Detonation of Meta-stable Clusters

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ABSTRACT

We consider the energy accumulation in meta-stable clusters. This energy can be much larger than the typical chemical bond energy (~1 ev/atom). For example, polymeric nitrogen can accumulate 4 ev/atom in the $N_8$ (fcc) structure, while helium can accumulate 9 ev/atom in the excited triplet state $He_2^*$. They release their energy by cluster fission: $N_8 \rightarrow 4N_2$ and $He_2^* \rightarrow 2He$. We study the locus of states in thermodynamic state space for the detonation of such meta-stable clusters. In particular, the equilibrium isentrope, starting at the Chapman-Jouguet state, and expanding down to 1 atmosphere was calculated with the Cheetah code. Large detonation pressures (3 & 16 Mbar), temperatures (12 & 34 kilo-K) and velocities (20 & 43 km/s) are a consequence of the large heats of detonation (6.6 & 50 kilo-cal/g) for nitrogen and helium clusters respectively. If such meta-stable clusters could be synthesized, they offer the potential for large increases in the energy density of materials.

INTRODUCTION

Small meta-stable clusters and molecules are able to accumulate energy that is orders of magnitude larger than the usual exothermic chemical reactions. Energy accumulated in the meta-stable state is released upon transition to the stable equilibrium state with lower energy. Depending on the type of chemical elements of the clusters, the system may transfer into a new stable configuration by either fusion of the clusters into a bulk sample (Type-I clusters), or fission of the large ensemble into small molecules or atoms (Type-II clusters)—analogous to energy release in nuclear matter.

Carbon Clusters

Carbon is a typical representative of the chemical elements that form Type-I clusters. The energy accumulated in carbon clusters increases as the number of atoms in the cluster decreases.
The smallest meta-stable carbon cluster found experimentally is the cage $C_{20}$ discovered by Prinbach [1]. Elesin and co-workers [2,3] have predicted the existence of meta-stable eight-atom carbon cluster $C_8$: Prismane. They have shown that isolated prismanes accumulate energy of $E_{acc} = 2.3 \text{ ev/atom}$ [2,3]. The lifetime of prismane is macroscopically long at room temperature. Prismanes can form quasi-one-dimensional meta-stable ensembles (see Fig. 1).

Prismane ensembles accumulate energy of $E_{acc} = 2.1 \text{ ev/atom}$ [3, 4]. However, the energy released upon fusion of prismanes does not exceed 20% of the accumulated energy $E_{rel} = 0.5 \text{ ev/atom}$ [5]. This is due to the transformation of the ensemble into other meta-stable long-lived configurations. This is a common feature of all Type-I clusters.

**Nitrogen Clusters**

Nitrogen is the typical representative of chemical elements that form Type-II clusters. The most stable form of nitrogen is diatomic molecule $N_2$. Theoretical calculations by Chen [7, 8] have shown that nitrogen clusters $N_4$, $N_8$, $N_{20}$, etc. are meta-stable and accumulate energy up to $E_{acc} = 3 \text{ ev/atom}$. This energy is released by fission of clusters into isolated $N_2$ molecules. Isolated nitrogen clusters have not been synthesized yet.

Elesin and co-workers have shown that $N_8$ boats form meta-stable ensembles while $N_8$ cubanes do not [6,9]. They have shown that $N_8$ boat ensembles (Fig. 2) accumulate the energy of $E_{rel} = 2 \text{ ev/atom}$ [6,9]. Their lifetime is macroscopically long at room temperature. The entire
accumulated energy is released upon fission of the boat ensemble into isolated $N_2$ molecules [6, 9]. This is a common feature of all Type-II clusters.

![Figure 2. Meta-stable $N_8$ boat structure [6,9].](image)

**Helium Clusters**

Helium atoms in the ground singlet state with spin $S = 0$ are chemically inactive and do not form even diatomic molecules. The helium atom $He^*$ in the excited triplet state with spin $S = 1$ is known to be meta-stable with a long lifetime: $\tau \approx 8,000$ seconds [10]. Recently it has been conclusively demonstrated experimentally by McKinsey et al. [11] that two helium atoms can form an excited meta-stable molecule: $He_2^*$ with at total spin $S = 1$. Measured lifetime of $He_2^*$ is $\tau \approx 18$ seconds—in agreement with theoretical predictions ($\tau = 10$ seconds). As shown both theoretically and experimentally, the energy accumulated in the $He_2^*$ molecule is $E_{acc} = 10$ ev/atom, and is released by electron spin flip. Two meta-stable helium quasi-molecules can form the meta-stable cluster $He_4^*$ (Fig. 3) with a total spin of $S = 2$ [6, 12].

