ABSTRACT

We present a methodology for the efficient calculation of the shock Hugoniot using standard molecular simulation techniques. The method is an extension of an equation of state methodology proposed by Erpenbeck [J. J. Erpenbeck, Phys. Rev. A 46, 6406 (1992)] and is considered as an alternative to other methods that generate Hugoniot properties. We illustrate the methodology for shocked liquid N$_2$ using two different simulation methods: (a) the Reaction Ensemble Monte Carlo method for a reactive system; and (b) the molecular dynamics method for a non-reactive system. The method is shown to be accurate, stable and generally independent of the algorithm parameters. We find excellent agreement with results calculated by other previous simulation studies. The results show that the methodology provides a simulation tool capable of determining points on the shock Hugoniot from a single simulation in an efficient, straightforward manner.

1. INTRODUCTION

The behavior of materials under conditions of extreme temperature and pressure is of significant interest in many fields of physics and fluid science [1-4]. Of special interest are energetic materials, a class of materials of critical industrial and military importance. These materials exhibit chemically and physically interesting behavior when exposed to extreme temperatures and pressures. In particular, when subjected to shock, energetic materials often undergo rapid reactions that produce a heterogeneous mixture of chemical species that are accompanied by huge energy releases and can produce pressures up to several hundred GPa and temperatures exceeding 10,000 K. For a sufficiently strong shock, a supersonic, self-propagating reaction wave known as a detonation can be initiated. Unfortunately, the extreme conditions along with the short time and length scales over which a detonation occurs poses considerable experimental challenges in characterizing the material behind the detonation front. Therefore, a concerted effort that combines experimental, theoretical, and simulation approaches is essential for furthering our understanding of such shocked systems. Advances in experimental capabilities provide us with crucial property data, while the continuing development of accurate equations of state have allowed reasonable predictions of various shock properties [5, 6]. Similarly, the development of novel methods to simulate these complex systems has been the focus of research efforts and has recently led to the invention of some uniquely effective simulation tools [7-11]. These classical simulation methods can be implemented irrespective of rate limitations, the production of huge energy releases, or extreme thermodynamic conditions.

The Hugoniot curve, a commonly calculated property in shock and detonation science, reveals many properties of shocked materials and knowledge of which is critical to the design of new materials and application platforms. This curve consists of the set of \((PVT)\) points for which the Hugoniot expression:

\[
H_g = E - E_o - \frac{1}{2}(P + P_o)(V_o - V)
\]  

(1)

is zero. In Equation (1), \(E\) is the specific internal energy, \(P\) is the pressure, and \(V=1/\rho\) is the specific volume (\(\rho\) is the specific density). The term specific refers to the quantity per unit mass, while the subscript “o” refers to the quantity in the initial unshocked state.

Presently, three approaches exist for calculating the shock Hugoniot states from classical molecular simulation, each with their own advantages and disadvantages. The first approach, which we term here the Erpenbeck equation of state method (E-EOS), is the most indirect of the approaches. The original version of the method involves performing several separate simulations at appropriately chosen temperatures and pressures. Each simulation generates an equation of state point for subsequent evaluation of the Hugoniot expression followed by interpolation to locate the point at which the expression is zero. The molecular dynamics (MD) method has been implemented in the Erpenbeck approach using reactive potentials that mimic chemical bond breaking and forming between species [12, 13].

An approach for calculating the shock Hugoniot properties from classical molecular simulation that is more direct than the E-EOS method is the piston-driven molecular dynamics method [7, 8]. Piston-driven MD generates a point on the Hugoniot curve from a single simulation, thus avoiding the need to calculate several EOS points (as in the E-EOS approach) in order to obtain the desired result. The method mimics the laboratory system by calculating properties behind the shock discontinuity in a shock wave simulation. Shock waves are produced by hitting the free edge of the molecular solid with a rigid layer of atoms that are moving at a constant velocity. Different shocked states
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are obtained by starting with different initial piston velocities [8].

A third approach, termed the uniaxial Hugoniotstat method [9], is a molecular dynamics method that utilizes modified equations of motion that constrain the system to states that correspond to points on the shock Hugoniot curve. This method is computationally more efficient than both MD implementations of the E-EOS method and the piston-driven shock wave simulations. The E-EOS method requires a system of only a few hundred atoms (with periodic boundary conditions imposed), but several simulations are required to generate a single shock Hugoniot point. The piston-driven MD method will produce results from a single simulation, however, the system size must be sufficiently large so that the properties behind the propagating shock wave can be averaged over time. The uniaxial Hugoniotstat method, on the other hand, can generate a point on the Hugoniot curve from a single simulation whose system size is relatively small.

