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Matrix Isolation Spectroscopy and Nuclear Spin Conversion of Propyne Suspended in Solid Parahydrogen

A. I. Strom,¹ A. Gutiérrez-Quintanilla,^{2,3a} M. Chevalier,² J. Ceponkus,⁴ C. Crépin,² and D. T. Anderson^{1b}

¹Department of Chemistry, University of Wyoming, Laramie, WY 82071-3838, USA

²Université Paris-Saclay, CNRS, Institut des Sciences Moléculaires d'Orsay, 91405, Orsay, France

³Instituto Superior de Tecnologías y Ciencias Aplicadas (InSTEC), Universidad de La Habana, Ave. Salvador Allende No. 1110, Quinta de los Molinos, La Habana 10400, Cuba

⁴Institute of Chemical Physics, Vilnius University, Sauletekio ave. 9 III, LT-10222 Vilnius, Lithuania

^aPresent address: Aix-Marseille Université, Laboratoire PIIM, Team ASTRO, Service 252, Saint Jérôme, Ave. Escadrille Normandie Niemen, 13013 Marseille, France.

^bAuthor to whom correspondence should be addressed: danderso@uwyo.edu

ABSTRACT: Parahydrogen (pH₂) quantum solids are excellent matrix isolation hosts for studying the rovibrational dynamics and nuclear spin conversion (NSC) kinetics of molecules containing indistinguishable nuclei with nonzero spin. The relatively slow NSC kinetics of propyne (CH₃CCH) isolated in solid pH₂ is employed as a tool to assign the rovibrational spectrum of propyne in the 600 – 7000 cm⁻¹ region. Detailed analyses of a variety of parallel ($\Delta K=0$) and perpendicular ($\Delta K=1$) bands of propyne indicate that the end-over-end rotation of propyne is quenched, but K rotation of the methyl group around the C_3 symmetry axis still persists. However, this single-axis K rotation is significantly hindered for propyne trapped in solid pH₂ such that the energies of the K rotational states do not obey simple energy level expressions. The NSC kinetics of propyne follows first-order reversible kinetics with a 287(7) min effective time constant at 1.7 K. Intensity-intensity correlation plots are used to determine the relative line strengths of individual *ortho*- and *para*-propyne rovibrational transitions, enabling an independent estimation of the ground vibrational state effective A'' constant of propyne.

1. Introduction

Spectroscopy of dopant molecules isolated in quantum condensed matter affords unique opportunities to probe the properties of weakly interacting, light mass media at the molecular level. In particular, inherently large amplitude dopant rotational motion is an incredibly useful probe of the quantum host dynamics determined by both kinetic and potential energy contributions, as one of the main qualifications of quantum condensed matter is the significance of the zero-point energy relative to the strength of the intermolecular forces.¹⁻⁴ Rovibrational spectroscopy of molecules and clusters embedded in ^4He nanodroplets has long been used to probe the microscopic dynamics of this unique quantum fluid, where rotational fine structure is usually observed due to the Bose-Einstein statistics associated with ^4He .⁵⁻⁹ The rovibrational spectra of dopants dissolved in liquid He nanodroplets are representative of the gas-phase molecular or cluster symmetry and yet the rotational constants are reduced compared to the isolated molecule. Solid parahydrogen (pH_2) provides another opportunity to study dopant rotation. However, stronger $\text{H}_2\text{-H}_2$ intermolecular forces relative to the intermolecular forces in He and the hexagonal close packed crystal structure in the solid pH_2 limit the propensity for observing dopant rotation in pH_2 matrices. Molecules that occupy single substitutional sites in the pH_2 crystal lattice (CH_4 ,¹⁰⁻¹³ NH_3 ,¹⁴ H_2O ,^{15,16} HCl ,¹⁷ and CO ¹⁸ to name published examples) undergo nearly free rotation because the potential barriers to rotation are extremely small in this highly symmetric solvation environment. When bigger molecules are confined in solid pH_2 that occupy larger substitutional sites (double and larger), the end-over-end tumbling motions of the dopant are typically quenched.^{19,20} Yet molecules containing methyl rotors (CH_3) that behave as symmetric tops, such as CH_3F ,²¹ or internal rotors, like CH_3OH ,²² can continue to show spinning

