Molecular Dynamics Simulation of the Order-Disorder Phase Transition in Solid NaNO₂

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Molecular dynamics simulation of the order-disorder phase transition in solid NaNO₂

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We present molecular dynamics simulations of solid NaNO₂ using pair potentials with the rigid-ion model. The crystal potential surface is calculated by using an a priori method which integrates the ab initio calculations with the Gordon-Kim electron gas theory. This approach is carefully examined by using different population analysis methods and comparing the intermolecular interactions resulting from this approach with those from the ab initio Hartree-Fock calculations. Our numerics show that the ferroelectric-paraelectric phase transition in solid NaNO₂ is triggered by rotation of the nitrite ions around the crystallographical c axis, in agreement with recent x-ray experiments [Gohda et al., Phys. Rev. B 63, 14101 (2000)]. The crystal-field effects on the nitrite ion are also addressed. An internal charge-transfer effect is found.

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I. INTRODUCTION

Sodium nitrite is a ferroelectric at room temperature. It has the orthorhombic structure, space group C\textsubscript{2h\textsubscript{2}}-Im\textsubscript{2}m\textsubscript{2}, with the dipole vector of the V-shaped nitrite anions aligned parallel to the crystallographic b direction, as shown in Fig. 1. The ferroelectric-paraelectric phase transition takes place at about 437 K, where the high-temperature phase is orthorhombic, space group D\textsubscript{2h\textsubscript{2}}-Im\textsubscript{2}mm, with the dipoles disordered with respect to the b axis. In a narrow temperature range from 435.5 K to 437 K, there exists an incommensurate antiferroelectric phase. The melting temperature is 550 K. Distinguished from displacive ferroelectrics in which the ferroelectric transition is driven by soft phonon modes, NaNO₂ offers a model system for research of the order-disorder structural phase transition and any associated ferroelectric instability.¹-³

Extensive experimental work on NaNO₂ has been devoted to probing the mechanism of the NO₂⁻ polarization reversal that triggers the order-disorder transition. The majority of studies support the c-axis rotation model, but there were also results favoring the a-axis rotation model.⁴ Recently, refined x-ray studies over a wide temperature range reinforced the c-axis rotation model.⁵,⁶ On the theoretical side, the microscopic model calculations done by Ehhardt and Michel supported the c-axis rotation mechanism,⁶ whereas mixed double rotations around the a axis and the c axis were suggested by Kinase and Takahashi.⁷ It has long been desirable to apply computer molecular dynamics (MD) simulations to NaNO₂ in order to achieve unambiguous understanding of the polarization reversal mechanism. Earlier MD simulations with empirical Born-Mayer pair potentials detected the c-axis rotation in above-room-temperature NaNO₂.⁸-¹⁰ Unfortunately, the low-temperature structure produced by those simulations was antiferroelectric and apparently disagreed with the experimental observations.

Lu and Hardy pointed out that the overall phase behavior of NaNO₂ could be simulated by using an a priori approach to construct the crystal potential surface (PES).¹¹ The Lu-Hardy (LH) approach was originally designed to deal with molecular crystals such as K\textsubscript{2}SeO₄, where there exists a mix of bonding types, that is, the intermolecular interactions are mostly ionic, but the constituent atoms in a molecule (SeO\textsubscript{4}²⁻ in K\textsubscript{2}SeO₄) bond covalently. In the LH approach, the intramolecular interactions were treated by applying the ab initio self-consistent field method to the gas-phase molecules, while the intermolecular pair potentials were computed within the Gordon-Kim (GK) electron-gas theory.¹² The crux of their application of the GK theory is how to partition the ab initio molecular charge density among the constituent atoms. Since there is no unique way to separate the charge density of a highly covalently bonded molecule, Lu and Hardy suggested equal separation in a spirit similar to the Mulliken population analysis (MPA). By using this atomic-level method, we could successfully describe the phase transitions in fluoroperovskites¹³ and ionic crystals with polyatomic molecules including SeO\textsubscript{4}²⁻,¹⁴ ClO\textsubscript{4}⁻,¹⁵ SO\textsubscript{4}²⁻,¹⁶ SiO\textsubscript{4}⁴⁻,¹⁷ and NO₃⁻¹⁸-⁲⁰ Note that the MPA happens to preserve the (zero) dipole moment of these molecules.

However, several problems appear when we moved on to deal with NaNO₂ where the NO₂⁻ radical has nonzero dipole moment and stronger chemical bonding. First, it is well known that the MPA, while certainly the most widely employed, is also somewhat arbitrary and the most criticized.²¹ In particular, the MPA overestimates the dipole moment of the free NO₂⁻ ion by about 120%. Other difficulties involved the free-ion approximation. Unlike in monatomic ionic crystals, there may exist considerable internal charge-transfer effects in molecular ionic crystals. Electronic band-structure calculations²² indicated that within a nitrite entity, the nitrogen atom and two oxygen atoms bond covalently, leading to high charge transferability between these constituent atoms. Therefore, in solid NaNO₂ the NO₂⁻ group will feel different crystal-field environments as it rotates and responds by redistributing the charge density among its three constituent atoms.

