Theoretical and experimental study of the inelastic neutron scattering spectra of β-5-Nitro-2,4-dihydro-3H-1,2,4-triazol-3-one

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Received 27 June 2005; revised 15 July 2005; accepted 15 July 2005

Abstract

The inelastic neutron scattering (INS) spectra of β-5-Nitro-2,4-dihydro-3H-1,2,4-triazol-3-one (β-NTO) are presented to 1400 cm\(^{-1}\). The β-NTO vibrational frequencies observed differ considerably from the α-NTO vibrational frequencies and normal mode frequency calculations for the isolated molecule. The INS spectrum contains detail unobserved in the previous IR studies, including combinations and overtones of the phonon and internal modes of β-NTO. The INS spectra are compared with periodic DFT calculations to show that the periodic DFT results correctly predict the solid-state molecular vibrational frequencies.

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Keywords: β-NTO; Inelastic neutron scattering; NTO; 5-Nitro-2,4-dihydro-3H-1,2,4-triazol-3-one

1. Introduction

5-Nitro-2,4-dihydro-3H-1,2,4-triazol-3-one (NTO) was first characterized as an insensitive energetic material in 1988 [1]. The insensitivity of NTO to heat, friction, and impact makes it suitable for a wide variety of applications, such as explosive formulations, plastic-bonded explosives, and as a sodium azide replacement in automobile air bag systems [2]. The insensitivity of NTO has been attributed to the broad range of decomposition mechanisms, such as nitrogen–hydrogen bond cleavage, auto-catalysis, and NO\(_2\) elimination with the formation of amide fragments [3]. The exact decomposition mechanism has yet to be elucidated, but a wide range of activation energies suggests that thermal decomposition is highly dependent on environmental conditions [4]. Hydrogen bonding has also been proposed play a key role in the insensitivity of other energetic materials, such as TATB [5] and ANPZ [6]. Evidence of hydrogen bonding in the solid-state of NTO has been observed in both X-ray diffraction [7] and spectroscopic studies [4]. Crystals of NTO have been examined in some detail. Two polymorphic phases are known to exist. The α-polymorph is the most stable, but an accurate crystal structure has yet to be determined because it undergoes a significant degree of twinning [2].

β-NTO forms a monoclinic (\(P2_1/c\)) unit cell with each molecule linked by four hydrogen bonds, which follow a helical 2₁-screw axis. The crystal structure of β-NTO at 100 K is presented in Fig. 1. X-ray diffraction investigations into the variable temperature crystal structure were performed in the range of 100–298 K [7]. Both thermal expansion, observed in the plane perpendicular to the molecules, and thermal contraction, noted along the hydrogen bond axis, was found, which indicates the anharmonic character of β-NTO [7].

An intermolecular packing potential was developed to describe the β-NTO structure within the approximation of the rigid molecule [8]. The potential was based on the earlier derivation of intramolecular force fields of β-NTO in both the gas and solid-state phases. These force fields were partially constructed from experimental and calculated vibrational frequencies. A close correlation was observed between the experimental vibrational frequencies and a partial IR spectrum of α-NTO published by Lee and Gilardi [9]. The positions of the vibrational mode frequencies were found to vary considerably between the infrared spectra of pure NTO films and NTO molecules isolated in an argon...
matrix, indicating the considerable hydrogen-bonding interaction in the solid-state [8]. In this paper, we bring to evidence that the vibrational frequencies of the two polymorphs of NTO differ considerably and that the vibrational frequencies of β-NTO can be adequately described with solid-state density functional theory (DFT).

