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Publisher's version / Version de l'éditeur:
https://doi.org/10.1021/ic0203771
Inorganic Chemistry, 41, 26, pp. 7064-7071, 2002-11-26
Structure and Dynamics of ND₃BF₃ in the Solid and Gas Phases: A Combined NMR, Neutron Diffraction, and Ab Initio Study

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Received June 3, 2002

The decrease in D→A bond lengths, previously reported for some Lewis acid/base complexes, in going from the gas to the solid phases is investigated by obtaining an accurate crystal structure of solid ND₃BF₃ by powder neutron diffraction. The B–N internuclear distance is 1.554(3) Å, 0.118 Å shorter than the most recent gas-phase microwave value and 0.121 Å shorter than the single molecule geometry optimized (1.672 Å, CISD/6-311++G-(d,p)) bond length. The crystal structure also shows N–D=F–B hydrogen bonds. The effects of this change in structure and of intermolecular hydrogen-bonding on nuclear magnetic shielding (i.e., chemical shifts) and the nuclear quadrupolar coupling constants (QCC) are investigated by ab initio molecular orbital and density functional theory calculations. These calculations show that the nitrogen ([¹⁵N and ¹⁴N] and boron ([¹¹B and ¹⁰B]) chemical shifts should be rather insensitive to changes in rBN and that the concomitant changes in molecular structure. Calculations on hydrogen-bonded clusters, based on the crystal structure, indicate that H-bonding should also have very little effect on the chemical shifts. On the other hand, the [¹¹B and ¹⁴N] QCCs show large changes because of both effects. An analysis of the ¹⁴N{¹⁹F} line shape in solid ND₃BF₃ yields a ¹¹B QCC of ±0.130 MHz. This is reasonably close an earlier value of ±0.080 MHz and the value of ±0.050 MHz calculated for a [NH₄BF₄]₄ cluster. The gas-phase value is 1.20 MHz. Temperature-dependent deuteron T₁ measurements yield an activation energy for rotation of the ND₃ group in solid ND₃BF₃ of 9.5 ± 0.1 kJ/mol. Simulations of the temperature-dependent T₁ anisotropy gave an Eₜ of 9.5 ± 0.2 kJ/mol and a preexponential factor, A, of 3.0 ± 0.1 × 10¹² s⁻¹. Our calculated value for a [NH₄BF₄]₄ cluster is 16.4 kJ/mol. Both are much higher than the previous value of 3.9 kJ/mol, from solid-state proton T₁ measurements.

Introduction

The charge transfer, Lewis acid–base complex, NH₃BF₃, was first prepared in the early 19th century and has since been synthesized by countless students in introductory inorganic laboratories. An early (1951) X-ray crystal structure provides what could best be described as estimates of rBN (1.60 Å), rBF (1.38 Å), and θNBF (107°). A more recent, and higher quality, structure of the methylene chloride solvated 18-crown-6 ether complex of NH₃BF₃ has been reported. To date no high quality structure, including accurate hydrogen positions, is available for solid NH₃BF₃. Determination of the gas-phase structure of NH₃BF₃ by rotational spectroscopy eluded researchers until the past decade. This was due to the tendency of the vapor of the solid adduct to form NH₂BF₂. It is evident from the X-ray and microwave structures that the BN bond length in NH₃BF₃ is shorter in the solid state. This puts NH₃BF₃ in a class of donor–acceptor complexes in which the D→A bond length decreases significantly in going from the gas to the solid phases. Other examples where this bond length decrease is observed are as follows: NH₃BH₄, N(CH₃)₃BF₃, NH₃-SO₃, N(CH₃)₂SO₃, N(CH₃)₂SO₄, CH₃CNBF₃, and HC-NBF₃. In the last case the decrease is particularly large at 0.835 Å!

