Multidimensional hydrogen tunneling dynamics in the ground vibrational state of the ammonia dimer

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(Received 20 March 1992; accepted 27 April 1992)

We have measured and assigned more than 800 new far-infrared absorption lines and 12 new microwave absorption lines of the ammonia dimer. Our data are analyzed in combination with all previously measured far-infrared and microwave spectra for this cluster. The vibration-rotation-tunneling (VRT) states of the ammonia dimer connected by electricdipole-allowed transitions are separated into three groups that correspond to different combinations of monomer rotational states: A + A states (states formed from the combination of two ammonia monomers in A states), A + E states, and E + E states. We present complete experimentally determined energy-level diagrams for the $K_a=0$ and $K_a=1$ levels of each group in the ground vibrational state of this complex. From these, we deduce that the appropriate molecular symmetry group for the ammonia dimer is G_{144} . This, in turn, implies that three kinds of tunneling motions are feasible for the ammonia dimer: interchange of the "donor" and "acceptor" roles of the monomers, internal rotation of the monomers about their C_3 symmetry axes, and quite unexpectedly, "umbrella" inversion tunneling. In the $K_a = 0 A + E$ and E + E states, the measured umbrella inversion tunneling splittings range from 1.1 to 3.3 GHz. In $K_a = 1$, these inversion splittings between two sets of E + E states are 48 and 9 MHz, while all others are completely quenched. Another surprise, in light of previous analyses of tunneling in the ammonia dimer, is our discovery that the interchange tunneling splittings are large. In the A + A and E + E states, they are 16.1 and 19.3 cm⁻¹, respectively. In the A + E states, the measured 20.5 cm⁻¹ splitting can result from a difference in "donor" and "acceptor" internal rotation frequencies that is increased by interchange tunneling. We rule out the possibility that the upper state of the observed far-infrared subbands is the very-low-frequency out-of-plane intermolecular vibration predicted in several theoretical studies [C. E. Dykstra and L. Andrews, J. Chem. Phys. 92, 6043 (1990); M. J. Frisch, J. E. Del Bene, J. S. Binkley, and H. F. Schaefer III, ibid. 84, 2279 (1986)]. In their structure determination, Nelson et al. assumed that monomer umbrella inversion tunneling was completely quenched and that "donor-acceptor" interchange tunneling was nearly quenched in the ammonia dimer [D. D. Nelson, G. T. Fraser, and W. Klemperer, J. Chem. Phys. 83, 6201 (1985); D. D. Nelson, W. Klemperer, G. T. Fraser, F. J. Lovas, and R. D. Suenram, *ibid.* 87, 6364 (1987)]. Our experimental results, considered together with the results of sixdimensional calculations of the VRT dynamics presented by van Bladel et al. in the accompanying paper [J. Chem. Phys. 97, 4750 (1992)], make it unlikely that the structure proposed by Nelson et al. for the ammonia dimer is the equilibrium structure.

I. INTRODUCTION

Over the past decade, a considerable variety of hydrogen-bonded dimers has been investigated by high-resolution spectroscopic methods¹⁻³ and in parallel by high-level *ab initio* techniques.⁴ The goal of this work has

been (1) to establish a predictive model for the dimer equilibrium structures and ultimately for the internal dynamics of small hydrogen-bonded clusters, and (2) to connect this detailed molecular knowledge with the behavior of the constituents in condensed phases. For the most part, good agreement has been obtained between theory and experiment for these prototypical systems. The most well-known exception is the singular case of the ammonia dimer.

For 23 years, since the matrix studies of Pimentel, Bulanin, and van Thiel,⁵ chemists had generally assumed that the minimum of the ammonia dimer intermolecular potential-energy surface corresponded to a linearly hydrogen-bonded geometry,⁶ similar to that determined for the water dimer.⁷ Figure 1, which shows a set of intermo-

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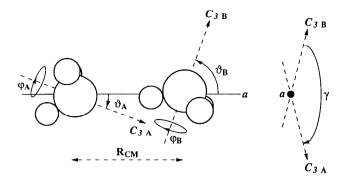


FIG. 1. The six intermolecular coordinates of the ammonia dimer. For clarity, we show the coordinate system used in the following paper (Ref. 22). Most important in our discussion are the angles θ_A and θ_B between the ammonia monomer symmetry axes and the **a** inertial axis of the dimer. These angles define the roles of the inequivalent ammonia monomers as "acceptor" and "donor." Donor-acceptor interchange tunneling, a large-amplitude motion which reverses these roles is shown to be a very facile motion in the ammonia dimer. The angles ϕ_A and ϕ_B describe internal rotations of the monomers about their symmetry axes, labeled C_{3A} and C_{3B} . Internal rotation also occurs readily in the ammonia dimer. The dihedral angle γ is the angle between the two planes that each contain one monomer symmetry axis and the **a** inertial axis of the complex. In this paper, we assume that the ammonia dimer contains a plane of symmetry; that is, we assume that $\gamma = 180^\circ$ (or 0°).

lecular coordinates for the ammonia dimer, illustrates one possible hydrogen-bonded configuration. However, Nelson, Fraser, and Klemperer's analysis of their molecularbeam electric resonance (MBER) rotational spectra^{8,9} indicated that the vibrationally averaged structure of the ammonia dimer was not a hydrogen-bonded one. Rather, they argued that while the complex was clearly asymmetric, the equilibrium structure was closer to being cyclic and centrosymmetric than linearly hydrogen bonded. In the coordinate system of Fig. 1, this proposed structure has $\theta_A = 48.6^\circ$ and $\theta_B = 115.5^\circ$, while ϕ_A , ϕ_B , and γ are not determined. This unexpected result was determined in a straightforward way from the observed $J=1 \leftarrow 0$ and $J=2 \leftarrow 1$ pure rotational transitions in two distinct "vibrational" states of the ammonia dimer. The measured ${}^{14}N$ quadrupole coupling constants eqQ_{aa} for each ammonia monomer, together with the μ_a dipole moment component of the complex, were inverted to determine the two angles θ_A and θ_B between the ammonia symmetry axes and the **a** inertial axis of the dimer.

Subsequently, Nelson and Klemperer¹⁰ presented an elegant group-theoretical description of some of the possible large-amplitude hydrogen tunneling motions that might occur within the ammonia dimer. We will discuss the group-theoretical analysis of Nelson and Klemperer in more detail later in this paper, since their analysis provides a starting point for our own. Nelson and Klemperer considered the possible effects of a "donor–acceptor" interchange tunneling motion, similar to that found in the $(HF)_{2}$,^{11,12} $(HCI)_{2}$,¹³ and $(H_2O)_{2}$,^{14,15} along with the possible effects of the internal rotation of the monomers about their symmetry axes on the rovibrational states of the ammonia dimer. They assumed that the umbrella inversion of the monomers would be totally quenched by anisotropy in

the ammonia dimer potential. Under these assumptions, two sets of pure rotational transitions are predicted to occur within the lowest $K_a=0$ manifold of the ammonia dimer. The two tunneling states that give rise to these transitions are shown to be separated by the difference in internal rotation frequencies of the "donor" and "acceptor," and in second order, their separation can be increased by interchange tunneling. Applying their group-theoretical model to the MBER data, Nelson et al.⁹ argued that the donor-acceptor interchange tunneling frequency was very small in the ammonia dimer because the two tunneling states accessed in their spectra appeared to lie very close in energy. In addition, since they measured different ¹⁴N quadrupole coupling constants for each monomer, they deduced that the donor-acceptor interchange tunneling had no pronounced averaging effect on the angles derived, in the usual way, from these measured ¹⁴N guadrupole coupling constants. Furthermore, careful isotopic substitution experiments revealed that similarly determined structures for a variety of ammonia dimer isotopomers were nearly identical. Nelson et al. inferred from these results that the equilibrium structure of the ammonia dimer was not very different from the vibrationally averaged structure that they determined. It is this claim that defines the apparent discrepancy between theory and experiment that has existed since the appearance of the MBER work.

Several extensive ab initio theoretical investigations¹⁶⁻²⁰ have followed the work of Klemperer group. In these studies, vibrational averaging, both from the tunneling motions, and from the floppy intermolecular vibrations of the ammonia dimer complex, is cited as the cause of the difference between the predicted equilibrium structure and the experimental results. Indeed, the intermolecular potential-energy surface of the ammonia dimer is generally found to be rather flat in some angular coordinates. Both Frisch et al.¹⁶ and Hassett, Marsden, and Smith²⁰ found a linearly hydrogen-bonded equilibrium structure, and a cyclic, centrosymmetric transition state for the interchange motion between two such potential minima that was only 0.1-0.2 kcal/mol higher in energy. Both of these groups also found that the harmonic zero-point energy of the intermolecular vibration in the interchange tunneling coordinate was approximately equal to the interchange barrier height.

The first high-resolution experiment to probe the other types of ammonia dimer tunneling states, which do not support pure rotational spectra, was performed by Havenith *et al.*,²¹ who discovered ammonia dimer spectra between 21 and 28 cm⁻¹, using tunable far-infrared laser spectroscopy. Although this work dramatically increased the number of known ammonia dimer vibration-rotationtunneling (VRT) states from 2 to 12, none of the farinfrared (FIR) transitions shared common upper or lower states. Thus, relatively little information on the groundstate tunneling energetics or dynamics could be extracted from this subset of the FIR spectrum. The results of Havenith *et al.* were interpreted as support for the conclusions of Nelson *et al.*, since no inconsistencies with the microwave work were revealed at that time.

In this paper, we present a far more complete experimental characterization of the ground vibrational state of the ammonia dimer than has been possible before. From our extensive new data set, we are able to show that the appropriate molecular symmetry group for the ammonia dimer complex is G_{144} . Thus, the donor-acceptor interchange tunneling, internal rotation of the NH₃ monomers about their C_3 symmetry axes, and monomer "umbrella" inversion tunneling are all feasible. We have precisely measured these interchange and inversion tunneling frequencies. Furthermore, we find that the two VRT states sampled by the pure rotational spectra are separated by the inversion tunneling frequency, not the interchange tunneling frequency. Indeed, the assumptions of Nelson et al.⁸⁻¹⁰ that umbrella inversion tunneling does not occur and that the interchange tunneling frequency is therefore very small are clearly invalid. Quite the opposite tunneling scheme is appropriate for this complex: we find that the interchange tunneling frequency is likely to be the largest of all hydrogen tunneling frequencies in the ammonia dimer.

The first explicit calculation of the VRT states on one of the *ab initio* ammonia dimer surfaces¹⁹ is described by van Bladel et al.²² in the accompanying paper. The qualitative picture obtained from the wave functions and relative energies thus computed has led these authors to anticipate the results found experimentally in our own work. More specifically, based on their calculations, they suggest assignments of the previously observed subbands near 483 and 614 GHz which are the same as the assignments that we have deduced by careful consideration of our data alone. Although it was not included in their calculations, van Bladel et al. also suggest that the ammonia monomer umbrella inversion tunneling motion may be occurring in the ammonia dimer complex. This result is unambiguously proven in our own analysis presented below. van Bladel et al. confirm the hypothesis of Nelson et al. that the wave functions of tunneling states that can be sampled by pure rotational transitions are to some extent "localized" with respect to interchange tunneling, and that one can therefore expect to measure two different ¹⁴N quadrupole coupling constants for the two different ammonia monomers in such states. However, they also show that the assumption of Nelson, Fraser, and Klemperer⁸ that the quadrupole coupling constants, measures of $\langle P_2(\cos \theta_A) \rangle$ and $\langle P_2(\cos \theta_B) \rangle$, can be used in conjunction with the dipole moment, a measure of $\langle P_1(\cos \theta_A) \rangle$ and $\langle P_1(\cos \theta_B) \rangle$, to deduce structural information may not be appropriate.

We emphasize that, even though some of our results and some of the results of van Bladel *et al.*²² are strikingly similar, each group performed its own analysis and drew corresponding logical conclusions before learning of the other's results. This agreement constitutes additional strong support for the independent analyses presented in both papers.

Here, we present three separate spectroscopically determined energy-level diagrams for the $K_a=0$ and $K_a=1$ levels of the ammonia dimer, corresponding to three different kinds of internal rotor states. These incorporate our new data set, the spectra previously reported by Havenith et al.,²¹ and the spectra reported by Zwart.²³ First, we describe our rotational analysis of these data and the observed Coriolis interactions. Then we discuss in more detail the physical and group-theoretical nature of the tunneling states of the ammonia dimer. Finally, we present our tunneling and vibrational assignment of the observed far-infrared and microwave transitions.

II. EXPERIMENT

The Berkeley tunable far-infrared laser spectrometers have been described previously.²⁴ Since the work of Havenith et al.²¹ on the $(NH_3)_2$ spectrum, we have added the following improvements: (1) we have optimized production of $(NH_3)_2$ by varying the concentration of ammonia in the argon carrier gas using mass flow controllers, (2) we have increased the effective sample path length of the tunable far-infrared laser sidebands (fixed-frequency farinfrared laser radiation mixed with tunable microwave radiation) through the planar jet expansion by multipassing²⁵ (usually 10 times) the sidebands in front of a longer (10 cm) slit nozzle,²⁶ (3) we have increased the far-infrared transmission to the detector by purging the entire beam path with dry nitrogen, and (4) we have increased our "routine" (but not "ultimate") detection sensitivity by subtracting the base-line variation due to microwave transmission resonances, and also the resonances of the sidebands inside the far-infrared laser cavity, from every scan. We have found it fruitful to rescan much of the region originally covered by Havenith et al.²¹ The frequencies and lasing mediums (given in parentheses) of the fixed-frequency far-infrared lasers used in the present study 527.9260 GHz (DCOOD), 584.3882 GHz were (HCOOH), 639.1846 GHz (CH₃OH), 692.9513 GHz (HCOOH), 716.1568 GHz (HCOOH), 761.6083 GHz (HCOOH), 768.8820 GHz (DCOOD), 787.7555 GHz (DCOOD), 850.4118 GHz (CH₃OD), 939.4940 GHz (CH₃OD), 980.5916 GHz (CH₃OD), 1016.8972 GHz (CH₃OD), 1042.1504 GHz (CH₂F₂), 1101.1594 GHz (CH₂DOH), 1110.3199 GHz (CH₂F₂), 1145.4302 GHz (CH₂F₂), and 1193.7273 GHz (CH₃OH).²⁷ The 584–768 GHz lasers were scanned at the highest sensitivity.

Our group has measured a total of nearly 4000 farinfrared absorption lines due to Ar–NH₃, $(NH_3)_2$, and other as-yet unanalyzed Ar_n– $(NH_3)_m$ clusters. Over 950 of these are assigned to $(NH_3)_2$. The signal-to-noise ratio observed on the strongest $(NH_3)_2$ absorption approaches 10 000:1, while the weakest assigned lines have a signal-tonoise ratio of only 3:1. (For comparison, Havenith *et al.*²¹ report a signal-to-noise ratio of 100:1 for the strongest lines they were able to assign to the ammonia dimer.)

The Caltech microwave direct absorption spectrometer has also been described previously.²⁸ Continuous searches were conducted in the regions 29.000–33.140 GHz, 40.800–57.735 GHz, and 60.900–62.286 GHz. Typical scanning conditions allowed observation of the $(NH_3)_2$ pure rotational spectrum, the " G_a " and " G_β " states discovered by the Klemperer group,^{8,9} at a signal-to-noise ratio of several hundred to one. Most of the microwave lines that we observed were part of the well-studied microwave spectrum of the *E* states of Ar-NH₃,²⁹ the pure rotational progressions of (NH₃)₂,^{8,9} as well as a few high-*J* lines in the pure inversion spectrum of the ammonia monomer.³⁰ We have measured 20 microwave absorption lines due to other species or series, and assign 12 of these to a new rotationtunneling subband of the ammonia dimer, as described later in this paper. The measured intensities of these new lines were about 2 orders of magnitude less that the observed intensities of the $K_a=0$ pure rotational transitions.