Unfortunately, none of these MD methods can be applied to the calculation of the shock Hugoniot locus over a wide range of temperatures and pressures unless certain conditions are fulfilled. Most energetic materials respond to shock by decomposing into a complex (sometimes heterogeneous) mixture of many different chemical species. Thus, for these multi-component systems, the implementation of the uniaxial Hugoniotstat method, the MD E-EOS method or the piston-driven MD method requires either: (1) a priori knowledge of the relative concentrations of each chemical species in the shocked state; or (2) a reactive potential that simulates bond breaking and bond formation. Typically, the relative species concentrations of the shocked state are lacking, moreover, knowledge of these quantities is desired. Furthermore, although significant advances have been made in developing reactive potentials for shocked materials, the potentials are presently limited to idealized representations of the chemistry that occurs [e.g., 13]. The most accurate interaction potentials for energetic materials available at this time are non-reactive.

Monte Carlo methodologies circumvent some of the restrictions associated with the MD methods for calculating shock Hugoniot states. The Reaction Ensemble Monte Carlo method (RxMC) [10] and the Composite Monte Carlo method [11] have both been used to calculate Hugoniot properties through the E-EOS approach. These two closely-related methods do not require a reactive potential or a priori knowledge of species concentrations for each Hugoniot state. They also do not require the specification of species chemical potentials or chemical potential differences to determine chemical equilibrium states of the reactive mixtures. Both methods have been applied to simple, spherically-averaged intermolecular potentials [10, 11] but can readily be applied to complex potentials that include multi-site and/or electrically-charged species as well as multi-phase mixtures. Therefore, in the absence of reactive potentials or a priori knowledge of species concentrations, the only applicable approaches for simulating the Hugoniot properties of a shocked material are the Erpenbeck EOS method performed using either the RxMC or Composite MC methods.

As previously mentioned however, the original E-EOS method requires simulations of several equation of state points to generate a single point on the shock Hugoniot curve. Each separate simulation requires sufficient equilibration and data collection steps. In an effort to reduce the number of steps and to minimize associated computational costs, we have implemented a numerical approach within the framework of the E-EOS approach. The resulting method requires only a single simulation to determine a point on the Hugoniot curve. The fitting procedure used to determine the root of the Hugoniot expression (Hg=0) in the original version of the E-EOS approach is replaced by an iterative numerical procedure built into the framework of the simulation.

A brief illustration of the method using isothermal-isobaric ensemble (NPT) simulations is given. The simulation is initiated at the specified temperature and pressure, and the Hugoniot expression is evaluated (using instantaneous values of V, P and E that depend only on the current configuration) at periodic intervals during the simulation run and accumulated for averaging. The averaged Hugoniot value is then used in a numerical root-finding algorithm (e.g., Newton-Raphson [14]) to provide an estimate of the Hugoniot pressure. The imposed pressure for the simulation is subsequently changed to correspond to the new estimate of the Hugoniot pressure. The simulation continues using this new imposed pressure constraint. This process is repeated until the Hugoniot function converges to zero (more precisely, within a desired tolerance of zero). The value of the pressure and corresponding volume averaged over the entire simulation run characterize the Hugoniot state at that temperature.

The method is akin to the phase equilibria methods that utilize thermodynamic integration to determine coexistence behavior [15-17]. In these calculations, a finite-difference algorithm is used to numerically integrate the differential equations, the Clausius-Clapeyron [15, 16] or the Gibbs-Duhem expressions [17], which govern the changes in thermodynamic parameters along the phase coexistence curve. Similarly, for the method introduced here, a numerical estimate of the root of the Hugoniot expression is made. Both approaches are iterated until the desired convergence has been reached.

