Initial Decomposition Reactions of Bicyclo-HMX [BCHMX or cis-1,3,4,6-Tetranitrooctahydroimidazo-[4,5-d]imidazole] from Quantum Molecular Dynamics Simulations

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ABSTRACT: We investigated the initial chemical reactions of BCHMX [cis-1,3,4,6-tetranitrooctahydroimidazo-[4,5-d]imidazole] with the following procedure. First we used density functional theory molecular dynamics simulations (DFT-MD) on the periodic crystal to discover the initial reaction steps. This allowed us to determine the most important reactions through DFT-MD simulations at high temperatures. Then we started with the midpoint of the reaction (unimolecular or bimolecular) from the DFT-MD and carried out higher quality finite cluster DFT calculations to locate the true transition state of the reaction, followed by calculations along the reaction path to determine the initial and final states. We find that for the noncompressed BCHMX the nitro-aci isomerization reaction occurs earlier than the NO2-releasing reaction, while for compressed BCHMX intermolecular hydrogen-transfer and bimolecular NO2-releasing reactions occur earlier than the nitrous acid (HONO)-releasing reaction. At high pressures, the initial reaction involves intermolecular hydrogen transfer rather than intramolecular hydrogen transfer, and the intermolecular hydrogen transfer decreases the reaction barrier for release of NO2 by ~7 kcal/mol. Thus, the HONO-releasing reaction takes place more easily in compressed BCHMX. We find that this reaction barrier is 10 kcal/mol lower than the unimolecular NO2 release and ~3 kcal/mol lower than the bimolecular NO2 release. This rationalizes the origin of the higher sensitivity of BCHMX compared to RDX (1,3,5-trinitrohexahydro-1,3,5-triazine) and HMX (octahydro-1,3,5,7-tetranitro-1,3,5,7-tetrazocine). We suggest changes in BCHMX that might help decrease the sensitivity by avoiding the intermolecular hydrogen-transfer and HONO-releasing reaction.

1. INTRODUCTION

cis-1,3,4,6-Tetranitrooctahydroimidazo-[4,5-d]imidazole was designed theoretically to have superior physical and energetic properties.1,2 This molecule, which can be considered as bicyclo-HMX, is denoted as BCHMX. Following a very intense effort in the 1980s to find efficient and economic synthetic procedures to obtain larger amounts of BCHMX,3,4 Zeman et al. in 2009 synthesized BCHMX and investigated the properties. The ignition temperature of BCHMX is 214–224 °C, compared to 204 °C for RDX (1,3,5-trinitrohexahydro-1,3,5-triazine) and 275 °C for β-HMX. The heat of explosion of BCHMX is 5.758 MJ/kg, which is approximately the same as pentaerythritol tetranitrate (PETN) (5.730 MJ/kg), but it is higher than for RDX (5.481 MJ/kg), and HMX (5.530 MJ/kg). In addition, the BCHMX melting point (268 °C) is much higher than PETN (141 °C) and RDX (204 °C) but slightly lower than HMX (275 °C).5 BCHMX has a predicted maximum density of 1.86 g/cm³, calculated detonation velocity of 9050 m/s, detonation pressure of 37 GPa, and maximum explosion heat of 6.518 MJ/kg.5–7 Thus, BCHMX has the potential to be a high-energy material. However, it has higher impact and electric-spark sensitivities, compared with RDX and β-HMX. In order to understand the origin of its high sensitivity we undertook to determine the mechanism with the expectation that this might provide clues to improve this material.

Although the synthesis8,9 properties, initiation reactivity,10 and thermal behaviors11 of BCHMX have been reported, the complex chemical behavior of BCHMX leaves many fundamental and practical issues to be understood. For instance, for high-energy materials, the initial thermal decomposition and reaction processes are essential in determining the energy release and detonation properties, as well as its sensitivity.11–17 But the initial decomposition reaction of BCHMX has not been reported. For such normal nitro-based explosives as HMX and RDX, the early chemistry involves decomposition via a branching chain of radical reactions, with the activation energy determined by the...
unimolecular NO₂ cleavage.18-22 In addition, intramolecular hydrogen transfer to produce nitrous acid (HONO) which decomposes to form HO and NO can compete with NO₂ cleavage as the first step of decomposition.11,13,23,24 Since BCHMX has a molecular structure similar to RDX or HMX, but is much more sensitive, we were particularly interested in examining the first step of the decomposition mechanism of BCHMX.

