must be considered by the designer in terms of their effects upon safety, reliability, and performance.

2) Crystal Size

The crystals of TNT in cast expls may vary from microscopic size to a substantial fraction of the size of the charge, depending upon casting conditions and procedure. The approach known as creaming results in very fine crystals. In mixed expls, which usually are cast in the form of slurries, the solid particles tend to inhibit crystal growth, although TNT crystals sometimes apparently grow around the particles of the slurry. The effects of particle size on initiation sensitivity, failure diameter, and performance characteristics also have been observed to apply to crystal size in cast TNT.

3) Uniformity of Composition.

Most castable expls are poured as slurries of RDX, Al, etc., in molten TNT. The instant a charge is poured, the particles of higher density than TNT start to settle, and those that are lighter start to rise. As a result, by the time the material solidifies, its compn varies from point to point within the charge. Another cause of nonuniformity of compn is the tendency of TNT to form essentially pure crystals, leaving other components of the mixt at grain boundaries and in the center of the charge that usually solidifies last. The most serious production problem of this kind is the settling of Al in larger charges of aluminized expls. The use of Al and other additives in very fine particle sizes can help to alleviate this problem but also tends to increase pouring difficulties because of the higher viscosities of the melts.

Standard Casting Procedure

The most common procedure for filling a proj or bomb case is to do so in a single pouring (Ref 7). In loading projs, a funnel or sprue provides a reservoir of molten expl to fill the volume left by the shrinkage. The expl in the funnel must, of course, remain liq and in communication with the center of the charge. When the filling hole is large enough, convective heat transfer maintains such conditions. In other instances, however, such conditions can be maintained only by means of steam heated funnels, steam finger, or hot probes.

Where the maintenance of a clear channel between sprue and the slowest freezing part of a charge is impractical, cavitation is avoided by casting charges in layers, each of which is allowed to "crust over" before pouring the next.

TNT melts at 81°. It forms eutectics with RDX, Tetnyl (68°), PETN (76°), and other "impurities" in the mix and makes these materials more soluble at higher temps. Thus, there is a general tendency for the solid content and, hence, the apparent viscosity of most castable mixts to decrease as the temp is increased. However, a reversal of the tendency toward the reduction in viscosity has been noted in Comp B when it heated above 100°.

From the eutectic or mp, the composition of the liq portion and its viscosity vary as heat is removed. It has been recommended that the heat content of any expl be reduced, before pouring, to the minimum compatible with the avoidance of air entrainment.

TNT may be cast after it has cooled to a point where a fairly large fraction of it has solidified to form a slurry of very small crystals. Such a slurry is obtained by stirring it as it cools. TNT cast in this manner is labeled by some as creamed TNT. Some have applied the term creamed to all expls that are cast after stirring until the last possible instant. Extreme caution must be taken to avoid air entrainment during stirring. This technique has the advantage of resulting in less shrinkage on cooling and solidification because a large portion of the TNT is already solid before casting.

Some Special Casting Techniques

1) Pellet Casting.

For very large charges, cooling time is reduced and shrinkage minimized by use of precast pellets. The best pellet casting technique is that of pouring a quantity of molten expl into the case, and then pouring in pellets, slowly enough so that they are not in contact with one another to avoid entrapping interstitial air (Ref 7). Although pellet casting
reduces the total amount of shrinkage voids, it makes it nearly impossible to maintain channels to the pockets of molten material. The most important advantages of pellet casting is the reduction of cooling time and minimizing the shrinkage in large charges. Pellet casting is not used in loading artillery props because of the development of cavities

2) Vacuum Melting and Casting.

Entrainment of air may be avoided by melting and casting under a vacuum. Vacuum melting is a fairly straight-forward procedure in the vacuum kettles that are maintained by many loading facilities. Vacuum casting requires specially designed molds or a vacuum chamber large enough to contain both kettle and mold. The definite increase in the cast density should indicate without question the advantage of vacuum melting, namely, an increase in viscosity of the vacuum melted material. Nevertheless, a divergence of opinion exists regarding the value of vacuum melting followed by pouring in air. Some investigators report results nearly as good as those obtained with complete vacuum melting and casting. Others maintain that so much air is entrained in the casting process that the value of vacuum melting is negligible. A possible explanation for this difference of opinion is the difference in techniques that can be applied in various types of operation

3) Vibration, Jolting and Centrifugal Casting.

Accelerating of a cast charge after pouring, but before solidification will often expedite the movement of air bubbles to the surface (Ref 4). Vibration and jolting often break the surface tension that causes bubbles to cling to surfaces (Ref 7). Centrifugal acceleration, of course, also accelerates the settling of denser components of mixtures (Ref 7). This has been used to advantage in loading HEAT ammo where it is desirable to have a richer composition of the more energetic compounds (RDX and HMX) around the cone in cyclotol or octol.

4) Controlled Cooling.

If an expl charge can be induced to cool from the bottom up, maintaining a nearly plane interface between liq and solid, densities well in excess of 99% of maximum theoretical are attainable. In a complicated programmed cooling, the thermal cycle of preheating the mold, pouring, and cooling takes over forty hours (Ref 2). At the other extreme is the use of strategically placed insulation to cause a charge to cool in the approximate desired pattern.

5) Extrusion

Extrusion may be considered a form of casting under pressure. In applications where cylindrical charges are required, some plastic bonded explosives can be extruded into the desired shape and then placed or pressed into the ammunition housing. Conventional extrusion tools are employed for this process.

6) Liquid Explosives

From the standpoint of casting, the pouring of liq or slurry expls is handled in the same manner as that of molten expls. The process is simpler in that pouring takes place at ambient temp. However, care must still be taken to avoid entrapment of air. Liq filling can be speeded up by pumping. If slurries are to be gelled in the ammo, the gelling agent is introduced just ahead of the cavity (Ref 12)

Pressing.

Standard Procedures.

1) Measurement of Explosive Charges

For small test quantities or for some premium quality production, direct reading one-pan balances are used. They are faster than analytical balances and provide an accuracy within one percent. Automatic weighing machines are also available.

The desire is always to load a specific weight of expl. This objective can be achieved to a sufficient degree of accuracy for many purposes by volumetric control, as in commercial blasting caps and squibs. The two most common volumetric measuring devices are scoops and charging plates. Scoops (Fig 1)

![Fig 1 Scoop Loading](image-url)
are filled and leveled against a rubber band. Careful scooping is accurate within 4%. Charging plates (Fig 2) lend themselves to production rates. After filling holes in the top plate and scraping off the excess, plates are aligned with cup holes. Since expl quantities are usually specified by weight, it is left to the loading plant to adjust the volume measured so as to take into account bulk density. There are now several automatic volumetric loading devices for production loading of primer mixes and lead azide used in initiators.