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There have been three recent ab initio calculations on NH$_3$-BF$_3$. Fujiiang et al. report an MP2/TZ2P geometry optimization of NH$_3$-BF$_3$ and calculate the BN bond length dependence of the energy, dipole moment, and electric field gradients at N and B.$^{15}$ Jonas and Frenking give $r_{B-N}$ for optimized geometries at various levels, including MP3/6-31(d,p) and QCISD/6-31G(d,p).$^{14}$ Jonas et al. have produced a comprehensive theoretical study of several Lewis acid—base complexes including NH$_3$-BF$_3$. Aside from a geometry optimization of NH$_3$-BF$_3$ at several levels of theory, they showed that $r_{B-N}$ decreases by about 0.05 Å upon dimerization.$^{15}$

The dynamics of the NH$_3$ and BF$_3$ groups in solid NH$_3$-BF$_3$ have been investigated by variable temperature $^1$H and $^{19}$F spin—lattice time ($T_1$) measurements.$^{16}$ This study yields activation energies for rotation of the BF$_3$ and NH$_3$ groups of 25.02 and 3.91 kJ/mol. This latter value seems unusually small considering the possibility of N—H···F—B hydrogen-bonding. Turning to the gas phase, an A,E splitting in the rotational spectrum of about 60 kHz led Legon and Warner to estimate an internal rotational barrier, $V_3$, of approximately 7 kJ/mol.$^5$ This is nearly twice the $E_3$ found in the solid.

NMR parameters are known to be sensitive probes of molecular structure and dynamics. The isotropic $^{11}$B and $^{19}$F chemical shifts of NH$_3$-BF$_3$ have been measured in solution.$^{17}$ In addition, the $^{11}$B nuclear quadrupolar coupling constant has been reported for solid NH$_3$-BF$_3$.$^{18}$ The spectrum was recorded at low field (1.41 T) in CW mode without $^{19}$F decoupling. The rather poor quality spectrum gave an $^{11}$B QCC of 80 ± 8 kHz. This value is over an order of magnitude smaller than that measured in the gas phase (about 1.2 MHz, depending on the torsional state).$^5$

The purpose of this study is to determine the molecular structure of solid NH$_3$-BF$_3$ including accurate hydrogen positions, to determine the activation parameters for ammonia group rotation in solid ND$_3$-BF$_3$ by deuterium $T_1$ and $T_2$ anisotropy measurements, and to redden the boron quadrupolar coupling constant in solid NH$_3$-BF$_3$. We will also employ ab initio molecular orbital and density functional theory (DFT) methods to calculate the barrier to rotation of the NH$_3$ group, the N and B nuclear quadrupolar coupling constants, and the chemical shielding (i.e., shift) parameters. These will be compared to experimental values available in the gas and solid phases.

### Theoretical Background

**Deuterium Line Shapes.** The solid-state deuterium NMR spectral line shape is dominated by the quadrupolar interaction. This is the interaction between the nuclear quadrupole moment, $eQ$, and the largest component of the electric field gradient (egf) tensor at the nucleus, $e_{zz}$. The quadrupolar coupling constant, $\chi$, which is a measure of the strength of the quadrupolar interaction, is given as

$$\chi = \frac{e^2 q Q}{h}$$

The line shape is also influenced by the asymmetry parameter of the efg tensor, $\eta$, which is defined as the difference between the two smaller principal components of the diagonalized efg tensor, $e_{xx}$ and $e_{yy}$, with respect to the largest component, $e_{zz}$,

$$\eta = \frac{e_{xx} - e_{yy}}{e_{zz}}$$

For the molecule studied here, the $\eta$ values are very small (<0.1), and it will be assumed that the efg is axially symmetric along the N−D bond. In that case, the spectrum from a deuterium nucleus located in a static N−D bond consists of a doublet with spacing

$$\Delta \nu = \frac{3e^2 q Q}{4h} (3 \cos^2 \alpha - 1)$$

where $\alpha$ is the angle between the X−D bond and the static magnetic field, $B_0$. In powdered samples, the line shape is determined by the random distribution of the X−D bond orientations with respect to $B_0$, yielding a typical Pake doublet with a characteristic splitting between peaks ($\alpha = 90\degree$ of $3\gamma/4$ and shoulders ($\alpha = 0\degree$) separated by $3\gamma/2$ (see Figure 4). The effects of motion on the deuterium powder spectrum are 2-fold. First the width of the spectral line is narrowed by the motion. In the case of a jumping motion of the X-D bond between three or more equally populated sites, as is expected for ND$_3$ and BD$_3$ groups, $\eta$ is averaged to zero and, if the motion is fast as compared to the quadrupolar interaction (i.e., $>10^6$ s$^{-1}$), the quadrupolar splitting, $\Delta \nu$, is reduced to