In order to elucidate the initial reaction of BCHMX, we report here DFT-MD simulations using the forces from the PBE-ulg25 flavor of DFT to examine the decomposition reaction of BCHMX. These studies on the initial reaction of BCHMX explain the relationship between initial reaction and sensitivities, and we suggest ways to improve its property.

2. METHODOLOGY

2.1. Density Functional Theory Molecular Dynamics Simulation (DFT-MD). Our DFT-MD simulations use interatomic forces calculated from quantum mechanics based on the PBE flavor of density functional theory (DFT),26,27 in which exchange and correlation are described using the generalized gradient approximation (GGA), and London dispersion (van der Waals attraction) is treated by the low gradient correction.25

These periodic QM calculations were performed using the VASP package.28-30 We found that a kinetic energy cutoff of 500 eV for the plane wave expansions gives excellent convergence of the total energies, energy differences, and structural parameters in structure optimization. The same energy cutoff was applied in the DFT-MD calculations. Only the gamma point of reciprocal space was sampled in the DFT-MD simulations. The convergence criteria were a $1 \times 10^{-6}$ eV energy difference for solving the electronic wave function and $1 \times 10^{-3}$ eV/Å force for structure optimization. For DFT-MD simulations, they were set to $1 \times 10^{-5}$ eV energy difference for solving the electronic wave function and $1 \times 10^{-3}$ eV/Å force. The spin-restricted KS solvers were used in the DFT-MD simulations, and the symmetry was not constrained.

The simulated system with eight molecules per periodic cell was obtained by replicating the unit cell (as shown in Figure 1) twice along the y and z directions. The structures were optimized before the DFT-MD simulations, as shown in Figure S1(a) of the Supporting Information (SI). To investigate the pressure effects on the initial reactions, we also performed simulations with the volume of the BCHMX unit cell compressed by 30%. This decreased the lattice constants $a \times b \times c$ from 8.59 Å × 7.07 Å × 8.75 Å to 7.86 Å × 6.73 Å × 7.03 Å with an initial external pressure of 20.0 GPa. This eight-molecule supercell is shown in Figure S1(b) (SI).

The DFT-MD simulation procedures were as follows. First the systems were heated at a constant rate from 20 to 300 K over 2 ps. Then the systems were equilibrated at 300 K for 1 ps using the NVT (constant volume, constant temperature, and constant number of atoms) ensemble. Finally, we heated the system from 300 to 3000 K uniformly over 20 ps. The time constant for the Nose–Hoover thermostat was 0.1 ps. The time step 1 fs was used for integrating the equation of motion. The high temperatures used in these simulations allowed many reactions to be observed within 20 ps of simulation. To analyze the fragments during the simulation, we used a bond length cutoff of 1.5 times of the normal bond length. These bond length cutoffs are shown in Table S1 of the SI.

2.2. Finite Cluster Calculation. To obtain accurate energetics, we extracted the molecules undergoing reactions from the trajectories of the periodic DFT-MD simulations. Then we located the nearby transition state, reactant, and product geometries in the gas phase at the level of PBE-ulg/6-31++G**(see), using Jaguar 8.2 transition states (TS) which were confirmed to have exactly one negative eigenvalue of the Hessian, which we followed by intrinsic reaction coordinate (IRC) scans to connect to the reactant and product structures. Thermodynamic properties were evaluated at 298.15 K and 1 atm. All gas-phase calculations were carried out using the Jaguar 8.2 package.31

3. RESULTS AND DISCUSSION

The DFT-MD simulations in this paper provide a very detailed, molecular-level description of the decomposition and reaction of condensed-phase BCHMX under various conditions. This information allowed us to extract valuable information about the complex chemistry involved, including unimolecular and multimolecular reactions.25 Our goal is to elucidate the initial reaction pathway as BCHMX decomposes and describe the events in this high-energy material as it evolves to form
intermediates that react with each other and with reactant to form eventually the final products observed theoretically. In this work, we focus on thermal decomposition of the condensed phase of BCHMX crystals, noncompressed BCHMX, and 30% compressed BCHMX and examine the initial reaction details under various conditions.