In summary, we demonstrate the accuracy and stability of a method to calculate the shock Hugoniot properties of materials when implementing the E-EOS method either in an MD or RxMC framework. (Implementation of the
modified E-EOS method using the Composite MC method is analogous to the RxMC method and will not be demonstrated here.) The method, which we term the adaptive Erpenbeck equation of state method (AE-EOS), is intended to be a tool that is an alternative to the existing methodologies in order to overcome some of their limitations. We demonstrate the validity of the AE-EOS method for calculating the Hugoniot properties of liquid N\textsubscript{2} in the non-reactive regime using molecular dynamics and in the reactive regime using Reaction Ensemble Monte Carlo. The outline of the paper is as follows. The formalism and practical details of the methodology are presented in Section II. Applications of the method are given in Section III, while assessments of the results are given in Section IV.

II. METHODOLOGY

A. Formalism

The Hugoniot function \( H_g\) is defined using averaged instantaneous ensemble, for example, when simulating with an isothermal-isobaric simulation and calculating the remaining variable. For instance, several simulations must be performed using various choices of the independent variables to generate sufficient points so that the Hugoniot state can be obtained through interpolation. (Detailed outlines of the original E-EOS approach using the molecular dynamics and the RxMC methods can be found in Refs. [12] and [10], respectively.) The adaptive Erpenbeck equation of state method presented in this work eliminates the interpolation procedure by using a root-finding algorithm to determine \( H_g=0 \). For example, the expression for finding the root of a function using the Newton-Raphson method is [14]

\[
H_{g} = \sum_{\text{i}=1}^{n} y_{i} H_{i}^{0} + U_{\text{conf}} - RT, \tag{4}
\]

then Eq. (1) can be rewritten as

\[
H_{g} = \left( \sum_{\text{i}=1}^{n} y_{i} H_{i}^{0} + U_{\text{conf}} - RT \right) - E_{0} - \frac{1}{2} \left( PV_{0} - PV + P_{0} V_{0} - VP_{0} \right) \tag{5}
\]

where \( E_{0}, P_{0}, \) and \( V_{0} \) are the values of the initial state (and thus constant) and \( U_{\text{conf}} = \sum_{\text{i}=\text{j}}^{n} U_{\text{ij}}(r_{\text{ij}}) \) where \( U_{\text{ij}} \) is the pair potential energy [18]. All quantities in Eqs. (4) and (5) are used here in the context of instantaneous quantities that depend only on the current configuration. The derivative term required in Eq. (3) is then

\[
\frac{dH_{g}}{dP} = \frac{d}{dP} \left( \sum_{\text{i}=1}^{n} y_{i} H_{i}^{0} + U_{\text{conf}} - RT \right) - \frac{1}{2} \frac{d}{dP} (PV_{0} - PV + P_{0} V_{0} - VP_{0}) \tag{6}
\]

The first and third terms on the r.h.s. of Eq. (6) are not functions of \( P \) and can be eliminated. Further since \( U_{\text{conf}} \) is calculated as an instantaneous value, \( U_{\text{conf}} = U_{\text{conf}}(r) \) only and thus can be eliminated. Finally, the last term is readily solved, so that Eq. (6) reduces to

\[
\frac{dH_{g}}{dP} = \frac{1}{2} (V - V_{0}) \tag{7}
\]

The algorithm for the adaptive Erpenbeck equation of state method with \( P \) chosen as the independent variable is as follows:

\begin{itemize}
  \item **Step 1**: Set the temperature for the Hugoniot state \( T_{\text{Hug}} \).
  \item **Step 2**: Guess the pressure for this Hugoniot state \( P_{\text{current}} \).
  \item **Step 3**: Perform an isothermal-isobaric ensemble simulation (MD or RxMC) at \( T_{\text{Hug}} \) and \( P_{\text{current}} \).
\end{itemize}
Step 4: After allowing the system to relax to \( P_{\text{current}} \), accumulate instantaneous values of \( H_g \) and \( dH_g/dP \) during the simulation using Eqs. (5) and (7), respectively.

Step 5: After a prescribed number of steps, calculate averaged values of \( H_g \) and \( dH_g/dP \) and predict the Hugoniot pressure using Eq. (3). (The averaged values of \( H_g \) and \( dH_g/dP \) can be determined by several methods, which are considered in the next section.)

Step 6: Repeat steps (3)-(6) until the results converge to the desired statistical uncertainty.

B. Practical Details

Next, we consider a few practical details of implementing the AE-EOS method. Our intent is to generalize the method for implementation into any of the standard molecular simulation techniques (MD, RxMC, or Composite MC). We consider the effect of several parameters on the accuracy and stability of the method. Below we address these issues, the logic behind our choices, and the tradeoffs involved.