2) Direct Pressing in Case

A large proportion of expl charges are loaded by direct pressing of expl charges in cases (Fig 3). Fits and tolerances of expl charge cases and loading tools are determined by reconciliation of three opposing factors: 1) production costs of components rise sharply as tolerances are reduced; 2) powdered expls tend to flow into the clearance between ram and case. In addition to creating a hazard, the expls wedged in this space can increase the frictional resistance to ram movement, and substantially decrease real loading press; and 3) interference between ram and case results in binding (which may be so severe as to prevent any pressing of the expl), damage to the case, inclusion of chips of case material in the expl, or all of these (Ref 1).

The cost of a set of loading tools may be distributed over a large number of items. For this reason, they are often made to fits and tolerances similar to those used for gages. Where cases are made by processes such as forging, drawing, and extrusion, which use most of the tolerance in lot-to-lot variation, some loading activities have found it worthwhile to maintain a series of loading tools of graduated dimension, using those giving the best fit possible with each lot of cases.

Production loading tools should be hardened (60 Rockwell C is common). The die should be ground, honed, and lapped or polished to an 8 or 16 rms micro-inch finish. Some claim better results if the final operation involves longitudinal rather than rotary motion.

The friction between the expl and the walls causes a gradient of press, and hence density, decreasing from the face of the ram. The slope of this press gradient, of course, is proportional to the coeff of friction between the expl and the walls, which varies with both expl and case material and also with the interior finish of the case. As a general rule, the density variations due to these gradients are kept within reasonable bounds by adherence to the general
rule-of-thumb that the length of an increment after consolidation should not exceed the diameter of the cavity (Ref 2).

The usual loading press of about 10,000 psi is well beyond the bursting strength of charge cups of any material that can be economically deep drawn. Hence, cups are supported by close fitting loading tools while being pressed. Most of the difference between the cup diameter before and after loading is accounted for by the expansion of the expl component, relieving residual stresses, as it is pushed out of the tool. For this reason, loading tools should be made to fit the maximum outside diameter of the cup, within a few ten-thousandths of an inch.

Standard dimensions and tolerances of cups are listed in MIL-STD-320 (Ref 11). Bore finish and hardness of the bushing are important factors in trouble-free ejection of finished cups. Lapped or honed bores are often specified. Where cases are heavier or where expls are to be loaded directly into fuze cavities, the interactions of case and tool tolerances, which may be sufficient to cause interference between the ram and any of the bores through which it passes, should be considered carefully. In some situations, where expls are to be loaded directly into fuze holes, the most practical way to attain alignment is to use a pin or dowel, similar to the loading ram, to hold the component in alignment with the ram guide while it is being clamped in place. It is best to use an alignment pin a thousandth of an inch or so larger than the loading ram. Fig 4 shows a set-up for hand loading of leads making use of an alignment ram and a mandrel.

3) Stop vs Pressure Loading

In production, it is possible either to press a controlled quantity of expl to a controlled height (called stop loading) or to apply a given load to a loading ram of a given diameter (called pressure loading). The inherent variations in production material introduce a certain amount of error in the density obtained by either method.

The relationship between loading press and charge density for commonly pressed expls is given in Table 1 (Ref 1). An approximation of the loading densities of six commonly used explosives is shown in the nomograph, Fig 5 (Ref 3). The pressure-density relationship varies somewhat from lot to lot. In addition, loading density is affected by such factors as ram clearance and increment length.

From the usual cup tolerances, it has been calculated that the cross-sectional area of the expl column of a detonator may vary by two or three percent. In normal production, a reasonable weighting tolerance for initiator charges is three or four percent. Thus, in stop loading, assuming that the height of an increment is exactly reproduced, the density may vary as much as seven percent.

The implication of the foregoing, that densities are more readily controlled by the control of loading pressure than by stop loading, has been borne out by experience. However, the production advantages of stop loading are sufficient to outweigh any theoretical disadvantages. It is important for stop loading to specify dimensions, quantities, and tolerances such that the max press is within limits imposed by tool strength. When items so loaded are used, safety and reliability determinations should take the effects of variable charge density into account. In either type of loading, a check of the loading density for each production lot is highly desirable.

When density is determined by pressure loading, variation in press, cross-sectional area, and charge weight each has an effect upon the column-height. Usually, the length tolerances specified cannot be held merely by holding the various quantities mentioned within their tolerances. The weight of expl must be adjusted to compensate for the other variables. Commonly, the last charge loaded is adjusted to fit the space.
### TABLE 1

**LOADING DENSITY OF VARIOUS EXPLOSIVES**

<table>
<thead>
<tr>
<th>Explosive</th>
<th>3</th>
<th>5</th>
<th>10</th>
<th>12</th>
<th>15</th>
<th>20</th>
<th>Cast</th>
<th>Crystal Density</th>
</tr>
</thead>
<tbody>
<tr>
<td>Composition A-3</td>
<td>1.47</td>
<td></td>
<td>1.61</td>
<td>1.65</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Composition B</td>
<td></td>
<td>1.59</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cyclonite (RDX)</td>
<td>1.46</td>
<td>1.52</td>
<td>1.60</td>
<td>1.63</td>
<td>1.65</td>
<td>1.68</td>
<td></td>
<td>1.82</td>
</tr>
<tr>
<td>EDNA (Hafnite)</td>
<td></td>
<td>1.39</td>
<td>1.46</td>
<td></td>
<td>1.51</td>
<td>1.55</td>
<td></td>
<td>1.71</td>
</tr>
<tr>
<td>Explosive D</td>
<td>1.33</td>
<td>1.41</td>
<td>1.47</td>
<td>1.49</td>
<td>1.61</td>
<td>1.64</td>
<td></td>
<td>1.72</td>
</tr>
<tr>
<td>Lead Azide</td>
<td>2.46</td>
<td>2.69</td>
<td>2.98</td>
<td>3.05</td>
<td>3.16</td>
<td>3.28</td>
<td></td>
<td>4.68</td>
</tr>
<tr>
<td>(2 specimens)</td>
<td>2.62</td>
<td>2.71</td>
<td>2.96</td>
<td></td>
<td>3.07</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lead Styphnate (Norm.)</td>
<td>2.12</td>
<td>2.23</td>
<td>2.43</td>
<td>2.47</td>
<td>2.57</td>
<td>2.63</td>
<td></td>
<td>3.1</td>
</tr>
<tr>
<td>Pentolite, 50-50</td>
<td></td>
<td></td>
<td>1.59</td>
<td></td>
<td></td>
<td></td>
<td>1.65</td>
<td></td>
</tr>
<tr>
<td>PETN</td>
<td></td>
<td>1.48</td>
<td>1.61</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1.76</td>
</tr>
<tr>
<td>Picric Acid</td>
<td>1.4</td>
<td>1.5</td>
<td>1.57</td>
<td>1.59</td>
<td>1.61</td>
<td>1.64</td>
<td>1.71</td>
<td>1.76</td>
</tr>
<tr>
<td>Picratol, 52/48</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1.62</td>
<td></td>
</tr>
<tr>
<td>Tetrylene</td>
<td>1.05</td>
<td>1.22</td>
<td>1.33</td>
<td>1.37</td>
<td>1.41</td>
<td>1.48</td>
<td></td>
<td>4.72</td>
</tr>
<tr>
<td>Tetryl</td>
<td>1.40</td>
<td>1.47</td>
<td>1.57</td>
<td>1.60</td>
<td>1.63</td>
<td>1.67</td>
<td></td>
<td>1.73</td>
</tr>
<tr>
<td>TNT</td>
<td>1.34</td>
<td>1.40</td>
<td>1.47</td>
<td>1.40</td>
<td>1.52</td>
<td>1.55</td>
<td>1.59</td>
<td>1.65</td>
</tr>
</tbody>
</table>