$$\Delta \nu_{eff} = \frac{3e^2 q Q}{8} (3 \cos^2 \theta - 1)$$

where $\theta$ is the angle between the X−D bond and the rotational axis. If the motional rate, $k$, is of the order of $P$ ($\sim 10^4$−$10^5$ s$^{-1}$) the spectral line shape is severely distorted because of the large $T_2$ anisotropy associated with the.
dynamic process. The distorted line shapes can readily be simulated for a given combination of $\chi$, $\theta$, and $k$.

**Deuterium Relaxation Times.** The spin–lattice relaxation time, $T_1$, characterizes the return of the bulk magnetization to its equilibrium state after a perturbing pulse. Relaxation is stimulated by time-dependent nuclear spin interactions and is most efficient when these fluctuations occur at the resonance frequency of the nucleus. For deuterons, the primary source of these fluctuating fields is the time-dependent nuclear spin interactions and $\chi$ is stimulated by time-dependent nuclear spin interactions and $\chi$ is characterized by the return of the bulk magnetization to its equilibrium state after a perturbing pulse. Relaxation is stimulated by time-dependent nuclear spin interactions and is most efficient when these fluctuations occur at the resonance frequency of the nucleus.

When the reorientational motion can be described by an exponential correlation function and the temperature dependence of the correlation time follows Arrhenius behavior, the rate of relaxation may be fit to the following equations:

$$\frac{1}{T_1} = K \left( \frac{\tau_c}{1 + \omega_0^2 \tau_c} + \frac{4\tau_c}{1 + 4\omega_0^2 \tau_c^2} \right)$$

$\tau_c = \tau_\omega \exp\left(\frac{E_a}{RT}\right)$

where $\tau_c$ is the correlation time for the motion ($\tau_c \propto k^{-1}$), $\tau_\omega$ is its value at infinite temperature, and $\omega_0$ is the Larmor precession frequency. In addition, $E_a$ is the activation energy for the motion influencing $T_1$, and $K$ depends on the strength of the dominant nuclear spin interaction (in this case $\chi^2$) and the exact nature of the reorientational motion. For jumps of the $X$–$D$ bond between three or more equivalent positions oriented at an angle $\theta$ about the rotational axis, $K$ is given by

$$K = \frac{9\tau_\omega^2 \chi^2}{20} (3 \cos^2 \theta - 1)$$

In this case, if a plot of $\ln(T_1)$ versus inverse temperature is made, one often sees a $V$-shaped curve, with a $T_1$ minimum at $\tau_\omega \omega_0 = 0.62$. There is, in general, much information to be extracted from this $T_1$ plot. The slope of the curve on either side of the $T_1$ minimum is characteristic of the activation energy for the motion influencing $T_1$. The intercept at $1/T = 0$ yields a value for $\tau_\omega$, while the depth of the minimum can be used to determine a value for the effective quadrupolar coupling constant. Note that knowledge of $\tau_\omega$ and $E_a$ allows calculation of the motional rate at any temperature through eq 6.

**Boron-10 Line Shapes.** For the case where $\eta = 0$ the $^{10}$B line shape consists of three pairs of Pake doublets, for the $0 \leftrightarrow \pm 1$, $\pm 1 \leftrightarrow \pm 2$, and $\pm 2 \leftrightarrow \pm 3$ transitions (compare to the deuterium case that has only one, $0 \leftrightarrow \pm 1$, set of transitions, hence one doublet). A simulation of such a set of doublets is shown in Figure 11. The separation between the peaks or horns of the Pake doublets is $\chi/20$, $3\chi/20$, and $5\chi/20$, respectively.

**Experimental Section**

Three different isotopomers were prepared for this study. Initially a sample of $^{11}$ND$_3$BF$_3$ (hereafter referred to as ND$_3$BF$_3$) was used for the deuterium line shape and $T_1$ work. Later a sample of $^{11}$ND$_3$BF$_3$ was required for the neutron diffraction structure determination.

