Explosion Temperatures of Metals and Other Elemental Dust Clouds

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Abstract

The Pittsburgh Research Laboratory of the National Institute for Occupational Safety and Health conducted a study of the explosibility of various metals and other elemental dusts dispersed in air, with a focus on the experimental explosion temperatures. The data are useful for understanding the basics of dust cloud combustion, as well as for evaluating the explosion hazards in the minerals and metals processing industries. The dusts studied included boron, carbon, magnesium, aluminum, silicon, sulfur, titanium, chromium, iron, nickel, copper, zinc, niobium, molybdenum, tin, hafnium, tantalum, tungsten, and lead. The dusts were chosen to cover a wide range of physical properties – from the more volatile materials such as magnesium, aluminum, sulfur, and zinc to the highly "refractory" (very low volatility) elements such as carbon, niobium, molybdenum, tantalum, and tungsten. These flammability studies were conducted in a 20-L chamber, using strong pyrotechnic igniters. The experimental data obtained included the minimum explosible concentrations, maximum explosion pressures, and maximum explosion temperatures. A unique multiwavelength infrared pyrometer was used to measure the temperatures. For the elemental dusts studied, all ignited and burned as air-dispersed dust clouds except for nickel, copper, molybdenum, and lead. The measured maximum explosion temperatures ranged from ~1550 K for tin and tungsten powders to ~2800 K for aluminum, magnesium, and titanium powders. The measured temperatures are compared to the calculated, adiabatic flame temperatures. In general, the dusts whose experimental temperatures were closer to the adiabatic values were those with the finer particle sizes, those that were more easily vaporized, and/or those that were intrinsically more reactive.

Disclaimer: The findings and conclusions in this report are those of the authors and do not necessarily represent the views of the National Institute for Occupational Safety and Health.

1. Introduction

This paper will concentrate on the explosion characteristics of metal and nonmetal elemental dusts. These dusts are important from both the practical and fundamental points of view. These metal and nonmetal elements are often used or occur in industry as powders or dusts, and an evaluation of their possible explosion hazards is important. In addition, the elemental dusts are ideal for studying the fundamental physicochemical processes occurring during combustion because they are pure, uniform substances with well-characterized phase transitions and thermodynamic properties. This is in marked contrast to the complex structures and heterogeneous chemistry associated with the combustion of carbonaceous materials such as coal and grain dusts. The elemental dusts also display a marked variation of reaction

\[\text{Kashdollar, CDC} \]
exothermicities in air and an enormous range in vapor pressures at their respective flame temperatures.

In the 1960's and earlier, the U.S. Bureau of Mines at Pittsburgh studied the explosibility of metal dusts using a cylindrical 1.2-L Hartmann chamber (Jacobson, Cooper, & Nagy, 1964) with an electric spark ignition source. More recent Pittsburgh Research Laboratory (PRL) studies used a nearly-spherical 20-L chamber and much stronger pyrotechnic ignitors to study the explosibility of metal and other elemental dusts (Hertzberg, Zlochower, & Cashdollar, 1991; Hertzberg, Zlochower, & Cashdollar, 1992; Cashdollar, 1994). The current paper is a continuation of these PRL studies, with an emphasis on experimental explosion temperatures measured with a six-channel infrared (IR) pyrometer. The current paper is also complementary to an earlier experimental study of coal dust explosion temperatures (Cashdollar & Hertzberg, 1983), using the same IR pyrometer. Other explosion characteristics measured for the metal and other elemental dusts included minimum explosible concentrations (lean flammable limits), maximum explosion pressures, and rates of pressure rise. In this paper, the terms "flammability" and "explosibility" are used interchangeably to refer to the ability of an airborne dust cloud to propagate an explosion after it has been initiated by a sufficiently strong ignition source. The explosions observed were rapid deflagrations, not detonations. The infrared temperatures and other experimental explosion data provide evidence to evaluate the general mechanisms of homogeneous versus heterogeneous combustion of these elemental dusts. Some preliminary metal explosion temperature data for iron dust are in Cashdollar (1994; 2000).

2. Experimental Equipment and Test Procedures

The dust flammability experiments in this paper were conducted in the PRL 20-L laboratory chamber (Cashdollar & Hertzberg, 1982a; Cashdollar, 1994), which has been used extensively to study the explosibility of various dusts. The chamber (Fig. 1) is near-spherical in shape and made of stainless steel, with a pressure rating of 21 bar. The vertical cross section is merely a schematic that is meant to show the chamber itself and the vertical positioning of the instrumentation; the positions of the instrumentation around the circumference are shown in the horizontal cross section. The chamber top is hinged and opens across the full chamber diameter. The hinged top is attached by six 3-in (19-mm) diameter bolts which are not shown on the drawings. Two PRL optical dust probes (Liebman, Conti, & Cashdollar, 1977; Cashdollar, Liebman, & Conti, 1981) were used to measure the uniformity of the dust dispersion at the positions shown in Fig. 1. The optical probes measured the transmission through the dust cloud. Thin jets of air kept the probe windows dust-free. For the majority of the tests, one dust probe with a 38-mm path length and a second dust probe with a 95-mm path length were used. The longer path length is more suitable for dusts with higher densities and larger particle sizes. The strain gauge pressure transducer measured the absolute explosion pressure and rate of pressure rise.