There exists a wide range of root-finding algorithms, including the Newton-Raphson method, the Secant method, the Bi-section method, and Halley’s method [14]. Newton-Raphson (Eq. 2) is a rather straightforward method but can be unstable near a horizontal asymptote or local minimum. A similar algorithm is Halley’s method that includes an additional term \((df(x))/dx\) from the Taylor series in the derivation of the method. When the pressure is chosen as the independent variable, \( d^2H_g/dP^2=0 \) (see Eq. 7) so Halley’s method reduces to the Newton-Raphson method. Another root finding method is the Secant method, which estimates the derivative term using \( f'(x) = (f(x_n) - f(x_{n-1})/x_n - x_{n-1}) \). However, use of the method in the AE-EOS method requires two recent points along the Hugoniot curve as opposed to only one for the Newton-Raphson method. Finally, if we can be certain that the solution of the Hugoniot expression lies within a known interval, then we can iteratively converge to the solution using the Bi-section method. However, a balance must be established between statistical uncertainty and the desired convergence when implementing the Bi-Section method in a molecular simulation, since statistical fluctuations cannot be greater than the size of the interval. In an effort to keep the method proposed here as general and straightforward as possible, we have implemented the Newton-Raphson method. The well-known problems of this method near a local minimum or asymptote have not been encountered for the Hugoniot expression in this work as well as for other work [10, 12, 13], but one should be mindful of its limitations.

In any molecular simulation, it is necessary to design a starting configuration so that the relaxed system is physically reasonable and computationally consistent. For example, consider an isothermal-isobaric ensemble simulation where periodic boundaries are imposed and where the potential energy function has a limited interaction range. In such a case, an appropriate number of molecules must be chosen so that the relaxed box size is consistent with the minimum image convention, i.e., one-half the box size must be greater than or equal to the potential cutoff distance [18]. Similarly, appropriate starting conditions for the AE-EOS method are required, particularly for the initial guess of the imposed pressure. Since the converged result (i.e., \( H_g=0 \)) will produce a Hugoniot pressure that is equal to the imposed pressure (within some specified tolerance), it is desirable to choose an initial pressure that is a good estimate of the actual Hugoniot pressure. Although we will show in Section III that the AE-EOS method is largely insensitive to the initial pressure guess, extremely poor initial guesses could result in numerical failures. For example, consider the simulation of a Hugoniot state such that \( T_{Hg}>T_o \) (recall that \( T_{Hg} \) is the temperature of the chosen Hugoniot state and \( T_o \) is the temperature of the unshocked material). If the initial guess of the pressure, \( P_{\text{initial guess}} \), causes an expansion of the simulation cell such that the specific volume is larger than the specific volume of the unshocked material, a negative pressure value will be predicted. From a practical standpoint, this is an unphysical occurrence since it implies that the material has expanded upon shock rather than being compressed. Furthermore, from a computational standpoint, the simulation cell will never converge to a negative imposed pressure. Such an occurrence, however, is analogous to choosing a starting configuration that relaxes to a physically unreasonable and computationally inconsistent state.

Consider the following ad hoc approach to choosing a reasonable initial guess of the pressure for Eq. (3). First, assume that the shocked material does not decompose (i.e., chemically react). This is a reasonable approximation at low shock pressures and reduces the first term on the r.h.s of Eq. (5) to \( U^\text{starting material} \). Next, neglect the contribution of \( U^\text{conf} \) and thus eliminate the second term on the r.h.s of Eq. (5). This approximation has no physical justification, however, a short simulation could be performed to calculate \( U^\text{conf} \) (although probably unnecessary given the lack of sensitivity of the final result on the initial guess of the pressure). Finally, estimate the amount of compression the starting material will undergo, e.g., \( V=0.7V_o \). This estimate presumably can be predicated on previous studies of the material or similar materials. With these approximations, an initial estimate of the Hugoniot pressure can be determined. Furthermore, as points along the Hugoniot curve are determined, better estimates of the initial pressure can be made by using these Hugoniot states.
For completeness of study, we consider three different schemes for averaging the instantaneous values of $H_g$ and $dH_g/dP$, which are then used in Eq. (3): (1) block averages; (2) running averages; and (3) block-to-running averages. Block averages are taken from a limited number of configurations immediately preceding the pressure adjustment step, while running averages are taken continuously over all configurations generated during the simulation run. The block-to-running averages scheme uses a block-averaging scheme for the equilibration period of the simulation run and then continues with a running average scheme for the production period. This scheme may most effectively remove the effects of a poor initial guess, while we expect the running average scheme to be the most effective alternative since fluctuations in the pressure will become increasingly damped as the simulation proceeds. Block averaging methods will likely be more slowly converging at best, and unstable at worst. Moreover, running average schemes have been the most successful scheme in the finite-difference algorithms used in the phase coexistence methods mentioned previously [15-17].