Vibrational spectroscopy has been extensively used to investigate the structural details and dynamical interactions in energetic materials [4,5,9–24]. Dlott’s theory of ‘vibrational up-pumping’ through ‘doorway’ modes [21] has generated interest in the low-frequency spectral region below 700 cm\(^{-1}\). Doorway modes are vibrations that arise because of weak anharmonic coupling between the phonons and intermolecular modes [21–23]. Previous work has indicated that energy transfer into vibrational modes below 700 cm\(^{-1}\) may be rate determining [24]. Nevertheless, vibrational information in this spectral region is often incomplete due to instrumental limitations or optical based selection rules. In these cases, inelastic neutron scattering (INS) spectroscopy is an attractive alternative to conventional vibrational spectroscopic methods. The INS intensity of all lattice and molecular vibrations is proportional to the displacements of the hydrogen atoms for a normal mode. The INS spectrum can be directly correlated to computational methods through calculation of the rms displacements of the mass-weighted normal mode hydrogen eigenvectors. This combined INS/DFT approach has been developed extensively over the last decade and has generated considerable success in interpreting vibrational spectra [14,15,25–32]. INS is limited, in a sense, by the resolution of the spectrometers, but this is offset by the wealth of information available in the accessible spectral region [32].

2. Method

2.1. Experimental methods

The β-NTO experiment was performed at the NIST Center for Neutron Research [33] using the Filter Analyzer Neutron Spectrometer (FANS) and the Fermi-Chopper time-of-flight (TOF) instrument. Detailed descriptions of the FANS instrument [34,35] and the TOF instrument [36]
are available. Polycrystalline samples of ca. 1 g were held at 15 K for these experiments. β-NTO was obtained from Picatinny Arsenal and used without further purification. The sample was found to be slightly hygroscopic and attempts were made to dry the sample in a vacuum oven. The INS spectra of β-NTO were normalized for the background contribution by using the DAVE program [37].

2.2. Computational methods

The β-NTO crystal structure used for the solid-state DFT was taken from Ref. [7] and is shown in Fig. 1. The crystallographic data for the unit cell at 100 K is as follows: Space group $P2_1/c$, $a = 9.3129$ Å, $b = 5.4458$ Å, $c = 9.0261$ Å, $\beta = 101.464^\circ$, $Z = 4$. A periodic lattice geometry optimization and vibrational analysis was performed with the DMol3 [38] package on the SGI Origin Array located at the Aeronautical Systems Major Shared Resource Center. The BLYP [38] functional was used for all DMol3 calculations in conjunction with a dnd numerical basis set. ‘Fine’ grid spacing was employed in all calculations.

Inelastic neutron scattering spectra (including overtones, combinations and phonon wings) were calculated from the BLYP/dnd normal mode eigenvectors using the A-Climax program v.5.1.3 [39]. In this program, the normal vibrational coordinates are mass-weighted and the scattering cross sections of the different atoms are taken into account. A Debye–Waller factor is accounted for in all fundamentals, overtone, and combination vibrations, which attenuates the vibrational intensities at higher frequencies. Overtone and combination intensities are calculated with their frequencies being assumed as the sum of the component fundamentals.

3. Results

3.1. Geometry

The β-NTO geometry optimized at the BLYP/dnd level is presented in Table 1. The numbering in Table 1 is consistent with the numbering scheme employed in Fig. 1a. The corresponding experimental X-ray diffraction values of β-NTO are shown at both 100 and 273 K. Previously published geometric parameters of a generic isolated NTO molecule are presented at the MP2/6-311G**, B3LYP/6-311G** and B3LYP/6-311 + G** theory levels. It is interesting to note that all methods produce a very similar geometry. The BLYP/dnd bond lengths differ from the crystal averages with rms deviations of $0.017$ Å (298 K) and $0.014$ Å (100 K). The rms deviations in the MP2/6-311G** bond lengths are $0.018$ Å (298 K) and $0.017$ Å (100 K). In the isolated molecule B3LYP/6-311G** results, the rms deviations increase to $0.019$ Å at both 298 and 100 K. The corresponding values for the larger basis set B3LYP/6-311 + G** calculation are $0.018$ Å (298 K) and $0.019$ Å (100 K). The largest deviations between the BLYP/dnd and generic isolated molecule calculated structures are observed primarily between the bond lengths of atoms involved in the hydrogen-bonding network, such as N9–O11 and C3–O7. We believe that the inclusion of the effect of hydrogen bonding on the structure of the molecules in the solid is an