(Densities are in g/cc)

---

**Fig 5** Nomograph of Loading Pressure and Density

---

A straight line through the point shown for a particular explosive will intersect the two scales to show the loading density resulting from any given loading pressure.

Example: The density of lead azide, pressed at 10,000 psi is about 2.9 g/cm³

Use with caution at high pressures.
remaining for it and the weight is specified as "approximate".

In addition to the pressing properties of the expl as such, the relationship between loading press and density is affected by such factors as ram movement, clearances, increment size, and the coefficient of friction between expl and case. The movement of the ram affects the relationship in two ways.

The first effect may be very serious. Where a balance or dead weight is used to determine the loading press, a rapid ram movement can result in a force due to acceleration of the masses moved which may vary from a substantial fraction to several times the force due to gravity. Analyses of some loading operations have revealed that the true loading press was three or four times that intended.

The second effect of ram movement usually works in the opposite direction (slower ram speeds plus a dwell at the peak press may cause an increase in density). This effect is due to the fact that, at loading pressures usually used, expls are stressed beyond their yield points and creep or flow plastically. This effect, of course, becomes more important at very high pressures, such as those used for delays. In addition to increasing the density, slower speeds plus dwell of the ram result in a more uniform density.

4) Pelletizing

Most powdered expls that are to be pressed are prepressed into pellets. The die of the loading tool permits closer tolerances and better finishes than are reasonable for cases that are loaded by direct pressing. Exceptions are primer mixes, PETN, and LA, although LA is pelletized in Canada on a production basis.

Although pellets for expl use are loaded by single operation methods in which weighed charges are pressed either by stop loading or by controlled pressure techniques, quantity production of pellets is accomplished in automatic pelleting machinery, in which the explosive is metered volumetrically by the controlled movements of punches. Single stroke presses of the types used for expls produce about 90 pellets per minute while rotary presses have rates of about 700 pellets per minute.

The density gradient resulting from wall friction, in addition to its effects on expl performance, may adversely affect the handling properties of pellets. Pellets consolidated from powders at low densities tend to be weak in two ways; their resistance to body fractures is often less than desirable, and they may crumble at corners and chalk off at surfaces. On the other hand, some materials become brittle and develop residual strains at high densities.

The effect of density variation on mechanical properties of pellets may cause difficulties even though the variation in expl properties is tolerable. On the other hand, the general superiority of the finishes of pelleting molds over those of charge cases and the use of double acting loading equipment result in somewhat smaller density gradients in pellets than in direct loaded explosives. The result of these counterbalancing trends is that the one-to-one limiting ratio of length to diameter which applies to increment loading also applies to pellets. For some materials, somewhat shorter pellets are desirable, particularly in larger sizes.

The diameter of a pelleting die may be maintained to almost any tolerance specified. Similarly, the distance between the top and bottom punches of an automatic pelleting machine, or the punch-to-heel distance in a stop pressing tool, can be held to any desired tolerance. Thus, the dimensional variations are essentially the variations in expansion of the material, during and after ejection from the die. The immediate expansion upon ejection for a typical expl used for pressed pellets is about 0.3%. Pellet-to-pellet variations are usually less than 0.1% but the expansion continues with storage at a rate that varies appreciably with conditions as well as with the comp of the expl. Pellets of an expl of known expansion characteristics, which are to be inserted into cups within a few hours after pelleting, may be held to dimensional tolerances of the order of 0.1% or less. However, tolerances of 0.3 to 0.5% are more practical.

Variations in density reflect variations in dimensions with those of the bulk density and flow characteristics of the expl, and those of the measured volume. With frequent pellet density determinations and occasional adjustment of the pelleting press, expls with good flow properties can be pressed into pellets reproducible in density to 1% in an automatic pelleting press.
5) **Reconsolidation**

Frequently, when it is desirable to attain the close confinement and continuity characteristic of expls loaded directly into their cases, it is difficult or inconvenient to do so. In such instances, pellets are inserted into the cavities and reconsolidated by pressing. In designing for reconsolidation, consideration must be given to the tolerances and variations of hole dimensions, pellet weight, and pressure-density relationship that enter into the determination of the relative location of the surface thru which the reconsolidation pressure is applied. Where this dimension is critical, the reconsolidation is done to a stop so that the tolerances appear in the density of the reconsolidated pellet. When reconsolidation is specified, the effects of these variations upon performance should be considered.

### Special Procedures

1) **Vacuum Pressing**

In the usual pressing operation, in which a granular expl is pressed from a bulk density of about half the crystal density to about 95% of the crystal density, the press rise in the interstitial gases (assuming isothermal compression and no leakage) may be in the neighborhood of 200 psi. The air may be presumed to diffuse out of the pellet, thru the continuous pores, quite rapidly after the pellet is ejected or the ram is removed, if it has not already leaked thru the clearance between ram and cavity during pressing.

When densities reach 99% of crystal density, the cycled press of the interstitial gases rises rapidly, limiting attainable densities. When under conditions of pressing, the expl or some component of it is caused to flow plastically, the pores may be closed into individual bubbles in which the compressed gases are retained to cause excessive growth after press removal or pellets that pop open when ejected from the die. In an open pore material, the relatively mobile gases tend to increase density gradients by distributing press without a correspondingly even distribution of the solid expl. For these three reasons, vacuum pressing is used where very high or uniform densities are required, or where significant plastic flow is anticipated during pressing.