III. ANALYSIS

A. Rotational analysis

A stick spectrum indicating the positions and estimated relative intensities of all the far-infrared absorption lines currently assigned to the ammonia dimer is presented in Fig. 2(a). (The intensities shown are calculated for 5 K, with adjustments made for obvious intensity perturbations.) Figure 2(b) is a chart indicating the new observed microwave transitions. (The symmetry assignments indicated in this chart are discussed later in this paper.) The relative energies of the vibration-rotation-tunneling (VRT) states are diagrammed in Figs. 3(a), 3(b), and 3(c), and the subband origins of the electric dipole transitions that connect the VRT states shown in these figures are listed in Table II. [The complete list of all the observed transitions (Table I) can be obtained from PAPS.³¹] These states are separated into three groups because of internal rotation of the ammonia monomers within the complex. For clarity, we label these groups by the symmetry types of the constituent ammonia monomer states: A + A, A + E, and E+E. The relative energies of all states within each group are now completely determined for $K_a=0$ and $K_a = 1$, and all of the transitions between states in the same group are fit simultaneously. The rotational parameters determined in each fit are listed in Table III and the lowest resulting calculated VRT energy levels (up to J=5) are listed in Table IV. (The frequencies of all the allowed low-J transitions can easily be calculated from the data in Tables IV to within 1 MHz.) One purpose of performing these fits is to quantitatively establish that the observed transitions have the common upper and lower states indicated in the energy-level diagrams. The simultaneous fits are also necessary to treat the strong Coriolis interations between certain $K_a = 0$ levels and nearby $K_a = 1$ levels with the same symmetry. Strong Coriolis perturbations occur in the A+E states about 600 GHz above the lowest A+Estate and even stronger Coriolis perturbations are present in the lowest E+E states. The VRT subbands which involve these affected states cannot be fit without this full Coriolis analysis.

For clarity and convenience, we have arbitrarily numbered the vibration-tunneling (VT) manifolds in each of the three groups of states. This numbering is indicated in the energy-level diagrams and identifies each transition in the line lists. $K_a \neq 0$ manifolds have two numbers, and the smaller number labels lower-energy component of each K-

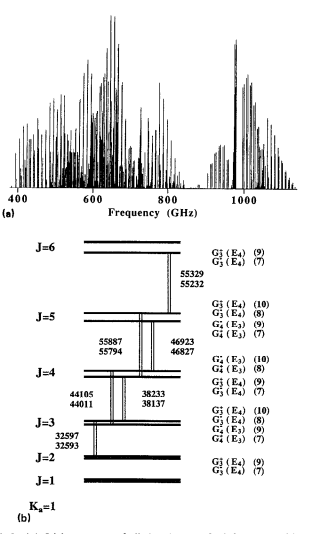
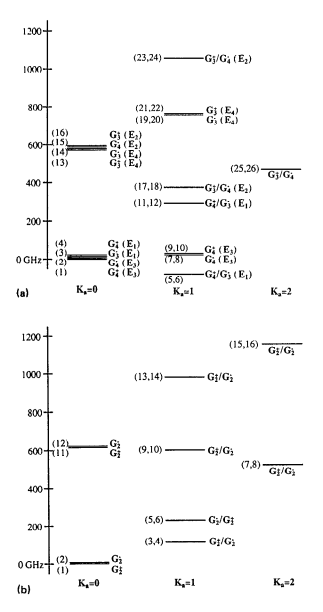


FIG. 2. (a) Stick spectrum of all the observed far-infrared transitions that have been assigned to the ammonia dimer between 13 and 38 cm⁻¹. The transitions assigned by Havenith (Ref. 21) and by Zwart (Ref. 23) are included. These lines represent about one-fourth of all the absorptions we have measured in the 16-41 cm⁻¹ frequency region in Ar/NH₃ expansions. Since these transitions were recorded over the course of several years under varying experimental conditions, this figure shows intensities calculated for a 5 K supersonic expansion, rather than experimentally observed signal strengths. In certain $|\Delta K_a| = 1$ subbands that involve strongly perturbed $K_a=0$ or $K_a=1$ manifolds, the observed R (or P) branch transitions are apparently an order of magnitude or more stronger than the weaker P (or R) branch transitions. We have scaled the calculated intensities of these branches to approximately preserve this unexplained effect. (b) Rotation-tunneling energy-level diagram for two of the $K_a = 1 E + E$ manifolds of the ammonia dimer. The 12 new microwave transitions are indicated. The small 48 MHz splittings are the nearly quenched ammonia monomer umbrella inversion tunneling frequency. The asymmetry doubling, which is larger than the tunneling splitting, is especially large due to Coriolis coupling of the upper components of the asymmetry doublets to nearby $K_a=0$ states. The numbers (7), (8), (9), and (10) are used for identification of these states in the text. The symmetry assignments are also discussed in the text.

type doublet. For each VT level n, the corresponding endover-end rotational states are fit to the phenomenological expressions:

$$E = E_n + B_n J(J+1) - D_n [J(J+1)]^2 + H_n [J(J+1)]^3 + \cdots,$$
(1)



$$E^{\pm} = E_n + B_n J(J+1) - D_n [J(J+1)]^2 + H_n [J(J+1)]^3 + \dots \pm [qB_n J(J+1) + qD_n [J(J+1)]^2 + \dots].$$
(2)

We use E_n to refer to the J=0 rotational state of each VT level n. Since we treat levels with different K_a values as separate "vibrations," E_n for $K_a \neq 0$ levels is the energy of a fictitious rotational state. We use expression (1) for all $K_a=0$ manifolds and we use expression (2) for the $K_a\neq 0$ levels that appear to have a true "asymmetry" doubling in the absence of perturbations. Some of the $K_a = 1$ states are split only by Coriolis interaction with nearby $K_a = 0$ states. When this effect is strong, we use expression (1) for both the $K_a = 0$ and $K_a = 1$ manifolds and determine the Coriolis coupling constant as described below. The Coriolis interaction pushes the $K_a=0$ level and one of the initially degenerate $K_a = 1$ levels apart. When this effect is weak, the Coriolis model still fits the data very well; however, the values of B and the Coriolis coupling constants are totally correlated and also not very well determined. In these weakly interacting cases we choose to present fits in which

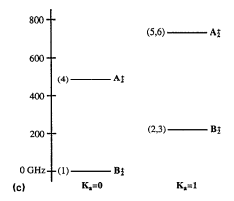


FIG. 3. (a) Complete energy-level diagram for the E + E tunneling states of the ammonia dimer. The observed tunneling levels (horizontal lines) are labeled by the arbitrary numbers (n) used for clarity in the text and are positioned according to the relative energies E_n determined in our fit. Table II(a) lists all of the observed VRT subbands that connect these E+E states. The indicated tunneling symmetries in G_{144} are taken to be the symmetries of the even rotational states. The symmetries of the G_{36} states that were split by the monomer inversion to give the G_{144} states are shown in parentheses. Table IV(a) tabulates the assigned symmetries and relative energies of the lowest few rotational levels in each of the observed E+E tunneling states. (b) Complete energy-level diagram for the A+Etunneling states of the ammonia dimer. The observed tunneling levels (horizontal lines) are labeled by the arbitrary numbers (n) used for clarity in the text and are positioned according to the relative energies E_n determined in our fit. Table II(b) lists all of the observed VRT subbands that connect these A+E states. The indicated tunneling symmetries in G_{144} are taken to be the symmetries of the even rotational states. Table IV(b) tabulates the assigned symmetries and relative energies of the lowest few rotational levels in each of the observed A + E tunneling states. (c) Complete energy-level diagram for the A+A tunneling states of the ammonia dimer. The observed tunneling levels (horizontal lines) are labeled by the arbitrary numbers (n) used for clarity in the text and are positioned according to the relative energies E_n determined in our fit. Table II(c) lists all of the observed VRT subbands that connect these A + Astates. The indicated tunneling symmetries in G_{144} are taken to be the symmetries of the even rotational states. Table IV(c) tabulates the assigned symmetries and relative energies of the lowest few rotational levels in each of the observed A + A tunneling states.

we use expression (1) for the $K_a=0$ manifold and expression (2) for the $K_a=1$ manifold with no Coriolis coupling included. Still other $K_a=1$ states are split both by Coriolis and asymmetry doubling effects.

We find that the rotational states within some VT manifolds cannot be fit well using only B and a small value of D (less than 0.1 MHz) in expression (1) or (2). Usually these perturbed manifolds come in pairs of one $K_a=0$ VT level near one $K_a = 1$ VT level, or one $K_a = 1$ VT level near one $K_a=2$ VT level. The two levels in each pair have second- and third-order distortion constants with similar magnitudes but opposite signs. We model the interaction between such pairs of manifolds as Coriolis coupling. In such cases, we first check that each pair of interacting levels can be connected by two successive electric-dipoleallowed transitions. This assures that the states in question have the same symmetry. We then fit off-diagonal matrix elements of the form $c[J(J+1)]^{1/2}$ between the appropriate unperturbed VRT levels of the same J and the same symmetry in $K_a=0$ and $K_a=1$. Matrix elements appropriate for coupling $K_a = 1$ states to $K_a = 2$ states are

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			Number	
Tunneling symmetry	$K'_a \leftarrow K'_a$	$(n') \leftarrow (n'')$	of lines	$E_{n'}-E_{n''}$
		(a)		
		(4)		
$G_4^-(E_3) \leftarrow G_4^+(E_3)$	1 ← 1	$(7,8) \leftarrow (9,10)$	12	48.2 MHz
$G_3^+/G_4^-(E_2) \leftarrow G_4^+/G_3^-(E_1)$	1 ← 1	(17,18) ← (5,6)	26	453.6 GHz ^a
$G_3^+(E_4) \leftarrow G_4^+(E_3)$	0 ← 1	(13) ← (9,10)	19	548.7 GHz
$G_3^-(E_4) \leftarrow G_4^-(E_3)$	0 ← 1	(14) ← (7,8)	18	551.0 GHz
$G_3^+(E_2) \leftarrow G_4^+(E_3)$	0⊷1	$(16) \leftarrow (9,10)$	15	554.7 GHz
$G_3^+(E_4) \leftarrow G_4^+(E_1)$	0⊷0	(13) ← (4)	14	575.5 GHz
$G_3^+(E_4) \leftarrow G_4^+(E_3)$	0→0	$(13) \leftarrow (1)$	23	577.9 GHz
$G_3^-(E_4) \leftarrow G_4^-(E_3)$	0⊷0	(14) ← (2)	21	579.0 GHz
$G_4^-(E_2) \leftarrow G_3^-(E_1)$	0⊷0	$(15) \leftarrow (3)$	19	580.3 GHz
$G_3^+(E_2) \leftarrow G_4^+(E_1)$	0⊷0	$(16) \leftarrow (4)$	23	581.4 GHz
$G_3^+(E_2) \leftarrow G_4^+(E_3)$	0 ← 0	$(16) \leftarrow (1)$	17	583.8 GHz
$G_3^+(E_4) \leftarrow G_4^+/G_3^-(E_1)$	0←1	$(13) \leftarrow (5,6)$	23	655.5 GHz
$G_4^-(E_2) - G_4^+/G_3^-(E_1)$	0 ← 1	$(15) \leftarrow (5,6)$	25	659.2 GHz
$G_3^+(E_2) - G_4^+/G_3^-(E_1)$	0 ← 1	$(16) \leftarrow (5,6)$	21	661.4 GHz
$G_3^+(E_4) \leftarrow G_4^+(E_3)$	1 ← 1	$(21,22) \leftarrow (9,10)$	53	730.78 GHz ^b
$G_3^-(E_4) \leftarrow G_4^-(E_3)$	1 ← 1	(19,20) ← (7,8)	53	730.82 GHz ^b
$G_3^+/G_4^-(??) \leftarrow G_4^+/G_3^-(E_1)$	2 ← 1	$(25,26) \leftarrow (5,6)$	24	746.4 GHz
$G_3^+(E_4) \leftarrow G_4^+(E_1)$	1←0	$(21,22) \leftarrow (4)$	4	757.6 GHz
$G_3^-(E_4) \leftarrow G_4^-(E_3)$	1 ←0	(19,20) ← (2)	18	758.8 GHz
$G_3^+(E_4) \leftarrow G_4^+(E_3)$	1 ←0	$(21,22) \leftarrow (1)$	17	759.9 GHz
$G_4^-/G_3^+(E_2) \leftarrow G_3^-/G_4^+(E_1)$	1 ← 1	$(23,24) \leftarrow (11,12)$	21	765.1 GHz
$G_4^-/G_3^+(E_2) \leftarrow G_4^+(E_3)$	1 ←0	$(23,24) \leftarrow (4)$	19	1051.8 GHz
$G_4^-/G_3^+(E_2) \leftarrow G_3^-(E_1)$	1 ← 0	(23,24) ← (3)	20	1052.9 GHz
$G_4^-/G_3^+(E_2) \leftarrow G_4^+(E_1)$	1 ← 0	(23,24) ← (1)	4	1054.2 GHz
		(b)		
$G_2^+/G_2^- \leftarrow G_2^+/G_2^-$	1 ← 1	(9,10) ← (3,4)	45	486.8 GHz ^a
$G_2^+ \leftarrow G_2^+ / G_2^-$	0 ← 1	$(11) \leftarrow (3,4)$	20	500.7 GHz
$G_2^- \leftarrow G_2^+ / G_2^-$	0 ← 1	$(12) \leftarrow (3,4)$	22	503.1 GHz
$G_2^+/G_2^- \leftarrow G_2^-$	1←0	(9,10) ← (2)	26	597.4 GHz
$\tilde{G_2^+}/\tilde{G_2^-} \leftarrow \tilde{G_2^+}$	1 ⊷0	(9,10) ← (1)	31	600.7 GHz
$G_2^+ \leftarrow G_2^-$	0 ← 0	(11) ↔ (2)	9	611.3 GHz ^c
$\tilde{G_2} \leftarrow \tilde{G_2}$	0⊷0	(11) ← (1)	24	613.7 GHz ^b
$\tilde{G_2^+} \leftarrow \tilde{G_2^+}$	0 ← 0	(12) ← (2)	23	614.6 GHz ^b
$G_2^- \leftarrow G_2^+$	0⊷0	$(12) \leftarrow (1)$	8	617.0 GHz ^c
$G_2^+/G_2^- \leftarrow G_2^+/G_2^-$	2 ← 2	$(15,16) \leftarrow (7,8)$	16	638.2 GHz
$G_2^+/G_2^- \leftarrow G_2^-/G_2^+$	1 ← 1	(13,14) ← (5,6)	60	747.2 GHz ^b
$G_2^+/G_2^- \leftarrow G_2^-$	1 ← 0	$(13,14) \leftarrow (2)$	28	975.9 GHz
$G_2^+/G_2^- \leftarrow G_2^+$	1←0	(13,14) ← (1)	31	979.2 GHz
$G_2^+/G_2^- \leftarrow G_2^+/G_2^-$	2 ← 1	(15,16) - (3,4)	49	1041.6 GHz
		(c)		
$A_2^+ \leftarrow B_2^+$	0→0	(4) ← (1)	22	483.3 GHz ^a
$A_2^+ \leftarrow B_2^+$	1 ← 1	$(5,6) \leftarrow (2,3)$	36	518.5 GHz
$A_2^+ \leftarrow B_2^+$	1⊷0	(5,6) ← (1)	31	729.0 GHz ^b

^aTransitions in this subband were observed independently by Zwart (Ref. 23).

^bThis subband was previously reported by Havenith et al. (Ref. 21).

^cQ branch only (due to Coriolis mixing of $K_a = 0$ upper states with $K_a = 1$ states).