![Figure 3. Meta-stable $He_4^*$ cluster [6,12].](image)
The energy accumulated in $\text{He}_4^*$ cluster is $E_{acc} = 9$ ev/atom. This energy is released upon spin-flip of one of the electrons occupying the excited energy level.

Meta-stable structures of $\text{He}_n^*$ clusters based on molecules (quasi-atoms) $\text{He}_2^*$ with the number of atoms up to $n=12$ have been found [6,10]. The energy accumulated in meta-stable clusters of $\text{He}_n^*$, based on molecules (quasi-atoms) $\text{He}_2^*$, is $E_{acc} = 9$ ev/atom for $6 \leq n \leq 12$. It has been shown that such $\text{He}_n^*$ clusters are ordered ferro-magnetically, with the total spin of the cluster $S = n/2$. Also, meta-stable structures of fully-spin polarized $\text{He}_n^{**}$ clusters have been found [6,10], based on excited triplet atoms of $\text{He}_1^*$ with the number of atoms up to $n=3$. The energy accumulated in meta-stable clusters $\text{He}_n^{**}$ based on excited triplet atoms $\text{He}_1^*$ is calculated to be $E_{acc} = 19$ ev/atom for $2 \leq n \leq 4$. It has been shown that $\text{He}_n^{**}$ based on exited triplet atoms $\text{He}_1^*$ are ordered ferro-magnetically, the total spin of the cluster being equal to $S = n$. Results of these studies are summarized in Table 1.

### Table 1. Energy content of meta-stable systems [6, 10]

<table>
<thead>
<tr>
<th>Meta-stable Systems</th>
<th>$E_{stored}$ (ev/atom)</th>
<th>$E_{released}$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Carbon Clusters</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$C_{20}$ Cage</td>
<td>1†</td>
<td>—fission—</td>
</tr>
<tr>
<td>$C_8$ Prismane</td>
<td>2.1</td>
<td>&lt;20%</td>
</tr>
<tr>
<td><strong>Nitrogen Clusters</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$N_4$ Tetrahedron</td>
<td>1.5—3</td>
<td></td>
</tr>
<tr>
<td>$N_8$ Cubanes</td>
<td>2—3.5</td>
<td></td>
</tr>
<tr>
<td>$N_8$ Boats</td>
<td>1.5—2.5</td>
<td>100%</td>
</tr>
<tr>
<td>$N_8$ Boats + $H$ bridges</td>
<td>1.5</td>
<td>100%</td>
</tr>
<tr>
<td>$N_8$ Cubane + $C$ bridges</td>
<td>2—2.5</td>
<td></td>
</tr>
<tr>
<td>$N_{20}$ Cages + $C$ bridges</td>
<td>2—2.5</td>
<td></td>
</tr>
<tr>
<td><strong>Helium Clusters</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\text{He}_2^<em>$ — $\text{He}_8^</em>$</td>
<td>9†</td>
<td>—fission—</td>
</tr>
<tr>
<td>$\text{He}_2^*$ ($n = 2$-12)</td>
<td>9</td>
<td></td>
</tr>
<tr>
<td>$\text{He}_n^{**}$ ($n = 2$-4)</td>
<td>19</td>
<td></td>
</tr>
</tbody>
</table>

† confirmed experimentally
THERMODYNAMICS

In this section we explore the thermodynamic properties of the detonation products gases resulting from the explosion of Type-II meta-stable clusters. In particular, we consider the detonation of $N_8$ clusters and $He_2^*$ clusters.

$N_8$ Detonation

The internal energy ($u$)—temperature ($T$) diagram for the detonation of $N_8$ clusters is depicted in Fig. 4. It shows the locus of equilibrium states of the detonation products gases, starting at the Chapman-Jouguet (CJ) point, expanding along the CJ isentrope: $S_{CJ} = 1.649 \text{ cal/g}$, and terminating at room temperature, as calculated by the Cheetah code [13]. Point $R$ denotes the initial state of the Reactants ($N_8$ clusters); the value $u_R = 6,652 \text{ cal/g}$ is based on ab initio local-energy-density-functional calculations for the fcc-form of polymeric nitrogen, as predicted by Mailhoit et al [14] and synthesized by Eremets et al [15] in a heated diamond anvil. The heat of detonation is $\Delta H_d = u_R(291K) - u_P(291K) = 6,672 \text{ cal/g}$. A Constant Volume Explosion (CVE) from point $R$ to point $P$ results in a temperature of $T_{CVE} = 6,400 \text{ K}$.