We also consider the effect of the frequency of re-setting the pressure during the simulation. Less frequent updates are expected to cause the results to converge more slowly while more frequent updates could possibly cause the root-finding scheme to become unstable or to fluctuate too greatly.

A final note on the convergence of the system to the predicted pressure is considered. Step (4) in the algorithm outline allows the system to converge to the predicted pressure value (within a few % of the predicted pressure for the most recent simulation steps) before re-evaluating the Hugoniot expression and it’s derivative ($dH_g/dP$) in the equilibration period only. This ensures that even for large changes in the predicted pressure, equilibrated information is still used in the $H_g$ and $dH_g/dP$ calculation. Typically, these large changes will only occur during the earliest stages of the simulation. At later times during the production cycles, this criterion is nearly always satisfied.

### III. APPLICATION

For demonstrative purposes, several shock Hugoniot states of liquid nitrogen are considered. The shock Hugoniot properties were predicted based on the initial states calculated previously: $T=77.0$ K; $p=0.808$ g/cm$^3$; $P=50.49$ MPa; $E=0.441$ KJ/g [10]. At pressures higher than ~30 GPa along the Hugoniot curve, the dissociation reaction of molecular nitrogen ($N_2 \leftrightarrow 2N$) occurs. Therefore, we demonstrate the AE-EOS method using the molecular dynamics technique only at pressures below 30 GPa while we demonstrate the AE-EOS method using the RxMC method at a wider range of pressures. Particles interact through an exponential-six potential, where potential parameters are given in Ref. [19]. A spherical cutoff for the particle-particle interactions was applied at $2.5r_{nm,N2}$ with long-range corrections added to account for interactions beyond this distance [20]. Electrostatic interactions between species were ignored. The unlike interactions between species $i$ and $j$ were approximated by the Lorentz-Berthelot combining rules [21]. 3375 $N_2$ molecules were used with all calculated quantities reduced by the exponential-6 potential energy ($\phi$) and size ($r_{nm}$) parameters of $N_2$. Periodic boundary conditions were imposed for all dimensions. Thermochemical reference data were used in calculating the ideal-gas enthalpies ($H_i^0$) required in Eq. (5) [22, 23].

#### A. Molecular dynamics

Molecular dynamics simulations in the isothermal-isobaric ensemble were performed using the Leap-Frog Verlet algorithm [18, 20] and the Melchionna modification of the Hoover-Nose equations of motion [24]. A thermostatting rate of 50 ps$^{-1}$ was used to maintain the imposed temperature while a barostatting rate ranging from 0.032-0.042 ps$^{-1}$ was used to maintain the imposed pressure. Initial configurations were generated from a face-centered-cubic (fcc) lattice structure with initial particle velocities selected from a Boltzmann distribution that corresponded to the imposed temperature. Preceded by an equilibration period of 0.127-0.254 ns during which the pressure was not re-set using Eq. (3), trajectories were followed for 1.32 ns with time steps ranging from 0.00763-0.0102 ps. All pressure values reported were determined using the virial theorem [18].

Three state points along the shock Hugoniot curve were determined: $T=883.9$; 3912.4; 6778.1 K. These state points are below the regime in which $N_2$ dissociates into atomic nitrogen. For each point, the Hugoniot pressure ($P_{Hg}$) was predicted in two simulations, one of which the initial pressure was much lower than the Hugoniot pressure and one in which the initial pressure was too high. The effect of the frequency of re-setting the imposed pressure was also studied. Two cases were considered, re-setting at: (a) every 100 steps; and (b) every 500 steps. Following the initial equilibration period used to relax the system from the fcc crystal to the imposed thermodynamic condition, an additional 0.305 ns of the trajectory was used to further equilibrate the system after the AE-EOS algorithm is implemented (i.e. the pressure is re-set at specified intervals). All quantities calculated during this time interval were not included in the final averages. A tolerance value of +/-5% was used in Step (4) for the pressure (see Section II.A), i.e., the calculated pressure was required to be within +/-5% of the most recent $P_{Hg}$ prediction before re-evaluating $H_g$ and $dH_g/dP$ and re-setting of the imposed pressure.
Table I: Predicted shock Hugoniot states of liquid N₂ using molecular dynamics in the AE-EOS method.¹