Our goals in this paper are twofold. First, we show that our atomistic level simulation methods involving pair potentials with the rigid-ion model is capable of correctly describing the phase behavior of NaNO₂. Second, we systematically examine the LH approach to understand why it works so well in molecular ionic crystal systems by the following
steps: (i) we develop another population analysis method that preserves the molecular dipole moment by directly fitting the *ab initio* charge density of a molecule; (ii) we carry out *ab initio* Hartree-Fock (HF) calculations of the intermolecular interactions and find that the pair potentials from the rigid-ion model can correctly reproduce the *ab initio* results; and (iii) we investigate the crystal-field effects on the NO$_2$ ion by embedding the ion and its first shell of neighbors in a lattice of point charges and find a remarkable internal charge-transfer effect. 23 Several MD simulations based on these modifications of the LH approach are also performed. The ferroelectric-paraelectric transition triggered by the c-axis rotation of the nitrite ions is observed in all versions of the LH approach. However, the transition temperatures predicted by these simulations are lower than the experimental values. Furthermore, the transition temperatures obtained from the original version are higher than those predicted by modified versions and closer to the experimental values. After careful examination, we notice that in the original LH approach, the NO$_2^-$ dipole moments were generally enhanced by about 20%. Such enhancement reinforces the ferroelectric state by raising the rotational barriers of NO$_2^-$, thus mimicking the anion polarization effect in the mean-field sense. Therefore, we conclude that the anion polarization effect is particularly important for the quantitative study of NaNO$_2$.

This paper is organized as follows. Section II describes the method used to obtain the PES of ionic molecular crystals. Section III analyzes the resulting intermolecular potentials in comparison with those obtained from *ab initio* calculations. Section IV presents the results of our MD simulations. The crystal-field effects on NO$_2^-$ are discussed in Sec. V. Concluding remarks are made in Sec. VI.

II. METHODOLOGY FOR PES CONSTRUCTION

Our MD simulation technique originates from the GK model for simple ionic crystals such as alkali halides, allowing that (molecular) ions in a crystal environment can be described as free ions. 24–26 Then it was extended to deal with molecular ionic crystals such as K$_2$SeO$_4$ in which strong intramolecular covalency exists. 12,14 The main idea is that the molecular ion (SeO$_4^{2-}$ in K$_2$SeO$_4$) is treated as a single entity, and intramolecular and intermolecular interactions are treated separately: first we perform *ab initio* quantum chemistry calculations for the whole molecular ion to obtain the optimized structure, the force constants, and the whole electron density $\rho(\mathbf{r})$. The intramolecular interactions are described by force constants within the harmonic approximation. As for the intermolecular interactions, we have to carry out electron population analysis to separate $\rho(\mathbf{r})$ onto each individual atom in the molecular ion, then use the Gordon-Kim electron-gas model to calculate the intermolecular pair potentials. This approach provides a parameter-free description for the crystal potential-energy surfaces, which allow structural relaxation, MD simulation, and lattice dynamics calculations.

In calculating the intermolecular forces, there are three major approximations as discussed in the following.

(1) We assume that the geometries and electronic densities of the separate ions remain unchanged once they have been obtained under given circumstance, such as in the equilibrium state of the gas or crystal phases. This approximation is the fundamental basis for the GK electron-gas theory. Generally speaking, we found that in an ionic crystal there is no strong chemical bond between ions, hence this approximation is reasonable.

(2) When dealing with the intermolecular interaction, we assume that the charge density of a rigid ion can be separated into its atomic constituents.

(3) We assume that the crystal potential energy is composed of the intermolecular and the intramolecular interaction, where the intramolecular interaction is expressed in terms of force fields and the intermolecular interaction is a sum of interatomic pair potentials.

Atomistic level simulations utilizing pair potentials and the rigid-ion model have achieved great success in describing many ionic systems. 25 We showed that this scheme can correctly describe the phase-transition behaviors of alkali halide fluoroperovskites, 13 and molecular crystals with tetrahedral 12,14–17 and equilateral triangular 18–20 radicals. We shall discuss this scheme and its modification in more detail in the remaining part of this section and the following section.

A. Pairwise additive approximation

In the GK model, we evaluate the interaction between two molecules based on the electron density, which is approximated as the sum of component densities taken from HF calculations. That is, if $\rho_A$ and $\rho_B$ are the component densities, then the total density is $\rho_{AB} = \rho_A + \rho_B$, and interaction potential is computed as the sum of four terms: Coulombic, kinetic, exchange, and correlation energies which are expressed in terms of the charge densities.
and the short-range part,

\[ V^\text{C} = V^\text{C}_A - V^\text{C}_B. \]  

Equation (2) is essentially the electrostatic interaction energy when the charge densities of the molecules are distributed as point charges on the constituent atoms, which is known as the Madelung potential energy.