The patented six-channel IR pyrometer (Cashdollar, Hertzberg, & Litton, 1979; Cashdollar & Hertzberg, 1982b) measured the explosion temperatures by observing the flame radiation through the sapphire window in the top of the 20-L chamber. The pyrometer uses room

2The majority of this research was conducted in the early 1990s when the PRL was part of the U.S. Bureau of Mines before its transfer to the National Institute for Occupational Safety and Health (NIOSH) in 1996.
temperature lead selenide photoconductive detectors and infrared interference filters to measure flame radiation simultaneously at six wavelengths: 1.57, 2.30, 3.84, 4.42, 4.57, and 5.00 μm. The pyrometer was calibrated over the temperature range 1200 to 1450 K using a large blackbody furnace. The linearity of the detectors was confirmed at even higher radiation levels, allowing the extrapolation to higher temperatures. The pyrometer measured the continuum radiation from the particles in the flame. The experimental explosion temperature was calculated by fitting the measured flame radiation data to the Planck equation as modified for non-blackbodies (Menzel, 1955; Cashdollar & Hertzberg 1982b):

$$H = \frac{2\alpha \varepsilon \hbar c^2}{\lambda^5} \left[ \exp\left(\frac{\hbar c}{\lambda kT}\right) - 1 \right]^{-1},$$  \text{ eqn (1)}

where $H$ is the observed radiation, $\alpha$ is an adjustable scale factor, $\varepsilon$ is the gray emissivity, $\hbar$ is Planck’s constant, $c$ is the speed of light, $\lambda$ is the wavelength, $k$ is Boltzmann’s constant, and $T$ is the absolute temperature. The combined scale factor $\alpha\varepsilon$ is dependent on the flame emissivity, the fraction of the view field filled by the flame, the transmission through any unburned dust on the sapphire window, and the detector sensitivity which varies with ambient temperature. Additional details on the infrared pyrometer and the Planck curve fitting procedure are in Cashdollar & Hertzberg (1982b). The advantages of this multiwavelength pyrometer are that temperatures can be calculated without knowing the flame emissivity and that observations can be made through windows that are partially obscured by unburned dust. This pyrometer has been previously used to measure explosion temperatures of coal dust clouds in an 8-L chamber at PRL (Cashdollar & Hertzberg, 1983), and it was also used to measure temperatures of coal dust flames on a flat-flame burner at Brigham Young University (Smoot & Horton, 1978). A similar
pyrometer has been used by Mackowski, Altenkirch, Peck, & Tong (1983) to measure
temperatures of coal dust flames on a burner.

For each 20-L chamber explosion test, the data from the various instruments were
collected by a custom, high-speed personal computer (PC) based data acquisition system (DAS)
that has the capability of smoothing the data and searching for peaks. The DAS is programmed
to conduct an iterative, nonlinear, least square fit of the pyrometer radiation data to the Planck
equation in order to calculate the explosion temperature versus time. The number of pyrometer
wavelengths used for the temperature calculation can be varied from three to six. After each
explosion test, the pressure, dust probe transmission, and IR temperature data were displayed
versus time.

The test procedures for the 20-L chamber are briefly described here. Additional details
of the 20-L chamber and test procedures are in Cashdollar & Hertzberg (1982a) and Cashdollar
(1994). For most of the tests, the dust was placed on top of the dispersion nozzle rather than in
the dust reservoir. After the dust and ignitor were placed in the chamber, the hinged top was
closed and bolted, and then the chamber was partially evacuated to an absolute pressure of
0.14 bar. Then a 0.3-s blast of dry air (from a 16-L reservoir at 8 to 9 bar pressure) dispersed
the dust and raised the chamber pressure to about 1 bar at ignition. The experimental dust
concentration reported for the 20-L chamber is the mass of dust divided by the chamber volume.
The ignition sources used for the 20-L tests were electrically activated, pyrotechnic ignitors
manufactured by Fr. Sobbe of Germany. For the 20-L tests reported in this paper, 2500-J
ignitors were used. This is the energy recommended in ASTM E1515 (2005b) for the
measurement of minimum exploisible dust concentration. The energy is the nominal calorimetric
value based on the mass of pyrotechnic powder in the ignitor. The 2500-J ignitor by itself
produces a pressure rise of about 0.3 bar in the 20-L chamber.