Fig 6 is a diagram of a vacuum loading tool. First, lower and top punches are advanced to a prepress position to compact the powder slightly. After evacuating to 1 mm Hg, full press is applied. Production of extremely high quality charges of TNT (pressed at elevated temp) and Composition A-3 (both at elevated and room temp) has been reported. Density spreads within 6-in diameter charges are 0.005 g/cc (Ref 2).

![Vacuum Pressing Apparatus](image-url)
2) Hot Pressing

The unique props of plastic bonded expls
are realized most fully if they are pressed at
elevated temps. Appropriate temps, of course,
are detd by the props of the plastic bonding
agents used and limited by the thermal insta-
bility of the expls. Temps as high as 130°
have been used (Ref 9). When heated to temps
approaching their melting points, expls and ad-
ditives used in expls, like most solids, are more
prone to plastic flow. Equipment required for
hot pressing of PBX has been found useful in
the production of high density charges of con-
ventional expls. TNT is pressed routinely to a
density of 1.62g/cc at 70° in the vacuum press-
ning process previously described, whereas cast
densities this high are unusual (Ref 2). Pre-
heating of the expl is more efficient than
waiting for it to heat in the mold but cannot
be used when thermoetting resins serve as
binder

3) Hydrostatic and Isostatic Pressing

When an expl is pressed in a die by means
of a ram, the fraction of the walls tends to
cause press and density gradients. In addition,
the one-dimensional compression can result in
an anisotropic structure and produce pellets
with residual strains. Where dimensional
stability, uniformity and high density are es-
tential to performance, hydrostatic pressing
and isostatic pressing have been used. In both
of these processes, the expl is compressed by
the action of a fluid, from which it is separated
by a rubber (or other elastomer) film

In hydrostatic pressing, the expl is placed on
a solid surface and covered with a rubber dia-
aphragm (Fig 7). Although this process elimi-

ates the gradients which result from wall
friction, some directionality of compression
remains which can result in anisotropic struc-
ture and residual strains. In isostatic pressing,
the explosive is placed in a rubber bag (Fig 8)
that is surrounded by the pressurizing fluid so
that the compression is essentially three di-

mensional

In addition to the production of high
quality charges, hydrostatic pressing and iso-
static pressing can be used to consolidate expls
which are so sensitive that frictional contact
with the walls of a conventional mold creates
a hazard. Materials like pure RDX, of which
it is difficult to make firm pellets except in
small sizes, can often be pressed hydrostatically
or isostatically (Ref 9)

Hydrostatic pressing and isostatic pressing
are usually applied to expls that have been
evacuated, frequently at elevated temps. Temps
up to 130° and pressures up to 30000 psi
have been used. The surfaces where pressure
is applied thru elastic membranes are, of course,
of relatively poorly defined form and dimensions.
Hence, these pressing processes must almost in-
variably be followed by machining

4) Pulsating Pressures

Expts have shown that pressures which
pulsate with an amplitude of a few percent
of the static press and at a frequency of about
60 Hz, when used with conventional molding
tools, make it possible to produce pellets four
or five diameters long with negligible density
gradients. The interesting possibilities of this
technique in production of explosive charges
have not yet been exploited
Finishing Operations

1) Machining

It has been found that the most uniform densities and compositions are attained by pressing or casting relatively large charges, and machining the charges needed from selected segments. Similarly, high quality charges can be made by isostatic or hydrostatic pressing, which also must be followed by machining operations. All standard machine shop operations including milling, drilling, sawing, boring, and turning — are applied in this work.

Many cast loaded items are filled thru the same hole as that into which the fuze is to be inserted. After casting, the sprue is broken off. Although it is a good plan to design the funnel to form a core for the fuze cavity, the problem of funnel extraction limits this practice to some extent. At best, then, the bottom of the fuze cavity is a rough, broken off surface and, generally, the cavity is not as deep as desired. The boring of fuze cavities to the specified depth and surface finish is a routine operation of production.

Profile lathes and forming tools may be used to form almost any desired surface of revolution. The special forms required for detonation wave shaping and other specialized output are often generated by such means. Expls may be machined to the same tolerances as metals. Turning and milling to a thousandth of an inch is not difficult with a good machine. However, the practical applicability of such precision is limited by the dimensional instability of most expol materials.

Safety is an important aspect in machining explosives. Since the sensitivity of an expol has meaning only in terms of the specific initiating impulse, the practice of machining each expol material by remote control is most desirable (Ref 2). On the basis of test data it is considered safe to machine Composition A-3, Composition B, and TNT at 200 ft/min surface speed.

Cut-off tools and small drills are more hazardous because of the poor cooling conditions. These operations, if necessary, should be performed at low speeds with intermittent cutting and frequent flushing. Water should be used wherever practical as a coolant, although tests at high speed under dry conditions are considered justification for dry machining where needed. The water keeps expol dust out of the air and cools the cutting operation.

2) Cementing of Compound Charges

Expol charges made of more than one expol, in which the contour of the boundary is an important design parameter, often are fabricated from cast, pressed, or machined components that are cemented together.

Cements that harden by the loss of solvent generally are to be avoided because the solvent can be lost only by diffusion thru the expol. Diffusion may be slow and the solvent may modify the properties of the expol. Two types of cement that have been used for this purpose are catalytic setting cements, like epoxy resins, and contact cements. Compatibility of the materials to be used should be checked. Compatibility of epoxy resins with most explosives depends upon the catalyst or hardener used (Ref 8). Data regarding bond strengths and other pertinent properties also have been compiled (Refs 5 & 6).

Both surfaces to be cemented should be clean and fit accurately to one another. A minimum continuous layer of cement should be applied to each surface to be joined. Where catalytic resins are used, provision should be made to hold the members in firm contact for the curing period. When contact cements are used, mechanical means are desirable to assure that the elements are in the proper position and orientation when they make the first contact.