 $c[J(J+1)-2]^{1/2}$. The *c* constants so determined are labeled by the numbers of the manifolds they couple. The form of the Coriolis coupling matrix elements is chosen to be equivalent to that of the Coriolis term given by Brocks *et al.*³² in their quantum prescription for modeling the intermolecular dynamics of van der Waals's dimers:

$$\delta_{\Omega',\Omega+1}B[j(j+1)-\Omega(\Omega+1)]^{1/2}$$
$$\times [J(J+1)-\Omega(\Omega+1)]^{1/2},$$

where B is the rotational constant of the complex [the same B used in expression (1) and (2) above], j is the angular momentum from rotation of the monomers, J is the total angular momentum, and Ω is the projection of total angular momentum along the **a** inertial axis. In this formalism Ω is equivalently the projection of both j and J onto the **a** axis, and when this Coriolis analysis is appropriate, we treat our quantum number K_a as Ω . This matrix element couples pure free internal rotor basis functions³² of the

	(a)		
		E_{13}	577 864.45 (53)
Bi	5115.17 (184)	B ₁₃	5050.422 (19)
D_1	0.0643 (16)	D ₁₃	0.051 10 (12)
E ₂	1094.55 (107)	E_{14}	580 081.15 (104)
B ₂	5115.64 (126)	B ₁₄	5049.887 (31)
D_2	0.07182 (38)	D_{14}	0.049 97 (19)
£3	1267.00 (73)	E_{15}	581 583.88 (73)
8 <u>,</u>	5115.52 (120)	B ₁₅	5050.786 (28)
D ₃	0.077 45 (38)	D ₁₅	0.051 36 (24)
54	2369.53 (81)	E_{16}	583 800.33 (56)
B ₄	5115.91 (172)	B ₁₆	5050.370 (19)
D4	0.0832 (16)	D ₁₆	0.051 25 (13)
E _{5.6}	- 77 628.40 (62)	$E_{17,18}$	375 971.81 (73)
9 _{5.6}	5096.55 (120)	B _{17,18}	5002.913 (24)
D _{5,6}	0.024 66 (20)	$D_{17,18}$	0.046 21 (16)
		<i>qB</i> _{17,18}	0.0175 (44)
E _{7.8}	29 088.00 (89)	$E_{19,20}$	759 910.56 (93)
B _{7.8}	5078.29 (63)	$B_{19,20}$	4933.472 (27)
D _{7.8}	0.039 34 (22)	D _{19,20}	0.011 26 (14)
1B7,8	24.67 (63)	$qB_{19,20}$	1.8319 (91)
$D_{7,8}$	0.005 01 (18)	$qD_{19,20}$	-0.002 133 (92)
E _{9.10}	29 136.21 (61)	$E_{21,22}$	759 919.60 (60)
B _{9,10}	5078.37 (61)	B _{21,22}	4933.530 (20)
D _{9,10}	0.037 66 (33)	$D_{21,22}$	0.011 45 (13)
7B _{9,10}	24.75 (61)	$qB_{21,22}$	1.8508 (94)
$qD_{9,10}$	0.003 41 (30	$qD_{21,22}$	-0.002 250 (92)
E _{11,12}	289 065.24 (74)	$E_{23,24}$	1054 170.59 (64)
B _{11,12}	5065.411 (33)	B _{23,24}	5141.999 (48)
D _{11,12}	0.063 47 (31)	D _{23,24}	0.2319 (11)
		H _{23,24}	0.000 2798 (65)
		$E_{25,26}$	668 760.09 (78)
		B _{25,26}	5038.885 (28)
		D _{25,26}	0.049 25 (21)

TABLE III. Rotational constants and relative energies as defined in Eqs. (1) and (2) for the (a) $E+E$, (b)
A+E, and (c) $A+A$ states.

Coriolis coupling constants

C _{1,5} C _{4,5}	-2117.2 (116) 2276.3 (129)	$(c_{3,6})^2 = (c_{1,4})^2 + (c_{4,5})^2$
C _{1,10} C _{4,10}	1519.4 (175) 1313.0 (158)	$(c_{2,8})^2 = (c_{1,10})^2 + (c_{4,10})^2$

		E_{11}	614 635.52 (43)
B	5110.410 54 (47)	$B_{11}^{}$	5035.139 (52)
D	0.052 683 (49)	D_{11}	0.050 93 (22)
<i>E</i> ₂	3309.41 (26)	E_{12}	617 027.53 (45)
B_2	5110.561 16 (48)	B ₁₂	5034.914 (47)
D_2	0.052 478 (51)	D_{12}	0.050 90 (24)
E _{3.4}	113 938.33 (38)	$E_{9.10}$	600 730.81 (41)
B _{3.4}	5125.420 (12)	$B_{9,10}$	5042.463 (76)
D _{3,4}	0.060 599 (80)	$D_{9,10}$	0.056 19 (35)
$qB_{3,4}$	0.4920 (20)	-,	
E _{5.6}	232 026.98 (54)	$E_{13,14}$	979 190.39 (39)
B5,6	5117.218 (19)	B _{13,14}	5044.414 (12)
D _{5.6}	0.053 93 (14)	D _{13,14}	0.033 893 (81)
$qB_{5,6}$	0.1536 (33)	$qB_{13,14}$	0.0302 (22)

(b)

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TABLE III. (Continued.	BLE III. (Continued	.)
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E _{7,8} B _{7,8}	517 333.29 (114) 5125.30 (17)	$E_{15,16}$ $B_{15,16}$	1155 578.12 (60) 5048.445 (17) 0.046 66 (11)
D _{7,8}	0.055 18 (88)	$D_{15,16}$	0.048 88 (11)
Coriolis coupling constants			
$c_{9,11} = c_{10,12}$	1270.09 (25)		
$c_{7,9} = c_{8,10}$	1597.7 (30)		
	(c)		
		E_4	483 299.54 (38)
<i>B</i> ₁	5136.646 (31)	B_4	5030.566 (29)
$\dot{D_1}$	0.058 92 (29)	D_4	0.048 93 (22)
<i>E</i> _{2,3}	210 526.08 (49)	$E_{5,6}$	729 026.77 (37)
$B_{2,3}^{-2,3}$	5140.986 (34)	B _{5,6}	5031.691 (28)
$D_{2,3}$	0.058 14 (33)	D _{5,6}	0.048 49 (24)
$qB_{2,3}$	2.731 (13)	<i>qB</i> _{5,6}	12.3862 (91)
$qD_{2,3}$	0.001 10 (15)	$qD_{5,6}$	-0.001 176 (92)

same internal rotor quantum numbers but different values of Ω . The actual VRT states of the ammonia dimer are expected to be mixtures of many free internal rotor states, so that the Coriolis matrix element between any two ammonia dimer states should be smaller than the free internal rotor limit given above. This is consistent with the magnitudes of the coupling coefficients determined in our fits.

We determined one of the Coriolis coupling coefficients in the lowest E+E states to be negative. This nonphysical result is an unresolved point of inconsistency in the analysis. This problem involves four of the E+E VT states (states 1, 4, 5, and 10) that are observed to have the same symmetry in the lowest E+E states. To clarify this discussion, we present a "close-up" diagram of only the lowest J=1 E+E states in Fig. 4. (The symmetry labels indicated in Fig. 4 will be discussed later in this paper.) Dashed lines show all possible $K_a = 0 - K_a = 1$ Coriolis interactions between the states 1, 4, 5, and 10. We determined all of the corresponding coupling constants in our fit. It is easy to show that all four of these states involved do mutually interact, as indicated by the dashed lines in Fig. 4, by comparing them to the pair of states 2 and 8 and the pair of states 3 and 6. States 8 and 10 are both upper components of $K_a = 1$ K-type doublets, and their rotational energy-level spacings are nearly indistinguishable at our experimental accuracy. State 8 is strongly perturbed through a Coriolis interaction with state 2; however, neither state 1 nor state 4 couples to state 10 in the same way. Similarly, states 6 and 5 are the upper and lower components of a $K_a = 1$ K-type doublet that is just barely split at high J. State 6 is strongly coupled to state 3, yet neither state 1 nor state 4 couples to state 5 in the same way. Further, we note that the deviation of the rotational spacings in state 1 from the rotational spacings in state 2 is equal and opposite to the deviation of the rotational spacings in state 4 from the rotational spacings in state 3. (This can be verified from Table III using simple arithmetic.) We know that this effect is not due to the direct coupling of state 1 to state 4 because they shift towards each other rather than further apart, and because states 13 and 16,

two higher $K_a=0$ states that are also very close in energy and have the same symmetry, do not interact. Therefore, we suggest that states 1 and 4 are coupled indirectly via interactions with the nearby $K_a=1$ states 5 and 10. In our analysis, we assume that Coriolis interactions are responsible for this coupling.

After we remove the effects of the strongest Coriolis interactions between $K_a=0$ and $K_a=1$, we find that the fitted B_n constants consistently cluster around different values for the three groups of tunneling states, and also that the B_n values of the "upper" states are consistently about 1% smaller than the "lower" state B_n values. Without this explicit Coriolis treatment, variations in the observed values of B_n would appear to be random. The rotational constants we determined after including the Coriolis interactions between $K_a=0$ and $K_a=1$ are still not pure "deperturbed" constants. Our data show evidence of strong Coriolis interactions between $K_a=1$ and $K_a=2$ as well, and we have not yet assigned all the $K_a=2$ states involved.

B. Molecular symmetry group and symmetry assignment

The appropriate choice for the molecular symmetry group of the ammonia dimer depends on which hydrogen tunneling motions occur on the time scale of our experiment. The most complete possible molecular symmetry group for the ammonia dimer is the permutation-inversion group G_{144} ,³³ which includes all operations that permute the three protons on each ammonia subunit, operations that permute the two ammonia subunits, and spatial inversion. Only permutations that correspond to breaking chemical N-H bonds are excluded from G_{144} . We assume that the ammonia dimer equilibrium structure contains a plane of symmetry, because there is no evidence that contradicts this simplifying assumption. Three kinds of tunneling motions can then be represented by the operations in G_{144} : interchange of the two inequivalent ammonia

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TABLE IV. Relative energies (MHz) of observed (a) E+E, (b) A+E, and (c) A+A states up to J=5.

(<i>n</i>)	Sym.	J	Ka	Rel. energy	(n)	Sym.	J	Ka	Rel. energy
				(;	a)				
(1)	G4 + (E3)	0	0	0.0	(13)	G3 + (E4)	0	0	577 864.4
(1)	G3 - (E4)	1	0	10 159.7	(13)	G4 - (E3)	1	0	587 965.1
(1)	G4 + (E3)	2	0	30 345.0	(13)	G3 + (E4)	2	0	608 165.1
(1)	G3 - (E4)	3	0	60 442.9	(13)	G4 - (E3)	3	0	638 462.2
(1)	G4 + (E3)	4	0	100 466.2	(13)	G3 + (E4)	4	0	678 852.4
(1)	G3 - (E4)	5	0	150 473.1	(13)	G4 - (E3)	5	0	729 331.1
(2)	G4-(E3)	0	0	1094.5	(14)	G3 - (E4)	0	0	580 081.2
(2)	G3 + (E4)	1	0	11 040.1	(14)	G4+(E3)	1	0	590 180.7
(2)	G4-(E3)	2	0	30 944.5	(14)	G3-(E4)	2	0	610 378.7
(2)	G3 + (E4)	3	0	60 830.9	(14)	G4+(E3)	3	0	640 672.6
2)	G4 - (E3)	4	0	100 726.3	(14)	G3 - (E4)	4	0	681 058.9
(2)	G3 + (E4)	5	0	150 657.3	(14)	G4+(E3)	5	0	731 532.8
3)	G3 - (E1)	0	0	1267.0	(15)	G4-(E2)	0	0	581 583.9
3)	G4 + (E1)	1	0	11 741.8	(15)	G3 + (E2)	1	0	591 685.2
3)	G3 - (E1)	2	0	32 684.6	(15)	G4 - (E2)	2	0	611 886.7
3)	G4 + (E1)	3	0	64 081.7	(15)	G3 + (E2)	3	0	642 185.9
3)	G3-(E1)	4	0	105 914.4	(15)	G4 - (E2)	4	0	682 579.1
3)	G4 + (E1)	5	0	158 159.6	(15)	G3 + (E2)	5	0	733 061.2
(4)	G4 + (E1)	0	0	2369.5	(16)	G3 + (E2)	0	0	583 800.3
4)	G3 - (E1)	1	0	12 630.9	(16)	G4-(E2)	1	0	593 900.9
(4)	G4 + (E1)	2	0	33 294.2	(16)	$G_{3}+(E_{2})$	2	0	614 100.7
4)	$G_{3}-(E_{1})$	3	0	64 481.8	(16)	G4-(E2)	3	0	644 397.4
4)	G4 + (E1)	4	0	106 189.3	(16)	G3 + (E2)	4	0	684 787.2
4)	G3 - (E1)	5	0	158 361.9	(16)	G4-(E2)	5	0	735 265.3
					(17)	G3 + (E2)	1	1	385 977.4
5,6)	$G_{3}-,G_{4}+(E_{1})$	1	1	-67 679.5	(18)	G4-(E2)	1	1	385 977.5
5)	G4 + (E1)	2	1	-47 777.3	(17)	G4-(E2)	2	1	405 987.5
6)	$G_{3}-(E_{1})$	2	1	-47 777.2	(18)	G3 + (E2)	2	1	405 987.7
5)	$G_{3}-(E_{1})$	3	1	-17 913.1	(17)	$G_{3}+(E_{2})$	3	1	435 999.9
6)	G4 + (E1)	3	1	- 17 912.9	(18)	G4 - (E2)	3	1	436 000.3
5)	G4 + (E1)	4	1	21 924.5	(17)	G4-(E2)	4	1	476 011.2
6)	G3 - (E1)	4	1	21 924.9	(18)	G3 + (E2)	4	1	476 011.9
5)	G3 - (E1)	5	1	71 749.0	(17)	G3 + (E2)	5	1	526 017.1
6)	G4 + (E1)	5	1	71 749.5	(18)	G4-(E2)	5	1	526 018.1
7)	G4 - (E3)	1	1	39 195.0	(19)	G4+(E3)	1	1	769 773.8
8)	G3 + (E4)	1	1	39 579.2	(20)	G3 - (E4)	1	1	769 781.1
7)	G3 + (E4)	2	1	59 408.1	(19)	G3 - (E4)	2	1	789 500.1
8)	G4 - (E3)	2	1	60 545.8	(20)	G4 + (E3)	2	1	789 521.9
7)	G4 - (E3)	3	1	89 724.9	(19)	G4 + (E3)	3	1	819 088.9
8)	G3 + (E4)	3	1	91 959.5	(20)	$G_{3}-(E_{4})$	3	1	819 132.3
7)	G3 + (E4)	4	1	130 142.5	(19)	$G_{3}-(E_{4})$	4	1	858 539.7
8)	G4 - (E3)	4	1	133 785.7	(20)	G4 + (E3)	4	1	858 611.3
7)	G4 - (E3)	5	1	180 656.4	(19)	G4 + (E3)	5	1	907 851.6
8)	G3 + (E4)	5	1	185 987.6	(20)	G3-(E4)	5	1	907 957.6
9)	G4 + (E3)	1	1	39 243.3	(21)	G4 + (E3)	1	1	769 782.9
10)	G3 - (E4)	1	1	39 626.9	(22)	G3 - (E4)	1	1	769 790.3
9)	G3 - (E4)	2	1	59 456.5	(21)	G3 - (E4)	2	1	785 509.3
10)	G4+(E3)	2	1	60 592.6	(22)	G4+(E3)	2	1	789 531.4
9)	G4 + (E3)	3	1	89 773.7	(21)	G4 + (E3)	3	1	819 098.4
10)	G3 - (E4)	3	1	92 005.2	(22)	G3 - (E4)	3	1	819 142.2
9)	G3 - (E4)	4	1	130 192.2	(21)	G3-(E4)	4	1	858 549.5
10)	G4 + (E3)	4	1	133 830.0	(22)	G4+(E3)	4	1	858 621.7
9) 10)	G4 + (E3)	5	1	180 707.8	(21)	G4 + (E3)	5	1	907 861.7
10)	G3 - (E4)	5	1	186 030.4	(22)	G3 - (E4)	5	1	907 968.7
	G4+,G3-(E1)	1	1	299 195.8	(23,24)	G4 - ,G3 + (E2)	1	1	1 064 453.7
		2	1	319 455.4	(23,24)	$G_{3}+,G_{4}-,(E_{2})$	2	1	1 085 014.3
11,12)	$G_{3-},G_{4+}(E_{1})$	~		A 4 A A A A A A			-		
11,12) 11,12)	G4+,G3-(E1)	3	1	349 841.0	(23,24)	G4-,G3+(E2)	3	1	
11,12) 11,12) 11,12)	G4+,G3-(E1) G3-,G4+(E1)	4	1	390 348.1	(23,24)	G3+,G4-(E2)	4	1	1 115 841.7 1 156 920.1 1 208 229 5
(11,12) (11,12) (11,12) (11,12) (11,12) (11,12)	G4+,G3-(E1)								

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TABLE IV. (Continued.)