3. Experimental Results and Discussion

The metal and nonmetal elemental dusts studied are shown in Fig. 2, highlighted at their
positions within the periodic table. Nineteen elements were studied, including boron (B),
carbon (C), magnesium (Mg), aluminum (Al), silicon (Si), sulfur (S) titanium (Ti),
chromium (Cr), iron (Fe), nickel (Ni), copper (Cu), zinc (Zn), niobium (Nb), molybdenum (Mo),
tin (Sn), hafnium (Hf), tantalum (Ta), tungsten (W), and lead (Pb). Table 1 lists the pertinent
physical data for the dusts. The first columns list the atomic number, elemental symbol, and
name, with a sample number listed for elements that were tested at more than one particle size.
The next column lists the density for each element. The size data are listed in three ways.
D(SEM) is an estimate of the size range for each dust as observed with a scanning electron
microscope. Next is the surface mean diameter as calculated (Cashdollar, Liebman, & Conti,
1981) from transmission measurements made with the optical dust probes in the 20-L chamber,
using the following equation:

\[
\tau = \exp \left( \frac{3OC_m}{2\rho D_s} \right),
\]

**eqn (2)**

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\(^3\)Mention of any company or product does not imply endorsement by NIOSH.
where $\tau$ is the transmission, $Q$ is the extinction coefficient, $C_m$ is the mass concentration, $\rho$ is the density of a particle, and $D_s$ is the surface mean diameter of the dust particles. The last column lists the estimated median diameter from a combination of SEM, dust probe, and other size data (sieving and Coulter Counter analyses).

Fig. 2. Periodic table of the elements, highlighting the dusts tested.

Scanning electron photomicrographs of six of the elemental dusts are shown in Fig. 3. A 10 $\mu$m size marker is shown at bottom center of each photomicrograph. The magnesium (Mg) in Fig. 3A is a flake. The aluminum (Al-3) in Fig. 3B, the iron (Fe-1) in Fig. 3D, and the tin (Sn) in Fig. 3F contain rounded particles, with many approximately spherical. The titanium (Ti) in Fig. 3C contains elongated or rodlike particles. The niobium (Nb) in Fig. 3E contains particles with sharp edges. The iron is the finest in size of these six dusts, as shown in the SEM photomicrographs and the listings in Table 1.

Examples of the pressure and radiation (at 2.3 $\mu$m wavelength) traces versus time are shown in Fig. 4A and 4B. These are from an iron dust (Fe-1 in Table 1) explosion in the 20-L chamber at a concentration of 600 g/m$^3$. The radiation reaches its maximum value before peak pressure because the flame front reaches the top window before combustion is completed in the lower parts of the chamber. The calculated temperature and standard deviation in temperature are shown in Fig. 4C and 4D. The maximum in the temperature trace (1740 K) occurs at the same time as the maximum in radiation, as expected. The standard deviation in the calculated temperature is fairly high at the start of the explosion while the amount of radiation is low. The standard deviation drops to a value of 25 K when the temperature reaches its maximum value.
Table 1. - Physical data for elemental dusts.