Written by S. M. KAYE

Refs:
Skeottini, “Ammunition Loading Techniques”, ExplosDevSectRept 43, PicArsn (1958)
8) M.C. St. Cyr, “Compatibility of Explosives with Polymers”, PATR 2595 (1959) 9) C.C.
Miser, “Capabilities of Explosive Loading Group of the Chemical Engineering Division
(WE)”, NAVORD Rep 6873, NavOrdLab, Silver Spring, Md (April 1960) 10) D. Kite Jr, A.K.
Behlert & E. Jerecerzewski, “Plastic Bonded Explosives for Use in Ammunition”,
ments for Use in Fuzes, MIL-STD-320 (July 1962) 12) S. Levmore, “Principal Charac-
teristics of the Gelled Slurry Explosive DBA-22M”, PATR 4237 (1971) 13) Anon, Eng-
Added Refs: a) C. Claessen, GerP 243981 (1910) (Double-walled steam heated funnel
for shell loading with molten expls) b) C.E. Bishel, GerP 227635 (1910) (Cast loading of
expls under press to increase their d)
c) Anon, GerP 244034 (1911) (to Deutsche Sprengstoff A-G) (High d is achieved by main-
taining a feeding head full of liq expl and subjecting it to press during charge solidification)
d) Anon, GerP 255096 (1911) (to Dynamit A-G) (A cold rod of HE is introduced into
the main axis of a molten charge to increase the rate of cooling, and thus prevent cavity
formation) e) W. Fairweather, BritP 24960 (1911) (Molten HE is poured into a shell
and subjected to mechanical press while the upper part is kept liq by heat, and the lower part is
allowed to solidify by cooling with w)
f) Anon, GerP 279526 (1913) (to Deutsche Sprengstoff A-G) (Cavity-free HE charges of
high d are obtained by casting into cases and subjecting them to centrifugal forces) g)
Colver (1918), 10, 319; 442–483, 671 (Vari-
ous methods of obtaining high d casts of HE are reviewed, eg, direct or press casting, chilled
molds or shells, cold core expl, and centrifuga-
gation) h) Anon, ChemAgc 1, 385–6 & 407–8 (1919) & CA 13, 3318 (1919) (Loading
of TNT & Amatol by melting and pouring, pouring shaped charges into blocks with sub-
sequent loading into shells, pressing block charges and fitting into shells, stemming, and
direct hydraulic pressing) i) C.A. Woodburry,
USP 1329566 (1920) (Loading of shells with a mixt prepd by dissolving PA in TNT and then
adding TNX while maintaining the temp of the mixt above the mp of TNT) j) F.
Hawkes, ArOr 2, 208–11 (1922) (Loading of shells with TNT and Amatol) k) J.P.
Madden & L. Fisher, USP 1420637 (1922) & CA 16, 2993 (1922) (A portion of TNT is
melted and then mixed with about 2.5 parts of solid TNT to form a plastic mixt for filling)
l) Anon, BritP 181030 (1922) & CA 16, 4349
(1922) (To facilitate loading of shells with
TNT or like expl under press, part of the TNT
is first melted and heated to about 85°, and
the remainder is then added dry and well stirred
to form a plastic dough, which is forced into
the shell) m) W.A. Gibbons, USP 1453933
(1923) (Cast loading by partially filling a shell
with solid HE, followed by pouring the HE in
liq form to fill the cavities) n) G.C. Hale,
ArOr 4, 838–40 (1925) (Ger methods for
loading HE shells) o) J.P. Harris, ArOr 7,
40–8 (1926) (Loading of ammo at PicArsn)
p) H.H. Olmstead, USP 1670689 (1928) & CA
22, 2467 (1928) (Molten expls are poured into
shells in successive layers, each about 1/2” in
thickness, and allowed to cool for about 30
minutes before adding the next layer)
q) O. Matter, USP 1903594 (1933) (A soln of
HE is gradually added to a liq in motion which is
miscible with the solvent in any proportions,
but does not dissolve the expl. The pptd expl is
melted to form a fluid mass contg unmelted
granules in suspension, and the resulting broth-
like mass is cast) r) C.R. Dutton, ArOr 20,
389–93 (1940) (Filling of bombs with a
molten PA/TNT mixt) s) H. Shaler, USP
2195429 (1940) & CA 34, 5283 (1940)
(Loading of expls into ammo is done by intro-
ducing an increment of molten expl, and
rotating the ammo on its axis until the increment
has solidified. This operation is then repeated
with successive expl increments) t) M.A.
Cook & C.O. Davis, USP 2353147 (1944)
(High d HE charges are obtained by heating
a mixt of two solid ingredients, one being a
nitrated organic compd, to a temp at which
one of the compds melts. A dispersing agent
is included in the mixt) u) L.A. Quayle,
MechEng 67, 599–606 (1945) (Volumetric
pouring machine for filling mines, bombs &
that deal with (1) design and development, acquisition, storage, movement, distribution, maintenance, evacuation, and disposition of materiel; (2) movement, evacuation, and hospitalization of personnel; (3) acquisition or construction, maintenance, operation, and disposition of facilities and (4) acquisition and furnishing of services. It comprises both planning, including determination of requirements, and implementation.

Ref: OrdTechTerm (1962), 180

**Long Tom.** Popular name for US 155mm self-propelled gun used during WWII. Extremely accurate, it fired a 95-lb proj a distance of 25700 yds. It weighed 15 tons and had a barrel length of about 23 ft. During the assault on the Siegfried Line these guns assisted in the destruction of 120 pillboxes, obliterating one with each round fired from a range of 300 yds.

The name has also been applied to the US 155mm self-propelled gun M-53, currently used by the Army and Marine Corps. It is full tracked to provide mobility for the gun and protection for the crew of six in offensive combat.


**Loading Factor.** Expressed as percentage, is the weight of HE fill of an ordnance item divided by the total weight of the item, multiplied by 100.

**Lobbe Explosive.** Mixt patented in Eng in 1861 contg Na nitrate, sawdust and lime

Ref: Daniel (1902), 411

**Locust-gum.** Proposed as a waterproofing coating for AN grains in so-called “water-resistant” expls

Ref: W.J. Taylor, USP 2654666 (1953) & CA 48, 3692 (1954)

**Logistics.** Those aspects of military operations

---

shells) v) H. Bernstorff & A. Allendörfer, USP 2390052 (1945) (Device for stirring expmets) w) H. Graham et al, Canad Chem & Process Indus 30, 37–41 (1946) (Bomb filling with 80/20 Amatol) x) A.N. Campbell & E.J. Pritchard, CanJRes 25B, 183–97 (1947) & CA 41, 4647 (1947) (It was hypothesized that cavitation & shrinkage in expl charges could be eliminated if it were possible to slow crystallization so that the melt would remain liq for a while at RT. Attempts to prepare expl mixts with such properties were unsuccessful) y) I.E. Aske, USP 2439443 (1948) (Casting app for vibrating molds) z) P. Campbell & W.H. Maxwell, USP 2439450 (1948) (App compensating for shrinkage of cast expls) za) L.F. Audrieth & D.D. Sager, USP 2482089–91 (1949) (A cast expl obtained by pouring a suspension of RDX or EDNA into a liq melt of TNT contg Tetryl or PETN) zb) A.R.V. Steele et al, BritP 765210 (1957) (A head of liq expl held in a hollow former provides a flow of expl to compensate for any contraction as solidification occurs) cc) J.G. Holmes, Ordn 42, 193–4 (1957) (Application of single-pour controlled cooling to filling bombs and other expl ordnance) dd) S.D. Stein & M.J. Margolin, “Proposed Shell Loading Standards Based on a Statistical Study of Setback Sensitivities”, PATR 2563 (1958)
Lovelace. Invented a priming expl consisting of K chlorate, PA & MF, and an expl powd contg K chlorate, PA & charcoal
Ref: Daniel (1902), 411

Low Detonation Pressure Explosives. Most expl materials in wide use today may be characterized by deton pressures ranging from approx 150–350 kilobars. PropInt materials, on the other hand, exhibit comparatively low press typical of defgrn reactions. The difference in pressures exhibited by these two classes of materials leaves an interesting gap, the exploration of which may yield valuable information on the propagation and kinetic limitations of deton materials.