The non-Coulombic energy terms are expressed in the uniform electron-gas formula,

\[ V_i = \int d\mathbf{r} [\rho_{AB}(\mathbf{r}) \epsilon_i(\rho_{AB}) - \rho_A(\mathbf{r}) \epsilon_i(\rho_A) - \rho_B(\mathbf{r}) \epsilon_i(\rho_B)], \]  

where \( \epsilon_i(\rho) \) is one of the energy functionals for the kinetic, exchange, and correlation interactions. 28 Note that Eq. (4) is not composed of pair potentials. In order to obtain the effective pairwise potentials, we approximate Eq. (4) using

\[ V_i = \sum_{m=A} \sum_{n=B} \int d\mathbf{r} [\rho_{mn}(\mathbf{r}) \epsilon_i(\rho_{mn}) - \rho_m(\mathbf{r}) \epsilon_i(\rho_m) - \rho_n(\mathbf{r}) \epsilon_i(\rho_n)], \]  

where \( \rho_{mn} = \rho_m + \rho_n \). \( \rho_m \) and \( \rho_n \) are the charge densities of individual atoms in the A and B molecules, respectively, which are obtained by a population analysis as described in the following section.

Even though the non-Coulombic forces as determined by Eq. (4) are not strictly additive, the above approximation appears to be adequate except at very short distances. As pointed out by Waldman and Gordon, 29 the main reason as to why this approximation is valid is because the Coulombic force, the largest contribution to the potentials, is additive. Based on our calculations, we find additivity of \( V_i \) holds only to within about 50%; however, the overlap contribution to the electrostatic energy dominates \( V_i \) and renders additivity to within 10%. One final remark is in order, for the sake of simplifying the two-electron integral in Eq. (1), the charge densities \( \rho_m \) and \( \rho_n \) are taken as their spherical average. As a result, the Coulombic interaction is not exactly evaluated. Nevertheless, as we shall show in Figs. 2 and 3, this error is compensated by those due to the pairwise additive approximation.

To summarize this section, we have demonstrated that it is possible to analytically express the intermolecular potentials \( V^C + V^C_i + V_i \) using Eqs. (2), (3), and (5) once the charge density of each individual atom is obtained by an electronic population analysis.

B. Electronic population analysis and intramolecular interactions

In this section, we discuss the ways to separate the electron density \( \rho(\mathbf{r}) \) of a molecule into its atomic constituents.
Then the electronic density of the molecule is

$$\rho(\mathbf{r}) = |\psi(\mathbf{r})|^2 = \sum_{ijkl} d_{ijkl} \chi_i(\mathbf{r} - \mathbf{R}_i) \chi_j(\mathbf{r} - \mathbf{R}_j),$$

(8)

where $d_{ijkl} = 2 w_{ijkl}$, which can be divided into two parts, namely, the net ($i = j$) and overlap ($i \neq j$) populations. The latter cannot be ignored in the presence of strong intramolecular covalency. Therefore, separating $\rho(\mathbf{r})$ into its atomic constituents is to split the overlap population. However, the way to achieve that is not unique. For example, we can introduce a set of weights $w_{ijkl}$ due to different criteria such that

$$d_{ik,ik} = d_{ik,ik} + \sum_{j \neq i,k} w_{ijkl} d_{ijkl} \int \chi_i(\mathbf{r} - \mathbf{R}_i) \chi_j(\mathbf{r} - \mathbf{R}_j) d\mathbf{r},$$

$$d_{jl,jl} = d_{jl,jl} + \sum_{i \neq j,k} (1 - w_{ijkl}) d_{ijkl} \int \chi_j(\mathbf{r} - \mathbf{R}_j) \chi_i(\mathbf{r} - \mathbf{R}_i) d\mathbf{r},$$

$$d_{ik,il} = d_{ik,il},$$

then we can rewrite Eq. (8) as

$$\rho(\mathbf{r}) = \sum_i \rho_i(\mathbf{r}) = \sum_{ijkl} d_{ijkl} \chi_i(\mathbf{r} - \mathbf{R}_i) \chi_j(\mathbf{r} - \mathbf{R}_j),$$

(10)

where $\rho_i(\mathbf{r})$ is the atomic density of atom $i$.