<table>
<thead>
<tr>
<th>Element</th>
<th>Dust</th>
<th>Density, g/cm³</th>
<th>D(SEM), μm</th>
<th>Dₜ(%τ), μm</th>
<th>Dₚ, μm</th>
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<tbody>
<tr>
<td>5 B</td>
<td>Boron</td>
<td>2.37</td>
<td>0.5-10</td>
<td>2-6</td>
<td>-3</td>
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<tr>
<td>6 C</td>
<td>Carbon-1</td>
<td>~2.1</td>
<td>0.2-2</td>
<td>#2</td>
<td>#1</td>
</tr>
<tr>
<td></td>
<td>-2</td>
<td>*</td>
<td>4-10 f</td>
<td>3-8</td>
<td>-1</td>
</tr>
<tr>
<td></td>
<td>-3</td>
<td>*</td>
<td>4-30 f</td>
<td>5-7</td>
<td>-4</td>
</tr>
<tr>
<td>12 Mg</td>
<td>Magnesium</td>
<td>1.74</td>
<td>10-50 f</td>
<td>30-80</td>
<td>-16</td>
</tr>
<tr>
<td>13 Al</td>
<td>Aluminum-1</td>
<td>2.70</td>
<td>3-30 f</td>
<td>-2-4</td>
<td>-1</td>
</tr>
<tr>
<td></td>
<td>-2</td>
<td>*</td>
<td>5-15</td>
<td>-10-14</td>
<td>-7</td>
</tr>
<tr>
<td></td>
<td>-3</td>
<td>*</td>
<td>3-60</td>
<td>-40-70</td>
<td>-15</td>
</tr>
<tr>
<td></td>
<td>-4</td>
<td>*</td>
<td>20-100 r</td>
<td>-60-180</td>
<td>-40</td>
</tr>
<tr>
<td>14 Si</td>
<td>Silicon</td>
<td>2.33</td>
<td>1-50</td>
<td>3-8</td>
<td>-4</td>
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<tr>
<td>16 S</td>
<td>Sulfur</td>
<td>2.07</td>
<td>~10-50</td>
<td>-60-100</td>
<td>-35</td>
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<tr>
<td>22 Ti</td>
<td>Titanium</td>
<td>4.54</td>
<td>10-60 r</td>
<td>~30-120</td>
<td>~25</td>
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<tr>
<td>24 Cr</td>
<td>Chromium</td>
<td>7.20</td>
<td>4-40</td>
<td>10-18</td>
<td>-10</td>
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<tr>
<td>26 Fe</td>
<td>Iron-1</td>
<td>7.87</td>
<td>1-8</td>
<td>4-11</td>
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<td></td>
<td>-2</td>
<td>*</td>
<td>10-40</td>
<td>-60-160</td>
<td>-45</td>
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<tr>
<td>28 Ni</td>
<td>Nickel-1</td>
<td>8.90</td>
<td>3-8</td>
<td>-30-45</td>
<td>-6</td>
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<tr>
<td>29 Cu</td>
<td>Copper</td>
<td>8.96</td>
<td>10-50</td>
<td>-30-60</td>
<td>-30</td>
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<tr>
<td>30 Zn</td>
<td>Zinc-1</td>
<td>7.13</td>
<td>6-50 f</td>
<td>3-8</td>
<td>-4</td>
</tr>
<tr>
<td></td>
<td>-2</td>
<td>*</td>
<td>10-100 r</td>
<td>-100-200</td>
<td>-45</td>
</tr>
<tr>
<td>41 Nb</td>
<td>Niobium-1</td>
<td>8.57</td>
<td>4-50</td>
<td>-30-55</td>
<td>-20</td>
</tr>
<tr>
<td></td>
<td>-2</td>
<td>*</td>
<td>10-60</td>
<td>-50-100</td>
<td>-30</td>
</tr>
<tr>
<td>42 Mo</td>
<td>Molybdenum</td>
<td>10.2</td>
<td>3-6</td>
<td>10-23</td>
<td>-5</td>
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<tr>
<td>50 Sn</td>
<td>Tin</td>
<td>5.75</td>
<td>2-20</td>
<td>14-30</td>
<td>-8</td>
</tr>
<tr>
<td>72 Hf</td>
<td>Hafnium</td>
<td>13.31</td>
<td>4-30</td>
<td>-7-13</td>
<td>-10</td>
</tr>
<tr>
<td>73 Ta</td>
<td>Tantalum</td>
<td>16.6</td>
<td>4-30</td>
<td>-20-60</td>
<td>-15</td>
</tr>
<tr>
<td>74 W</td>
<td>Tungsten-1</td>
<td>19.3</td>
<td>#1</td>
<td>1-3</td>
<td>#1</td>
</tr>
<tr>
<td></td>
<td>-2</td>
<td>*</td>
<td>5-25</td>
<td>-30-100</td>
<td>-10</td>
</tr>
<tr>
<td>82 Pb</td>
<td>Lead</td>
<td>11.3</td>
<td>20-60 r</td>
<td>~100-1000</td>
<td>~40</td>
</tr>
</tbody>
</table>

Note: For the SEM size listing, f denotes flake or plate-like particles and r denotes rod-like particles.