![Figure 4. Le Chatelier diagram for the detonation of $N_8$ (fcc) meta-stable clusters.](image)
At temperatures below 6,000 K, the composition of the detonation products consists of diatomic nitrogen (concentration of 35.7 mol/kg). As the energy increases, nitrogen starts to dissociate: at \( \nu = 4,000 \text{ cal/g} \) the equilibrium composition is 4.87 mol/kg of \( N \) and 33.3 mol/kg of \( N_2 \). At the \( CJ \) point, the composition is 66.3 mol/kg of \( N \) and 2.6 mol/g of \( N_2 \), in other words, the gas is almost fully dissociated. Thus, it is the change in composition that causes the non-monotonic behavior of the equilibrium \( u-T \) curve. Comparison of theoretical calculations of nitrogen in this energy regime prove that it is the equilibrium composition results, not the frozen composition results, that agree with experimental data [16].

The corresponding locus of states of the \( N_8 \) detonation products gases in the pressure (\( p \))—volume (\( v \)) plane is presented in Fig. 5.

![p-v diagram for the detonation of \( N_8 \) (fcc) meta-stable clusters.](image)
The computed points are fit with the JWL function:

\[ p_{\text{JWL}}(\rho, T) = A \left[ 1 - \frac{\omega}{R_1 v} \right] \exp(-R_1 v) + B \left[ 1 - \frac{\omega}{R_2 v} \right] \exp(-R_2 v) + \rho R_{DP} T \]  

The constants are listed in Table 2.

Table 2. JWL constants for the products of meta-stable cluster detonations

<table>
<thead>
<tr>
<th>Case</th>
<th>A (M-bar)</th>
<th>B (M-bar)</th>
<th>( R_1 ) (—)</th>
<th>( R_2 ) (—)</th>
<th>( \omega = \gamma - 1 ) (—)</th>
<th>( R_{DP} ) (bar-cc/g-K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>( N_8 ) (fcc)</td>
<td>8.04</td>
<td>0.1557</td>
<td>2.2496</td>
<td>0.5</td>
<td>0.1945</td>
<td>2.969</td>
</tr>
<tr>
<td>( He_2^* )</td>
<td>80.06</td>
<td>1.9985</td>
<td>2.5484</td>
<td>0.5</td>
<td>0.2483</td>
<td>20.785</td>
</tr>
</tbody>
</table>

**\( He_2^* \) Detonation**

The Le Chatelier diagram for the detonation of \( He_2^* \) clusters is depicted in Fig. 6. It shows the locus of equilibrium states of the detonation products gases, starting at the Chapman-Jouguet (CJ) point, expanding along the CJ isentrope: \( S_{CJ} = 5.536 \) cal/g, and terminating at room temperature, as calculated by the Cheetah code [13]. Point \( R \) denotes the initial state of the Reactants (\( He_2^* \) clusters). The value \( u_R = 50,039 \) cal/g is based on Elesin’s *ab initio* local-energy-density-functional calculations for \( He_2^* \), where the accumulated energy was \( E_{acc} = 8.7 \) ev/mol [10, 12, 17]. This gives a heat of detonation of \( \Delta H_d = 50,039 \) cal/g. A constant volume explosion from point \( R \) to point \( P \) results in a temperature of \( T_{CVE} = 27,550 \) K. The computed points are well fit by the following cubic function of temperature:

\[ u(T) = -824 + 0.767 \cdot T + 5.33 \times 10^{-5} \cdot T^2 - 5.204 \times 10^{-10} \cdot T^3 \]  

The corresponding locus of states of the \( He_2^* \) detonation products gases in the pressure \((p)\) —volume \((v)\) plane is presented in Fig. 7. Computed points are fit by the JWL function (1), whose constants for this case are listed in Table 2.
Figure 6. Le Chatelier diagram for the detonation products of $He_2^*$ meta-stable clusters.

Figure 7. $p$-$v$ diagram for the detonation products of $He_2^*$ meta-stable clusters.
Detonation properties of the selected meta-stable clusters are summarized in Table 3. The extraordinarily large detonation pressures (3 & 16 mega-bars), temperatures (12 & 34 thousand K), and wave velocities (20 & 43 kilometers per second) are a consequence of the large energies (and densities) stored in the Reactants state $R$ of the meta-stable clusters of $N_8$ and $He^*_{2}$.