In our previous studies, the overlap electronic density is equally separated, i.e., $w_{ijkl} = 1/2$ in Eq. (9), similar to the MPA. In Table I, we present the electronic multipole moments of $\text{SnCl}_6^{2-}$, $\text{SeCl}_4^{2-}$, $\text{SeO}_2^{2-}$, $\text{SO}_2^{2-}$, $\text{ClO}_4^{2-}$, $\text{CO}_3^{2-}$, and $\text{NO}_3^{2-}$ calculated using the MPA, and compare them with the \textit{ab initio} values. We note that for these symmetrical molecules, the MPA preserves total charge and zero dipole moment. However, for a molecular ion such as $V$-shaped $\text{NO}_2^{2-}$ or linear $\text{CN}^-$, the Mulliken population

<table>
<thead>
<tr>
<th>$AB_n$</th>
<th>$\mu$</th>
<th>$\vartheta_{zz}$</th>
<th>$\Omega_{zzz}$</th>
<th>$\Phi_{zzzz}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{CN}^-$</td>
<td>0.14 (0.17)</td>
<td>-27.35 (-29.41)</td>
<td>7.99 (10.68)</td>
<td>-143 (-166)</td>
</tr>
<tr>
<td>$\text{NO}_2^-$</td>
<td>0.57 (0.26)</td>
<td>-19.64 (-21.64)</td>
<td>-2.60 (-5.87)</td>
<td>-60 (-76)</td>
</tr>
<tr>
<td>$\text{NO}_3^{-}$</td>
<td>0 (0)</td>
<td>-15 (-16)</td>
<td>0 (0)</td>
<td>-38 (-41)</td>
</tr>
<tr>
<td>$\text{CO}_2^{-}$</td>
<td>0 (0)</td>
<td>-17 (-17)</td>
<td>0 (0)</td>
<td>-48 (-48)</td>
</tr>
<tr>
<td>$\text{ClO}_4^{-}$</td>
<td>0 (0)</td>
<td>-111 (-111)</td>
<td>0 (0)</td>
<td>-548 (-548)</td>
</tr>
<tr>
<td>$\text{SO}_2^{-}$</td>
<td>0 (0)</td>
<td>-119 (-120)</td>
<td>0 (0)</td>
<td>-620 (-627)</td>
</tr>
<tr>
<td>$\text{SeO}_2^{-}$</td>
<td>0 (0)</td>
<td>-143 (-143)</td>
<td>0 (0)</td>
<td>-823 (-816)</td>
</tr>
<tr>
<td>$\text{SiO}_2^{-}$</td>
<td>0 (0)</td>
<td>-157 (-158)</td>
<td>0 (0)</td>
<td>-1011 (-1019)</td>
</tr>
<tr>
<td>$\text{ScCl}_6^{2-}$</td>
<td>0 (0)</td>
<td>-319 (-318)</td>
<td>0 (0)</td>
<td>-5112 (-5071)</td>
</tr>
<tr>
<td>$\text{SnCl}_6^{2-}$</td>
<td>0 (0)</td>
<td>-845 (-844)</td>
<td>0 (0)</td>
<td>-19834 (-19668)</td>
</tr>
</tbody>
</table>

*In the HF calculations, basis set D95* were used for $AB$ and $AB_2$, 6-31G* for $AB_3$ and $AB_4$, and 3-21G* for $AB_5$.

*The electrostatic moments $\mu$ (dipole), $\vartheta$ (quadrupole), $\Omega$ (octapole), and $\Phi$ (hexadecapole) refer to the center of mass of the molecule with the standard orientation defined in \textit{GAUSSIAN} 98 (Ref. 30).
seems inadequate. Given total charge and the dipole moment 0.26 a.u. of NO₂ obtained from the \textit{ab initio} calculations (Table I), a population analysis which preserves these values would give rise to -0.092e on N and -0.454e on O. Whereas, using the MPA ($w_{ijkl}=1/2$), the charges on the N and O atoms are 0.1624e and -0.5812e, respectively, which renders the dipole moment 0.57 atomic unit, overestimated by 120%.

Therefore, it is desirable to determine $w_{ijkl}$ in such a way that the calculated multipole moments of the molecule are consistent with the \textit{ab initio} values. One possible way is to evaluate $w_{ijkl}$ by fitting the \textit{ab initio} charge density, as shown in Eqs. (9) and (10), with the values of multipole moments as constraints. An alternative way is to directly fit the shape density, Eq. (10), with $\tilde{d}_{ik,jl}$ being the parameters and $\chi_{kl}(r-R_i)\chi_i(r-R_j)$ being the dependent variables. To simplify the computation, only the radial parts of $\chi_{kl}(r-R_i)\chi_i(r-R_j)$ were kept. This fitting population analysis (FPA) is similar to that proposed by Parker and his co-workers as an alternative implementation of the GK model. 32

To obtain the atomic electron densities on the nitrogen and oxygen atoms by the above MPA or FPA of the electron density of the nitrite anion, \textsc{gaussian} 98 program package 30 is employed to solve the Hartree-Fock-Roothan equations for NO₂ in the gas phase; the in-crystal ions will be discussed in Sec. V. The atomic-orbital basis sets used are a double-zeta basis with polarization functions (D95*) for the nitrogen and oxygen atoms. The optimized geometry of NO₂ with the N-O bond length $R_{NO}=1.233$ Å and the bond angle $\theta_{ONO}=116.6^\circ$ is comparable to the experimental geometry of NO₂ in the ferroelectric phase of NaNO₂ with $R_{NO}=1.236$ Å and $\theta_{ONO}=115.4^\circ$. 34

