Observation of a Minimum Reaction Initiation Threshold in Ball-Milled Ni + Al Under High-Rate Mechanical Loading

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HOWARD G. WHITE, PhD  JEFFREY D. KUHN, MAJ, PhD  JENNIFER L. JORDAN, PhD
Technical Advisor  Branch Chief  Project Manager
Ordnance Division  Energetic Materials Branch  Energetic Materials Branch

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Observation of a Minimum Reaction Initiation Threshold in Ball-Milled Ni + Al Under High-Rate Mechanical Loading

Eric B. Herbold, Naresh N. Thadhani, Jennifer L. Jordan

School of Materials Science and Engineering
Georgia Institute of Technology
Atlanta, GA 30332

Air Force Research Laboratory
Munitions Directorate
Ordnance Division
Energetic Materials Branch
Eglin AFB, FL 32542-5910

AFRL-RW-EG-TP-2011-015

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Two types of microstructurally distinct ball-milled Ni+Al powder compacts are characterized for the investigation of reaction initiation threshold under high-rate mechanical loading using a modified rod-on-anvil Taylor impact-test setup. It is observed that the kinetic energy threshold for reaction decreases to a minimum then increases with milling time. It is also observed that the kinetic energy required for reaction initiation is lower for the 95% theoretical maximum density (TMD) ball-milled powder compacts than for the 65% theoretical maximum density (TMD) compacts. The results are discussed on the basis of competing effects of reactivity enhancement and deformability reduction caused by prior ball-milling of the powder mixtures. © 2011 American Institute of Physics.

Nickel powder, aluminum powder, ball mill, reaction
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Chemical reactions in powder materials have been an active area of investigation in recent years for synthesis of novel compounds by various methods of dynamic loading or processing techniques involving relatively “fast” or “moderate” reaction rates. Investigations of “fast” reactions include laser-induced desorption ionization in metal-mo-oxidic12 and those resulting from shock-compression of powder mixtures.1–11 The latter are characterized by “shock-induced” chemical reactions occurring within the time-scale of the shock duration: typically on the order of several microseconds. In contrast, “shock-assisted” chemical reactions initiate following the peak pressure state during the thermal equilibra-tion time scale2,3,7,9 and are being investigated for applications relevant to energetic materials.17 Mechanisms leading to shock-induced chemical reactions in powders, whether mechanochemical or thermomechanical in nature, vary as a function of the intrinsic and extrinsic properties of materials and their loading conditions.18 For example, pore collapse,19 localized melting,20,11 and disparate material properties may affect the occurrence of reaction. In contrast, the synthesis of intermet-allic compounds at longer time-scales resulting from gradual and explosive reactions at the grain level has been observed in powders processed using high-energy ball-milling.13–16

Self-sustaining high-temperature synthesis (SHS) reactions in intermetallic-forming systems result in the release of large amounts of heat and occur during ball-milling of Ni and Al powders mixed in an equiatomic ratio.13,14 Arrested reactive milling (ARM) techniques halt the milling process prior to SHS reaction resulting in a fine blend of constituent powders.16 Typical ARM materials have a layered microstructure at length-scales on the order of tens to a few hundred nm. During thermal analysis the apparent temperature threshold for reaction of these powders have been shown to decrease with increased milling time at the expense of total energy output.9

A recent investigation of shock induced chemical reactions in ball-milled Ni+Ti powder shows that increased milling times raises its crush-strength due to strain hardening.9 Correspondingly, the extent of reaction observed during shock-compression decreased with increasing ball-milling time. These results illustrate reduced reactivity of ball-milled powder mixtures during shock-compression (uniaxial-strain loading) despite a more intimately mixed powder.6

Here, the reaction/combustion characteristics of ball-milled Ni+Al (each at 50 at.%) powders are investigated using modified rod-on-anvil Taylor impact experiments with pressed pellets that vary in microstructure and density, mounted onto Cu rods, and impacted against a high-strength anvil. Mechanisms related to impact initiated initiation of gasless chemical reactions are important for applications involving novel energetic materials. The high strain-rates (102–105 s−1) and combined loading conditions indicate whether the reaction occurs during compaction, shear or severe plastic straining in the axial and lateral directions.