Fig. 5 shows the curve fits to the Planck equation for four dust explosions at high dust concentrations where the flame would be optically thick. The flame radiation, $H/ae$, is plotted versus the wavelength. The respective dust concentrations for the explosions were: 850 g/m³ for the magnesium (Mg), 800 g/m³ for the niobium (Nb-I), 900 g/m³ for the iron (Fe-I), and 1200 g/m³ for the tin (Sn). The pyrometer radiation data in Fig. 5 were measured at the time of peak flame radiation at the 2.3 μm wavelength for each explosion. All four curves show a good fit of the data points to the Planck curve at a particular temperature. As shown in the figure, the relative radiation values at the two shortest wavelengths have the most effect on the temperature calculation. Of the four dusts, the magnesium explosion had the highest temperature - 2740 K, and the niobium had the next highest temperature - 2180 K. The iron had a temperature of 1840 K, and the tin had the lowest temperature - 1590 K.
Explosibility data for three aluminum dusts (Al-2, Al-3, and Al-4 from Table 1) as a function of dust concentration are shown in Fig. 6. The data were measured in the 20-L chamber, using 2,500-J igniters. The explosion pressures, rates of pressure rise, and measured explosion temperatures are shown as a function of dust concentration. The explosion pressure data in Fig. 6C are the maximum measured explosion pressures (with the pressure rise due to the ignitor subtracted) divided by the starting pressure (approximately 1 bar). This corrects for small variations in the starting pressure. Fig. 6B shows the size normalized maximum rate of pressure rise, (dp/dt)$^{1/3}$, for each explosion test. Note that the turbulence level is lower in the PRL 20-L chamber than that recommended in ASTM E1226 (2005a) or NFPA 68 (2002), and therefore these (dp/dt)$^{1/3}$ data are not recommended for vent sizing calculations. The rate of pressure rise data are, however, useful as a relative measure.
of dust reactivity. For each dust size in Fig. 6, the explosibility data show that explosions are not observed below a certain dust concentration. This is the minimum exploisible concentration (MEC) or lean flammable limit (LFL), which is measured by ASTM E1515 (2005b). The criterion used to determine the MEC value under these test conditions is an absolute explosion pressure of 2 bar, or, equivalently, a pressure rise of 1 bar. From the data in Fig. 6, the MEC's with the 2,500-J ignitors for the Al-2, Al-3, and Al-4 dusts are about 90, 90, and 120 g/m³, respectively. Additional MEC data at 5,000-J ignitor energy for these and other dusts are in Cashdollar (1994). For aluminum, the stoichiometric concentration (for formation of Al₂O₃) is \( C_{\text{stoich}} = 310 \text{ g/m}^3 \). The explosion pressure and \( (dp/dt)^{1/3} \) do not reach their maximum values until concentrations well above \( C_{\text{stoich}} \) for the aluminum dust. At even higher dust concentrations, both \( P_{\text{max}} \) and \( (dp/dt)_{\text{max}} V^{1/3} \) level off as all of the oxygen in the chamber is consumed. However, the dusts show no evidence of a "normal" rich limit as would be observed for flammable gases. It should be noted that the efficiency of the dust dispersion is more uncertain at the higher dust concentrations. The maximum explosion pressure and rate of pressure rise are slightly higher for the Al-3 dust than for the larger Al-4 dust. The even finer sized Al-2 dust has a significantly higher maximum explosion pressure and much higher rate of pressure rise than the other two aluminum dusts. The data (not shown in the figure) for the finest sized Al-1 dust were even higher.

![Fig. 6. Explosibility data for three aluminum dusts.](image)

![Fig. 7. Explosibility data for titanium dust.](image)
The explosion temperatures shown in Fig. 6A were measured with the six-wavelength IR pyrometer. The pyrometer observed the continuum radiation from the particles, and temperatures were calculated from the best Planck curve fit to the IR radiation data. The temperature data shown in Fig. 6A were calculated at the time of maximum radiation. This was generally also the time when the standard deviation of the fit to the Planck curve was smallest and the time when the measured temperature was highest. The time of maximum radiation was usually earlier than the time of maximum pressure and probably corresponded to the time when the flame front reached the top sapphire window. The maximum measured particle temperatures for the Al-3 dust were ~2,800 K, well below the maximum calculated adiabatic temperature, $T_{ad,max} = 4,060$ K, for ideal combustion at constant volume. The maximum measured particle temperatures for the Al-4 dust were even lower, although there were only a limited number of measurements for this dust. These experimental temperatures are only those of the particles in the explosion, and the gas temperatures may have been different. There were no experimental temperatures measured for the two finest sizes of aluminum dust (Al-1 and Al-2) because a sapphire window cracked during early testing of the Al-1 dust, and the windows were replaced with steel blanks for later tests of both of these dusts.

Explosibility data from the 20-L chamber for titanium dust as a function of concentration are shown in Fig. 7. The MEC for this dust is about 70 g/m$^3$, using the 2,500-J ignitors. As with the aluminum dusts, the maximum explosion pressures and rates of pressure rise for the titanium dust are at dust concentrations far above the stoichiometric value of 420 g/m$^3$. The maximum measured particle temperatures for the titanium dust were ~2,800 to 3,000 K, well below the maximum calculated adiabatic temperature, $T_{ad,max} = 3,990$ K, for ideal combustion. The maximum experimental explosion temperatures for the titanium dust are comparable to those of the Al-3 dust in Fig. 6.