The intramolecular interaction within the harmonic-oscillator approximation can also be obtained from frequency analysis in \textsc{gaussian} 98.30 The lowest vibration frequency (1192 K) of NO₂ is considerably higher than the highest vibration frequency (318 K) obtained from Raman spectroscopy as well as the order-disorder transition temperature (437 K). Therefore, it is justified to treat the internal motion of the nuclei in the NO₂ group within the harmonic approximation, or even as a rigid rotor.

Note that the polarizability of NO₂ at its optimized geometry is highly anisotropic, that is, with $\alpha_{xx}=7.820$, $\alpha_{yy}=10.823$, and $\alpha_{zz}=23.825$ in atomic units (see Fig. 1 for coordinate convention). Thus, one would expect this polarization to seriously affect the intermolecular pair potentials, and thus render the rigid-ion approximation in question. We shall examine this question in the following section by comparing the intermolecular potentials obtained from this approach with those from \textit{ab initio} HF calculations.

III. INTERMOLECULAR POTENTIALS

The \textit{ab initio} HF calculations are performed by using \textsc{gaussian} 98 program package 30 to scan the potential-energy surface of NO₂ : Na⁺ and NO₂ : NO₂ dimer. The D95* basis set is used for the nitrogen and oxygen atoms. As for the sodium atoms, we used both the standard 6-31G* basis and the Slater-type orbitals for Na⁺ taken from the Clementi and Roetti table.36 It turned out that the difference between them is small. In these calculations, the geometrical variables of NO₂ are frozen at their equilibrium values, since we showed previously that the NO₂ group in NaNO₂ could be treated as a rigid rotor. However, in the HF calculations, the electronic structure is allowed to vary in order to minimize the total energy, thus the electronic polarization effects are included.

In our calculations of intermolecular interactions, one NO₂ is fixed with its center of mass being the origin of the coordinate system, the dipole vector pointing to the y axis and the O-O line being aligned parallel to the z axis (see Fig. 1). Then, Cartesian coordinates ($x$, $y$, $z$) are the position of Na⁺ or the center of mass of another NO₂.

In order to study the effects of different population analysis schemes on the pair potentials, we performed three different calculations based on our rigid-ion models: (i) MPA with pair potentials [Eq. (5)], (ii) FPA with pair potentials, and (iii) FPA with nonpair potentials [Eq. (4)], referred to as models I, II, and III, respectively. We shall show in the following that the electronic polarization effect could be revealed from examining the differences between models I and II, while errors due to the pairwise additive approximation could be analyzed from the differences between models II and III.

In Figs. 2 and 3, we compare the intermolecular potentials obtained from models I–III and those from the \textit{ab initio} HF calculations. The results for the NO₂ : Na⁺ dimer are shown in Fig. 2: we find in both models I and II that the overall shapes of the GK potentials as a function of molecular separation agree with the \textit{ab initio} results, with the lowest NO₂ -Na⁺ potential energy emerging in the (0,−R,0) configuration; whereas model III predicts incorrectly the lowest potential energy in (R,0,0). It thus appears that in model II the errors caused by the pairwise additive approximation are compensated by the errors due to FPA.

On the other hand, the electronic polarization effect also manifests itself in Fig. 2 based on the following two observations. First, notice that our rigid-ion models I and II fit best to the \textit{ab initio} results for the (R,0,0) configuration [Fig. 2(d)]. We attribute that to the anisotropic polarizability of NO₂ ($\alpha_{xx}<\alpha_{yy}<\alpha_{zz}$), thus the electronic cloud of NO₂ is most unlikely to be polarized along the x direction. Second, for (0,R,0) [Fig. 2(a)], the minimum potential energy in model II is closer to the \textit{ab initio} values than model I, whereas the result reverses for (0,−R,0) [Fig. 2(c)]. To understand this, we observe that for (0,R,0), Na⁺ is closer to O than to N, thus in the \textit{ab initio} calculations the electrons were attracted toward the N atom, leading to a smaller dipole moment. Therefore, the results yielded by model II, which produced smaller dipole moment than model I, tend to be closer to the HF results for (0,R,0). Obviously, the situation reverses for (0,−R,0).