3.1. Reaction Mechanisms from DFT-MD. 3.1.1. Noncompressed BCHMX. We first examined the initial decomposition reaction of noncompressed BCHMX. The molecular fragments during the cook-off simulation are plotted in Figure 2 as a function of temperature. The starting supercell consists of eight BCHMX molecules (208 atoms). The first reaction occurs as the temperature increases to ~1700 K at which point one BCHMX molecule experiences unimolecular decomposition, releasing the first NO₂ molecule, while the instantaneous pressure increased to ~1.0 GPa. Later at ~1900 K, more NO₂ molecules are released, as shown in Figure 2. Thus, we consider that unimolecular NO₂ cleavage is the initial reaction for noncompressed BCHMX. We will discuss the reaction mechanisms in more detail in the section describing the DFT-MD as above. This leads to the results in Figure 3. Here we find that BCHMX molecules start to react at ~2050 K (12 ps, leading to an instantaneous pressure increase to ~22.0 GPa), involving both intermolecular hydrogen transfer and NO₂-releasing mechanisms. One hydrogen atom in a CH bond in BCHMX reacts with the N atom of the nearby BCHMX to form a N–H bond simultaneously with the first NO₂ released. Thus, the initial reaction in the compressed system is different from that for the noncompressed case. Then at ~2150 K (about 1 ps later), BCHMX molecules decompose one by one, releasing more NO₂ and HONO molecules, which is similar to the mechanism in RDX and HMX in which hydrogen transfer to/from HONO takes places at the beginning.

3.2. Reaction Mechanism from Finite Cluster Calculations on Either the Single Molecule or the Dimer. To understand the nature of the initial decomposition reactions of noncompressed and compressed BCHMX, we extracted the activated intermediates from the condensed-phase simulation and analyzed the reaction mechanism as a gas-phase reaction. The crystal structure for BCHMX has one long NN bond (1.427 Å) and three short NN bonds (1.355, 1.352, and 1.365 Å). We were surprised by the big differences and optimized the crystal structure using PBE-ulg where we find excellent agreement (bonds of 1.412, 1.364, 1.369, and 1.372 Å). Apparently these variations are related to the dihedral CNNO angle which should be 0 for the best resonance stabilization of the NO₂ with the ring N. Indeed the dihedral angle for the long bond is O₃–N₂–N₆–C₁ = 28.0°, whereas the other three are O₆–N₃–N₇–C₂ = 11.0°, O₇–N₄–N₈–C₂ = 12.5°, and O₂–N₁–N₅–C₁ = 10.5°. However, for the gas phase our calculations (PBE-ulg) lead to a symmetrical BCHMX molecule with C₃ symmetry where two N–N bonds (N₁–N₅ and N₃–N₇) are equal distance of 1.430 Å and the other two (N₂–N₆ and N₄–N₈) of 1.450 Å. The dihedrals O₇–N₄–N₈–C₂ and O₃–N₂–N₆–C₁ are equal to 26.4° and the other two dihedrals O₂–N₁–N₅–C₁ and O₆–N₃–N₇–C₂ to 20.6°. These results are compared in Table 1 with results for B3LYP

**Figure 2.** Species analysis for decomposition of noncompressed BCHMX heated from 300 to 3000 K over 20 ps. The first decomposition reaction for BCHMX occurs at ~1700 K (9.5 ps), releasing one NO₂ molecule. The second reaction occurs at 1910 K (11.0 ps) releasing two NO₂ dissociating from the decomposed BCHMX fragment.

**Figure 3.** Species analysis for the decomposition of 30% compressed BCHMX heated from 300 to 3000 K over 20 ps. At ~2050 K (12 ps), two BCHMX molecules interact and release one NO₂ molecule, and then after another 1 ps (at ~2150 K), BCHMX molecules start decomposing one by one, releasing more NO₂ and HONO molecules.
which are similar. This big increase in NN is apparently due to the larger dihedrals in the gas phase.

First we considered the noncompressed periodic system, BCHMX-TS1-Int1-TS2-Int2, is the pathway extracted from the DFT-MD trajectory and then optimized as a gas-phase reaction. We found (see Figure 4) that the nitro-aci isomerization reaction to make the aci isomer of BCHMX Int1 (at $+37.7$ kcal/mol) via TS1, in which one H atom of BCHMX transferred from the C atom to the nearby O atom, has a barrier of 38.5 kcal/mol, making it the rate-determining step (RDS). Starting with Int1 the easiest pathway to eliminate the NO$_2$ molecule is breaking the N–N bond via TS2 (at $+41.2$ kcal/mol).