(<i>n</i>)	Sym.	J	Ka	Rel. energy	(<i>n</i>)	Sym.	J	Ка	Rel. energy
					(25,26)	G4-,G3+	3	2	729 219.6
					(25,26)	G3+,G4-	4	2	769 518.1
					(25,26)	G4-,G3+	5	2	819 882.3
					(b)				
(1)	<i>G</i> 2+	0	0	0.0	(11)	G_2+	0	0	614 635.t
(1)	G2 —	1	0	10 220.6	(11)	G2-	1	0	624 934.1
(1)	G2+	2	0	30 660.6	(11)	G_{2+}	2	0	645 515.4
(1)	G2	3	0	61 317.3	(11)	G2	3	0	676 352.6
(1)	G2+	4	0	102 187.1	(11)	<i>G</i> 2+	4	0	717 414.8
(1)	G2-	5	0	153 264.9	(11)	G2-	5	0	768 671.3
(2)	G2-	0	0	3309.4	(12)	G2-	0	0	617 027.5
(2)	G^{2+}	1	0	13 530.3	(12)	G2+	1	0	627 293.0
(2)	G2-	2	0	33 970.9	(12)	G2	2	0	647 813.8
(2)	G2 + G2	3 4	0 0	64 628.6 105 499.6	(12)	G2+ G2-	3 4	0 0	678 572.3
(2) (2)	$G_2 = G_2 + G_2$	4 5	0	156 579.0	(12) (12)	62- 62+	4 5	0	719 546.1 770 711.7
(3)	G2 -	1	1	124 187.9	(9)	G2-	1	1	610 587.0
(3)	62- 62+	1	1	124 187.9	(10)	G_2^{-}	1	1	610 619.7
(3)	G_2^+	2	1	144 685.7	(9)	62+ 62+	2	1	630 435.8
(4)	G2 -	2	1	144 691.6	(10)	G2-	2	1	630 528.1
(3)	G2-	3	1	175 428.7	(9)	G2 -	3	1	660 238.8
(4)	G_2^+	3	1	175 440.6	(10)	G2+	3	1	660 408.5
(3)	G_2^+	4	1	216 412.7	(9)	$G^2 +$	4	1	700 021.3
(4)	G2	4	1	216 432.3	(10)	G2-	4	1	700 277.4
(3)	G2-	5	1	267 631.6	(9)	G2	5	1	749 806.0
(4)	G2+	5	1	267 661.2	(10)	G2+	5	1	750 150.8
(5)	<i>G</i> 2+	1	1	242 260.9	(13)	G2-	1	1	989 279.0
(6)	G2	1	1	242 261.5	(14)	G2+	1	1	989 279.1
(5)	G2	2	1	262 727.4	(13)	$G_2 +$	2	1	1 009 455.5
(6)	G2+	2	1	262 729.3	(14)	G2 —	2	1	1 009 455.8
(5)	G2+	3	1	293 424.0	(13)	G2 —	3	1	1 039 718.1
(6)	G2	3	1	293 427.7	(14)	G2+	3	1	1 039 718.8
(5)	G2-	4	1	334 346.7	(13)	G2+	4	1	1 080 064.5
(6)	G_2^+	4	1	334 352.8	(14)	G2-	4	1	1 080 065.7
(5)	G2+	5	1	385 490.4	(13)	G2	5	1	1 130 491.4
(6)	G2	5	1	385 499.6	(14)	G2+	5	1	1 130 493.2
(7,8)	$G_{2+,G_{2-}}$	2	2	547 960.0	(15,16)	$G_{2}+,G_{2}-$	2	2	1 185 867.1
(7,8)	$G_{2}^{-},G_{2}^{-}+$	3	2	578 519.6	(15,16)	$G_{2}^{2}-,G_{2}^{2}+$	3	2	1 216 152.7
(7,8)	G_{2}^{+},G_{2}^{-}	4	2	619 256.6	(15,16)	$G_{2+,G_{2-}}$	4	2	1 256 528.3
(7,8)	G2-,G2+	5	2	670 163.3	(15,16)	G2-,G2+	2	2	1 306 989.5
					(c)				
(1)	A2+	0	0	0.0	(4)	<i>B</i> 2+	0	0	483 299.5
(1)	B2	1	0	10 273.1	(4)	A2 —	1	0	493 360.5
(1)	A2+	2	0	30 817.8	(4)	B2+	2	0	513 481.2
(1)	<i>B</i> 2 –	3	0	61 631.3	(4)	A2-	3	0	543 659.3
(1)	A2+	4	0	102 709.4	(4)	B2+	4	0	583 891.3
(1)	B2	5	0	154 046.6	(4)	A2-	5	0	634 172.5
(2)	A2+ B2-	1 1	1 1	220 802.4 220 813.3	(5)	B2+A2-	1	1 1	739 065.2
(3) (2)	B2 B2	1 2	1	220 813.3 241 353.5	(6) (5)	A2— A2—	1 2	1	739 114.7 759 140.9
(2)	A2 +	2	1	241 333.3	(6)	B2+	2	1	759 140.9
(2)	A2+ A2+	3	1	272 176.6	(5)	B2+B2+	2 3	1	739 289.4
(2)	B2	3	1	272 242.5	(6)	A2-	3	1	789 548.5
(2)	B2	4	1	313 267.5	(5)	A2-	4	1	829 393.9
(2)	A2 +	4	1	313 377.6	(6)	B2+	4	1	829 888.4
(2)	A2+	5	1	364 620.4	(5)	B2 + B2 +	5	1	879 563.3
(3)	B2	5	1	364 786.2	(6)	A2-	5	1	880 304.4

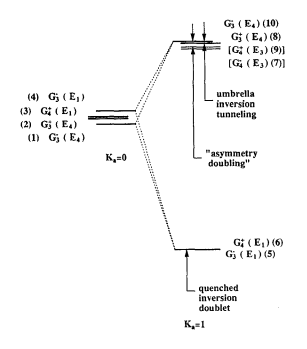


FIG. 4. Enlarged energy-level diagram for the lowest J=1 E+E states. The observed Coriolis interactions within the G_3^- states (1), (4), (5), and (10) are indicated with dashed lines. We also observed Coriolis coupling between the G_4^+ states (3) and (6) as well as between the G_3^+ states (2) and (8). The $K_a = 1$ G_4^- state (7) and G_4^+ state (9) are unperturbed by $K_a=0$ states. The Coriolis perturbations are discussed in the rotational analysis section of the text and the symmetry assignments are discussed in the E+E symmetry analysis section.

monomers (the donor and the acceptor), internal rotation of the monomers about their C_3 symmetry axes, and umbrella inversion tunneling of each monomer. A subgroup of G_{144} , the permutation inversion group G_{36} , was used in the analysis of Nelson and Klemperer.¹⁰ This group does not account for the umbrella inversion tunneling, and was used because it was assumed that the umbrella inversion tunneling would be totally quenched by the anisotropy in the ammonia dimer potential-energy surface. Perhaps surprisingly, we deduce from our data that G_{144} is, in fact, the appropriate molecular symmetry group for the ammonia dimer. Nevertheless, it is useful to examine the data in terms of both G_{36} and G_{144} descriptions.

In order to interpret the ammonia dimer VRT spectrum, we must determine how many tunneling states arise from each vibration-rotation state of the dimer, and we must label them with irreducible representations of the molecular symmetry group. This will allow us to predict the number of VRT states that will be spectroscopically "connected" for each vibration-rotation state of the ammonia dimer. The molecular symmetry group of the rigid, nontunneling complex is C_s . The correlation between C_s and G_{36} has been derived previously by Nelson and Klemperer.¹⁰ They showed that, in G_{36} , there are eight tunneling sublevels associated with each rotational level of the rigid $C_{\rm s}$ complex,

$$A' = A_1(66) \oplus A_4(78) \oplus G(144) \oplus G(144) \oplus E_1(36)$$
$$\oplus E_2(36) \oplus E_3(30) \oplus E_4(42),$$
$$A'' = A_2(78) \oplus A_2(66) \oplus G(144) \oplus G(144) \oplus E_1(36)$$

$$\oplus E_2(36) \oplus E_4(42) \oplus E_2(30)$$
 (3)

The nuclear-spin statistical weights for the VRT levels of $(^{14}NH_3)_2$ are given in parentheses after each irreducible representation. Also, Eqs. (3) are aligned so that the symmetry species of the even- and odd-J end-over-end rotational levels of each of the eight tunneling sublevels are shown in the vertical columns. The selection rules for electric-dipole-allowed transitions, assuming molecular symmetry G_{36} , were found to be $A_1 \leftrightarrow A_3$, $A_2 \leftrightarrow A_4$, $G \leftrightarrow G$, $E_1 \leftrightarrow E_2$, $E_3 \leftrightarrow E_3$, and $E_4 \leftrightarrow E_4$. Inspection of Eqs. (3) with these selection rules in mind reveals that transitions between either the A or E symmetry levels must involve a change in tunneling state while transitions between the Gsymmetry levels may or may not involve a change in tunneling state. That is, for example, since A_1 and A_3 appear in different vertical columns in Eqs. (3), the frequency of a transition between a $J=0A_1$ rotation-tunneling state and a $J=1 A_3$ rotation-tunneling state will include a contribution from tunneling as well as the usual rotational spacing. Even though rotation-tunneling states of A_1 and A_2 symmetry are not connected by electric dipole transitions, the energy separations between these states are purely rotational and are thus determined in the rotational fits. In G_{36} , the electric dipole selection rules separate the observed VRT levels of the ammonia dimer into four groups: (1) the A_1 , A_2 , A_3 , and A_4 levels (A+A states, we explain this nomenclature later in this section), (2) the G levels (A+E)states), (3) the E_1 and E_2 levels (E+E states), and (4) the E_3 and E_4 levels (also E+E states). All of the tunneling sublevels within each group are connected by electricdipole-allowed transitions, but transitions between tunneling sublevels in different groups are forbidden. We emphasize that, in G_{36} , there is only one tunneling sublevel of each E symmetry type. Also, we note that there are two groups of E+E states in G_{36} and electric dipole transitions are not allowed between these two groups.

When umbrella inversion tunneling is considered feasible in the ammonia dimer, G_{144} becomes the appropriate molecular symmetry group. Now, there are 22 tunneling sublevels associated with each rotational level of the rigid complex having C_s symmetry, but only 14 tunneling sublevels have nonzero nuclear-spin statistical weight for $(^{14}NH_3)_2$. We expect that the umbrella inversion tunneling motion has the highest barrier of all the tunneling motions we are considering for the ammonia dimer. Thus, we imagine that the more widely spaced G_{36} tunneling states are further split by small umbrella inversion tunneling frequencies into the G_{144} tunneling states. The correlation from G_{36} to G_{144} is

$$A_{1} = A_{1}^{-}(0) \oplus E^{-}(0) \oplus B_{2}^{+}(66),$$

$$A_{2} = B_{1}^{-}(0) \oplus E^{+}(0) \oplus A_{2}^{-}(78),$$

$$A_{3} = A_{1}^{-}(0) \oplus E^{+}(0) \oplus B_{2}^{-}(66),$$

$$A_{4} = B_{1}^{+}(0) \oplus E^{-}(0) \oplus A_{2}^{+}(78),$$

$$G = G_{1}^{+}(0) \oplus G_{1}^{-}(0) \oplus G_{2}^{+}(72) \oplus G_{2}^{-}(72),$$

$$E_{1} = G_{3}^{-}(21) \oplus G_{4}^{+}(15),$$

$$E_{2} = G_{3}^{+}(21) \oplus G_{4}^{-}(15),$$

$$E_{3} = G_{4}^{+}(15) \oplus G_{4}^{-}(15),$$

$$E_{4} = G_{3}^{+}(21) \oplus G_{3}^{-}(21).$$
(4)

Again, the nuclear-spin statistical weights for VRT levels of $({}^{14}\text{NH}_3)_2$ are shown in parentheses after the G_{144} irreducible representations. So, the complete correlation between each C_s rotational level and the G_{144} tunneling sublevels is

$$A' = A_1^+ \oplus B_1^+ \oplus 2E^- \oplus A_2^+ \oplus B_2^+ \oplus 2G_1^+ \oplus 2G_1^- \oplus 2G_2^+ \oplus 2G_2^- \oplus 2G_3^+ \oplus 2G_3^- \oplus 2G_4^+ \oplus 2G_4^-, A'' = B_1^- \oplus A_1^- \oplus 2E^+ \oplus B_2^- \oplus A_2^- \oplus 2G_1^- \oplus 2G_1^+ \oplus 2G_2^- \oplus 2G_2^+ \oplus 2G_4^- \oplus 2G_4^+ \oplus 2G_3^- \oplus 2G_3^+.$$
(5)

Equations (5) are also aligned so that the vertical columns show the alternation of symmetry label with even and odd J. The electric dipole selection rules in G_{144} are $A_1^+ \leftrightarrow A_1^-$, $B_1^+ \leftrightarrow B_1^-$, $E^+ \leftrightarrow E^-$, $A_2^+ \leftrightarrow A_2^-$, $B_2^+ \leftrightarrow B_2^-$, $G_1^+ \leftrightarrow G_1^-$, $G_2^+ \leftrightarrow G_2^-$, $G_3^+ \leftrightarrow G_3^-$, and $G_4^+ \leftrightarrow G_4^-$. Pure rotational transitions are allowed in the G_1^+ , G_1^- , G_2^+ , G_2^- , E^+ , and $E^$ levels, but transitions between all other rotation-tunneling levels require a change in tunneling state. In G_{144} , the A + Astates are the A_1^+ , A_1^- , B_1^+ , B_1^- , E^+ , E^- , A_2^+ , A_2^- , B_2^+ , and B_2^- levels, the A+E states are the G_1^+ , G_1^- , G_2^+ , and $G_2^$ levels, and the E+E states are the G_3^+ , G_3^- , G_4^+ , and $G_4^$ levels. Since some of these have zero statistical weight for $(^{14}NH_3)_2$, we expect to observe only three groups of ammonia dimer VRT levels in G_{144} : (1) the A_2^+, A_2^-, B_2^+ , and B_2^- levels (A+A states), (2) the G_2^+ and G_2^- levels (A+E states), and (3) the G_3^+ , G_3^- , G_4^+ , and G_4^- levels (E+E states)states). We emphasize that, in G_{144} but not in G_{36} , there are two tunneling sublevels of each E+E symmetry type, and all the E+E tunneling manifolds can be connected together by electric-dipole-allowed transitions. Later, we exploit these important differences between the grouptheoretical description of the E+E states in G_{36} and that of the E+E states in G_{144} to assign the appropriate molecular symmetry group for the ammonia dimer.