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<tr>
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</thead>
<tbody>
<tr>
<td>1.56</td>
<td>883.9(1.2)</td>
<td>4.82(9)</td>
<td>19.7(2)</td>
<td>-2.356E-4(0.013)</td>
<td>883.9(1.1)</td>
<td>4.8(1)</td>
<td>19.9(3)</td>
<td>8.901E-3(0.16)</td>
</tr>
<tr>
<td>7.92</td>
<td>883.9(1.2)</td>
<td>4.8(1)</td>
<td>19.74(3)</td>
<td>1.001E-4(0.023)</td>
<td>883.9(1.2)</td>
<td>4.8(1)</td>
<td>20.0(3)</td>
<td>1.600E-2(0.19)</td>
</tr>
<tr>
<td>5.97</td>
<td>3912(52)</td>
<td>18.4(3)</td>
<td>15.6(2)</td>
<td>-1.550E-3(0.15)</td>
<td>3912(52)</td>
<td>18.4(2)</td>
<td>15.6(2)</td>
<td>-3.101E-5(0.043)</td>
</tr>
<tr>
<td>30.23</td>
<td>3912(53)</td>
<td>18.4(1)</td>
<td>15.56(8)</td>
<td>2.790E-3(0.063)</td>
<td>3912(54)</td>
<td>17.8(6)</td>
<td>16.0(4)</td>
<td>1.841E-1(1.1)</td>
</tr>
</tbody>
</table>

¹Hugoniot states taken from [10].

Table II: Predicted shock Hugoniot states of liquid N₂ at $T=7963.0$ K using the Reactive Monte Carlo method. Values determined previously [10] by the E-EOS method are $P=36.0$ GPa; $V=13.35$ cm³/mole N₂; $x(N₂)=0.975$.²

<table>
<thead>
<tr>
<th>$P_{\text{initial}}$ [GPa]</th>
<th>$P$ [GPa]</th>
<th>$V$ [cm³/mol N₂]</th>
<th>$x(N₂)$</th>
<th>$H_g$ [kJ/g]</th>
<th>$P$ [GPa]</th>
<th>$V$ [cm³/mol N₂]</th>
<th>$x(N₂)$</th>
<th>$H_g$ [kJ/g]</th>
</tr>
</thead>
<tbody>
<tr>
<td>11.9</td>
<td>36.1(1)</td>
<td>13.33(6)</td>
<td>0.975(1)</td>
<td>2.129E-5(0.013)</td>
<td>36.2(1)</td>
<td>13.32(3)</td>
<td>0.975(1)</td>
<td>-5.957E-4(0.16)</td>
</tr>
<tr>
<td>60.1</td>
<td>36.1(1)</td>
<td>13.33(6)</td>
<td>0.975(1)</td>
<td>1.118E-4(0.023)</td>
<td>36.1(1)</td>
<td>13.33(4)</td>
<td>0.975(1)</td>
<td>2.937E-5(0.19)</td>
</tr>
</tbody>
</table>

²Mole fraction of N₂, so $x(N₂)=N_{N₂}/N_{\text{total}}$ and $x(N)=\frac{1}{2}N_{N}/N_{\text{total}}$ where $N_{\text{total}} = 3375$. A comparison between the Hugoniot properties predicted using the original E-EOS and AE-EOS methods is shown in Table I. Good agreement is found for all cases considered, with pressure and specific volume values well within statistical uncertainty. Table I also shows that no dependence on the initial pressure guess is found, with values within a few percent of the E-EOS method results.
B. Reaction Ensemble Monte Carlo

The Reaction Ensemble Monte Carlo method was used to assess the accuracy of the AE-EOS method at a wider range of conditions than considered using the molecular dynamics technique including conditions under which \( \text{N}_2 \) dissociates (\( \text{N}_2 \leftrightarrow 2\text{N} \)). Details of the methodology can be found in the original papers [25-27] as well as in recent applications of the technique, which implemented the E-EOS method [10]. In addition to intermolecular potentials that describe non-reactive interactions between species \( \text{N}_2 \) and \( \text{N} \) in the equilibrium mixture, RxMC also requires inputting the ideal-gas internal modes (vibration, rotation, electronic). The vibrational and rotational contributions to the ideal-gas partition functions of both species were calculated using a standard source [28], and supplemented with electronic level constants that included the ground state and six excited electronic states for \( \text{N}_2 \) [22], along with electronic energy levels for \( \text{N} \) taken from Moore and Gallagher [23].