To maximize the signal-to-noise ratio in the $^{10}$B study, a sample of ND$_3$BF$_3$ was prepared. Deuterium was included in this sample to reduce the $^{10}$B–$^{1}$H dipole line broadening in the $^{10}$B spectrum. This allows us to perform a double resonance $^{10}$B($^3$F) experiment rather than the more demanding $^{10}$B($^1$H,$^3$F) experiment.

All three isotopomers were prepared by bubbling ammonia-$d_1$ gas (MSD isotopes) through a solution of $^{11}$BF$_3$ or $^{10}$BF$_3$ in ether (Aldrich). The solid product was precipitated out of solution, filtered, and washed with dry ether. The sample is stable in air, but the deuterium will eventually be replaced by hydrogen when exposed to moisture. The purity of the samples was checked by $^{19}$F and $^{10}$B (or $^{11}$B) spectra of the solid dissolved in D$_2$O. Some samples contained an impurity that was determined to be NH$_4$BF$_3$.

Structural data for ND$_3$BF$_3$ were determined by powder neutron diffraction measurements at 5 K. The powder samples were contained in vanadium tubes of 5 mm i.d. and wall thickness of 0.18 mm, which were filled with helium gas. The samples were rotated about one axis throughout data collection to improve averaging over crystallite orientations. The experiments were done on the C2 Dualspec high-resolution powder diffractometer at the NRU reactor at Chalk River, and data were collected from 5 to 120$^\circ$ with a step size of 0.025$^\circ$. The monochromator was Si(311), and the takeoff angle was $2\theta = 92.7^\circ$. The neutron wavelength of 1.3284 Å was calibrated using a separate Si powder standard. The sample temperature was controlled by a commercially available open cycle design helium cryostat, with separate silicon diodes for the measurement and control of temperature. Structure refinement was carried out using the GSAS Rietveld refinement code.\(^{19}\)

Deuterium spectra, obtained at 30.74 MHz on a Bruker ASX-200, of a 100 mg sample of solid ND$_3$BF$_3$ employed a quadrupolar echo pulse sequence.\(^{20}\) The spectra were obtained at temperature intervals of about 20 K from room temperature (291 K) down to 110 K. Deuterium spin–lattice relaxation times were measured at temperatures down to 110 K using an inversion–recovery quadrupolar echo pulse sequence. For each temperature 16 different variable delays were used. The $T_1$ at each temperature was determined using the TONE program. For all deuterium experiments a $\pi/2$ pulse of 2.6 $\mu$s, a pulse spacing of 40 $\mu$s, and a relaxation delay of at least 5$T_1$ were used. After each temperature change a period of 15 min was allowed for the sample to reach thermal equilibrium. A Eurotherm B-VT2000 was used for temperature regulation, and the sample temperature was checked before each run by lowering a thermocouple down the bore of the magnet into the sample compartment.

Solid-state $^{10}$B spectra were obtained on a Chemagnetics CMX Infinity ($B_0 = 4.7$ T, $v_0(10B) = 21.5$ MHz) NMR spectrometer. Single pulse experiments were used with pulses of 2 $\mu$s (90$^\circ$ pulse was 5.5 $\mu$s). The spectra were acquired with $^{19}$F high power decoupling.

Theoretical calculations were performed with Gaussian 98.\(^{21}\) Nuclear quadrupole coupling constants were obtained from the calculated electric field gradients via the relationship $\chi = -235Qq_{zz}$, where $\chi$ is the QCC in kiloHertz, $Q$ is the nuclear quadrupole moment in millibarns, and $q_{zz}$ is the largest calculated electric field gradient at the nucleus of interest in atomic units. The $Q$ values recently published by Pykkö were used.\(^{22}\) The values used in this study are as follows: $Q(^1H) = 2.86$, $Q(^1B) = 40.59$, $Q(^{10}B) = 84.59$, and $Q(^{11}B) = 20.1$.