Explosibility data for two iron dusts (Fe-1 and Fe-2 from Table 1) as a function of concentration are shown in Fig. 8. The data were measured in the 20-L chamber, using 2,500-J ignitors. The explosion pressures, rates of pressure rise, and measured explosion temperatures are shown as a function of dust concentration as in the previous two figures. From the data in Fig. 8, the MEC's for the Fe-1 and Fe-2 dusts are about 220 and 500 g/m$^3$, respectively, using the 2,500-J ignitors. However, there is considerable uncertainty in these values, especially for the Fe-2 dust, due to the scatter in the data. For iron, the stoichiometric concentration (for formation of Fe$_2$O$_3$) is $C_{stoich} = 650$ g/m$^3$. The explosion pressure is close to its maximum value at slightly above $C_{stoich}$ for both iron dusts. However, $(dP/dt)^{1/3}$ is considerably less than its maximum at $C_{stoich}$ and reaches its maximum value at almost twice $C_{stoich}$ for the Fe-1 dust. At the higher dust concentrations, the maximum explosion pressure, $P_{max}$, and $(dP/dt)_{max}V^{1/3}$ level off as all of the oxygen in the chamber is consumed and the dusts show no evidence of a "normal" rich limit. It should be noted that the efficiency of the dust dispersion is more uncertain at the higher dust concentrations, particularly for the dusts with higher density. The explosion temperatures shown in Fig. 8A were measured with the six-wavelength IR pyrometer at the time of maximum radiation. The maximum measured particle temperatures for the Fe-1 dust were ~1,800 K, well below the maximum calculated adiabatic temperature, $T_{ad,max} = 2,490$ K, for ideal combustion at constant volume. The maximum measured particle temperatures for the Fe-2 dust were even lower - about 1,500 to 1,700 K. Again, these experimental temperatures are only those of the particles in the explosion, and the gas temperatures may have been different. For all three
Explosibility characteristics shown in Fig. 8, the Fe-1 dust has higher values than the Fe-2 dust, showing that it is more reactive, due to its finer particle size.

Explosibility data from the 20-L chamber for the two niobium dusts as a function of concentration are shown in Fig. 9. The data were measured in the 20-L chamber, using 2,500-J ignitors. The explosion pressures, rates of pressure rise, and measured explosion temperatures are shown as a function of dust concentration as in the previous figures. From the data in Fig. 9, the MEC's for the Nb-1 and Nb-2 dusts are about 250 and 420 g/m$^3$, respectively, using the 2,500-J ignitors. However, there is considerable uncertainty in these values, especially for the Nb-2 dust, due to the scatter in the data. For both niobium dusts, the maximum explosion pressure and rates of pressure rise are found at dust concentrations at or above the stoichiometric value of C$_{\text{stoich}} = 650$ g/m$^3$. The explosion temperatures shown in Fig. 9A were measured with the six-wavelength IR pyrometer at the time of maximum radiation. The maximum measured particle temperatures for the Nb-1 dust were ~2,100 K, well below the maximum calculated adiabatic temperature, T$_{\text{ad, max}} = 3,540$ K, for ideal combustion. There were no temperature measurements for the Nb-2 dust explosions. As for the other metal dusts, the finer sized niobium dust (Nb-1) appears to be more reactive than the larger Nb-2 dust.

Table 2 summarizes the explosibility data for the elemental dusts tested, based on the data curves of Figs. 6 through 9 and similar data for the other elemental dusts. The first column of the table lists the dust by elemental symbol and sample number. The second column lists the median size from the last column of Table 1. The next column lists the MEC as measured in the 20-L chamber using the 2500-J ignitor. Additional MEC data using a 5000-J ignitor are in
maximum adiabatic explosion pressure is expressed as the ratio of the
rise of the ignitor and normalized to a starting pressure of
Mo, W-2, and Pb dusts were actually determined using the stronger
JANAF tables (Chase, Davies, Downey, Frurip, & McDonald, 1985) and the Thermodynamic Properties of Individual Substances (TPIS) tables (Gurvich, Veyts, & Alcock, 1991). As
discussed for Fig. 6, the listed experimental explosion pressure, \( P_{\text{max}} \), is corrected for the pressure rise of the ignitor and normalized to a starting pressure of 1 bar.a. For the experimental explosion pressure, the values listed in Table 2 are the average of the highest three to six data points for each dust. Note that some of the experimental MEC and \( P_{\text{max}} \) values have been revised

<table>
<thead>
<tr>
<th>Dust</th>
<th>( D_{\text{med}} )</th>
<th>MEC</th>
<th>( P_{\text{ad, max}} )</th>
<th>( P_{\text{max}} )</th>
<th>( C_{\text{ad, max}} )</th>
<th>( T_{\text{ad, max}} )</th>
<th>( P_{\text{equil}} )</th>
<th>( \beta ) fraction</th>
<th>( T_{\text{pyrometer}} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>B</td>
<td>~3</td>
<td>~110</td>
<td>10.5</td>
<td>7.0</td>
<td>130</td>
<td>3300</td>
<td>0.027</td>
<td>&gt; 1</td>
<td>1700×100</td>
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Notes: NF means the dust was nonflammable or nonignitable.
F means the dust was flammable but MEC could not be determined.
from those reported in Hertzberg, Zlochower, & Cashdollar (1992) and Cashdollar (1994), based on additional data.