<table>
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<th>BLYP/dnd</th>
<th>MP2 6-311G**</th>
<th>B3LYP 6-311G**</th>
<th>B3LYP 6-311 + G**</th>
<th>Exp(b) (298 K)</th>
<th>Exp(b) (100 K)</th>
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<td>117.500</td>
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<td>118.460</td>
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<td>125.700</td>
<td>126.200</td>
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</table>

a Values are as reported in Ref. [4].

b Reported in Ref. [7].
important aspect of a proper description of the vibrations of these species. There is some evidence of electron rearrangement in the BLYP/dnd bond lengths, presumably due to hydrogen bonding in the solid-state. The C=O bond is 0.03 Å longer in the solid-state calculations than in the isolated molecule. The BLYP/dnd calculated bond length is in close agreement with the X-ray diffraction results at 100 K [7]. Another possible indication of electron rearrangement is revealed in the calculated bond length of N4–H8. This bond lengthens from 1.009 Å in the isolated molecule MP2/6-311G** calculations to 1.050 Å in the BLYP/dnd solid-state calculations. Unfortunately, we are unable to confirm this experimentally. The N4–H8 distance was not reported in the most recent structural analysis of β-NTO [7] and the positions of the hydrogen atoms were not precisely determined in the 1993 X-ray structural determination by Lee et al. [9]. A neutron diffraction study is currently underway in our laboratory, but will be discussed elsewhere [40].

3.2. Molecular vibrations

The experimental INS spectra and BLYP/dnd theoretical spectrum are shown in Fig. 2. Table 2 presents the mode assignments and fundamental vibrational frequencies of β-NTO. Our mode assignments are in close agreement with those given by Sorescu et al. [4]; however, our assignments reveal the contribution of the hydrogen bonding. The INS frequencies are compared to the BLYP/dnd calculated frequencies, the previously published isolated molecule calculated frequencies, and the IR frequencies of α-NTO. There is good agreement between the BLYP/dnd calculated INS frequencies and the experimental INS frequencies. A comparison of the INS vibrational frequencies of β-NTO to the IR vibrational frequencies reported by Sorescu et al., which were thought to be from α-NTO, indicates significant differences between the vibrational frequencies of the two polymorphs. IR and Raman frequencies of β-NTO are not available at this time, but we hope our report stimulates interest in obtaining them.

The vibrational signals between ~10 and 1400 cm\(^{-1}\) can be compared with experiment on the basis of the BLYP/dnd calculated frequencies and intensities. As shown in Fig. 2, the INS and BLYP/dnd calculations are in good agreement with one another, and both the peak spacings and relative intensities are reproduced. In general, the BLYP/dnd calculations slightly overestimate the vibrational frequencies, a typical result of DFT. The BLYP/dnd calculation quantitatively gives the correct intensities to 1400 cm\(^{-1}\). The rms deviation between experimental and the BLYP/dnd results is 8.43 cm\(^{-1}\). The corresponding rms values for the isolated molecule calculations are, at 58.45 cm\(^{-1}\) (MP2/6-311G**), 61.88 cm\(^{-1}\) (B3LYP/6-311G**) and 61.09 cm\(^{-1}\) (B3LYP/6-311+G**), notably larger than the solid-state BLYP/dnd rms value. The large rms deviations of the isolated molecule calculations reflect the large effect hydrogen bonding has on the vibrational frequencies and indicate the necessity of including these interactions in a quantum chemical calculation.

The experimental INS spectrum in Fig. 2 shows a rather large background contribution from ~530 to 1400 cm\(^{-1}\). It is observed that, while the BLYP/dnd calculations indicate some contribution to the background from a phonon wing, this does not completely account for the large background present. We believe that there was a small fraction of water in the sample, which gives rise to the additional background contribution in this spectral region. This could be confirmed by the appearance of an O–H band between 3200 and 3500 cm\(^{-1}\).
There are six fundamental vibrational modes, namely $v_6$, $v_8$, $v_{12}$, $v_{14}$, $v_{15}$, and $v_{18}$, where frequency deviations of 10 cm$^{-1}$ or greater are observed between the experimental INS and BLYP/dnd frequencies. As can be seen from Table 2, these modes primarily involve large out-of-plane hydrogen atom motion. Based on the crystalline geometry shown in Fig. 1b, these deviations suggest that the hydrogen bond network is not modeled exactly in our solid-state calculations and that the hydrogen bonds are calculated to be longer than the corresponding crystallographic bond lengths. The solid-state calculations, however, represent a substantial improvement over the isolated molecule calculations, which have frequency deviations as large as 55 cm$^{-1}$ for these particular modes, even after scaling by the accepted scaling factor [41].