The energy level required for initiation and the amount released during reaction vary as a function of milling time. Here, ball-milled Ni+Al powder compacts are formed from powders milled in a hardened steel vial with Ar in a SPEX-8000 ball-mill with temperature control. Two different material types, distinguished by the initial size of the powder (designated SARM1 and SARM2), are employed to investi-gate the role of ball-milling time, powder grain microstruc-ture, and the density of the compacts. Differences in the single-grain microstructure are produced by milling 50 μm Al (H50, Valimet, Inc.) and 5–15 μm Ni (Alfa-Aesar) for SARM1 compared to 2 μm Al (H2, Valimet, Inc.) and 100 μm Ni (−150 + 200 mesh, Alfa-Aesar) for SARM2. Both powder types are milled with 7.94 mm stainless steel balls. The averaged time to reaction (t>0) for each powder is 9959 ± 862 s for SARM1 and 12632 ± 988 s for SARM2.
and was determined by repeating the ball milling procedure three times with 10 g of powder until reaction was detected by thermocouples attached to the outside of the vial. Subsequent batches of each mixture were produced under identical milling conditions for times corresponding to 35% and 65% of the averaged \( t_R \) for the impact tests.

The differences in microstructure between the two unsieved powders are compared in Fig. 1. Figure 1(a) shows SARM1 powder milled to 0.35\( t_R \). There are moderately deformed Ni particles within the Al matrix, as well as small amounts of Ni particles dispersed between sub-micron layers of Al. Figure 1(b) shows SARM2 powder milled to 0.35\( t_R \). Here the Ni particles are initially much larger than in (a) and are flattened with the soft Al particles resulting in a relatively coarse laminate particle. Figure 1(c) shows that the microstructure shown in Fig. 1(a) becomes a highly refined laminate structure when milled for 0.65\( t_R \). This is also apparent in Fig. 1(d) where the flakelike microstructure is refined when milled for 0.65\( t_R \) compared with Fig. 1(b). The microstructures of the individual grains in Fig. 1(c) and 1(d) look very similar in laminate thickness. However, comparing Fig. 1(a) and 1(b) the microstructural difference between individual grains suggests that a finer laminate structure (and higher contact surface area) is achieved with SARM2 powder as shown in (b) as opposed to the powder with small, hard Ni particles in SARM 1 as is apparent in (a).

The impact experiments are performed in air using a modified rod-on-anvil Taylor test setup. The 3.17 mm diameter pellets are mounted to a 7.62 mm diameter, 38.1 mm long high-strength steel anvil with hardness Rc 46 ground to 16G. The 3.17 mm diameter at the Cu sabot, \( d_2 \), is the diameter at the anvil, \( D = (d_2 - d_1)/2 \), and \( z \) is the averaged angle of the side of the cylinder. There are two distinct high-rate loading conditions following impact. Considering the measured shear (\( \dot{z} \)) and axial (\( \dot{e}_z \)) strain rates, the first stage of pellet deformation is dominated by compaction and a relatively high shear strain rate followed by the second stage involving mainly high axial and radial (\( \dot{e}_r \) and \( \dot{e}_r \)) strain rates. Figure 2(e) shows a bright “flash” of light as the Cu sabot contacts the anvil, which is an indication of the fast reaction in the ball-milled Ni+Al powder pellet. The reaction propagates into a powder dust cloud shown in Fig. 2(f). Each experimental image indicating reaction occurred at the latter stage where the particles have dynamically deformed by an averaged axial and lateral strain of approximately 0.8 (see the last column for \( \dot{e}_r \) and \( \dot{e}_z \) in Table I), though values of local strain at the scale of the particle are likely higher.

In Fig. 3 the kinetic energy of the Cu sabot and pellet is plotted as a function of (a) time of ball-milling and (b) the pressed density of the two types of sieved powders. In both (a) and (b) the open and closed markers (i and I for SARM1 and \( \nabla \) and \( \nabla \) for SARM2) indicate no reaction or a distinct ‘flash’ of light across the width of the pellet as seen in Fig. 2(e), indicating reaction before the sabot hit the anvil, respectively. The markers at \( t = 0 \) s correspond to as-blended (non-milled) 92% dense Ni+Al powder pellet based on the results of our prior work.