The sixth column lists the concentration, \( C_{\text{ad,max}} \), at which the calculated adiabatic temperature, \( T_{\text{ad,max}} \) (seventh column) is a maximum. The \( C_{\text{ad,max}} \) values are generally somewhat higher than the \( C_{\text{stoich}} \) values. The adiabatic temperature is also calculated from the NASA-Lewis computer code (McBride & Gordon, 1996) for constant volume combustion. The calculated adiabatic temperatures listed in Hertzberg, Zlochower, & Cashdollar (1992) and Cashdollar (1994) were for constant pressure combustion, which was more appropriate for studying the minimum explosible concentrations (lean flammability limits).

The eighth column lists the ideal equilibrium vapor pressures of the elements at the maximum adiabatic flame temperatures that are calculated using standard thermodynamic tables of the Gibbs free energy of the condensed phases and the product vapors. These values are from the JANAF tables (Chase et al., 1985) and the TPIS tables (Gurvich et al., 1991). That vapor pressure is given by:

\[
P_{\text{equil}} \text{ (bar)} = 1.01 \exp \left( - \frac{\Delta G_v - \Delta G_c}{RT} \right),
\]

where \( \Delta G_v - \Delta G_c \) refers to the free energy difference between the vapor and condensed phases, \( R \) is the gas constant, and \( T \) is the calculated adiabatic flame temperature in kelvins.

The ninth column gives an estimate of \( \beta \), the maximum fraction of the element that is evaporated in the flame at the calculated maximum temperature. That estimate is based on the ideal evaporation rate into an effective vacuum given by the Hertz-Knudsen equation (Nesmeyanov, 1963). The further assumption of a flame residence time of 0.01 sec gives:

\[
\beta = 2.6 \frac{\rho_{\text{equil}}}{d_p \rho} \left( \frac{M}{T} \right)^{1/2},
\]

where \( M \) is the gram-atomic or molecular mass of the vapor species, \( T \) is the flame temperature, \( d_p \) is the particle diameter and \( \rho \) is the particle density (Hertzberg et al., 1992).

The \( \beta \)-values so calculated are idealized estimates where the rate of evaporation in a reactive medium is taken to be equivalent to a vacuum in terms of an assumption of no vapor recondensation. The calculated flame temperature is also an idealized value which assumes perfect mixing, equilibrium conditions, and no heat loss to the walls. Because the assumed flame residence time of 0.01 sec is only an estimate, the calculated \( \beta \)-factor should be considered an order of magnitude estimate of the relative volatilities of the elements at the ideal maximum flame temperature. It is clear from the listed values in Table 2 that most of the elements should be totally volatilized at their maximum flame temperatures. The exceptions are carbon, niobium, molybdenum, tantalum, and tungsten - the most refractory elements, which have only a small fraction of material volatilized. Copper also has a very low volatility, but that is due to its low calculated flame temperature. The larger size of the iron (Fe-2) and the nickel are partially volatilized.

The last column in Table 2 lists the experimental IR temperature, \( T_{\text{pyrometer}} \), in kelvins. As for the experimental explosion pressure, the experimental temperature value listed in Table 2 is the average of the highest three to six data points for each dust. The uncertainty listed for the temperature values is based on the scatter in the data for multiple tests. The standard deviation
for the fit to the Planck equation for an individual test was generally about half that value. There may be some additional uncertainty in the experimental temperatures listed in Table 2 due to the extrapolation of the blackbody calibration of the pyrometer for the higher temperatures or to the possible nongray emissivity of the dust explosion.

As mentioned in the discussion of Fig. 6, experimental temperatures were not measured for the two finest sizes of aluminum dust (Al-1 and Al-2) because a sapphire window had cracked during early testing of the Al-1 dust, and the windows were replaced with steel blanks for later tests of these dusts. These two aluminum dusts had much faster rates of pressure rise than any of the other dusts (Cashdollar, 1994). The listed IR temperatures for the Al-3 and Al-4 in Table 2 are based on the temperatures of the aluminum metal particles in the flame because the combustion product, Al_2O_3, is transparent at all of the IR pyrometer wavelengths. For many of the metals, the observed infrared temperatures are those of the incompletely volatilized metal particles since the metals have higher emissivities than the product oxides, which are often transparent in part or all of the infrared spectrum. There is no listed IR temperature for the Nb-2 dust because there was insufficient material to measure the explosion temperature at high dust concentrations. Experimental particle temperatures were not measured for the sulfur dust because sulfur vaporizes at a low temperature and the combustion product is a gas.

Four of the elemental dusts (Ni, Cu, Mo, and Pb) listed in Table 2 could not be ignited, even with a 5,000-J ignitor. These four dusts showed almost no pressure rise beyond that of the ignitor. It is not surprising that the Cu and Pb did not ignite because their C_{ad,max} values are very high and their T_{ad,max} values are probably too low to sustain flame reaction. The Ni and Mo have reasonably high T_{ad,max} values, similar to that of Fe and higher than those of zinc and tin, but the rate of reaction may be too slow to sustain flame propagation. The larger sizes of several other dusts (C-3, Zn-2, and W-2) could not be ignited, although the Zn-2 showed some slight activity in some of the tests.