### 3.3. Phonon modes

The frequencies and assignments of the $\beta$-NTO phonons are provided in Table 3. Four well-defined peaks in the TOF $\beta$-NTO spectrum appear with maxima at $(P_1)\ 37$ cm$^{-1}$, $(P_2)\ 60$ cm$^{-1}$, $(P_3)\ 93$ cm$^{-1}$, and $(P_4)\ 235$ cm$^{-1}$. Fig. 2 shows
The phonon overtone and combination bands observed in the INS spectrum of β-NTO are summarized in Table 3. There are four phonon combination bands which are assigned to \( P_1 + P_2 \) (105 cm\(^{-1}\)), \( P_1 + P_3 \) (121 cm\(^{-1}\)), \( P_2 + P_3 \) (230 cm\(^{-1}\)) and \( P_1 + P_3 \) (272 cm\(^{-1}\)). Four instances of coupling between the phonon modes and the fundamental vibrations are found in the INS spectrum, which are assigned to \( P_2 + v_4 \) (246 cm\(^{-1}\)), \( P_4 + v_1 \) (313 cm\(^{-1}\)), \( P_1 + v_6 \) (375 cm\(^{-1}\)) and \( P_2 + v_1 \) (776 cm\(^{-1}\)). Although combination modes occur between all vibrational modes with a resulting intensity largely dependent on the intensity of the fundamental vibrations, it is interesting to note that the four most intense combination modes that are discussed above all contain a significant degree of N–H vibration. This suggests that these four modes may be considered as ‘doorway modes’ and offers a possible mechanism for bond breaking, which is induced by phonon excitation. Furthermore, an intense combination of a phonon band and fundamental vibration at 776 cm\(^{-1}\) suggests that vibrations up to 800 cm\(^{-1}\) or higher may also play a role in vibrational up-pumping.

### 3.4. Combinations and overtones

The overtone and combination bands of the fundamental vibrations observed in the INS spectra of β-NTO are summarized in Table 4. Combination and overtone vibrations typically display more anharmonic character than fundamental transitions because they have much stronger interactions. Therefore, the accurate representation

<table>
<thead>
<tr>
<th>INS BLYP/dnd</th>
<th>MP2/6-311G**</th>
<th>B3LYP/6-311G***</th>
<th>B3LYP/6-311 + G***</th>
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<tbody>
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<td></td>
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<td>Intensity</td>
<td>Frequency</td>
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<td>210</td>
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<td>( P_1 + P_1 )</td>
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<td>( P_1 + P_5 )</td>
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<td>( P_3 + P_3 )</td>
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<td>( P_2 + P_4 )</td>
<td>321</td>
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<tr>
<td>( 2 \times P_4 )</td>
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<td>378</td>
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<tr>
<td>( P_3 + P_4 )</td>
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All values are reported in cm\(^{-1}\). Intensities are reported in arbitrary units to facilitate comparison.

---

Table 3

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<th>INS BLYP/dnd</th>
<th>Frequency</th>
<th>Intensity</th>
<th>Frequency</th>
<th>Intensity</th>
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<td>( P_2 )</td>
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<td>( P_3 )</td>
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<td>( P_4 )</td>
<td>235</td>
<td>27</td>
<td>241</td>
<td>18</td>
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</table>

Combinations

| \( P_1 + P_2 \) | 105       | 29        | 105       | 20        |
| \( P_1 + P_3 \) | 121       | 58        | 124       | 35        |
| \( P_2 + P_3 \) | 230       | 32        | 231       | 25        |
| \( P_2 + P_4 \) | 246       | 27        | 248       | 16        |
| \( P_1 + P_4 \) | 272       | 19        | 278       | 13        |
| \( P_1 + P_5 \) | 313       | 33        | 308       | 25        |
| \( P_1 + P_6 \) | 375       | 24        | 373       | 18        |
| \( P_2 + P_5 \) | 776       | 78        | 787       | 65        |

All values are reported in cm\(^{-1}\). Intensities are reported in arbitrary units to facilitate comparison.