It is useful to explore the correlation between the tunneling sublevels of the ammonia dimer and the rotational states of two ammonia monomers. The ammonia monomer equilibrium geometry has a threefold axis of symmetry, so the rotational states are classified as A or E, for $k_c \pmod{3}$ =0 and $k_c \pmod{3} \neq 0$, respectively. There are also two proton spin states, I=3/2(ortho) and I=1/2(para) of A

and E symmetry, respectively, which pair with the A and E rotational states. This leads to the well-known fact that electric dipole transitions between the states of A and Esymmetry are forbidden. This selection rule also applies to the dimer complex when internal rotation tunneling is feasible: it leads to the three separate groups of G_{144} tunneling states that we discussed above. In the accompanying paper, van Bladel et al.²² refer to the A + A states as ortho-ortho states, the A + E states as ortho-para states, and the E + Estates as para-para states. If there is a very low barrier to internal rotation in the ammonia dimer, as in the more strongly hydrogen-bonded HOH-NH3 dimer where the barrier to internal rotation of the ammonia acceptor was determined by Stockman et al.³⁴ to be $\leq 10 \text{ cm}^{-1}$, a correlation between dimer internal rotation states and monomer rotational states is directly useful for estimating the relative internal rotation splittings in the ammonia dimer. Since the ammonia monomers in the ammonia dimer can interchange their donor and acceptor roles, the average barrier to internal rotation in the ammonia dimer is expected to be higher than in HOH-NH₃. If the ammonia rotors are substantially hindered so that a high barrier analysis of the tunneling splittings is appropriate-such a formalism has recently been described by Coudert and Hougen³⁵-the correlation between dimer tunneling states and monomer rotational states is still useful for establishing upper limits for the relative internal rotation tunneling splittings in the ammonia dimer.

In the calculations described in the accompanying paper, van Bladel et al.²² have used basis functions formed from linear combinations of coupled monomer symmetric rotor wave functions.³¹ These basis functions have been symmetrized so that they transform according to irreducible representations of G_{36} . The true VRT states of the ammonia dimer are not described by the pure (or "free") symmetrized basis functions; rather, they are mixed by anisotropy (long-range electrostatic, induction and dispersion interactions, short-range exchange forces, etc.) in the intermolecular potential-energy surface. The description of the true mixed VRT states of the ammonia dimer is the ultimate goal in the calculations of van Bladel et al. Nevertheless, the free internal rotor limit has been used by Nelson and Klemperer¹⁰ to illustrate the energetics and internal rotation dynamics due to the internal rotation of the ammonia monomers about their C_3 axes. We describe the free-rotor components of some of the lowest-energy G_{36} basis functions and the G_{144} analogs for total J=0 below.

The A_1 and A_4 tunneling states of G_{36} , or the A_1^+ , B_2^+ , B_1^+ , A_2^+ , and E^- tunneling states of G_{144} , are constructed from ammonia monomer A states. For example, the combination of two ammonia monomers in 0_0 rotational states transforms like A_1 in G_{36} , and the combination of one monomer in 0_0 with the other in 1_0 transforms as $A_1 \oplus A_4$. In these lowest A + A basis functions, there is no projection of internal angular momentum from internal rotation about the monomer C_3 symmetry axes. (Of course, the true VRT wave functions probably also contain contributions from 3_3 states, coupled in by electrostatic interactions involving the ammonia monomer electric octopole moment

as well as short-range repulsive contributions caused by steric effects from the three hydrogens on each ammonia monomer. However, the 33 contributions will not be dominant in the lowest A + A states.) To generalize from G_{36} to G_{144} , we must include the umbrella inversion states of the monomers. There is a symmetric and antisymmetric inversion component for every monomer rotational state. In the A states with $k_c=0$, only one component has nonzero statistical weight; in 0₀, this is the antisymmetric inversion state 0_0^- . The combination of two monomers in 0_0^+ transforms as A_1^+ in G_{144} , which of course has zero statistical weight. One monomer in 0_0^+ plus one monomer in 0_0^- gives E^- in G_{144} , which also has zero statistical weight. The combination of two monomers in 0_0^- transforms as B_2^+ in G_{144} , which we can observe spectroscopically.

The G states of G_{36} , and the G_1^+ , G_1^- , G_2^+ , and $G_2^$ states of G_{144} , are constructed from one monomer in an A state plus one monomer in an E state. The lowest-energy free monomer basis function of G symmetry in G_{36} involves one monomer in 0_0 (an A state) with the other in 1_1 (an E state). The fourfold degeneracy arises because this combination can be taken four ways: $(0_0+1_{+1}, 0_0+1_{-1}, 1_{+1})$ $+0_0$, $1_{-1}+0_0$). In the lowest A+E states, the internal rotation angular momentum (j=1, in this example) is projected onto the C_3 axis of just one of the ammonia monomers. (More classically, we might say that the E monomer is rotating about its symmetry axis, but the A monomer is not rotating at all.) The relative energy of this lowest A + Ebasis function with respect to the lowest A + A basis function is equal to the rotational energy of the free 1_1 ammonia monomer rotational state. In G_{36} , there are two A + Etunneling states for each C_s "vibrational" state of the complex. In the high barrier tunneling limit, it can be seen from the work of Coudert and Hougen,³⁵ that the energy separation of these two A + E tunneling states (in the absence of interchange tunneling) is equal to the difference in the hindered internal rotation frequencies of the inequivalent monomers. That is, the splitting between one A + E state and the A + A states is equal to the hindered internal rotation frequency of the donor and the splitting between the other A+E state and the A+A states is equal to the hindered internal rotation frequency of the acceptor. It is straightforward to generalize to G_{144} combinations. Both the symmetric and antisymmetric inversion levels of an ammonia monomer E rotational state have nonzero statistical weight. The combinations of one monomer in 0_0^+ (which has zero statistical weight) with the other in 1^+_1 or l_1^- transform like G_1^+ or G_1^- in G_{144} , and both of these symmetries of course have zero statistical weight. The combinations of one monomer in 0_0^- with the other in 1_1^+ or l_1^- transform like G_2^+ or G_2^- in G_{144} , and both have nonzero statistical weight and thus are observable.

The E_1 , E_2 , E_3 , and E_4 states of G_{36} , and the corresponding G_3^+ , G_3^- , G_4^+ , and G_4^- states of G_{144} , are constructed from combinations of two ammonia monomers in E states. The transformation properties of the corresponding basis functions are more complicated. For example, the lowest-energy E + E combination, formed from two monomers in 1_1 states, transforms like $2E_1 \oplus E_2 \oplus 3E_3$ in G_{36} .

The doubly degenerate combination of $(1_{+1}+1_{-1}, 1_{-1})$ $+1_{\pm 1}$) transforms as E_1 when the total internal angular momentum (j) is 0 or 2, while it transforms as E_2 for j=1. The doubly degenerate combination of $(1_{+1}+1_{+1}, 1_{-1})$ $+1_{-1}$) transforms like E_3 for all possible total internal angular momenta i=0, 1, or 2. In the E+E states, the internal rotation angular momentum comes from the vector sum of the angular momentum from both monomers and can have a nonzero projection onto both monomer C_3 symmetry axes. Although we may expect the E_1 and E_2 tunneling sublevels to behave perhaps differently from the E_3 and E_4 sublevels, we see that they will still have roughly the same energy in the free internal rotor limit. The relative energy of the lowest E+E basis functions with respect to the lowest A + A function is equal to twice the rotational energy of the 1₁ rotational energy of the free monomer. In the high barrier treatment of Hougen and Coudert,³⁵ the splitting between the $G_{36} E + E$ states and the A + A states is equal to the hindered internal rotation frequency of the donor plus the hindered internal rotation frequency of the acceptor. An additional splitting, which we expect to be small, between the (E_1, E_2) and (E_3, E_4) pairs can be related to the difference in frequencies of an internal rotation tunneling motion in which both monomers rotate together clockwise or counterclockwise about their C_3 symmetry axes and an internal rotation tunneling motion in which both monomers rotate together but in opposite senses.

Summarizing the above observations, we expect the internal rotor levels of the ammonia dimer to be separated into three equally spaced clumps: the lowest group would be the A + A states, next, the A + E states, and the highest group would be the E+E states. If there were no barrier to internal rotation, the spacing between these groups would be equal to the 1₁ rotational energy of the ammonia monomer. As the barriers to internal rotation become more appreciable, these spacings decrease to the hindered internal rotation frequencies of the complexed monomers, and the internal rotor tunneling states within the A+E and E+Egroups start to split apart.

In their group-theoretical analysis, Nelson and Klemperer¹⁰ divided the correlation of rigid molecule rotational levels with the tunneling-rotational levels of the ammonia dimer into two steps, considering two ways to arrive at G_{36} from C_s . One way, called the $2C_3 + I$ limit, assumes that the energy splittings caused by the internal rotation of the ammonia monomers about their C_3 symmetry axes are much larger that the splittings caused by the donoracceptor interchange tunneling. The pattern of ammonia dimer tunneling energy levels has previously been assumed to resemble this limit.^{9,21} In the $2C_3 + I$ limit, we would expect to find that the A_1 and A_4 (A+A), E_1 and E_2 (E+E), and the E_3 and E_4 (E+E) pairs of tunneling sublevels are each split by small interchange tunneling frequencies. The pair of G(A+E) sublevels can be split by a contribution from the internal rotation tunneling (the difference in internal rotation frequencies of the two inequivalent monomers) as well as the small interchange tunneling frequency. The pure rotational transitions discovered

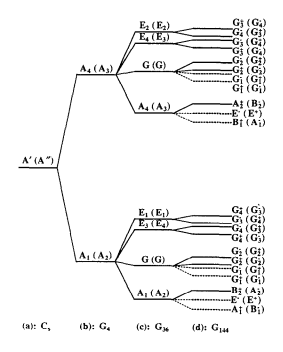


FIG. 5. Correlation diagram for the tunneling levels of the ammonia dimer. Proceeding from the left-hand column, (a) one rotational state of the rigid complex with a plane of symmetry is spit into rotation tunneling sublevels as (b) the "donor-acceptor" interchange, (c) C_3 internal rotation, and (d) monomer umbrella inversion tunneling motions become feasible. The symmetry species of each rotation tunneling state in the appropriate molecular symmetry group for each column is indicated for even J(odd J). See the text for the complete explanation.

by the Klemperer group were assigned to these G sublevels. Our microwave search was initially undertaken in order to measure the rotation-interchange tunneling subbands in the A+A and E+E tunneling sublevels.

The opposite way of arriving at G_{36} from C_s , called the $I+2C_3$ limit,¹⁰ assumes that the interchange tunneling is much faster that the internal rotation. This turns out to be a better description of the now vastly expanded experimental results, and we illustrate this qualitative scheme in Fig. 5. As in Nelson and Klemperer's analysis,¹⁰ we discuss each column in this diagram, proceeding from left to right. In column (a) we show one rotational level of a rigid ammonia dimer with an equilibrium structure that contains a plane of symmetry. The appropriate molecular symmetry group in this case is C_s . In column (b) the donoracceptor interchange tunneling splits this rotational state into a symmetric tunneling sublevel and an antisymmetric tunneling sublevel. This new symmetry operation doubles the size of the molecular symmetry group, so that it is now G_4 . In column (c) the internal rotation of each monomer about its C_3 symmetry axis causes an additional ninefold splitting of each symmetric or antisymmetric interchange tunneling sublevels into four distinct tunneling sublevels: an A+A, an A+E, and two E+E states. These internal rotation sublevels are sketched assuming the energy ordering as described above. The internal rotation symmetry operations increase the size of the molecular symmetry group by a factor of $3 \times 3 = 9$ times, to give G_{36} . Finally, in column (d), the umbrella inversion of both monomers becomes feasible, and increases the size of the molecular symmetry group by a factor of $2 \times 2 = 4$ times to give G_{144} .

The tunneling levels that we have accessed spectroscopically are connected by electric dipole transitions into three groups. It is straightforward, using the relative nuclear-spin statistical weights observed in the closely spaced K-type doublets, as well as the presence of the previously observed pure rotational transitions in the A+Estates, to assign these groups to the A+A states, the A+Estates, and the E+E states. Additional confirmation of the A+A and E+E assignment is simply the fact that more E+E states were more observed than could possibly be expected for the A+A states alone.

1. The E+E states

The VRT levels identified as the E+E states are shown in Fig. 3(a). A total of 509 observed electric dipole transitions connect these levels.³¹ Table II(a) lists all of the observed subband origins for the transitions connecting these E+E states and serves as a quick summary of this subset of the spectrum. The fitted rotational constants, relative J=0 energies, and Coriolis coupling constants are shown in Table III(a), and the calculated lowest VRT energy levels (J=0-5) from this fit are listed in Table III(a). We discuss the most interesting features of this data set below.

The most important observation to make about the observed E+E states is that two of the four lower $K_a=0$ E+E states have the same symmetry and two of the four upper $K_a = 0 E + E$ states have the same symmetry, as we mentioned in our description of the Coriolis analysis above. We learned this by discovering four sets of $|\Delta K_a| = 1$ transitions between two $K_a = 0$ VRT states and a single $K_a = 1$ VRT state. (These are the $K_a = 0 \leftarrow 1$ subbands near 550 GHz, the $K_a = 0 \leftarrow 1$ subbands near 660 GHz, the $K_a = 1 \leftarrow 0$ subbands near 760 GHz, and the $K_a = 1 \leftarrow 0$ subbands near 1050 GHz.) This leads to two important conclusions, as illustrated in the abbreviated correlation diagram shown in Fig. 6. First, G_{144} is the appropriate molecular symmetry group for the ammonia dimer. The E+E states of G_{36} are E_1, E_2, E_3 , and E_4 , and there is only one tunneling sublevel of each symmetry for each vibrational level. However, in G_{144} , the E+E states are comprised of two G_3^+ , two G_3^- , two G_4^+ , and two G_4^- sublevels, which is consistent with our observations. Second, the interchange tunneling splitting must be large.

Consider the possible correlations of four $G_{144} E + E$ states to two $G_{36} E + E$ states. If, as in the $(2C_3+I)$ limit shown in Fig. 6(a), the two $G_{36} E + E$ states were interchange tunneling partners such as the pair E_1 and E_2 or the pair E_3 and E_4 , then the resulting G_{144} states would be G_3^+ , G_3^- , G_4^+ , and G_4^- ; these all have different symmetries. On the other hand, as in the $I+2C_3$ limit shown in Fig. 6(b), if the two $G_{36} E + E$ states were E_1 and E_3 , which are both symmetric with respect to interchange tunneling, then the resulting G_{144} states would be G_3^- , G_4^+ , G_4^+ , and G_4^- ; two of these have the same symmetry G_4^+ , which is consistent with our observation of two pairs of *nearby* $K_a=0$ states with the same symmetry. If the four lowest $K_a=0 E+E$

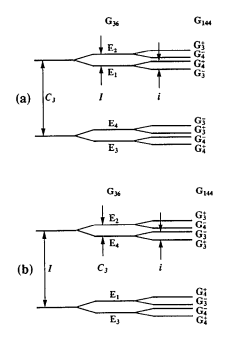


FIG. 6. (a) Correlation of G_{36} (no monomer umbrella inversion) E+E tunneling states in the $2C_3+I$ limit to the G_{144} (with monomer umbrella inversion) E+E tunneling states. In this case, the G_{144} states with the same symmetry are well separated. (b) Correlation of $G_{36} E+E$ tunneling states in the $I+2C_3$ limit to the $G_{144} E+E$ tunneling states. In this case, two pairs of G_{144} states with the same symmetry, the G_4^+ states and the G_3^+ states, lie close in energy. This tunneling scheme is consistent with our far-infrared spectra.

states are symmetric with respect to interchange tunneling, then four more $K_a=0$ E+E states which are their antisymmetric interchange partners should exist. We did not find these interchange partners for the lower four $K_a=0$ E+E states in our microwave search, nor have we assigned far-infrared subbands that might *originate* from the interchange partners of the lowest E+E states; this suggests that the interchange tunneling splitting in the E+E states is large. Later, we deduce that the antisymmetric E+Einterchange partners are actually the upper states of the $K_a=0+0$ subbands we measured near 580 GHz. This suggests that the $I+2C_3$ tunneling scheme described by Nelson and Klemperer¹⁰ is the most appropriate.