\({\text{Vol. 41, No. 26, 2002}}\) Penner et al.
The deuterium line shapes and $T_1$ anisotropies were simulated with the program MXET1.\textsuperscript{23} The $^{19}$B spectrum was simulated with the WSOILDS software.\textsuperscript{24}

**Results and Discussion**

**Neutron Diffraction Structure of ND$_3$BF$_3$.** The analysis of the neutron diffraction pattern was accomplished by starting with the published X-ray structure and refining the structural parameters, including the deuterium positions. The results of the refinement of the structure of solid ND$_3$BF$_3$ are summarized in Tables 1–6. Figure 1 shows the observed and calculated powder patterns. The unit cell, which contains eight molecules, is depicted in Figure 2. The crystal structure indicates that each fluorine atom on one molecule is hydrogen-bonded to one deuterium atom from each of three neighboring molecules, thus forming an infinite network of hydrogen bonds throughout the crystal lattice. This arrangement is shown in Figure 3a. Similarly, each deuteron of a ND$_3$BF$_3$ molecule is hydrogen-bonded to one fluorine from each neighboring molecule. This arrangement is shown in Figure 3B. Table 7 compares our neutron diffraction analysis with the earlier results of Geller et al.\textsuperscript{13} The most important difference is that the neutron diffraction analysis yields a B–N bond that is shorter by 0.046 Å.

**Deuterium NMR Spectra and $T_1$.** The $^2$H line shape of solid ND$_3$BF$_3$ does not change in the temperature region between 300 and 110 K. This spectral line shape (see Figure 4 for an example) is consistent with a ND$_3$ group rapidly rotating at a rate, $k$, greater than $10^8$ s$^{-1}$. If one assumes a rapid interchange between the three N–D deuterons and uses the neutron diffusion structure, eq 4 yields a $\chi(2H)$ of 184 kHz for the nonexchanging deuterons. Values for other ND$_3$ donor–acceptor complexes are $198 \pm 5$ kHz for ND$_3$BH\textsuperscript{25} and $162 \pm 5$ kHz for ND$_3$SO.\textsuperscript{26} The deuterium QCC is a local property and is relatively independent of the acceptor group. On the other hand, it is rather sensitive to N–D····X–A interactions. For example, gas-phase rotational spectra and molecular orbital calculations show that $\chi(2H)$ for isolated NH$_3$ is 290.6 kHz, whereas solid-state NMR experiments yield a $\chi(2H)$ value of 156 kHz.\textsuperscript{27} Similarly, our calculations of $\chi(2H)$ for single molecules of NH$_3$BF$_3$ give values of 252 (B3LYP/6-311++G**) and 255 kHz (MP2/6-311++G**). Evidently there is a significant intermolecular N–H····F–B interaction, as implied by the neutron diffraction structure, which lowers the deuterium QCC. Details of the calculations of hydrogen-bonding effects on $\chi(2H)$ in ND$_3$BF$_3$ will be presented later (vide infra).

A plot of the deuterium $T_1$ as a function of sample temperature is shown in Figure 5. A low-temperature $T_1$ minimum could not be reached with our apparatus. This is consistent with the ND$_3$ rotational rate being in excess of $10^8$ s$^{-1}$ at temperatures above 110 K. The slope of the ln $T_1$ versus 1/T plot gave an activation energy of $9.1 \pm 0.1$ kJ/mol. Since the $T_1$ minimum could not be reached, it was not possible to determine the temperature-dependent rates for ND$_3$ rotation. In this case the alternative is to measure the $T_1$ anisotropy.\textsuperscript{28} For a three-site exchange the anisotropy is given by $\chi^2(3H)$ in the rate of relaxation of the powder line shape is dependent on the exchange rate. The inversion recovery deuterium spectra for one temperature are shown in Figure 6. As can be seen, the shoulders of the spectral line shape relax faster than the peaks. A plot of ln $k$ versus 1/T is shown in Figure 7. A linear regression yields $E_a = 9.5 \pm 0.2$ kJ/mol and a preexponential factor, $A$, of $3.0 \pm 0.1 \times 10^9$ s$^{-1}$ for rotation of the ND$_3$ group. Earlier proton $T_1$ measurements provide $E_a$ and $A$ values of 3.91 kJ/mol and $8.99 \times 10^{10}$ s$^{-1}$, respectively.\textsuperscript{16} Although our results are not consistent with these numbers, there is agreement in $E_a$ between our independent $T_1$ and $T_1$ anisotropy measurements. The preexponential factor, $A$, also known as the frequency factor or attempt frequency, is often identified with the torsional frequency for that motion. The attempt frequency for rotational motion of the ND$_3$ group, $A$, is in good agreement with the torsional frequency, $v_T$, of $5.7 \times 10^2$ s$^{-1}$ obtained from the far-infrared spectrum of solid ND$_3$BF$_3$.\textsuperscript{29}