In Fig. 3, curves (1)–(2) are trendlines indicating the dependence of impact-initiated reaction energy on prior ball-milling time decreases to a minimum around \( t_{min} = 0.35 t_R \) and then increases with milling time for both powders. Such a trend is due to the competition of two effects. First, ball-milling creates fresh reactant surfaces to be brought into contact, thereby enhancing the reactivity of the powders. Further milling work-hardens the reactants and creates localized interfacial reactions making it more difficult for subsequent impact-initiated reactions. This indicates an optimum time exists for arrested reactive milled powders that can provide the highest sensitivity of impact-initiated reactions.
However, for increased milling times shown in Fig. 3(b), the apparent reaction threshold appears at the same level of energy at 0.65\(t_R\) though the powders were milled for different amounts of time, respectively, and may have different levels of strain-hardening given their initial sizes. With the same milling conditions (stoichiometric and charge ratio, milling temperature, and ambient conditions) the energy required for initiation of reaction of the SARM1 and SARM2 may tend to a constant value independent of milling time.

In Fig. 3(b) the kinetic energy is plotted against the degree of compaction energy as the density of the pellet increases, which may be linked to the amount of load carried by the pellet. The low-density sample may dissipate energy during the compaction process, reducing bulk distributed deformation, which requires higher threshold energy (see Table I at later times). The difference in threshold values between the two materials when comparing Figs. 3(a) and 3(b) suggests that the reaction threshold depends on the density and powder microstructure in these dynamic tests for similar milling conditions.

The dependence of air or vacuum environments on the impact initiation thresholds of long PTFE/Al composite rods show that first light occurs almost twice as fast in air as in vacuum at the same impact stress, suggesting that particle size effects are less important in experiments surrounded by air. However, the dependence of the threshold time on the milling time is strongly linked to microstructural differences and the results shown in Fig. 3 indicate that these differences are prevalent even in the presence of air.

Ball-milled Ni+Al powder compacts were characterized for the investigation of reaction threshold in modified Taylor impact tests. The kinetic energy (impact velocity) required for the observation of fast reaction decreases as fresh surfaces of the constituents are in close contact with minimal strain hardening and then increase due to the strain-hardening of the powder grains. There is an optimal milling time related to a minimum energy located around 0.35\(t_R\) for these materials, which is related to grain microstructure as indicated by its appearance. The energy required for reaction decreases with increasing pellet density, which is attributed to the balance of energy between the compaction process and bulk distributed deformation of the powder particles.

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**TABLE I.** Pellet measurements from Fig. 2. The parameter \(h\) is the pellet length (\(h\) is the engineering strain), \(d_1\) (\(r_1\)) is the diameter (eng. strain) at the Cu sabot, \(d_2\) (\(r_2\)) is the diameter (eng. strain) at the anvil, \(D = (d_2 - d_1)/2\), and \(z\) is the angle of the side of the cylinder \([z = \tan(D/h)]\).

<table>
<thead>
<tr>
<th>(t) ((\mu)s)</th>
<th>(h) (mm)</th>
<th>(d_1) (mm)</th>
<th>(d_2) (mm)</th>
<th>(D) (mm)</th>
<th>(z) (rad.)</th>
<th>(r_1)</th>
<th>(r_2)</th>
<th>(\dot{a}(10^6\text{ s}^{-1}))</th>
<th>(\dot{\epsilon}_1(10^4\text{ s}^{-1}))</th>
<th>(\dot{\epsilon}_2(10^4\text{ s}^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>(0 - 1.41)</td>
<td>1.27</td>
<td>3.17</td>
<td>3.17</td>
<td>0.425</td>
<td>0.045</td>
<td>0.241</td>
<td>0.297</td>
<td>0.0002</td>
<td>0.00012</td>
<td>0.00006</td>
</tr>
<tr>
<td>(1)</td>
<td>0.964</td>
<td>3.26</td>
<td>4.11</td>
<td>0.445</td>
<td>0.241</td>
<td>0.334</td>
<td>0.521</td>
<td>0.00174</td>
<td>0.00039</td>
<td>0.00017</td>
</tr>
<tr>
<td>(2)</td>
<td>0.590</td>
<td>4.23</td>
<td>4.82</td>
<td>0.555</td>
<td>0.535</td>
<td>0.669</td>
<td>0.767</td>
<td>0.00248</td>
<td>0.00057</td>
<td>0.00028</td>
</tr>
</tbody>
</table>

*This estimate is based on the velocity of 342 m/s of the projectile and a gap of 0.482 mm shown in Fig. 2(a). Spatial measurements are within 4 pixels or \(\pm 0.12\) mm.
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