We place G_3 and G_4 labels on all the E + E VRT states using nuclear-spin statistics, but we must make an educated choice between two possible overall assignments of + and - labels. The two lower $K_a = 0 E + E$ states that have the same symmetry are either both G_4^+ or both G_4^- . So the choices for the four lower E+E $K_a=0$ states are (1) G_4^+ (from E_1), G_3^- (from E_1), G_4^+ (from E_3), and G_4^- (from E_3); or (2) G_4^- (from E_2), G_3^+ (from E_2), G_4^- (from E_3), and G_4^+ (from E_3). We prefer assignment (1) because all of these states are symmetric with respect to interchange tunneling. When interchange tunneling is large, as in the $I + 2C_3$ limit, the correlation diagram shows unambiguously that the E_1 and E_3 G_{36} states must be the lowest-energy E+E states. In clusters such as $(HCl)_2$ and (HF)₂, in which interchange is the only possible tunneling motion, the symmetric interchange tunneling state is always lower in energy.

The pattern of $K_a = 1$ E + E tunneling states looks rather different from that in $K_a=0$. In addition, the differing $K_a = 1$ behavior of these states further subdivides them into two groups. The $K_a = 1$ VT manifolds labeled in Fig. 3 as (5, 6), (11, 12), (17, 18), and (23, 24) are spaced apart by 367, 87, and 678 GHz. But, as shown in Table II(a), the splitting between the two levels with the same J within each of these four manifolds, which is reflected in our rotational constants qB_n , is essentially zero to within our experimental resolution, or very small. If there were no Coriolis interactions involving these states, the qB_n values would be exactly zero. By contrast, the $K_a = 1$ manifolds labeled in Fig. 3 as (7,8), (9,10), (19,20), and (21,22) are spaced apart by 48 MHz, 731 GHz, and 9 MHz. The values of qB_n for these levels, listed in Table II(b), are close to the values of the asymmetry parameter $\frac{1}{4}$ (B-C) that can be predicted from the proposed experimental structure (21) MHz) or the proposed theoretical structure (15 MHz). The effects of Coriolis interactions have already been removed from these values of $qB_{7,8}$ and $qB_{9,10}$ in Table III(b). The two qualitatively different observed Coriolis interactions of these $K_a = 1$ E + E states show how they correlate to the two different groups of $G_{36} E + E$ states.

First, let us consider the very-lowest-energy $K_a = 1 E$ +E tunneling manifold (5,6). For this level, we found that $qB_{5.6}$ is zero. The J=1 $K_a=1$ states of this manifold have symmetry species G_3^- (5) and G_4^+ (6), and they are shown in Fig. 4. The $G_3^- 1_1$ (5) state is pushed down in energy by a Coriolis interaction with the $G_3^- l_0$ states (1 and 4) that are ~78 GHz higher in energy and the G_4^+ 1₁ state (6) is pushed down in energy by a Coriolis interaction with the G_4^+ 1₀ state (3) that is also ~78 GHz higher in energy. This is why the observed splitting of the l_1 states is so small. We might think of a pair of two 1_1 rotational wave functions as $2^{-1/2}(|K_a=+1\rangle+|K_a=-1\rangle)$ and $2^{-1/2}(|K_a=+1\rangle-|K_a=-1\rangle)$, of which only the "+" linear combination can interact with $K_a=0$ states. But the observed Coriolis interactions show that this does not describe the $K_a = 1$ states (5,6). Rather, the + rotational states (or even simpler rotational states such as $|K_a = \pm 1\rangle$) of two different tunneling states are present in this $K_a = 1$ pair. Since the two 1_1 states have alternating intensities (due to different nuclear-spin statistical weights), their symmetries must be G_4^+ and G_3^- , and the two states are the symmetric and antisymmetric umbrella inversion tunneling components of a $G_{36} E_1$ state. That is, the umbrella inversion tunneling splitting of the G_{36} E_1 sublevel is exactly zero in $K_a = 1$. The 367 GHz splitting of the (5,6) and (11,12) manifolds is not due to umbrella inversion tunneling; instead this large energy separation may be the result of coupling of a large internal angular momentum to the a inertial axis. This effect is also found in the $K_a = 1 E$ states of Ar-NH₃.^{29,36,37}

Now, let us consider the example of the $K_a=1 \ E+E$ tunneling manifold (9,10). We found that $qB_{9,10}$ is ~25 MHz, and that the separation between the (7,8) and (9,10) manifolds is only ~48 MHz. As shown in Fig. 4, the $J=1 \ K_a=1$ states in this manifold also have symmetry species G_4^+ (9) and G_3^- (10). The G_3^- 1 state (10) is

pushed up in energy by a Coriolis interaction with the $G_3^ 1_0$ states (1 and 4) ~ 30 GHz lower in energy, but the G_4^+ 1_1 state (9) does not interact with the G_4^+ 1_0 state (3) also \sim 30 GHz lower in energy. This leads to the very large observed value of $qB_{9,10}$. The observed Coriolis interactions confirm that the G_3^- state (10) can be thought of as a $[2^{-1/2}(|K_a=+1)]$ symmetric rotational state $+|K_a=-1\rangle$)] because it can interact with the $K_a=0$ state and that the G_4^+ state (9) can be thought of an antisymmetric rotational state $[2^{-1/2}(|K_a=+1\rangle - |K_a=-1\rangle)]$. For even J, the symmetric rotational states of manifold (9,10) have symmetry G_4^+ and the symmetric rotational states of manifold (7,8) have symmetry G_4^- . Therefore the $K_a = 1$ manifolds (7,8) and (9,10) are the symmetric and antisymmetric umbrella inversion tunneling components of a G_{36} E_3 tunneling state. And, the very small ~48 MHz energy difference between them is, in fact, the $K_a = 1$ umbrella inversion tunneling frequency.

We find that the electric dipole selection rules in G_{36} can be viewed as very approximate additional selection rules for transitions between the G_{144} VRT states. This means, for example, that we would expect to see $G_3^+(E_2) \leftarrow G_3^-(E_1)$ but not $G_3^+(E_4) \leftarrow G_3^-(E_1)$. We also find, however, that the Coriolis coupling between $K_a=0$ and $K_a = 1$ levels almost completely spoils the approximate G_{36} selection rules. That is, we observe subbands that would be forbidden by the G_{36} selection rules when either the lower or upper state is affected by strong Coriolis interactions. These extra observed subbands allow us to determine, for example, the energy difference between the G_{36} E_1 level and the $G_{36} E_3$ level. The observed transitions that defy the G_{36} selection rules are often weaker than similar transitions that obey the G_{36} selection rules. We use these observations to assign the G_{36} symmetries of the $K_a=0$ E+E states, since we know the G_{36} symmetries of the $K_a = 1$ states. We indicate both the G_{144} and G_{36} symmetries in Fig. 3(a) and in Ref. 31 and Tables II(a) and IV(a).

The tunneling rotation subband that we discovered in the microwave search also involves the E+E states. The observed transitions in this subband are indicated in Fig. 2(b). This $K_a = 1 \leftarrow 1$ subband with a band origin at 48 MHz, between states (7,8) and (9,10), directly measures the umbrella inversion tunneling splitting of the $K_a = 1 G_{36}$ E_3 level. Although these transitions obey the approximate G_{36} selection rules as well as the rigorous G_{144} selection rules, we found that they are approximately 2 orders of magnitude weaker than the strong pure rotational transitions found in the A+E states. The measurement of these relative intensities is possible because our microwave lines are measured by direct absorption. The relative intensity of our E + E microwave subband may be small partly because the population of the lower state cannot be fully relaxed and also partly because the E+E nuclear-spin statistical weights are smaller. We also note, however, that if K_{a} and K_c quantum numbers are assigned to the apparent "asymmetry" doublets (as usual, with 111 lower in energy than l_{10} , etc.), the observed microwave transitions do not strictly correspond to an a-, b-, or c-type subband. Rather,

the expected rotational selection rules for the Q and R branches of an *a*-type subband are reversed. The observed hyperfine pattern of these transitions is 2–3 MHz wide. When a suitable formalism in G_{144} is derived to interpret the quadrupole hyperfine structure in the ammonia dimer, these well-resolved triplets will undoubtedly provide important structural information.

2. The A+E states

The ammonia dimer A+E VRT levels are shown in Fig. 3(b). A total of 392 observed electric dipole transitions connect these states (see Ref. 31). Table II(b) summarizes all the subband origins for the observed transitions connecting the A+E states. The fitted rotational constants, relative J=0 energies, and Coriolis coupling constants are shown in Table III(b), and the calculated lowest VRT energy levels (J=0-5) from the fit are listed in Table IV(b).

There are two G_1^+ , two G_1^- , two G_2^+ , and two $G_2^$ tunneling sublevels in G_{144} . Pure rotational transitions that do not involve a change in tunneling states are allowed in all of these levels. However, for $({}^{14}NH_3)_2$, only the G_2^+ and G_2^- VRT states have nonzero statistical weight. Before obtaining our new results on the E+E states, we believed the analyses of Nelson et al.^{8,9} and Havenith et al.²¹ that the monomer umbrella inversion tunneling was quenched (i.e., that G_{36} was the appropriate molecular symmetry group), that the interchange tunneling splitting of the A + E states was very small, and that the observed far-infrared transitions involved an A" intermolecular vibration. Thus, consideration of the observed A + E states alone would not unambiguously reveal that the correct molecular symmetry group for the ammonia dimer is actually G_{144} . As in the E+E states, we must again choose between two possible overall assignments of + and - labels for the G_2 states in G_{144} . We will arbitrarily assume, throughout this paper, that the lowest-energy $K_a = 0 A + E$ state that we observed has G_2^+ symmetry.

Four sets of pure rotational transitions are expected for the $G_{144} A + E$ states, in $K_a = 0$. Nelson and Klemperer^{8,9} reported only two sets of pure rotational transitions. From far-infrared transitions that we observed, we can deduce that if the state Nelson and Klemperer called G_{α} actually has G_2^+ symmetry, then the state they called G_β really has G_2^- symmetry. These are the two umbrella inversion components of just one of the two G_{36} G states, and they are split 3.3 GHz apart. This partially quenched umbrella inversion frequency is about seven times smaller than that in a free ammonia monomer (~ 23.8 GHz).³⁰ Since they are the lowest-energy A + E states that we observed, they most probably correlate to the lower of the two G_{36} G states in the ground vibrational state of the ammonia dimer. The other allowed pair of pure rotational transitions in the next-highest G_2 states that would correlate to the upper of the two G_{36} G states, was not observed either by us or by Nelson et al.^{8,9} In our microwave search, we were able to observe the pure rotational transitions in the $K_a = 1$ manifold labeled (5,6) in Fig. 2, as well as very, very weak pure rotational transitions in the $K_a = 2$ manifold labeled (7,8).

These last transitions seemed weak because the (7,8) states, ~500 GHz in energy above the lowest A + E states, should not be well populated in a cold supersonic expansion. However, we could not observe the pure rotational transitions in the $K_a = 1$ manifold (3,4) which is only ~100 GHz above the lowest A + E states. These observations lead us to believe that the other two G_2 states (the upper of the two G_{36} G states in the ground vibrational state) are either located more than 500 GHz in energy above the lowest G_2 states or they have no dipole moment to provide intensity for the pure rotational transitions. We observed very strong $\Delta K_a = 0$ far-infrared transitions out of all the lower A + E states shown in Fig. 3(b), regardless of whether or not we could observe their pure rotational progressions in the microwave region, and the relative farinfrared intensities seemed to depend only on the relative populations in our supersonic expansion. Therefore, we assume that the population argument applies to the unobserved G_2 microwave transitions. Later, we deduce that the two G_2 states in question were found earlier in the farinfrared study by Havenith et al.²¹ near 614 and 617 GHz.

The behavior of the $K_a = 1 A + E$ states is similar to the E+E states that correlate to the $G_{36} E_1$ and E_2 tunneling sublevels. The energy differences between the four $K_a = 1$ A+E manifolds are much larger than those in $K_a=0$. In each $K_a = 1 A + E$ manifold, the observed splitting of levels with the same J is very small, as is evident in the fitted values of qB_n in Table III(b). Coriolis interactions between $K_a=0$ and $K_a=1$ states cause all the observed values of qB_n to be nonzero. However, the analysis presented here explicitly treats only the strongest Coriolis coupling which occurs ~ 600 GHz above the lowest A + E states. As listed in Table IV(b), the symmetries of the $J=1, K_a=1$ states in the affected manifold are G_2^+ (10) and G_2^- (9). These l_1 levels are both pushed down by the G_2^+ (12) and G_2^- (11) 1_0 states ~17 and ~15 GHz higher in energy. Again, this shows that both members of the $K_a = 1$ pair are symmetric rotational states, $2^{-1/2}(|K_a=+1\rangle+|K_a=-1\rangle)$, that can interact with $K_a = 0$ rotational states. Also, one state in the pair must be symmetric with respect to umbrella inversion while the other state must be antisymmetric, because the $K_a = 0$ states they interact with are symmetric and antisymmetric umbrella inversion tunneling states. That is, the umbrella inversion frequency in the $K_a = 1 A + E$ states is exactly zero. Furthermore, the observed Coriolis interaction between the $K_a = 1$ manifold (9,10) and the $K_a = 2$ manifold (7,8), we find that the umbrella inversion frequency in the $K_a = 2 A + E$ states is also exactly zero.

At this point it becomes possible to assign the two infrared bands observed by Fraser *et al.*³⁸ near 980 cm⁻¹ in double resonance with the lowest G_2 states. In their modified molecular-beam electric resonance experiment, they measured the strength of the pure rotational $J=1 \leftarrow 0$ transitions in the lowest G_2^+ (1) and G_2^- (2) tunneling levels while stepping an overlapping line-tunable CO₂ laser between 914 and 987 cm⁻¹. When monitoring the G_2^+ microwave transition, they observed an infrared peak near 977 cm⁻¹, and when monitoring the G_2^- microwave transition, they observed an infrared peak near 981 cm⁻¹. This approximate 4 cm⁻¹ splitting very likely corresponds to the umbrella inversion tunneling splitting in the upper state. This umbrella inversion frequency is approximately 9 times smaller than that in a free ammonia monomer in $v_2=1$ (~ 36.5 cm⁻¹),³⁰ so this result is consistent with our work on the ground state of the ammonia dimer in that both for the ground state and the $v_2=1$ state of ammonia the complexed umbrella inversion tunneling frequency is about an order of magnitude smaller than the freeumbrella inversion frequency.

3. The A+A states

The four A+A states we have observed are shown in Fig. 3(c). The 89 observed electric dipole transitions that connect these levels are available from PAPS.³¹ The band origins of the three observed subbands are given in Table II(c), the fitted rotational constants and relative J=0 energies are shown in Table III(c), and the calculated VRT energy levels (J=0-5) from the fit are listed in Table IV(c).

Only one-third of the $G_{144} A + A$ VRT states have nonzero statistical weight for $({}^{14}NH_3)_2$. It is not possible to distinguish between G_{36} and G_{144} by considering the A + Astates alone. The populated A + A tunneling sublevels in G_{144} are B_2^+ and A_2^+ . The only difference between these and the $G_{36} A_1$ and A_4 states is that the G_{144} states might be slightly shifted by the small umbrella inversion frequency, which we cannot measure. The observed statistical weights show that the lower A + A states have B_2^+ tunneling symmetry, which means that they are symmetric interchange tunneling states, while the upper A + A states have A_2^+ tunneling symmetry and are thus antisymmetric interchange tunneling states.

No Coriolis perturbations are observed among the A +A states. Yet, the $K_a = 1 A + A$ states still have nonzero qB_n values. Let us assume this $K_a = 1$ splitting is caused by structural asymmetry, so that we may extend the analysis of Havenith et al.²¹ to extract two pieces of information from the A + A states that we have observed. First, we can confirm that the ammonia dimer molecular plane of symmetry contains the **a** and **c** inertial axes, as we may anticipate by computing the moments of inertia of either the experimental or theoretical structure of the ammonia dimer. Second, we can confirm that the interchange tunneling path is the "trans" path described by Havenith et al. ²¹ [by analogy to (HF)₂, ¹²], which has the cyclic C_{2h} transition state predicted by the ab initio potential-energy surfaces. The other path considered by Havenith et al.²¹ is a "cis" path [with a $C_{2\nu}$ transition state, also by analogy to (HF)₂]; cis interchange tunneling results in a different equivalent rotation of the ammonia dimer complex than trans interchange tunneling, and thus different symmetries of the $K_a = 1$ asymmetry doublets.