### Table 1. N–N Bond Distance and CNNO Dihedral Angle for All Four N–N Bonds of the BCHMX Molecule in the Condensed Phase (Experiment and PBE-ulg Level) and the Gas Phase (B3LYP, PBE, PBE-ulg Levels)

<table>
<thead>
<tr>
<th>N–N bond and CNNO torsion</th>
<th>condensed phase</th>
<th>gas phase$^a$</th>
<th>PBE-ulg</th>
<th>PBE</th>
<th>B3LYP</th>
</tr>
</thead>
<tbody>
<tr>
<td>N1–N5 (Å)</td>
<td>1.365</td>
<td>1.364</td>
<td>1.430</td>
<td>1.430</td>
<td>1.400</td>
</tr>
<tr>
<td>N2–N6 (Å)</td>
<td>1.412</td>
<td>1.427</td>
<td>1.450</td>
<td>1.450</td>
<td>1.430</td>
</tr>
<tr>
<td>N3–N7 (Å)</td>
<td>1.355</td>
<td>1.369</td>
<td>1.430</td>
<td>1.430</td>
<td>1.400</td>
</tr>
<tr>
<td>N4–N8 (Å)</td>
<td>1.352</td>
<td>1.372</td>
<td>1.450</td>
<td>1.450</td>
<td>1.430</td>
</tr>
<tr>
<td>O2–N1–N5–C1 (deg)</td>
<td>16.2</td>
<td>10.5</td>
<td>20.6</td>
<td>20.2</td>
<td>16.9</td>
</tr>
<tr>
<td>O3–N2–N6–C1 (deg)</td>
<td>27.5</td>
<td>28.0</td>
<td>26.4</td>
<td>25.5</td>
<td>22.9</td>
</tr>
<tr>
<td>O6–N3–N7–C2 (deg)</td>
<td>14.8</td>
<td>11.0</td>
<td>20.6</td>
<td>20.2</td>
<td>16.9</td>
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<tr>
<td>O7–N4–N8–C2 (deg)</td>
<td>12.8</td>
<td>12.5</td>
<td>26.4</td>
<td>25.5</td>
<td>22.9</td>
</tr>
</tbody>
</table>

$^a$Data from ref 5. $^b$Data from crystal structure optimized by VASP in PBE-ulg level. $^c$Data from molecular structure optimized by Jaguar in PBE-ulg, PBE, and B3LYP levels.

Figure 4. Mechanism of the first nitro-aci isomerization event and for the NO$_2$-releasing reactions from finite cluster calculations starting from the trajectory from the cook-off simulation of noncompressed BCHMX. The first step (TS1 at $38.5$ kcal/mol) is intramolecular hydrogen transfer within the BCHMX molecule, from CH to OH, which is followed quickly by NO$_2$ release (TS2 at $41.2$ kcal/mol). Configurations TS1, Int1, and Int2 were extracted from the DFT-MD trajectory and then optimized as a gas-phase reaction, while Int1-TS3-Int3 is a proposed probable reaction path to release HONO. The DFT-MD simulations also did not find the TS4 to Int4 pathway to release HONO, although the DFT finds this to have a barrier (TS4) of 33.9 kcal/mol, which is a lower barrier HONO releasing pathway in the gas phase. Unit is in kcal/mol.
kcal/mol) to form an intermediate \( \text{Int}_2 \) (shown in Figure 4), which we find to have a barrier of 3.5 kcal/mol above \( \text{Int}_1 \).

Although BCHMX DFT-MD on the noncompressed period system did not find a HONO-releasing reaction in the first decomposition of BCHMX, the adiabatic QM calculations find two HONO-releasing pathways. One pathway stretches the N=N bond via \( \text{TS}_3 \) (at +40.5 kcal/mol) to release a HONO molecule, leading to a barrier of 2.8 kcal/mol above the aci isomer of BCHMX \( \text{Int}_1 \) (at +37.7 kcal/mol). The other pathway is for BCHMX to release HONO directly via \( \text{TS}_4 \), with a barrier of 33.9 kcal/mol, which is lower than the barrier of the nitro-aci isomerization reaction. In the DFT-MD simulation for the noncompressed condensed phase the HONO-releasing reaction seems to be hindered by the nearby BCHMX molecule with a contact distance of 2.76 Å, whereas the \(-\text{NO}_2\) that dissociates is \(~3.62\) Å from the closest BCHMX (see Figure S3, SI). This seems to explain why HONO release was not observed in the fast heating rate MD simulation.

The reaction energy (barrier) for unimolecular NO\(_2\) cleavage reaction without hydrogen transfer in BCHMX is calculated to be 39.0 kcal/mol (see Figure 4), which is 1.3 kcal/mol higher than the intramolecular hydrogen-transfer barrier, explaining why the intramolecular hydrogen-transfer first takes place before the unimolecular NO\(_2\) cleavage reaction in MD simulation.