Constant-pressure RxMC simulations of shocked \( \text{N}_2 \) were initiated from 3375 \( \text{N}_2 \) particles placed on an fcc-lattice structure. Simulations were performed in steps, where a step (chosen with equal probability) was either a particle displacement, a forward reaction step, or a reverse reaction step. A change in the simulation cell volume was attempted every 500 steps. Simulations were equilibrated for 1.5x10^6 steps after which averages do not correspond to the averaging used in the reverse reaction step. A change in the simulation cell volume was attempted every 500 steps. Simulations were equilibrated for 1.5x10^6 steps after which averages of the quantities were taken over 8x10^6 steps. Uncertainties were estimated using the method of block averages by dividing the production run into 10 equal blocks [20]. Reported uncertainties are one standard deviation of the block averages. (Note that these block averages do not correspond to the averaging used in the \( P_{\text{Hg}} \) prediction scheme.) The maximum displacement and volume change were adjusted to achieve an acceptance fraction of approximately 0.33 and 0.5, respectively. Depending on the system conditions, the acceptance fraction of the reaction steps ranged from 0.075-0.375.

Three points along the shock Hugoniot curve were considered: \( T=2008.4; 7963.03; 10935.2 \) K. Again for each point, \( P_{\text{Hg}} \) was predicted from two initial guesses that differ from the known value [10] by +/- 67%. We predicted the shock Hugoniot properties based on the same initial state used previously [10] and in the molecular dynamics study above. A tolerance value of +/-2.5% was used in Step (4) for the pressure (see Section II.A), a value slightly lower than used in the molecular dynamics simulations above. Analogous to the study above, the effect of the frequency of re-evaluating the Hugoniot pressure and subsequent re-setting of the imposed pressure was studied. Two cases were considered, re-setting at: (a) every 5000 steps; and (b) every 50,000 steps. Additionally, three different averaging schemes of the instantaneous values of \( \text{H}_g \) and \( d\text{H}_g/dP \) were assessed: (a) block averages; (b) running averages; and (c) block-to-running averages. A description of each type is given in Section II.B. For each of the initial pressure guesses, all six series (2 \( \text{H}_g \) frequency evaluations X 3 averaging schemes) were considered.

Comparisons between the Hugoniot properties predicted by the E-EOS and the AE-EOS methods for the 7963.03 K case are given in Table II. We find excellent agreement between the E-EOS and the AE-EOS results for all quantities calculated. Nearly all AE-EOS quantities fall within +/-0.5% of the E-EOS calculations with none greater than +/-0.9%. Consequently, no dependence on the initial pressure guess or the Hugoniot expression evaluation frequency is evident. Likewise, no apparent dependence on each of the averaging schemes is found.

IV. DISCUSSION

We have presented a computationally efficient methodology for calculating the shock Hugoniot properties of materials using classical molecular simulations. The method is an extension of the Erpenbeck EOS approach and allows for the determination of a point along the Hugoniot curve from a single simulation. The method, termed the adaptive Erpenbeck equation of state method (AE-EOS), uses a numerical estimate of the root of the Hugoniot expression to determine the corresponding thermodynamic state. AE-EOS is applicable for any simulation method that determines points along the Hugoniot curve by generating EOS points, which includes molecular dynamics [18,20], Reaction Ensemble Monte Carlo [25,26], and the Composite Monte Carlo method [11]. The method was demonstrated to be accurate and numerically stable. For the systems in this study, the method was not particularly sensitive to the algorithm parameters indicating the method’s robustness and ability to be readily implemented. Furthermore, a substantial savings in the computational requirements is gained through the implementation of the AE-EOS method. For comparable statistical uncertainty, 4-5 complete simulations are required to generate a single point on the Hugoniot curve for the original Erpenbeck method while only a single simulation is needed in the AE-EOS method. Hence, a computational gain of approximately 4-5 fold is made when implementing the AE-EOS method. Overall, the AE-EOS method appears to be slightly more accurate and stable when applied to the RxMC method as compared to the molecular dynamics method. This may be attributed to the relative stability of the Monte Carlo method over the molecular dynamics technique for constant-temperature and constant-pressure simulations.