**Barrier to Internal Rotation.** The gas-phase barrier to internal rotation in NH$_3$BF$_3$ has been estimated to be 7 kJ/mol.\textsuperscript{13} Fluorine-19 and proton $T_1$ measurements yield barriers

<table>
<thead>
<tr>
<th>Table 1</th>
<th>atomic positions</th>
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<td>0.0748(9)</td>
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<td>0.1016(11)</td>
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<tr>
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<th>Table 2</th>
<th>Intramolecular Distances (Å)</th>
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</tr>
<tr>
<td>B11–N15</td>
<td>1.554(5)</td>
</tr>
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account the decrease in the B–N rotation. The internal rotational barrier in isolated NH
anisotropy measurements give a value of 9.5 kJ/mol for NH
is exclusively due to intramolecular interactions between the
for rotation of the BF
3
and NH
3
groups in the solid as 25.0
and 3.91 kJ/mol, respectively, while our deuterium
rotational barrier in isolated NH
3
BF
3
is exclusively due to intramolecular interactions between the
N–H and the B–F bonds. In the solid one must take into
account the decrease in the B–N bond length and intermo-

Table 3

<table>
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<th>Internal Angles (deg)</th>
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<td>F12–B11–F13</td>
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<tr>
<td>F13–B11–F14</td>
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<tr>
<td>B11–N15–D16</td>
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Table 4

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<tr>
<td>D17···F84</td>
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<tr>
<td>D18···F32</td>
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\[ \frac{1}{2} - x, \frac{1}{2} + y, z \]

\[ -\frac{1}{2} + x, y, \frac{1}{2} - z \]

\[ -x, -y, -z \]

Table 5

<table>
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<th>Intramolecular Angles (deg)</th>
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<tr>
<td>N15–D17···F84</td>
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<tr>
<td>N15–D18···F32</td>
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\(^a\) Obtained using PLUTON\(^3\) and does not provide esd values.

Table 6

<table>
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<th>Lattice Parameters</th>
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<tr>
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</tr>
<tr>
<td>U(_{33}), Å(^2)</td>
</tr>
<tr>
<td>U(_{13}), Å(^2)</td>
</tr>
</tbody>
</table>

\(^a\) These values were refined but damped during the analysis.

Figure 1. Neutron diffraction intensity profile and calculated fit for solid
ND
3
BF
3
at 5 K. The upper part of the diagram shows the observed and
calculated profiles, and the lower part of the diagram shows the difference
between observed and calculated profiles.

for rotation of the BF
3
and NH
3
groups in the solid as 25.0
and 3.91 kJ/mol, respectively, while our deuterium
anisotropy measurements give a value of 9.5 kJ/mol for NH
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3
BF
3
is exclusively due to intramolecular interactions between the
N–H and the B–F bonds. In the solid one must take into
account the decrease in the B–N bond length and intermo-

Figure 2. Crystallographic unit cell for solid ND
3
BF
3n

\( V(θ) = 16.4 \, \text{sin}^2(3θ/2) \) (also shown in Figure 9). If the structure
were allowed to relax during the rotation of the NH
3
group, a B3LYP/6-311++G(\(^d,p\)) calculation yields a barrier of 16.4
kJ/mol. The relative energy as a function of rotational angle
is shown in Figure 9. The points nearly follow the equation
\( V(θ) = 16.4 \, \text{sin}^2(3θ/2) \). If a high
barrier is assumed, the potential function $V_3 \sin^2(3\theta/2)$ is replaced by $3V_3 \theta^2/2$, and a harmonic oscillator Schrödinger equation can be solved to give

$$V_3 = \frac{8}{9} I \nu^2$$

where $I$ is the reduced moment of inertia for the rotation of the ND$_3$ group; $I = 3m_r^2 \sin \theta$. The previously measured torsional frequency in solid ND$_3$BF$_3$ gives a $V_3$ value of about 19 kJ/mol. Since the model used to obtain this number assumes a very high barrier to rotation, this number is an overestimation of $V_3$. Either way, our calculated and experimental values and the value estimated from the torsional