Next we consider the compressed BCHMX case. Here the first reaction is an electrophilic substitution, leading to the intermolecular hydrogen-transfer and NO\(_2\)-releasing reaction, as shown in Figure 5. This reaction takes place between two BCHMX molecules in the compressed phase at about 2050 K, and in this reaction one hydrogen atom (H6) of the C3 atom in BCHMX reacts with the N atom (N5) of the nearby BCHMX to make a new N=H bond via \( \text{TS}_6 \), followed by forming the N=H bond and the first NO\(_2\) release. Configurations \( \text{TS}_6, \text{Int}_6, \) and \( \text{TS}_7 \) were extracted from the DFT-MD trajectory and then optimized as a gas-phase reaction. BCHMX-TS8-Int8 is another possible path to release HONO calculated in the gas phase but not observed in the DFT-MD simulations. Unit is in kcal/mol.

Figure 5. Mechanism of the first intermolecular hydrogen-transfer and NO\(_2\)-releasing reactions from finite cluster calculations starting from the trajectory from the cook-off simulation of compressed BCHMX. One hydrogen atom (H6) of the C3 atom in BCHMX (reacts with the N atom (N5) of the nearby BCHMX to make a new N=H bond via \( \text{TS}_6 \), followed by forming the N=H bond and the first NO\(_2\) release. Configurations \( \text{TS}_6, \text{Int}_6, \) and \( \text{TS}_7 \) were extracted from the DFT-MD trajectory and then optimized as a gas-phase reaction. BCHMX-TS8-Int8 is another possible path to release HONO calculated in the gas phase but not observed in the DFT-MD simulations. Unit is in kcal/mol. The Journal of Physical Chemistry C Article DOI: 10.1021/jp510328d J. Phys. Chem. C 2015, 119, 2290–2296

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barrier for RDX11 and 39.8 kcal/mol NO2 dissociation barrier. BCHMX explains the higher sensitivity of BCHMX.

initial decomposition reaction of TKX-50 38 and DTTO, 39 for HMX.17 Thus, the lower initial decomposition barrier for atom with a halogen atom such as F or Cl. Also synthesizing the

4. CONCLUSIONS

We examined the initial chemical reactions of the explosive BCHMX by DFT-MD simulations to discover the reaction steps at high temperature followed by full DFT calculations on a finite cluster to obtain the transition states for the reactions. Key points of the our simulations are

(1) For noncompressed BCHMX, DFT-MD finds that nitroacitiomerization reaction is first, followed by unimolecular NO2 release.
(2) For compressed BCHMX (20 GPa), DFT-MD finds that intermolecular hydrogen transfer is first, followed by bimolecular NO2 release.
(3) For compressed BCHMX, the intermolecular hydrogen-transfer decreases the reaction barrier for releasing NO2 by ~7 kcal/mol.
(4) The HONO releasing reaction is more favorable (29.9 kcal/mol barrier) for compressed BCHMX, with a reaction barrier 10 kcal/mol lower than the unimolecular NO2 releasing reaction and ~3 kcal/mol lower than the bimolecular NO2 releasing reaction.

These simulations rationalize the high impact sensitivity of BCHMX, compared to RDX and HMX. It arises because of much lower initial reaction barriers (29.9–33.5 kcal/mol) for BCHMX compared to the 39.0 kcal/mol NO2 dissociation barrier for RDX11 and 39.8 kcal/mol NO2 dissociation barrier for HMX.17 Thus, the lower initial decomposition barrier for BCHMX explains the higher sensitivity of BCHMX.

We also have used DFT-MD simulations to investigate the initial decomposition reaction of TXK-5058 and DTTO,39 which have been predicted to be insensitive energetic materials. Here we find that the first decomposition barrier is lower than 45.1 kcal/mol for TXK-50 and lower than 45.9 kcal/mol for DTTO. Thus, the lower decomposition barrier for BCHMX can explain the higher sensitivity of BCHMX.

In order to decrease the sensitivity of BCHMX, we suggest modifications to avoid the intermolecular hydrogen-transfer and HONO reaction. This can be done by replacing the H atom with a halogen atom such as F or Cl. Also synthesizing the cocrystal combining BCHMX with other EMs, such as HMX, might decrease the sensitivity of energetic materials.40,41

ASSOCIATED CONTENT

Supporting Information

Atomic coordinates of all intermediates and TS shown in this study, coordinates for structure of BCHMX, and the bond type and bond cutoff in the fragment analysis. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes
The authors declare no competing financial interest.

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