Similarly, we show in Fig. 3 the NO₂ -NO₂ intermolecular potentials using models I–III and the HF method. The configurations are chosen as follows: the two NO₂ molecules are initially parallel at their experimental low-temperature separation (1.82 Å, 2.83 Å, 2.69 Å) and then
one of them rotates around each of the $x$, $y$, $z$ axes through its center of mass, as shown in Figs. 3(a)–3(c). The results of models I–III agree reasonably well with the ab initio calculations. On closer examination, in Fig. 3(c) in angle ranges $0^\circ$–$90^\circ$ and $270^\circ$–$360^\circ$, model I fits better to the ab initio results than models II and III. This feature would be important when the rotation of $\text{NO}_2^-$ around the $c$ axis dominates its rotations around the $a$ and $b$ axes, as will be demonstrated in Sec. IV.

Summarizing this section, in spite of the presence of electronic polarization when two molecules are brought closer, the intermolecular potentials for the $\text{NO}_2^-$ : $\text{Na}^+$ and $\text{NO}_2^-$ : $\text{NO}_2^-$ dimers could be correctly reproduced by using models I and II.

### IV. MOLECULAR DYNAMICS SIMULATIONS

After the potential-energy surface for NaNO$_2$ has been obtained, we are prepared to undertake MD simulations. Long-range Coulombic interaction among point charges calculated from the population analysis, while each of the short-range pair potentials are fitted by using an exponential-polynomial function accurate within 0.1%. In the following description, the $x$, $y$, $z$ directions correspond to the crystallographic $a$, $b$, $c$ directions of NaNO$_2$, respectively, see Fig. 1.

#### A. Lattice relaxation

Before we proceed with the molecular dynamics simulations, we perform lattice relaxation for the ferroelectric structure of NaNO$_2$ both with and without the $Im\bar{2}m$ space-group symmetry constraints. This relaxation procedure provides the crystal structure with zero force on each atom, that is, an energy extremum; it also produces a test to the PES because the resultant structures have to agree reasonably with the experimental data for further simulations to be reliable. We perform both static and dynamic relaxations: the static one is an application of the Newton-Raphson algorithm and usually results in finding a local minimum of the energy, and the dynamic one is a simulated annealing calculation for overcoming that limitation. We start the static lattice relaxation with the experimental parameters. In Table II we present the lattice and basis parameters deduced from the experiments and static relaxation. In all cases, the static relaxation produced essentially the same structure that strongly resembles the experimental structure. Most of the lattice constants in the relaxed structure are shorter than the experimental values (by 3.7%, 1.5%, and 8.5% for $a$, $b$, and $c$, respectively, in model I, and by 0.5%, $-2.4\%$, and 10% for $a$, $b$, and $c$, respectively, in model II). Hence the calculated volume is smaller than the experimental one by 13% for model I and 10% for model II, a common feature for simulations using the GK model, which will be addressed in more detail in the following section.

Next, we go on to relax the statically relaxed crystal structure to zero temperature using a simulated annealing algorithm, in which the amount of kinetic energy in the molecules slowly decreases over the course of the simulation. We find that the (zero-temperature) ground states in models I and II are close to the statically relaxed structures, whereas there are substantial changes taking place in model III. By monitoring the orientations of the nitrite ions, we find that the ground structure in model III, still orthorhombic with $a = 3.90494$ Å, $b = 4.8441$ Å, and $c = 5.0770$ Å, is ferroelectric with the dipole moments of $\text{NO}_2^-$ aligned along the $a$ axis rather than the experimental $b$ axis. So we conclude that the PES given by models III did not reflect reality. This concurs with the previous discussion on the intermolecular potentials (Sec. III). In the following, we use only models I and II to simulate the phase transition in NaNO$_2$.

#### B. MD simulations

Using the isenthalpic, isobaric ensemble, our MD simulation is started with a zero-temperature zero-pressure orthorhombic cell ($4a\times4b\times4c$) consisting of 512 atoms. Periodic boundary conditions are imposed to simulate an infinite crystal. The MD calculations are carried out in the Parrinello-Rahman scheme which allows both the volume and the shape of the MD cell to vary with time. The calculation of the energies and forces was handled by the Ewald method. A time step of 0.002 ps was used to integrate the equations of motion. In our heating runs, we raise the temperature of the sample in stages, 20 K each time, up to 1000 K. At each stage, the first 2000 time steps were employed to equilibrate the system, then 10,000 time steps were collected for subsequent statistical analysis. Since our simulations employ periodic boundary conditions, we cannot distinguish the incommensurate structure (i.e., phase II of solid NaNO$_2$).