The reliable generation of deton press under 100 kilobars should offer advantages from an engineering viewpoint in applications where higher pressures are neither needed nor desired. Certain plastic/expl formulations described (Ref 1) offer these advantages in addition to others, such as the capability of being extruded or injection molded into difficult configurations and then polymerized in place. ExpIs included superfine PETN, acetone fine RDX, dextrinated LA and Thallous Azide

Low Energy Detonating Cord (LEDC). When detog cord is not required directly to initiate HE, but solely to transmit deton from one place to another, it is advantageous to use a cord with very low charge weight. LEDC consists of an extremely small core load of HE, one or two grains per ft, which is contained in a small continuous lead tube. The lead tube is protected by wrappings of paper or cotton cord and an outer jacket of plastic. The countering provides good tensile strength and unlimited w resistance. LEDC has a vel of deton of 24000 ft/sec, slightly faster than standard detong cord (Ref 2)

When used for laying above ground to connect shots in civil engineering, quarrying and open-pit operations, LEDC has the advantage of producing much less noise than the normal grade. This avoids any requirement of covering the cord with earth or sand when used in populous areas (Ref 1)

See also Cord, Detonating in Vol 3 of Encycl, p CS29-R

Low Explosive (LE). An expl which when used in its normal manner deflagrates or burns rather than detonates; that is, the rate of advance of the reaction zone into the unreacted material is less than the vel of sound in the unreacted material. Low expls include proplnts, certain primer mixts, BlkPdr, photoflash pdrs and delay comps. Whether an expl reacts as a HE or LE depends on the manner in which it is initiated and confined. For example, a double base proplnt when initiated in the usual manner is a LE. However, this material can be made to deton if the proplnt is initiated by an intense shock. Conversely, a HE like TNT, under certain conditions, can be ignited by flame and will burn without detonating
Ref: OrdTechTerm (1962), 181-R

Low-Explosive Devices for Performing Mechanical Functions. Known as actuators or cartridge actuated devices (See Encycl 2 (1962), C70–C72) low-explosive devices are small parts in which a proplnt is ignited to produce gas. The gas generated performs such mechanical functions as closing switches, actuating valves, cutting cables, severing bolts, dispensing fluids, inflating bags, or starting engines

Basically, the devices are of two types, small actuators like dimple, bellows, or piston motors and larger pressure cartridges and gas generators. They are compact, efficient, reliable, inexpensive, and deliver a burst of mechanical work in a short period of time.

Actuators provide a force over a small distance while gas generators exploit the energy of expanding gas
Ref: G. Cohn, "Low-Explosive Devices for Performing Mechanical Functions", Proceedings of the New Mexico Section of the ASME (Behavior and Utilization of Explosives in Engineering Design) (2–3 March 1972)
Low Freezing Dynamites (LF Dynamites). See under Dynamite (Freezing of Dynamites Containing NG) in Encycl 5 (1972), D1588-R to D1593-L.

Low Melting Ammonium Nitrate Explosives. See under Ammonium Nitrate Blasting Explosives, High Explosives and Propellants (AN High Explosives) in Encycl 1 (1960), A346-R to A348-L, A352-R.

Low Order Burst. Functioning of a proj or bomb in which the expl fails to attain a high order deton. Usually evidenced by the breaking of the container into a few large fragments instead of a large number of smaller fragments Ref: OrdTechTerm (1962), 181-R.


Low Temperature Effect on Explosive Properties. Tests conducted at temps of $-80^\circ$ and $-183^\circ$ with expls commonly used in detonators showed that the vel of deton and Trauzl lead block expansion values were practically unaffected, while brisance, as indicated by the lead plate test, was greatly reduced. The performance of detonators, as judged by an initiation test on unconfined cartridges contg LOX (qv) expls, was also greatly reduced. Only detonators contg Mannitol Hexanitrate, when immersed in LOX expls for 3 minutes at $-183^\circ$, resulted in complete deton (Ref 1).

Burning, and the initiation, growth, and propagation of expls are often retarded or prevented by very low temps. Tests of blasting caps at liq nitrogen temps showed much decreased sensitivity (Ref 3).

The effect of low temps upon the sensitivities of initiators is usually quite small because the change from room temp is only a small fraction of the rise associated with initiation. However, systems that are marginal with respect to growth or propagation of expl reaction will usually fail in low temp testing. (Ref 3)

The most noticeable effect of low temp upon stable deton results from a shrinkage in vol. Because of the higher d at low temps, deton velocities, and consequently deton pressures, are higher. These increases are too small to have practical significance. Where propagation time is critical, and must be synchronized with a process that is independent of temp, this effect, now accentuated by reduction in distance, can be a source of difficulty (Ref 2)


Lubrication

Fuzes. A lubricant is expected to perform the jobs of minimizing friction, wear, and galling between sliding or rolling parts. It must do these jobs under two types of conditions: (1) those which are inherent in the component element itself—such as load, speed, geometry, and frictional heat—and (2) those which are imposed from external sources—such as temp and comp of the surrounding atm, nuclear radiation, inactive storage, vibration, and mechanical shock. The imposed conditions are usually the more restrictive ones for lubricant selection.

Mechanical fuse components contain elements which undergo a variety of sliding and rolling motions, and combinations of the two. For example, a mass translating on guide rods involves linear sliding only, the balls in a ball bearing involve essentially all rolling motion, and meshing gear teeth surfaces experience both rolling and sliding motions. For any given type of motion, the lubricant found to be satisfactory in one case will not necessarily be suitable for another if loads, speeds, etc., are not similar.

Selection of the proper lubricant requires not only knowledge of the specific function which the lubricant is required to perform in the device being lubricated but also consideration of the interactions include chemical processes—such as corrosion of the metal parts.
by components of the lubricant, eg, corrosion due to oxidation of MoS2 in the absence of suitable inhibitors, or solution of copper alloys during lubricant oxidation processes; or physical interactions, eg, attack by active organic materials on synthetic elastomers and plastic structural members. In addition, the inherent stability of the lubricant must be considered. Stability is of particular importance if storage for long periods of time with or without elevated temps (which speeds up oxidation rate) is involved. (In general, lubricants are inhibited against oxidation by appropriate additives, but since temp is an important parameter, the oxidation stability characteristics of the lubricant should be taken into account in connection with the expected storage life and pertinent temps of the mechanism being lubricated). Oxidation of fluid or semi-fluid lubricants may lead to thickening of the lubricant with consequent increased forces being required for operation, or corrosive attack on the materials of construction.