Table 3. Detonation properties of Type-II meta-stable clusters

<table>
<thead>
<tr>
<th>Property</th>
<th>$N_8$ (fcc)</th>
<th>$He^*_{2}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\rho_0$ (g/cc)</td>
<td>3.186</td>
<td>3.164</td>
</tr>
<tr>
<td>$H_d$ (cal/g)</td>
<td>6,673</td>
<td>50,039</td>
</tr>
<tr>
<td>$p_{CJ}$ (M-bars)</td>
<td>3.14</td>
<td>16.3</td>
</tr>
<tr>
<td>$\rho_{CJ}$ (g/cc)</td>
<td>4.266</td>
<td>4.423</td>
</tr>
<tr>
<td>$T_{CJ}$ (K)</td>
<td>12,061</td>
<td>34,362</td>
</tr>
<tr>
<td>$W_{CJ}$ (km/s)</td>
<td>19.7</td>
<td>42.8</td>
</tr>
<tr>
<td>$U_{CJ}$ (km/s)</td>
<td>5.0</td>
<td>12.2</td>
</tr>
<tr>
<td>$a_{CJ}$ (km/s)</td>
<td>14.7</td>
<td>30.6</td>
</tr>
<tr>
<td>$\Gamma_{CJ}$ (-)</td>
<td>2.948</td>
<td>2.515</td>
</tr>
<tr>
<td>$p_{CVE}$ (M-bars)</td>
<td>1.14</td>
<td>8.97</td>
</tr>
<tr>
<td>$T_{CVE}$ (K)</td>
<td>10,750</td>
<td>27,550</td>
</tr>
<tr>
<td>$E_{CVE}$ (cal/g)</td>
<td>6,673</td>
<td>50,039</td>
</tr>
</tbody>
</table>

* see Appendix for density calculations

EXPLOSION SIMULATIONS

Here we use numerical simulations with our 3-D AMR code [18], to study the explosion environment created by the detonation of 1-g meta-stable clusters in a 6.6-liter chamber (right circular cylinder, $d = 20$ cm, $h = 21$ cm) at STP conditions (1 bar and 273 K). The thermodynamic properties of the detonation products ($u$-$T$ relation, JWL function, etc.) from the previous section were used to define the Equation of State (EOS) inputs to the AMR code. To handle the non-convex EOS effects, we used a generalized Godunov scheme of Bell et al [19].

$N_8$ Cluster Explosion

A 1-g spherical charge of $N_8$ clusters, with a density of 3.186 g/cc, was located at the center of the chamber filled with $N_2$ at STP. We assumed an initial value problem where the
energy was released as a constant volume explosion that deposited 6,673 calories instantaneously throughout the charge. With these initial conditions, the AMR code simulated the evolution of the explosion fields in the chamber. Pressure and impulse histories on the chamber lid at $r = 5$ cm are presented in Fig. 8. There they are compared with: (i) waveforms from the explosion of 1 gram of Al-SDF charge—releasing 7,400 calories via turbulent combustion with air, and (ii) the explosion of 1 gram of TNT charge in nitrogen—releasing 1,148 calories by the detonation wave (no after-burning). These are based on experimental measurements described by Kuhl and Reichenbach [20]. The $N_8$ blast had a peak value of 89 bars, compared to 23 and 14 bars for the TNT and Al-SDF explosions. The $N_8$ explosion created 4 times the impulse of the TNT explosion and twice the impulse of the Al-SDF explosion.

![Figure 8. Waveforms created by 1-g explosions in a 6.6-liter chamber. Numerical simulation of a $N_8$ explosion is compared with data from Al-SDF explosions in air and TNT explosions [20].](image)

**He$_2^*$ Clusters Explosion**

A 1-g spherical charge of He$_2^*$ clusters ($\rho_0 = 3.164$ g/cc) was located at the center of the 6.6-liter chamber filled with He at STP. An initial value problem was posed where the energy was released as a constant volume explosion that deposited 50,039 calories instantaneously
throughout the charge. Then, the AMR code simulated the evolution of the explosion in the chamber. Pressure and impulse histories on the chamber lid at $r = 5$ cm are presented in Fig. 9. The $\text{He}_2^*$ blast wave had a peak pressure of 743 bars, or more than 8 times the $N_8$ blast wave.

Figure 9. Waveforms created by 1-g explosions in a 6.6-liter chamber. Numerical simulation the explosion of a 1-g $\text{He}_2^*$ charge is compared with waveforms from Fig. 8.