Figure 4. Experimental (upper) and simulated (lower) deuterium NMR spectrum of solid ND$_3$BF$_3$. The narrow peak in the center of the experimental spectrum is due to ND$_3$BF$_4$.

Figure 5. Plot of the deuterium ln $T_1$ vs inverse temperature for solid ND$_3$BF$_3$.

Figure 6. Deuterium inversion recovery spectra for solid ND$_3$BF$_3$ at 290 K and simulated spectra for different values of the interpulse spacing.
frequency for the barrier to \( \text{NH}_3 \) rotation in solid \( \text{NH}_3\text{BF}_3 \) are all significantly higher than the earlier solid-state experimental values.\(^{16}\)

**Boron, Nitrogen, and Deuterium Quadrupolar Coupling Constants.** Calculations (B3LYP/6-311++G(d,p)) of \(^{14}\)N and \(^{11}\)B nuclear quadrupolar coupling constants were performed for boron–nitrogen bond lengths ranging from 1.4 to 1.8 Å, with optimization of all other structural parameters. The results are shown in Figure 10. The \(^{14}\)N QCC appears to be rather insensitive to changes in \( r_{BN} \), and no clear trend is discernible. On the other hand, the \(^{11}\)B QCC changes significantly and, in fact, passes through zero and changes sign at about 1.5 Å. This difference can be easily explained electrostatically. Isolated BF\(_3\) is planar and has a large \(^{11}\)B QCC. The calculated and experimental values are 2.53 (B3LYP/6-311++G**) and 2.64 MHz, respectively.\(^{31}\)

As a B–N bond is formed, the arrangement around the boron becomes tetrahedral with the fourth atom being the nitrogen. Since fluorine and nitrogen are rather electronegative atoms, there will be some arrangement where the electric field gradients along the three B–F bonds and along the B–N bond will cancel. For B–F bond lengths of about 1.4 Å this occurs when the B–N bond length is about 1.5 Å. In contrast, the geometry of the \( \text{NH}_3 \) moiety shows relatively little change as the B–N bond is being formed. This together with the lower electron density around the boron results in the \(^{14}\)N QCC being rather insensitive to \( r_{BN} \). Since the electric field gradient at the nucleus, and hence the QCC, is a local molecular electronic property, the presence of the fluorine atoms on the boron has very little effect on the \(^{14}\)N QCC. Now that we have determined the crystal structure of \( \text{NH}_3\text{BF}_3 \), it is possible to calculate the \( \chi^{(14)}(\text{N}) \) and \( \chi^{(11)}(\text{B}) \) for gas-phase and solid \( \text{NH}_3\text{BF}_3 \). These values are presented in Table 8 together with the previously reported gas-phase microwave and solid-state NMR values. The quality of the spectrum previously presented by Olliges et al. is rather poor.\(^{18}\) This is primarily due to strong \(^{11}\)B–\(^{19}\)F and \(^{11}\)B–\(^{1}\)H dipolar couplings exacerbated by the small \(^{11}\)B QCC (relative to the dipolar couplings). Our attempts to eliminate these dipolar couplings by \(^{19}\)F decoupling and replacement of the protons by deuteron resulted in very modest improvements. An alternative is to examine the \(^{10}\)B spectrum. A \(^{10}\)B spectrum of solid \( \text{ND}_3\text{BF}_3 \), with \(^{19}\)F decoupling, is shown in Figure 11. The most prominent feature is the so-called Pake doublet

of the central 0→1 transition. A simulation based on a fit of this feature is also shown. The satellite 1→2 and 2→3 transitions do not have the intensity in the experimental spectrum as they do in the simulation. We cannot provide an explanation for this.