A wide variety of fluid and semi-fluid lubricants is available covering a wide temp range of applicability, a range of compatibility with organic and inorganic structural materials, and a range of other properties which may be pertinent, eg, nonspreading, lubricity, etc. In addition, both dry powdered and bonded solid-film lubricants are available. The choice of a lubricant depends on the totality of functions which the lubricant must perform, and the structural and functional features of the mechanism being lubricated. For example, a very severe nonspreading and low vapor pressure requirement in connection with long term storage may lead to a choice of a solid lubricant; whereas adhesion problems with bonded lubricants at high loads or with thin films associated with low mechanical tolerances may complicate the use of dry film lubricants. In fuzes subject to high rates of spin (above 25000 rpm), fluid and semi-fluid lubricants tend to be displaced by centrifugal force causing loss of lubricant and possible contamination of other fuze parts. Requirements for corrosion protection may require additives not accessible with dry lubricants.

In simpler fuzes, choice of proper materials, plating, and finishes can obviate a separate lubricant.

Descriptions of available lubricants — oils, greases and solid — with summaries of their properties are contained in Ref 5.

_Space Applications._ All known liquid lubricants and fatty acids evaporate, and they are, therefore, unsuitable for space conditions. Tests showed that liquid lubricants do not even provide adequate lubrication in the lower vacuums of space simulators. Solid lubricants, such as molybdenum disulfide, tungsten disulfide, and the soft metals have given better results. However, the known data about space lubricants are results of simulator measurements made in the pressure range of 10^-5 to 10^-6 torr which does not simulate real space conditions, and therefore, these available data cannot be considered completely valid. It can be expected that definite data on lubricant performance in a vacuum will be obtained by conducting tests in a simulator that reaches the low 10^-10 torr range. At this pressure level, the monolayer formation time is increased to at least several hours which will result in a sufficient time span for observing the metal surfaces (Ref 6).

_Explosives._ Lubricants are utilized in HE's to minimize compn adherence to dies used for pelleting and press loading operations. Typical of materials used with Tetraly have been graphite, stearic acid, Ca, Ba and Mg stearates (Refs 1, 2 & 3). Forchielli (Ref 4) reported on the beneficial effects of the metallic stearates (Ca, Li, Co, Zn & Cr) and talcum pdr with Composition A-3.

Luck and Cross. Prep’d, in 1898, proplont powds by adding aq solns of Pb and Zn acetates to NC as stabilizers. This addition could be conducted in the presence of 30–50% acetone, after which the mix was dried
Ref: Daniel (1902), 412

Luck and Durnford. Proposed, in 1896, to prepare proplont powd from nitrated hydrocellulose which had been treated with aq starch soln. This allowed the material to be compacted into grains, blocks, leaflets, etc
Ref: Daniel (1902), 411-12.

Luck Powder. A proplont proposed, in 1900, for use in shotguns. It consisted of NC powd in which the NC was replaced by either cellulose acetate, butyrate or benzoate. This substitution permitted the prep of stocking powds with a slow rate of combustion
Ref: Daniel (1902), 413

Addnl Refs: 1) A. Michel-Lévy & H. Muraour, CR 206, 1566–8, 1938) & CA 32, 5629 (1938) (Luminosity of expls) 2) H. Muraour, A. Michel-Lévy & J. Rouvillois, CR 208, 508-10 (1939) & CA 33, 3157 (1939) (Luminosity of expls) 3) H. Muraour, Chim&Ind (Paris) 47, 3–15 (1942) & CA 37, 6461 (1943) [Luminosity has its origin in the shock waves generated by expls, and not in emitted gas as claimed by Laffite, CA 20, 1324 (1926)]
4) A. Michel-Lévy & H. Muraour, CR 224, 695–6 (1947) & CA 41, 5723 (1947) (Origin of luminosity which accompanies the firing of expls in untapped drill holes. It is not a function of the gas evolved, but of the amount of air in the hole) 5) L. Defret & P. vanWouwer, Chim&Ind (Paris) 69, 1086–7 (1953) & CA 49, 6608–9 (1955) (Luminosities produced by expl shock waves)

Lundholm and Sayers. Proposed in 1899 (BritP 10376) methods for the manuf of several expls, eg, mixts of NC with nitroaromatic compds, mixts of oxy- or hydro-NC with various organic compds, etc. They also proposed charging projectiles by introducing expls in layers
Ref: Daniel (1902), 413

Lunge, Georg. (1839–1923). Ger chemist, noted for work on technological analytical methods. Inventor of the nitrometer (See below); author of numerous publications, among them (in collaboration with E. Berl), “Chemisch-technische Untersuchungsmethoden”, a classic work on technical analysis
Ref: Hack’h (1944), 501-R

Lunge Nitrometer. An app designed by G. Lunge for the detn of nitrogen in either organic or inorganic nitrates. It is so designed that the nitrate or nitric ester is dissolved in conc sulfuric acid and the soln, without entrained gas, is afterwards admitted to a reaction vessel. The nitric oxide gas from the reaction is measured in a mercury eudiometer in cc at atm press, a barometer and thermometer are read, and the wt of nitrogen in the nitric oxide and the percentage of nitrogen in the sample are calc’d (Ref). This app was subsequently modified by the DuPont Co, and is known as the DuPont Nitrometer (See Encycl 1 (1960), A373-L to A377-L for description of use)
Ref: Davis (1943), 270–1

Lupersol DDM. A proprietary catalyst of Wallace & Tiernan, Inc, Buffalo, NY, contg 60% methylethylketone peroxide in dibutylylphthalate, used as the hardener in Lamincan (qv)
Ref: 1) OrdiTechTerm (June 1962), 183-R
LX Explosives. A code employed by the Lawrence Livermore Laboratory, Univ of California (Livermore), USA, to designate formulations in production. A specific code designation is assigned to an expl when the state of development of its formulation has reached the point where a set of reasonable manufg specifications can be written, the evaluation of the material’s chemical, physical, expl props and sensitivity is essentially complete, and the material has a definite application.

This code consists of the two letters LX followed by a dash, two digits, a second dash, and finally a single digit. The first pair of digits is merely an arbitrary serial number assigned in sequence, while the last digit denotes a subclass in the series. Thus, LX-01-0, LX-02-1, ..., LX-05-0, etc. The last digit provides for small changes in manufg specifications. For example, when LX-04-0 has undergone a revision of expl particle size, new lots, manufd under the revised specification, are identified as LX-04-1.