The dust with the lowest T_{ad,max} value that produced explosions was zinc with T_{ad,max} = 2070 K. However, the Zn is easily vaporized at flame temperatures. Considering its low T_{ad,max} value, it is not surprising that the larger size of Zn could not be reproducibly ignited with the 2,500-J ignitor.

For most of the experimental temperature calculations in Table 2, all six pyrometer channels were used (see Fig. 5), but in some cases only channels 1, 2, 3, and 6 or channels 1, 2, and 3 were used. For most of the elemental dusts, the emissivity was high. However, for the Si and Ta, the calculated emissivity was low (~0.1). For three dusts (B, Si, and Hf), only the first three pyrometer wavelengths were used to calculate the temperatures, because of higher emissivity at the three longer wavelengths. Because of this, the temperatures for the boron, silicon, and hafnium are more uncertain than those of the other dusts. For the silicon, the combustion product SiO_2 would be transparent at the shorter IR wavelengths and opaque at the longer IR wavelengths. Therefore, the temperature measurement from the three shorter wavelengths would be the temperature of the silicon metal. The radiation at the longer three wavelengths would be a combination of radiation from the Si metal and SiO_2 metal oxide.

4. Conclusions

A summary comparison of the experimentally measured explosion pressures (P_{max}) and the calculated adiabatic pressures (P_{ad,max}) from Table 2 is shown in Fig. 10. The data points for the dusts are identified by the elemental symbols. The lines identify values at 40%, 60%, 80%,
and 100% of the calculated adiabatic pressures. The dusts whose experimental pressures are closest to the calculated adiabatic values are B, Al-1, Al-2, Si, and Fe-1. These are also among the finest sized dusts, with median diameters below 10 μm.

A summary comparison of the experimentally measured explosion temperatures ($T_{pyrometer}$) and the calculated adiabatic temperatures ($T_{ad,max}$) from Table 2 is shown in Fig. 11. The data points for the dusts are identified by the elemental symbols. The lines identify values at 40%, 60%, 80%, and 100% of the calculated adiabatic temperatures. The two dusts whose experimental temperatures are closest to the $T_{ad,max}$ values are Zn-1 and Mg. These are also two of the dusts that are the most easily vaporized (see Table 2).

The aluminum, silicon, titanium, and iron (Fe-1) also have high volatilities at their flame temperatures. These dusts have measured explosion temperatures that are at an intermediate position in the figure, at about 70% $T_{ad,max}$. Two of the dusts with high volatilities, boron and hafnium, have experimental temperatures that are only about half of the $T_{ad,max}$ values. These were also dusts that had a larger than normal uncertainty in their temperature measurements, as

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**Fig. 10.** Comparison of calculated and experimental explosion pressures, with data points shown as element symbols

**Fig. 11.** Comparison of calculated and experimental explosion temperatures, with data shown as element symbols
discussed in the previous section. The flammable dusts (C, Nb, Ta, and W) that have very low volatilities are also those whose measured explosion temperatures are among the farthest from their $T_{ad,max}$ values.

For dusts that were tested at more than one particle size, the experimental temperatures of the finer sizes were closer to the calculated temperatures, as expected. The larger sizes of aluminum (Al-4) and iron (Fe-2) had measured temperatures that were farther from the adiabatic values than the temperatures for the finer sizes. The two sizes of carbon (C-1 and C-2) that produced explosions had about the same measured temperatures, but both were very fine sizes.

In general, the dusts whose experimental temperatures are closer to the adiabatic values are those with the finer particle sizes, those that are more easily vaporized, and/or those that are intrinsically more reactive. However, the boron and hafnium appear to be exceptions since they are both fine size and easily volatilized.

The experimental pressure and temperature data presented here will be useful in evaluating various models of metal dust combustion. Additional information on the mechanisms of metal combustion is in Hertzberg, Zlochower, & Cashdollar (1992) which discusses the volatility of the metals and in Kanury (1975) which has information on combustion properties of carbon and metals. The results reported in this paper provide information on the combustion characteristics of nineteen metal and nonmetal elemental dusts. The experimentally measured temperatures can help in the understanding of the combustion process. However, because of the effects of particle size, these data should not be considered to be representative of all dust samples of these elements. Ideally, explosion temperatures would be measured for identical, small particle sizes of the various elemental dusts in order to study their intrinsic combustion properties, but it would be difficult to obtain these. For the practical use of explosibility data for safety planning, it is important to test the particular size of dust that occurs at a particular industrial plant, rather than trying to extrapolate from other test data for a different size of the same material.

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References