### Table II. Experimental and theoretical structural parameters for the Im$\bar{2}m$ phase (III) of NaNO$_2$.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Experiment$^a$</th>
<th>Model I</th>
<th>Model II</th>
<th>Model III</th>
</tr>
</thead>
<tbody>
<tr>
<td>$a$</td>
<td>3.5024</td>
<td>3.3889</td>
<td>3.5013</td>
<td>3.7353</td>
</tr>
<tr>
<td>$b$</td>
<td>5.5209</td>
<td>5.4542</td>
<td>5.5485</td>
<td>5.4257</td>
</tr>
<tr>
<td>$c$</td>
<td>5.3789</td>
<td>4.9254</td>
<td>4.8403</td>
<td>4.9669</td>
</tr>
<tr>
<td>$\gamma/b$ of N (2$a$)</td>
<td>0.0781</td>
<td>0.0498</td>
<td>0.0433</td>
<td>0.0586</td>
</tr>
<tr>
<td>$\gamma/b$ of Na (2$a$)</td>
<td>0.5437</td>
<td>0.5537</td>
<td>0.5492</td>
<td>0.5437</td>
</tr>
<tr>
<td>$\gamma/b$ of O (4$d$)</td>
<td>$-0.0443$</td>
<td>$-0.0704$</td>
<td>$-0.0740$</td>
<td>$-0.0610$</td>
</tr>
<tr>
<td>$\gamma/c$ of O (4$d$)</td>
<td>0.1965</td>
<td>0.2111</td>
<td>0.2151</td>
<td>0.2098</td>
</tr>
</tbody>
</table>

$^a$From x-ray-diffraction experiments at 30 K, see Ref. 37.
the quadrupole moment be $q$, the lattice vectors associated with the $i$th anion rotate primarily about the $c$ axis. When $T>T_m$, $(\Theta_{xx} + \Theta_{yy})/2\Theta_{zz} \approx 1$, i.e., NaNO$_2$ becomes an orientational liquid.

Further, in Fig. 6 we show the mean-square atomic displacements $U_{ij} = \langle u_i^2 \rangle$ where $i,j = 1,2,3$ denotes the displacements along the $a,b,c$ axes, respectively. Different models of NO$_2^-$ reversal are expected to satisfy the following conditions. (1) Rotation around the $c$ axis: $U_{22}(N), U_{33}(N) < U_{11}(N)$ and $U_{22}(O), U_{33}(O) < U_{11}(O)$. (2) Rotation around the $a$ axis: $U_{11}(N), U_{22}(N) < U_{33}(N)$ and $U_{11}(O), U_{33}(O) < U_{22}(O)$. This figure relates to recent x-ray experiments which used the same quantities to investigate the polarization reversal mechanism. The experiments confirmed that the first condition holds for both ferroelectric and paraelectric phases. Another important feature revealed by the experiments is that $U_{22}(Na), U_{33}(Na) < U_{11}(Na)$ in the ferroelectric phase, whereas $U_{11}(Na), U_{33}(Na) < U_{22}(Na)$ in the paraelectric phase. That is, $U_{11}(Na)$ and $U_{22}(Na)$ are reversed across $T_C$. These features are reproduced in Fig. 6 with exception of $U_{11}(O), U_{33}(O) < U_{22}(O)$ in the paraelectric phase. This means the NO$_2^-$ motions in our simulations are more mobile than those in the real crystal, rendering the simulated transition temperatures lower than the experimental values of $T_C = 437$ K and the melting temperature 550 K. In other words, the barriers to NO$_2^-$ rotation in our models are too small.

![FIG. 4. Temperature variation of lattice constants $a$, $b$, $c$ (solid, dashed, and dotted lines, respectively; left scale) and volume of the unit cell (open circles; right scale) for (a) model I and (b) model II.](image)

![FIG. 5. Mean dipole moment $M_{ij}(T)$ and quadrupole moment $\Theta$ of the whole NaNO$_2$ crystal as a function of temperature for the MD runs for (a) model I and (b) model II.](image)
In addition, in Fig. 7, we show the average crystal structures of NaNO$_2$ at different temperatures. The ellipsoids in these pictures represent the root-mean-square deviations of the atoms from their average positions and thus indicate the thermal motions of these atoms. The $c$-axis rotation mode can be clearly seen, particularly in Fig. 7~$c$.

It is worth mentioning the less desirable agreement between theoretical and experimental volumes, namely, the 13% discrepancy for model I and 10% for model II. To address this we make one simple change: by following Waldman and Gordon, we increase the kinetic-energy term in the Gordon-Kim potentials by 5%, this reduces the discrepancy to 9% for model I and 6% for model II. Having done this we rerun the MD to obtain values of $T_c$ of 360 K for model I and 303 K for model II. While this change worsens the value for model I, the value for model II is virtually unchanged. And in both cases the transition mechanism is unaltered. Thus the slight hardening of the short-range potentials removes most of the volume discrepancies. However, there is no material change in the mechanism of the phase transition. This robustness of the results with respect to minor variations in the potential demonstrates that our basic conclusions remain valid.