To illustrate this, we first assign K_a and K_c quantum numbers to the observed A + A states. Since there is no way to rotate and displace two ammonia monomers and arrive at a structure in which the **b** and **c** inertial axes are switched with respect to the starting configuration, we will assume that the **b** and **c** axes are the same in the upper and

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lower states of the far-infrared transitions. Thus we find, for example, that the 1_{11} state in the $K_a = 1 A + A$ manifold (2,3) has B_2^+ symmetry, and the 1_{10} state in this manifold has A_2^- symmetry. Recall that the 0_{00} state in the $K_a=0$ A+A manifold (1) has B_2^+ symmetry and that the l_{01} state in the same manifold has A_2^- symmetry. Next, we extend Table II of Havenith et al.²¹ (which lists the symmetry species of the rotational wave functions in G_{36} , assuming that the a and b inertial axes lie in the plane of symmetry, for the cases of *trans* interchange tunneling only and *cis* interchange tunneling only) to the G_{144} molecular symmetry group and to include the possibility that the a and c inertial axes might lie in the plane of symmetry. In this way, we find that the symmetries of the rotational states that we stated above are only consistent having the a and c inertial axes in the plane of symmetry and interchange proceeding along a trans path.

C. Vibrational assignment

Several ab initio workers^{16,18,39} have calculated harmonic frequencies for the intermolecular "normal" vibrations of the ammonia dimer. These modes are classified in the C_s molecular symmetry group, but the grouptheoretical correlation tables presented earlier in this paper can be used to enumerate all the G_{144} tunneling sublevels that correlate to each of these C_s rovibrational states. The specific G_{144} symmetry of each vibration can be multiplied by the symmetries of the ground-state tunneling sublevels as an initial estimate of the ordering of the tunneling sublevels in the excited vibrational state. We have argued above that all of the lower states of the far-infrared subbands are symmetric with respect to the donor-acceptor interchange tunneling and all of the upper states are correspondingly antisymmetric. We now address the following question: Do the far-infrared transitions connect the symmetric and antisymmetric interchange levels within the ground vibrational state of the ammonia dimer complex, or does the upper state correspond to a low-frequency intermolecular vibration?

We have no data that rule out the first possibility. Hassett, Marsden, and Smith^{20} and $\text{Frisch et al.}^{16}$ calculate a low barrier to interchange tunneling. That is, they find that the cyclic transition state is only 0.1–0.2 kcal/mol above their calculated equilibrium structure. In addition, they find that the harmonic zero-point energy of the vibration in the same coordinate as the lowest-energy path for the interchange tunneling motion is approximately equal to the barrier height. Our observation of a large interchange tunneling splitting is consistent with this feature of high-quality theoretical surfaces.

Frisch *et al.*,¹⁶ Andrews and Dykstra,¹⁸ and Hassett, Marsden, and Smith³⁹ all predict a very-low frequency torsional mode with a harmonic frequency at 20, 27, and 35 cm⁻¹, respectively. All three groups give the C_s symmetry of this mode as A''. Assuming that the intermolecular vibrations of the ammonia dimer have one dimensional representations in G_{144} , the G_{144} symmetry of this torsional mode is A_1^- , B_1^- , A_2^- , or B_2^- . Our observations rule out each of these possibilities. The upper state of our far-

infrared transitions have a vibrational symmetry of either A_1^+, B_1^+, A_2^+ , or B_2^+ and correlates to a C_s mode with A' symmetry. That is, the $K_a = 0 \leftarrow 0$ transitions in the A + Astates can only be observed if the vibrational symmetry has a + label, the fact the two close $K_a = 0 E + E$ tunneling levels that have the same symmetry have + labels in both the lower and upper states also requires that the vibrational symmetry have a + label, and the fact that we always observe the $K_a=0$ G_2^+ level below the G_2^- level also suggests that the vibrational symmetry has a + label. The lowest-energy A' mode predicted by the various theoretical workers lies at 121, 81, and 72 cm^{-1} , respectively, and is the vibration along the interchange tunneling coordinate we mentioned above. We do not believe that the assignment of our spectra to this vibration is reasonable since our transitions occur at a factor of 4 lower in energy, and because we have not assigned any subbands that might originate from the antisymmetric interchange levels of the ground state in the event that we were observing this vibrational band.

IV. DISCUSSION

In this paper, we have presented extensive new information on the VRT levels of the ammonia dimer. The characterization of all the ground-state tunnelingrotational energy levels in $K_a=0$ and $K_a=1$ is complete. We have definitively assigned all previously reported farinfrared subbands^{21,23} (the new assignments are noted in Table II), and indeed, the information provided by any new measurements of far-infrared subbands within the ground state of the ammonia dimer will be redundant with the results presented here. There should be a few strong far-infrared subbands at lower frequency from where we scanned: in the A + A states, there should be a $K_a = 0 \leftarrow 1$ subband at 272.8 GHz [(4) \leftarrow (2,3)]; in the A+E states, there should be $K_a = 0 \leftarrow 1$ subbands at 382.6 and 385.0 GHz [(11) \leftarrow (5,6) and (12) \leftarrow (5,6)]; and in the E+Estates, there should be $K_a = 1 \leftarrow 0$ subbands at 373.6, 374.6, and 376.0 GHz (this last one might be weak) $[(17,18) \leftarrow (4), (17,18) \leftarrow (3), \text{ and } (17,18) \leftarrow (1)]$. The frequencies of the lowest J transitions in these subbands can be calculated to within 1 MHz from Table IV. The last five subbands involve upper or lower states that are perturbed by Coriolis interactions, and in $\Delta K_a = \pm 1$ subbands involving these perturbed states we have noticed intensity perturbations that made the R branch particularly strong and the P branch very weak (or vice versa).

Exploiting our new definitive characterization of the ground vibrational state of the ammonia dimer, we are able to improve on the interpretation of the infrared spectrum of the ammonia dimer measured in the monomer $v_2=1$ region by Huisken and Pertsch⁴⁰ and by Snels *et al.*⁴¹ Since line-tunable CO₂ lasers were used in these experiments, and since the upper state of these infrared spectra dissociates rather quickly, rotational resolution of these bands was not possible. The main features found in these spectra are a broader, more intense band, with obvious, but still unresolved structure near 979 cm⁻¹, and a sharper single band near 1004 cm⁻¹. These two bands were interpreted as

excitations of the two inequivalent ammonia subunits within the complex. However, we now know that it is not possible to excite either the donor or the acceptor by itself when donor-acceptor interchange tunneling is occurring within the complex. van Bladel et al. show in the accompanying paper that only the A+E states are "localized" with respect to interchange tunneling. Thus, it may be useful to think of this entire spectrum as components of a single transition from the ground vibrational state, that we have characterized here, to a single excited state. In the $K_a = 0 \leftarrow 0$ portion of this spectrum, four subbands involving the A+E states should be observable: two from the lower A+E states found by Nelson et al.^{8,9} to the lower A+E states in the excited vibrational states, and two from the lower A+E states to the upper A+E states in the excited vibrational state. Both pairs are expected to be strong transitions because the pure rotational transitions in the lower A + E states in the ground state and the transitions across the A+E tunneling splitting in the ground state are both strong. In their infrared-microwave doubleresonance experiment, Fraser et al.³⁸ identified one pair of A+E transitions under the 979 cm⁻¹ peak, but did not scan under the 1004 cm⁻¹ peak, because the normal isotope CO₂ laser that they were using does not provide that frequency. We believe that the second pair of A + E transitions lies under the 1004 cm⁻¹ infrared peak. This ~ 25 cm⁻¹ internal rotation and interchange tunneling splitting of the A+E states in $v_2=1$ is not very different from the value that we have measured in the ground state.

It had previously been thought that the anisotropy of the intermolecular potential completely "quenched" the umbrella inversion of the ammonia monomers, but our results clearly show that this motion still occurs in the ammonia dimer complex. This leads to the molecular symmetry group G_{144} , rather than the group G_{36} , which has guided earlier analyses. In this work, we have identified $K_a = 0$ umbrella inversion tunneling splittings of 3.3094(3) GHz and 2.3920(6) GHz in the "lower" and "upper" A +E states, respectively. In the E+E states, the umbrella inversion tunneling splittings are 1.094(1) GHz (splitting of the $G_{36} E_3$ level), 1.103(1) GHz (E_1), 2.217(1) GHz (E_4) , and 2.217(1) GHz (E_2) . These frequencies, are about 10 times less than the inversion tunneling splitting of \sim 23.8 GHz (Ref. 30) in the ground state of the free ammonia monomer. Thus, it is indeed true that monomer umbrella inversion tunneling is partially quenched in the ammonia dimer; it occurs on essentially the same time scale as end-over-end rotation in the low-J, $K_a=0$ states.

In Ar-NH₃, the umbrella inversion of the monomer is nearly free in the observed $|\Omega| = 0$ states, but completely quenched in the observed $|\Omega| = 1$ states, in which the ammonia lone pair tends to point either towards or away from the argon atom.^{36,37} However, the two $|\Omega| = 1 E$ states of Ar-NH₃ that correlate to the j=1, $|k_c|=1$ states of the ammonia monomer are split apart by 200 GHz.²⁹ This splitting would occur in the absence of monomer inversion and it arises largely because there is an energy difference between the localized ammonia-lone-pair-towards-argon and ammonia-hydrogens-towards-argon configura-

tions.^{36,37} These $|\Omega| = 1$ Ar-NH₃ E states show no true asymmetry doubling. However, small splittings (that look like "asymmetry doubling") are observed because each initially degenerate component of the $|\Omega| = 1 E$ states is Coriolis coupled to a different $|\Omega| = 0$ state, and these two nearby $|\Omega| = 0$ states are split ~22.7 GHz apart by monomer umbrella inversion.²⁹ In the ammonia dimer, the K_a =1 A + E states and the $K_a = 1 E + E$ states that correlate to the $G_{36}E_1$ and E_2 states behave like the $|\Omega| = 1 E$ states of Ar-NH₃. The umbrella inversion in all of these ammonia dimer $K_a = 1$ states is quenched and they show no true asymmetry doubling, as explained in the preceding sections. The $K_a = 1$ separations of these states (large energy separations which are not caused by monomer inversion) are 118.0887(7) and 378.4596(6) GHz in the lower and upper A+E states, respectively, and 366.694(1) and 678.199(1) GHz in the E+E states that correlate to the G_{36} states E_1 and E_2 , respectively. It is interesting to note that, by looking at the G_{144} symmetry labels alone, we cannot tell the difference between states that are members of an asymmetry doublet and states that are members of umbrella inversion doublets in the ammonia dimer A+Estates, in the ammonia dimer E+E states that correlate to the G_{36} E_1 and E_2 states, or in the Ar-NH₃ E states. By contrast, the ammonia dimer $K_a = 1$ states that correlate to G_{36} E_3 and E_4 states show both real asymmetry doubling and monomer umbrella inversion tunneling splittings. In these states, the G_{144} symmetry labels do distinguish the asymmetry doublets from the umbrella inversion tunneling doublets. For example, the asymmetry doublets in one umbrella inversion state are G_4^+ (E_3) and G_3^- (E_4), while in the other umbrella inversion state they are G_4^- (E₃) and G_3^+ (E₄). In these $K_a = 1$ states we have measured the very small umbrella inversion tunneling splittings of 48(1) MHz of the lower E_3 state (G_{36} tunneling symmetry) and 9(1) MHz of the upper E_4 state (G_{36} tunneling symmetry).

In the ammonia dimer, we have identified donoracceptor interchange tunneling frequencies of 483.2995(4) GHz $(K_a=0)$ and 518.5007(6) GHz $(K_a=1)$ in the A+Astates, 578.4255(9) GHz ($K_a=0$) and 730.8030(7) GHz $(K_a=1)$ between the E+E states that correlate to the G_{36} E_3 and E_4 states, and 580.8738(7) GHz ($K_a=0$) and 609.3528(7) GHz ($K_a=1$) between the E+E states that correlate to the $G_{36} E_1$ and E_2 states. The corresponding measured frequencies in the A+E states are 614.1768(3) GHz $(K_a=0)$ and 616.9779(4) GHz $(K_a=1)$, but this splitting is due to a difference in donor and acceptor internal rotation frequencies that has been increased due to interchange tunneling (i.e., interchange has a second-order effect on this splitting of the A + E states). Because of their initial energy separation caused by internal rotation, the interchange tunneling does not mix the lower and upper A+E states very much, so that they cannot be called symmetric or antisymmetric with respect to interchange tunneling. This is clearly illustrated in the lowest A + E wave functions by van Bladel et al.²² in the accompanying paper. Let us assume the upper limit of 400 GHz for the initial internal rotation separation of these A+E states, which

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corresponds to free internal rotation of the acceptor about its C_3 axis and no internal rotation of the donor. Let us further assume that interchange tunneling mixes only the lower and upper A+E states that belong to the ground vibrational state of the complex; this is probably the most important effect of interchange feasibility on the A+Estates. Then, we can solve for the off-diagonal "interchange coupling" of the A+E states in this two-state approximation. We find that the interchange matrix element is ~ 230 GHz for the A + E states. Thus, if there were initially zero internal rotor splitting (rather that the 400 GHz we assumed) between these A + E states, interchange tunneling would split them by ~ 460 GHz. Since we assumed the upper limit for the initial internal rotor splitting, 460 GHz is a lower limit for the interchange tunneling frequency in the two-state approximation. The upper limit for the interchange tunneling frequency in the A+E states is the measured ~ 615 GHz splitting.

Donor-acceptor interchange tunneling frequencies have also been measured in other hydrogen-bonded homodimers, namely (HCl)₂, (HF)₂, and (H₂O)₂. For (H³⁵Cl)₂, the donor-acceptor interchange tunneling splitting has been measured at 463.979(2) GHz (Ref. 13) by tunable far-infrared laser spectroscopy, and the barrier to this motion has been empirically estimated to be 28.4 $cm^{-1.42}$ For (HF)₂, the donor-acceptor interchange tunneling splitting has been measured in the microwave at 19.94704(2) GHz, 43 and an empirical potential-energy surface puts this barrier at 302 cm^{-1.44} And for $(H_2O)_2$, the donor-acceptor interchange tunneling splitting has also been measured in the microwave at 19.526 73 MHz¹⁴ and the *ab initio* barrier to this motion is 304 cm^{-1} .⁴⁵ The interchange tunneling frequency in the ammonia dimer A +A states (483.2 GHz) is quite similar to the interchange tunneling frequency in (HCl)₂ (464.0 GHz). In both complexes, the interchange tunneling motion is predicted to occur via rotation of both monomers, each about an axis perpendicular to the plane of the dimer, so that we can compare the reduced masses for interchange tunneling by comparing the monomer rotational constants. For the ammonia monomer, $A = B \approx 9.9$ cm⁻¹ and for H^{35} Cl monomer, $B \approx 10.6 \text{ cm}^{-1}$;⁴⁶ these values only differ by ~7%. Recently, Hassett, Marsden, and Smith²⁰ have calculated a barrier height of 29 cm^{-1} for donor-acceptor interchange tunneling in the ammonia dimer. Since both the interchange tunneling frequencies and the reduced masses for this motion are similar for the ammonia dimer and the hydrogen chloride dimer, we believe that our measurements support the 29 cm^{-1} barrier predicted by Hassett, Marsden, and Smith, which is also similar to the empirical barrier of 28.4 cm⁻¹ deduced for (HCl)₂. By contrast, the rotational constant for (HF)₂ is ≈ 20.9 cm^{-1.46} If the barrier to interchange tunneling in the ammonia dimer were at all similar to the $\sim 300 \text{ cm}^{-1}$ barrier to interchange tunneling in the hydrogen fluoride dimer, the interchange frequency in the ammonia dimer should be less than the ~ 20 GHz interchange frequency in the hydrogen fluoride dimer. This is clearly far from being the case.