Chamber Pressures

The impulse curves of Figs. 8 and 9 were fit with linear functions; their constants are listed in Table 4. The slope of the impulse history represents the mean chamber pressure: $\Delta \bar{p}_c$.

The helium cluster explosion created a mean chamber pressure of 86 bars; this is to be compared with 12 bars for the nitrogen cluster explosion in contrast with 8.8 bars and 2.4 bars for the Al-SDF and TNT explosions, respectively.

Table 4. Impulse curve fits: $I_i(t) = a_i + \Delta \bar{p}_{c,i} \cdot t$

<table>
<thead>
<tr>
<th>Case ($i$)</th>
<th>$a_i$ (bar-ms)</th>
<th>$\Delta \bar{p}_{c,i}$ (bars)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{He}_2^*$ cluster explosion in air</td>
<td>0.496</td>
<td>86.32</td>
</tr>
<tr>
<td>$N_8(fcc)$ cluster explosion in air</td>
<td>0.014</td>
<td>11.93</td>
</tr>
<tr>
<td>Al-SDF explosion in air</td>
<td>-1.65</td>
<td>8.829</td>
</tr>
<tr>
<td>TNT explosion in $N_2$</td>
<td>0.007</td>
<td>2.357</td>
</tr>
</tbody>
</table>
Flow Field Visualization

Figures 10 and 11 present a cross-section of the temperature field at different times during $N_8$ and $He^*_2$ cluster explosions, as computed by the AMR code. The Detonation Products (DP) interface is unstable and Taylor cavities (interpenetration of air into the DP), even before the first reflection from the chamber walls. The flow rapidly evolves into an intense mixing layer.

RÉSUMÉ

Theoretically, meta-stable clusters are able to accumulate energy that is much larger than those found in typical exothermic chemical reactions. For example, *ab initio* local-energy-density-functional calculations [10, 12, 14, 17] predict energy accumulation equivalent to heats of detonation of 6,673 cal/g and 50,0039 cal/g for $N_8$ and $He^*_2$ clusters, respectively. The Meta-stable Cluster State (MCS) serves as the Reactants state for the explosion. Detonation of meta-stable clusters creates Products gases such as $N_2$ molecules and He atoms, respectively. The locus of thermodynamic states of the Products gases in the $u-T$ plane and the $p-v$ plane have been calculated by use of the thermodynamic code Cheetah [13]. The appropriate assumption for the locus is that the Products are in thermodynamic equilibrium. For nitrogen, this results in a non-monotonic behavior of the $u(T)$ locus—due dissociation of $N_2$ into atomic nitrogen at large energies [16].

CONCLUSIONS

Meta-stable cluster explosions can produce pressures and impulses that are much larger than TNT detonations and even Al-air combustion systems. Meta-stable clusters of nitrogen and helium have been shown to exist theoretically. If they could be synthesized, then they would offer the potential of much higher energy density media.
Figure 10. Cross-section of the temperature field at different times during the $N_8$ cluster explosion in a 6.6-liter chamber (red corresponds to 6,000 K).
Figure 11. Cross-section of the temperature field at different times during the $He_2^*$ cluster explosion in a 6.6-liter chamber (red corresponds to 27,000 K).
ACKNOWLEDGMENTS

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APPENDIX

The initial density of the $N_8$ (fcc) cluster media was estimated from the relation:

$$\rho_0 = \frac{M}{V_0 \cdot A} = \frac{14 \text{ g/mol}}{7.3 \times 10^{-24} \text{ cc} \cdot 6.022 \times 10^{23} \text{ atoms/mol}} = 3.186 \text{ g/cc}$$

where the volume $V_0 = 7.30(A^0)^3$ was based on Mailhiot’s calculations for the fcc-form of polymeric nitrogen [14] and $A$ denotes Avogadro’s number. For the cubic gauche (cg) form, Mailhiot [14] predicts a volume of $V_0 = 6.67(A^0)^3$ which is close to the value $V_0 = 6.592(A^0)^3$ measured by Eremets [15]—thereby confirming the accuracy of his volume/density predictions for polymeric nitrogen. The initial density for $He_2^*$ was estimated from:

$$\rho_0 = \frac{M}{V_0 \cdot A} = \frac{8 \text{ g/mol}}{4.2 \times 10^{-24} \text{ cc} \cdot 6.022 \times 10^{23} \text{ atoms/mol}} = 3.163 \text{ g/cc}$$

where the volume $V_0 = 4.2(A^0)^3$ is based on an atom-to-atom separation of two angstroms, according to Elesin’s calculations for meta-stable $He_2^*$ clusters [10].

REFERENCES