\( \text{b} \) Values are as reported in Ref. [4].
of these transitions is an important challenge for the harmonic approximation.

One overtone transition \((2\nu_d)\) is observed in the INS spectrum at \(367\ \text{cm}^{-1}\). The BLYP/dnd frequency calculated with the A-Climax program deviates by \(11\ \text{cm}^{-1}\) from the experimental value. This deviation presumably reflects some degree of anharmonicity. The notable intensity difference between the calculated and experimental spectra can be justified by noting the slight difference in intensity between the calculated and experimental fundamental vibration, \(\nu_d\).

In many cases, the calculated vibrational frequencies of the combination transitions are adequately described with the BLYP/dnd calculations. The largest frequency deviations are observed for combination modes that have a significant contribution from hydrogen motion, primarily \(\nu_1\), \(\nu_4\), \(\nu_6\), and \(\nu_{11}\). This is consistent with the X-ray diffraction, which indicates the anharmonic nature of the hydrogen bonds of \(\beta\)-NTO. The isolated molecule frequencies deviate significantly, often up to \(200\ \text{cm}^{-1}\), from the INS experimental frequencies for combinations of these fundamental vibrations. The anharmonic nature of \(\beta\)-NTO is also reflected in the inaccurate calculated intensities of several combination bands.

4. Conclusions

The vibrational spectrum of the energetic material \(\beta\)-NTO was calculated using the BLYP/dnd full periodic representation employed in DMol3 and compared to the INS experimental frequencies. The simulated INS spectrum was constructed using the A-Climax program and included combinations and overtones up to four quanta. The degree of agreement between the theoretical and experimental vibrational spectra demonstrates the advantages of the A-Climax program as a useful method for calculations and analysis of the vibrational spectroscopy of polyatomic molecules.

Our analysis of the INS vibrational spectra verifies that the molecular packing of NTO in its crystalline lattice plays an important role in the general pattern of the vibrational spectrum of the molecule. By comparing the similarities between calculated and measured INS spectral plots, we attempted to establish a one-to-one correspondence between calculated and measured bands, thereby assigning the normal modes. In addition, four phonon modes were assigned. Four instances of intense vibrational coupling between the phonon and internal modes were found. All internal modes that were coupled to the phonon modes were found to have significant N–H character, suggesting a possible mechanism for bond breaking induced by excitation of phonons. Evidence was found of coupling between the phonon and fundamental vibrations up to \(776\ \text{cm}^{-1}\), indicating higher frequency modes might also play a role in vibrational up-pumping.

We found generally good agreement between the calculated geometry of the NTO molecule and the experimentally determined geometry of the \(\beta\)-NTO molecule at \(100\ \text{K}\). We also found good agreement between both the intensities and frequencies of the solid-state BLYP/dnd calculations and the experimental INS values. In general, this work indicates that the quantum chemical description of highly hydrogen-bonded systems requires a proper treatment of the number of degrees of freedom of the lattice, the effects of intermolecular interactions on the structure of the molecule and the effect of intermolecular interactions on the coupling of vibrations of the molecules in the lattice.

Acknowledgements

The NIST Center for Neutron Research is acknowledged for providing neutron beam access on the FANS and TOF instruments. The Wright-Patterson Aeronautical Major Shared Resource Center is thanked for access to the DMol3 program. Picatinny Arsenal is thanked for providing the sample of \(\beta\)-NTO. An NRC Fellowship with the Army Research Laboratory supported J.C. during the course of this research.

References

[33] Certain commercial software and instruments are identified in this paper to foster understanding. Such identification does not imply recommendation or endorsement by the National Institute of Standards and Technology, nor does it imply that the software or equipment identified are necessarily the best available for this purpose.