Some time ago, Frisch et al.¹⁶ also calculated the bar-

rier to interchange tunneling in the ammonia dimer, and arrived at a value of ~ 70 cm⁻¹. A similar value of 77 cm^{-1} is computed by van Bladel *et al.*²² for the site-site potential of Sagarik, Ahlrichs, and Brode.¹⁹ In the accompanying paper, van Bladel et al.²² show that a surface with a barrier of 167 cm^{-1} gives an interchange tunneling frequency that is three times lower than what is observed. Liu et al.¹⁸ have performed a high-level calculation of the electrostatic interaction surface for the ammonia dimer at fixed R_{CM} (including electric multipole-multipole interactions, induction, dispersion, but no short-range repulsion) and obtained an electrostatic contribution to this barrier of 250 cm^{-1} . This reveals that the *ab initio* values for this interchange barrier are the sums of large opposing contributions, so that this feature of the ammonia dimer potentialenergy surface is the most difficult to predict accurately.

The "local internal axis method (IAM)," a model developed by Hougen which assumes high barriers to tunneling, is a type of analysis that does not require knowledge of the entire potential-energy surface. This approach has been used quite fruitfully to characterize the tunneling motions and some of the energetics of (HF)₂ (Ref. 12) and $(H_2O)_2$ ¹⁵ In such a high barrier limit, Coudert and Hougen³⁵ have derived possible tunneling energy-level patterns for $(XY_3)_2$ molecules in the G_{36} molecular symmetry group. A general result of their work is that the interchange tunneling splitting of the A + A states must be larger than the interchange tunneling splitting of the E+Estates. This is because the splitting of the A + A states involves the sum of all the various contributions from different possible tunneling paths, while the splitting of the E+E states involves both sums and differences of the same contributions. In this work, we find that the E+E interchange frequency of the ammonia dimer is much larger than the A + A interchange frequency (by 100 GHz). This is more than can possibly be accounted for by the umbrella inversion tunneling not considered by Coudert and Hougen. The authors do mention that they suspect the high barrier analysis will not be appropriate for the ammonia dimer, and they are indeed correct.

Both Nelson et al.⁹ and Havenith et al.²¹ have argued that the internal rotation tunneling frequency is much faster than the interchange tunneling frequency in the ammonia dimer. That is, they believed that the $2C_3+I$ limit proposed by Nelson and Klemperer¹⁰ was an appropriate description of tunneling dynamics in this complex. Although nuclear-spin statistics prevent us from directly measuring the internal rotation tunneling frequency, we can use the measured rotational constants for the A + A, A+E, and E+E states to compare the internal rotation tunneling frequency to the other known tunneling frequencies. The $K_a = 0$ tunneling states that are separated by only umbrella inversion tunneling frequencies have nearly identical B_n rotational constants, which deviate from each other by less than 0.02%. [These are the A + E states (1) G_2^+ and (2) G_2^- , A + E states (11) G_2^+ and (12) G_2^- , E + Estates (1) G_4^+ (E_3) and (2) G_4^- (E_3), E+E states (3) $G_3^ (E_1)$ and (4) G_4^+ (E_1) , E+E states (13) G_3^+ (E_4) and (14) G_3^- (E₄), and E+E states (15) G_4^- (E₂) and (16)

 G_3^+ (E₂).] On the other hand, the $K_a=0$ tunneling states that are separated by the interchange tunneling frequency have B_n values that deviate from each other by 1%-2%. [These are the A + A states (1) B_2^+ and (4) A_2^+ , the A + Estates (1) and (11) both G_2^+ , the A + E states (2) and (12) both G_2^- , the E + E states (1) G_4^+ (E_3) and (13) G_3^+ (E_4), the E + E states (2) G_4^- (E_3) and (14) G_3^- (E_4), the E + Estates (3) G_3^- (E₁) and (15) G_4^- (E₂), and the E+E states (4) G_4^+ (E_1) and (16) G_3^+ (E_2).] The B_n rotational constants of the $K_a=0$ tunneling states that are separated by only the internal rotation tunneling frequency deviate from each other by 0.3%-0.4%. [These are taken as the average B_n values for the A+A, A+E, and E+E states that correlate to the lower (symmetric) interchange tunneling state and the average B_n values for the A + A, A + E, and E+E states that correlate to the upper (antisymmetric) interchange tunneling state.] When compared to the other two observed deviations in rotational constants, this implies that the internal rotation tunneling frequency is smaller that the interchange tunneling frequency, but larger than the umbrella inversion frequency; it is probably several wave numbers in magnitude. We emphasize, however, that the internal rotation frequency is not very small. The strong Coriolis coupling observed between manifolds with different K_a is evidence that the tunneling sublevels of the ammonia dimer do have some substantial internal angular momentum. The absolute upper limit for the internal rotation frequency would be 400 GHz or 13.3 cm⁻¹, the rotational energy of a free ammonia monomer in the 1, rotational state. However, even this upper limit is smaller than the observed donor-acceptor interchange tunneling frequencies. In other words, our observations lead us to conclude that the opposite $I + 2C_3 [+i]$ limit proposed by Nelson and Klemperer¹⁰ is the appropriate description of the tunneling motions in the ammonia dimer—not the $2C_3$ +I limit that has been assumed in previous analyses.

We can also compare the observed A rotational constants of the ammonia dimer to the A rotational constant predicted from the structure of Nelson *et al.*^{8,9} (115 GHz) and from a linearly hydrogen-bonded structure (105 GHz). The average energy difference between the $K_a=0$ and $K_a = 1$ levels ($\approx A$) of the A + A states is 228 GHz, for the A + E states it is 173 GHz, and for the E + E states it is 111 GHz. These values are closer to the predicted value for rotation of the entire ammonia dimer about its a inertial axis than to the value (400 GHz) for rotation of an ammonia monomer constituent about its c axis. By contrast, in the ammonia water complex, in which internal rotation of the ammonia monomer about its C_3 axis is nearly free, the structural value of A is ~150 GHz, but $K_a = 1 \leftarrow 0$ transitions in the ammonia water A states were measured near 620 GHz.³⁴ This comparison suggests that the internal rotation cannot be as free in the ammonia dimer as it is in the ammonia water complex. In the ammonia dimer, the internal rotation frequency must be smaller than the interchange frequency in order for the interchange tunneling motion to average out the effect of acceptor internal rotation on the A rotational constant.

Liu et al.¹⁸ have also calculated the electrical interac-

tion contributions to the internal rotation barriers in the ammonia dimer. They find, at a hydrogen-bonded configuration with $\theta_A = 12.8^{\circ}$ and $\theta_B = 102^{\circ}$ in Fig. 1, that the barrier to accepter internal rotation is $\approx 10 \text{ cm}^{-1}$, while the barrier to donor internal rotation is $\approx 650 \text{ cm}^{-1}$. At the cyclic transition state, with $\theta_A = \theta_B = 62.5^{\circ}$, the internal rotation barrier is $\approx 210 \text{ cm}^{-1}$. This suggests that there may be a considerable "interchange averaged" barrier to internal rotation in the ammonia dimer. However, these calculations did not include the possible effects of short-range forces, which could substantially change these barriers.

Finally, we turn to the structure of the ammonia dimer. In this paper, we have clearly demonstrated that a substantial amount of large-amplitude tunneling motion occurs on the time scale of our experiment. It makes little sense to speak of a rigid structure *per se* for the ammonia dimer, and it is particularly difficult to imagine the monomer inversion occurring in a rigid configuration. Thus, when we discuss the "structure" of the ammonia dimer, we refer to the minima of its intermolecular potential-energy surface, rather than to fixed positions of the constituent nuclei.

The structure proposed by Nelson et al.^{8,9} was based on the assumption that neither the donor-acceptor tunneling nor zero-point vibrational averaging affects the measured ¹⁴N quadrupole coupling constants in the two observed $K_a = 0$ A + E pure rotational progressions. In the accompanying paper, van Bladel et al.²² show that it is indeed true that these A + E wave functions are "localized" with respect to the interchange tunneling. This same qualitatively localized behavior is also characteristic of the water dimer E states, which are comprised of one ortho water monomer plus one para water monomer.¹⁴ This effect is unique to homodimers involving two monomers in different proton nuclear-spin states. By contrast, such "localization" is not expected for any of the tunneling states of $(HF)_2$ (Refs. 11 and 12) or $(HCl)_2$ —and, indeed, only one average ³⁵Cl or ³⁷Cl quadrupole coupling constant is measured in (H ³⁵Cl)₂ or (H ³⁷Cl)₂.¹³ Further examples are the A and B states of (H₂O)₂,¹⁴ and the A+A and E+E states of (NH₃)₂.²² Similar to the case of the E states of the water dimer, van Bladel et al.²² find that the localized A + E states of the ammonia dimer do not necessarily have to be close in energy, contrary to the stated expecta-tions of Nelson *et al.*^{9,10} This finding is confirmed by our experimental results. van Bladel et al.²² also find that the ab initio surface used in their quantum dynamical calculations is sufficiently flat that vibrational averaging causes the measured values of $eqQ_{aa}(NH_3^A)$ ($\langle P_2(\cos) \theta_A \rangle$) and $eqQ_{aa}(\mathrm{NH}_{3}^{B})$ ($\langle P_{2}(\cos \theta_{B}) \rangle$) to yield different values of θ_{A} and θ_B than those that seem consistent with the measured value of

$$\mu_a \propto (\langle P_1(\cos \theta_A) \rangle + \langle P_1(\cos \theta_B) \rangle),$$

such that all these observed quantities *cannot* be used together to deduce the structure, as Nelson, Fraser, and Klemperer⁸ have done. In addition, van Bladel *et al.*²² find that both of the above methods of estimating θ_A and θ_B

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yield values that deviate significantly from the equilibrium values of θ_A and θ_B for the surface used. van Bladel *et al.*²² are the first workers to actually quantify these deviations, thus removing all doubt that the effects of vibrational averaging *must* be considered in the determination of the ammonia dimer equilibrium structure.

It is now obvious that an appropriate analysis has not vet been performed to determine the equilibrium structure of the ammonia dimer. The proper way to approach this problem, which is beyond the scope of the present paper, is to fit a full six-dimensional (or eight-dimensional, if it is necessary to treat the monomer umbrella inversion in full detail) potential-energy surface to all of the spectroscopic data presented here, and then find the most-stable structure described by this fitted potential. This method would involve multiple iterations of the VRT state calculations described by van Bladel et al. as required by a nonlinear least-squares-fitting routine, and this would be computationally very demanding. Such analyses have recently vielded experimental potential-energy surfaces for Ar-H₂O (three dimensions),^{1,47} and for Ar-NH₃ (Ref. 37) (three dimensions plus an adiabatic treatment of monomer inversion), using the very efficient collocation method to calculate the VRT states of these complexes.

Theoretical investigators¹⁶⁻²⁰ who calculated *ab initio* intermolecular potential-energy surfaces for the ammonia dimer immediately following the experiments of Nelson et al.^{8,9} also suggest that "vibrational averaging" on their surfaces would lead to the experimentally observed molecular constants. Since we no longer need to judge theoretical surfaces by the criterion that they possess equilibrium structures that match the vibrationally averaged structure proposed by Nelson et al., we now reexplore the proposed ab initio structures. In general, the workers who used the largest basis sets, optimized all degrees of freedom, and accounted for electron correlation and basis-set superposition error, found that a nearly linearly hydrogen-bonded geometry was the equilibrium configuration for the ammonia dimer.^{16,20} In the coordinates of Fig. 1, Hassett et al.¹⁶ found $\theta_A \approx 24^\circ$ and $\theta_B \approx 94^\circ$, with ϕ_A rotated by 60° from the way we have drawn it. Frisch et al.¹⁶ found $\theta_A \approx 8^\circ$ and $\theta_B \approx 110^\circ$. Most the experimental information that has been directly compared with the ab initio results comes from the Klemperer laboratory. The theoretically predicted linearly hydrogen-bonded structure has a dipole moment of two dimensions.^{16,18,20} Vibrational averaging along the interchange tunneling coordinate could make the measured dipole moment smaller than two dimensions, and should affect the normal isotopomer more than the fully deuterated isotopomer. However, Nelson et al.^{8,9} found that the measured dipole moment of 0.5 D for (ND₃)₂ is actually smaller than the measured dipole moment of 0.8 D for $(NH_3)_2$. This result was thought to rule out the linearly hydrogen-bonded structure. Recently, however, Hassett, Marsden, and Smith²⁰ have finally pointed out the plausible explanation for the measured dipole moments consistent with their ab initio surface. Several competing effects are involved: while the zero-point vibration in the interchange tunneling coordinate leads toward a cyclic geometry which has no dipole moment, thus predicting a smaller expected dipole moment for $(NH_3)_2$, zero-point motion in the orthogonal in-plane vibrational mode leads to a head-to-tail geometry which has an approximately three-dimensional dipole moment, thus predicting a larger dipole moment for $(NH_3)_2$ than for $(ND_3)_2$.

Another calculation that favors the linearly hydrogenbonded structure is the empirical study by Liu *et al.*¹⁸ They computed highly accurate electrostatic contributions to the angular potential-energy interaction between two ammonia monomers fixed at the experimental center of mass distance. They, too, arrived at a configuration with $\theta_A = 12^{\circ}$ and $\theta_B = 101^{\circ}$. As we mentioned earlier, this calculation resulted in higher barriers to the various tunneling motions than the full *ab initio* potentials predict. Dykstra and Andrews¹⁸ later found that it is possible to fit atom-atom repulsive contributions to augment the electrical interaction surface that reduce the interchange barrier and can even shift the equilibrium structure towards the structure proposed by Nelson *et al.*^{8,9}

Other recent ab initio structural calculations which do not have a linearly hydrogen-bonded equilibrium structure are less convincing. Latajka and Scheiner¹⁷ used a smaller basis than either Frisch et al.¹⁶ or Hassett, Marsden, and Smith,²⁰ and found that a cyclic geometry with $\theta_A = 68^\circ$ and $\theta_{B} = 112^{\circ}$, which is close to the transition state of Frisch et al. and Hassett, Marsden, and Smith, was the equilibrium structure on their surface, and is 0.2 kcal/mol more stable than the linearly hydrogen-bonded form. Hassett, Marsden, and Smith²⁰ were able to obtain similar results when they reduced their basis size. Such an equilibrium structure is definitively ruled out by our experimental data. All of the electric-dipole-allowed transitions that are required to involve a change of interchange tunneling state in the C_s conformer would become pure rotational transitions in this proposed C_{2h} conformer, and it is unclear how such a structure can have a dipole moment, which is clearly required for such transitions to be observable. Sagarik, Ahlrichs, and Brode¹⁹ claim that they have predicted an equilibrium structure ($\theta_A = 61^\circ$ and $\theta_B = 105^\circ$) very similar to the one proposed by Nelson et al. However, van Bladel et al.²² have shown that the true minimum of that particular surface actually occurs at the linearly hydrogen-bonded geometry! The proposed structure of Sagarik, Ahlrichs, and Brode¹⁹ was originally preferred by Nelson et al.9 because a calculation of vibrational averaging effects was not necessary in order to achieve agreement between the theoretically and experimentally proposed structures.

ACKNOWLEDGMENTS

We thank Dr. Luca Dore, Nick Pugliano, and Jeff Cruzan for their help with the far-infrared search, and Sakae Suzuki and Paul Stockman for their help with the microwave experiments. We thank Professor Ad van der Avoird, Nick Pugliano, Paul Stockman, and Sakae Suzuki for providing useful comments on this manuscript. J.G.L. would like to thank Joyce Bumgarner for three weeks of hospitality in Pasadena, CA, while most of the microwave experiments were performed. This work was supported by the National Science Foundation, Grant No. CHE-8612296. G.A.B. would like to thank the David and Lucille Packard and Alfred P. Sloan Foundations for support.

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