LX-01 is a liq material, characterized by a wide liq range (−65° to +165°F), moderate energy release, and good stability and sensiti-vity props. Caution: The TNM component is moderately volatile and highly toxic.

LX-02 is a material of putty-like texture, characterized by ability to propagate in very small diameters.

LX-04 is a solid expl characterized by excellent mechanical and compatibility props, an energy release about 9% less than LX-09 and sensitivity props much superior to LX-09.

LX-07-2 is a modification of LX-04 with a higher energy release (5% less than LX-09), obtained at the expense of some degradation in mechanical props (less elongation, etc) and in sensitivity.

LX-08 is an extrudable, curable expl developed for use in Dautremond timing tests.

LX-09 is similar to the LASL expl PBX-9404, but with significantly improved thermal stability and slightly poorer physical props.

LX-10 is in the same energy class as LX-09 and PBX-9404, but utilizing HMX and Viton A like LX-04, and having excellent thermal characteristics. It also exhibits high creep resistance, but may be somewhat more sensitive than the others.

LX-11 is like LX-04, but intentionally degraded in energy by adding an addnl 5% binder.

LX-13 is a variant of the LASL expl XTX-8003.

Ref: B.M. Dobratz, “Properties of Chemical Explosives and Explosive Simulants”, Lawrence Livermore Laboratory, Univ of California (Livermore), UCRL-51319 (15 Dec 1972), pp 3-3, 3-4, 17-1, 17-2.

Typical LX formulations are given in the following table:

<table>
<thead>
<tr>
<th>Explosive</th>
<th>Ingredient</th>
<th>Weight %</th>
<th>Color</th>
</tr>
</thead>
<tbody>
<tr>
<td>LX-01</td>
<td>Nitromethane</td>
<td>51.7</td>
<td>Clear</td>
</tr>
<tr>
<td></td>
<td>Tetranitromethane</td>
<td>33.2</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1-Nitropropane</td>
<td>15.1</td>
<td></td>
</tr>
<tr>
<td>LX-02-1</td>
<td>PETN</td>
<td>73.5</td>
<td>Buff</td>
</tr>
<tr>
<td></td>
<td>Butyl rubber</td>
<td>17.6</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Acetylatedulcitrone</td>
<td>6.9</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cab-O-Sil *</td>
<td>2.0</td>
<td></td>
</tr>
<tr>
<td>LX-04-1</td>
<td>HMX</td>
<td>85.0</td>
<td>Yellow</td>
</tr>
<tr>
<td></td>
<td>Viton A **</td>
<td>15.0</td>
<td></td>
</tr>
<tr>
<td>LX-07-2</td>
<td>HMX</td>
<td>90.0</td>
<td>Orange</td>
</tr>
<tr>
<td></td>
<td>Viton A</td>
<td>10.0</td>
<td></td>
</tr>
<tr>
<td>LX-08</td>
<td>PETN</td>
<td>63.7</td>
<td>Blue</td>
</tr>
<tr>
<td></td>
<td>Silicone rubber</td>
<td>34.3</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cab-O-Sil</td>
<td>2.0</td>
<td></td>
</tr>
<tr>
<td>LX-09-1</td>
<td>HMX</td>
<td>93.3</td>
<td>Purple</td>
</tr>
<tr>
<td></td>
<td>p-DNPA</td>
<td>4.4</td>
<td></td>
</tr>
<tr>
<td></td>
<td>FEFO ***</td>
<td>2.3</td>
<td></td>
</tr>
<tr>
<td>LX-10-0</td>
<td>HMX</td>
<td>95</td>
<td>Bl-grn spots</td>
</tr>
<tr>
<td></td>
<td>Viton A</td>
<td>5</td>
<td>on white</td>
</tr>
<tr>
<td>LX-11-0</td>
<td>HMX</td>
<td>80</td>
<td>White</td>
</tr>
<tr>
<td></td>
<td>Viton A</td>
<td>20</td>
<td></td>
</tr>
<tr>
<td>LX-13</td>
<td>PETN</td>
<td>80</td>
<td>Green</td>
</tr>
<tr>
<td></td>
<td>Silicone rubber</td>
<td>20</td>
<td></td>
</tr>
</tbody>
</table>

* Cab-O-Sil — trademark for colloidal silica particles
** Viton A — trademark for a synthetic rubber derived from vinylidene fluoride and hexafluoropropylene
*** FEFO — bis (2-fluoro-2-dinitroethyl) formal
Lycopodium (Club-Moss or Vegetable Sulfur). Fine yellow powd consisting of flammable spores of Lycopodium clavatum, which grows in North America, Asia and Europe (Ref 3). It has been used in pyrotechnic formulations, eg, in green stars: Ba perchlorate 70.5, Mg powd 11.8, shellac 11.8 & lycopodium powd 5.9% (Ref 2). It is also a constituent of some expls (Ref 1), eg, in Leonard's smokeless powds (qv)
Refs: 1) Daniel (1902), 405 2) Davis (1943), 86 3) CondChemDict (1971), 530

Lyddite. Br expl, similar in comp to the Fr Melinite (qv). The name was derived from the town of Lydd, near which the expl was manufd and tested. It contains either PA alone, or in admixture with about 10p of other aromatic hydrocarbons, added principally to lower the mp of the PA. Lyddite was adopted in 1888 for charging torpedo warheads, and later during the South African war, for HE projectiles
Refs: 1) Daniel (1902), 414 2) Colver (1918), 14 3) Bebie (1943), 95 4) Davis (1943), 166

Lysol and Nitrated Derivative
Lysol. Brown, oily liq; creosote odor; poisonous. A mixt of alkali compounds of the higher phenols with fat and resin soaps. Obtained by boiling a mixt of heavy tar-oils, fats and resin with alkali. Sol in w, alc, eth, chl & benz (Ref 4). A registered trademark of Lehn & Fink Products Co, Bloomfield, NJ, for a cresvlic disinfectant and antiseptic (Ref 3). On nitr-ation it yields an expl
Nitrolysol (Nitrated lysol). Mp about 64°. Prepd by one-stage nitrination of lysol. It has been proposed for use as an additive to reduce the mp of other expls such as PA. It is also suitable for the manuf of plastic expls, and for coating hygroscopic substances such as Na nitrate, to protect against moisture (Refs 1 & 2)
Refs: 1) E. Raynaud, FrP 449785 (1912) & CA 7, 2687 (1913) 2) Colver (1918), 692 3) Hack's (1944), 503 4) CondChemDict (1950), 409; not found in later editions