The AE-EOS method is intended as an alternative tool to similar techniques, namely, piston-driven molecular dynamics [7,8] and uniaxial Hugoniot molecular dynamics [7,8].
dynamics [9], since AE-EOS offers some notable advantages when simulating reactive mixtures. The most striking advantage is that the AE-EOS method does not require either: (1) a priori knowledge of the relative concentrations of each chemical species in the shocked state; or (2) a reactive potential that simulates bond breaking and bond formation. At least one of these conditions must be met for the other methods to be applicable to the simulation of reactive mixtures.

REFERENCES
Novel simulations of energetic materials: circumventing limitations in existing methodologies

John K. Brennan, Betsy M. Rice and Edward F.C. Byrd

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Motivation

Need for:

1. Better performing energetic materials
2. Materials to keep pace with advancing technology (notional materials)

Goal:

Efficient design of energetic materials
Design Approaches

1. Laboratory experiments
   - Trial and error process
   - Years to develop \( \sim 20 \) years for M43 propellant
   - Costly & environmentally harmful
     - 50,000 lbs XM39 propellant
     - \( {\text{Millet et al Clean Prods. \\& Proc. 2000}} \)

2. Theoretical models (Ex: Cheetah)
   - Fast calculations
   - Limited predictive capabilities
   - Not practical for notional materials
   - require EOS

3. Computational methods

Computational Methodologies
Molecular Simulation

- Simulation of chemically reacting systems:
  1. Molecular Dynamics:
     - Requires reactive potential for each reaction
     - Not applicable to mixtures
  2. Reaction Ensemble Monte Carlo

Reaction Ensemble Monte Carlo

- Simulates chemical reaction equilibria
- Not limited by reaction rate
- Not limited by activation energy barrier

- W. Smith and B. Triska (1994)
### Reaction Ensemble Monte Carlo

<table>
<thead>
<tr>
<th>Input requirements:</th>
</tr>
</thead>
<tbody>
<tr>
<td>• specify reactions occurring</td>
</tr>
<tr>
<td>• intermolecular potential models</td>
</tr>
<tr>
<td>• ideal gas intra-molecular contributions (vibrational, rotational, electronic)</td>
</tr>
<tr>
<td>• predicts physical effects not chemical effects</td>
</tr>
</tbody>
</table>

**ideal gas reaction** $\Rightarrow$ **RxMC** predict chemical reaction behavior under non-ideal conditions

- reactive potential not needed
- multiple reactions, multiple phases possible
- reaction *equilibria* not *kinetics*
Example: \(2 \text{ A} \leftrightarrow \text{ B}\)

- particle displacement
- box volume change

**forward reaction**

**reverse reaction**

initial state

shocked state

\(E=E_0=\frac{1}{2}(P+P_0)(V_0-V)\)

\(V=1/\rho\)

**shock Hugoniot properties**
\[ \text{N}_2 \leftrightarrow 2\text{N} \]

**Graph 1:**
- **X-axis:** Volume [cm³/mol of N₂]
- **Y-axis:** Pressure [GPa]
- Data points represent:
  - RxMC (non-reactive)
  - RxMC (reactive)
  - Experiment

**Sources:**
- Zubarev and Telegin (1962)
- Nellis et al. (1991)

**Graph 2:**
- **X-axis:** Pressure [GPa]
- **Y-axis:** Species mole fraction
- Data points represent:
  - N₂
  - N
2 NO ⇌ N₂ + O₂

Schott, Shaw and Johnson (1985)

Nitromethane

- prototypical EM
- 27 reacting species
- 42 reactions

C. Mader (2002)
Nitromethane

![Graph showing experimental and predicted values of pressure (P) and volume (V) for nitromethane.](image)

**Capabilities of RxMC**

- Assess accuracy of theoretical models

- Predictive tool for energetic material development

- Supercritical fluid phase separation

*energy storage*

*polymeric nitrogen*
Capabilities of RxMC

- Predictive tool for other non-ideal environments

Capabilities of RxMC

- Sol-gel processing of nanostructured materials
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