**C. Rotational barriers**

Based on the above simulation results, the order-disorder phase transition in NaNO$_2$ involves the rotation of the nitrite ions. We devise a scheme to calculate the rotational barriers: Starting from the experimental ferroelectric structure taken to be the zero-energy state, we rotate one of the two nitrite ions in the unit cell around the $a$, $b$, and $c$ axes with the center of mass of the nitrite ion being fixed. The results are shown in Fig. 8. The bottom of each barrier is located at zero rotational angle that denotes the ferroelectric structure. For both models I and II, the rotation around the $c$ axis has an energy barrier 5–10 times smaller than those of the other rotations, which is a characteristic of nitrites. Hence, our calculations unambiguously reveal that the reorientation of NO$_2^-$ in the paraelectric phase occur essentially by rotations around the $c$ axis. Moreover, the barriers calculated in model I are higher than those in model II, confirming that the transition temperatures predicted by model I are higher than those predicted by model II.
V. CRYSTAL-FIELD EFFECTS

Generally speaking, our simulations predict lower transition temperatures than the experimental values. We believe that the main reason is due to underestimation of anion polarization effect in solid NaNO₂. In model II, NO₂⁻ possesses the same dipole moment as in its gas phase, thus the polarization effect is totally neglected. In model I, the dipole moment has been enhanced by Mulliken charges—such enhancement could be interpreted as taking into account the polarization effect in the mean-field sense—thus leading to higher transition temperatures than those obtained from model II. The above comparison indicates that more improvement for dealing with the anion polarization effect would be needed to raise the calculated transition temperatures closer to the experimental values.

To manifest the substantial crystal-field effects on the NO₂⁻ ions, we perform the following HF calculations: the crystal field of ferroelectric NaNO₂ is simulated by placing the nitrite ion and its six nearest Na⁺ cations at the center of a 4 × 4 × 4 orthorhombic point-charge lattice with spacings equal to the experimental lattice parameters. Charges in the faces of the lattice are scaled to maintain overall neutrality. All anions except the central NO₂⁻ are represented by single point charges on their centers of mass. Hence, there are 168 point charges surrounding the NO₂⁻ (Na⁺)₆ cluster. Calculations of this type were proposed by Fowler and co-workers in the studies of monatomic ions and cyanides. The same basis set D95* is employed for the in-crystal NO₂⁻ ion as for the free NO₂⁻ ion, while the minimal basis set STO-3G is used for the Na⁺ ions. The cations, however, are relatively insensitive to the crystal environment and they are included here only to account for their compressing effect on the NO₂⁻ wave functions. We find that adding extra basis functions to Na⁺ will not change the results significantly. The central NO₂⁻ initially points in the b direction as in the ferroelectric phase of NaN₂. Subsequently, we rotate the NO₂⁻ about the a, b, and c axes through its center of mass.

As shown in Fig. 9(a), the dipole moment of the central NO₂⁻ changes considerably as it rotates, indicating strong crystal-field effects on the reorientation of NO₂⁻. We also find that the dipole moment is sensitive to the location of the rotation center of NO₂⁻. In the context of population analysis, increase of the dipole moment of NO₂⁻ implies that more electrons are distributed on the O atom, i.e., electrons are flowing from the nitrogen atom to the oxygen atoms. Conversely, decrease of the dipole moment indicates a reversal in electron transfer. Therefore, we have demonstrated considerable intramolecular charge transfer, although the intermolecular charge transfer is usually small in ionic crystals.

Although strong crystal-field effects have been revealed by these ab initio calculations, the rotational barriers obtained from the polarizable-ion model [Fig. 9(b)] are in qualitative agreement with those from the rigid-ion models [Figs. 8(a) and 8(b)], confirming that the rigid-ion model is capable of describing the phase behavior in NaN₂.

VI. CONCLUDING REMARKS

We have presented MD simulations of NaN₂ using a hybrid a priori method consisting of ab initio calculations and Gordon-Kim electron-gas theory to analytically calculate the crystal potential surface. This method has been carefully examined by using different population analysis methods. We have carried out ab initio Hartree-Fock calculations of the intermolecular interactions for NO₂⁻:Na⁺ and NO₂⁻:NO₂⁻ dimers and concluded that the pair potentials of the rigid-ion model can correctly reproduce the ab initio results. We demonstrated that a rigid-ion model is capable of describing phase behavior in solid NaN₂.

We also addressed the crystal-field effects on the NO₂⁻ ion by performing Hartree-Fock calculations on a NO₂⁻ (Na⁺)₆ cluster embedded in a lattice of point charges. We conclude that the partial charges on the nitrogen and oxygen atoms are fluctuating in solid NaN₂ in response to changing crystal-field environments, which arise particularly from the rotation of the nitrite ions. Our MD simulations are based on two rigid-ion models using MPA and FPA, respectively. The model using MPA, which enhances the dipole moment of NO₂⁻ in the gas phase, gives rise to more comparable results with the experiments. Such enhancement stabilizes the ferroelectric structure by raising the rotational barriers of NO₂⁻, thus mimicking the anion polarization effect in the mean-field sense. To quantitatively simulate NaN₂, a more elaborate polarizable-ion model is needed.

ACKNOWLEDGMENTS

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