As shown in Table 8, our geometry-optimized structure for NH₃BF₃ gave ¹⁴N QCC that is in excellent agreement with the observed gas-phase value. Both of these values are about an order of magnitude larger than the two experimental values for the solid state. If a calculation is done on a single molecule with the neutron diffraction structure, the QCC drops to 257 kHz, much closer to the NMR values. If intermolecular N–H⋯F–B hydrogen-bonding is taken into account by calculating the QCC for the tetramer shown in Figure 3A, the value drops further by a factor of 5–50 kHz, in good agreement with experimental values.

Unfortunately there is no experimental value for the ¹⁴N QCC in solid NH₃BF₃. Our calculated single molecule value agrees well with the gas-phase microwave number. Using the solid-state molecular structure reduces the QCC slightly, putting it in better agreement with the microwave value; this is fortuitous. A calculation on the tetramer depicted in Figure 3B causes a significant drop in the ¹⁴N QCC. Therefore, one can predict that, because of hydrogen-bonding effects, the solid-state ¹⁴N QCC should be at least 1 MHz smaller than the microwave value.

Going from the gas-phase structure to that in the solid and on to tetramer B appears to have very little effect on the deuterium QCC. The result of the last calculation is surprising considering that the deuterons should be directly involved in intermolecular hydrogen-bonding. It has been shown experimentally that hydrogen-bonding reduces the QCC. As pointed out in a previous section of this discussion, the deuterium QCC for ammonia drops from 290.6 kHz in the gas phase to 156 kHz in the solid state.

**Chemical Shielding and Chemical Shifts.** Calculations (B3LYP/6-311++G(d,p)) of the components of the nuclear magnetic shielding tensor (δ and σ for boron and nitrogen) as a function of r_NB in Figure 12. The variation of shielding with bond length is rather small, being less than 8 ppm in all cases. In addition, the anisotropy in the shielding, Δσ = σ∥ − σ⊥, is also small. The isotropic shielding is given as σ_iso = 1/3(σ∥ + 2σ⊥). The nitrogen shielding values can be converted to the chemical shift scale with respect to liquid CH₃NO₂ according to the equation δ = −135.8 − σ (32). For a single molecule using the solid-state structure a B3LYP/6-311++G(d,p) yields a δ_iso of −353.0 ppm. A calculation on the tetramer (Figure 3B), where each NH₃ is surrounded by three hydrogen-bonded NH₃BF₃ molecules, gave a value of −353.6 ppm. As can be seen, the effect of N–H⋯F–B hydrogen-bonding on the ¹⁴N chemical shift is negligible. The components of the chemical shift tensor, δ∥ and δ⊥, have not been experimentally determined. Unfortunately, there is no scale available to convert boron nuclear magnetic shielding to chemical shifts.

**Conclusions**

Our calculations and measurements, together with earlier measurements, show that the nuclear quadrupolar coupling constant can be a very sensitive probe of electronic, hence molecular, structure. We have successfully solved the neutron diffraction structure for solid NH₃BF₃. The crystal structure shows two important features: a B–N bond length that is much shorter than that in the gas phase and N–H⋯F–B hydrogen bonds. This structure, together with that from the gas phase, proved very useful for setting up the ab initio calculations. These calculations clearly demonstrated that the changes in the boron and nitrogen quadrupolar coupling constants observed in going from the gas to solid phases is a result of two effects: the large change in the B–N bond length and the N–H⋯F–B hydrogen-bonding. Calculations employing the gas and solid-state structures also show that the higher barrier to rotation of the NH₃ moiety in the solid is due to these factors. The calculated gas-phase deuterium QCC is much larger than that measured in the solid. Including the abovementioned, effects in the calculations do not bring the two into closer agreement. As yet we have no explanation for this.

**Acknowledgment**. We thank the Natural Sciences and Engineering Research Council (NSERC) of Canada for financial support. The efforts of Guy Bernard (University of Alberta) and Chris Kirby (University of Western Ontario) toward our attempts to acquire ¹⁰B spectra are greatly appreciated.