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2
3 motion of the CH₃ rotor. This spinning motion can be used to probe quantum solvation in larger,
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5 more complex molecules, such as acetylacetone (CH₃(C=O)CH₂(C=O)CH₃).^{23,24}
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8 The accessible temperature range is an important aspect of probing the rotational motions
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10 of dopants trapped in quantum condensed matter. All of the molecules that demonstrate rotation
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12 while solvated in solid pH₂ have small moments of inertia (CO being the only exception in the
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14 literature¹⁸), and, thus, the rotational motion endures in spite of the rotational barriers induced by
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16 the pH₂ host. This means that the energy of the lowest excited rotational state of the dopant is
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18 also large with respect to the available thermal energy in the accessible temperature range in
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20 solid pH₂ (1.5 to 4.3 K), so there is very little population in excited rotational levels that
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22 significantly limits the number of rovibrational transitions that can be observed. Nonetheless,
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24 closer examination of the list of rotating molecules reveals another important attribute, namely
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26 indistinguishable hydrogen atoms. As a consequence of the symmetrization postulate of quantum
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28 mechanics,²⁵ excited rotational states connected to the different nuclear spin isomers of the
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30 dopant species can become kinetically trapped. Optimally, one would like to observe multiple
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32 rovibrational transitions to help tease apart different rotational and vibrational contributions to
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34 the measured transition energies in order to maximize the information content on the rotational
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36 dynamics of the dopant isolated in solid pH₂. Of course, nuclear spin constraints are active for
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38 methyl rotors in symmetric top molecules with two different moments of inertia, such as CH₃F.
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40 In this case, sufficient population of excited $K=1$ rotational states are kinetically trapped, such
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42 that transitions out of these levels are observable in the IR spectra of freshly deposited samples
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44 of CH₃F isolated in solid pH₂.²¹ With time, these transitions out of excited rotational states
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46 decrease in intensity due to condensed phase NSC. Thus, nuclear spin restrictions on the
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48 rotational wavefunctions provide the means to study dopant rotational motion in low temperature
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quantum hosts not possible in the fully equilibrated samples. This was successfully demonstrated for CH_3F^{21} and $\text{CH}_3\text{OH}^{22}$ doped pH_2 solids.

Our groups, in Wyoming and Orsay, became interested in investigating the infrared spectroscopy and solid state dynamics of methyl rotors trapped in solid pH_2 based on previous experiments conducted in Orsay.^{23,24} Preliminary results on the infrared spectroscopy of propyne in solid pH_2 , that are presented in the thesis of Gutiérrez,²⁶ were provided to the Wyoming group to stimulate further collaborative studies. The idea is to develop useful spectroscopic tools to gain information about how the quantum cavity around the methyl rotor perturbs the large amplitude K rotational motion. Specifically, we want to develop analytical methods that employ NSC phenomena to rigorously assign the measured infrared spectra for a number of molecular dopants containing methyl groups. For example, which types of vibrational modes provide the richest sources of information? If spectra contain significant fine structure, how can NSC be used to assign the different peaks? A number of researchers have already addressed some of these issues with much success,^{14,21-24,27} and we intend to refine the existing spectroscopic tools to better determine how solvation in quantum condensed matter perturbs rotational dynamics in dopant molecules. Furthermore, rovibrational spectroscopy can be used for NSC kinetics studies of methyl rotor molecules with different attributes. Ultimately, our goal is to measure signatures of NSC and quantitative nuclear spin populations of molecules for applications to photochemical and reactive studies conducted in solid pH_2 .

2. METHODS

Most of the spectroscopic data presented here was acquired in the laboratory in Wyoming. Some additional experiments were conducted in Orsay as this laboratory has the capability of performing prolonged studies of a single pH_2 sample. A detailed description of the experimental

methods used in Orsay is published elsewhere,²⁸ and here we point out some important differences between the two laboratories. In the laboratory in Orsay, samples are not obtained by co-deposition but rather are deposited through a single deposition tube. The samples are thinner and the concentration of dopant is higher than the samples grown in Wyoming. The lowest temperature achieved in the laboratory in Orsay is 2.8 K, the highest resolution of the FTIR spectrometer is 0.125 cm⁻¹, and samples can be kept frozen for several days at a time. As such, the spectroscopic data from Orsay was recorded at 2.8 and 4.0 K, for samples that were kept frozen for longer times and annealed, and thus could be used to complement and confirm the results from the Wyoming laboratory.

The experimental apparatus for pH₂ matrix isolation spectroscopy in the Wyoming laboratory has been discussed previously,^{14,29} so only details relevant to this study are presented below. Propyne-doped pH₂ solids are grown via the rapid-vapor-deposition (RVD) technique developed by Fajardo and Tam.^{30,31} This procedure involves the codeposition of independent pH₂ and propyne gas streams onto a precooled BaF₂ optical substrate held near ~2.5 K within a sample-in-vacuum /He cryostat, producing millimeters-thick propyne-doped pH₂-enriched samples in about ~30 mins. Enriched pH₂ gas (>99.97%) was rapidly introduced into the cryostat via the outlet tube of a variable-temperature *ortho*-to-*para* (*o/p*) converter packed with granular Fe(OH)₃ catalyst and backed by 800-1000 torr of normal H₂ gas (Linde, ≥99.999%) passed through a needle-valve set to achieve flowrates of $\Phi_{H_2} = 200\text{-}500 \text{ mmol}\cdot\text{hr}^{-1}$. In this work, the *o/p* converter is operated around 15 K producing pH₂ solids containing orthohydrogen (oH₂) concentrations ≈100 ppm (approximated via gas-phase H₂ rotational partition functions or determined experimentally using the oH₂-induced $Q_1(0)$ feature³¹ and sample thickness³²). Room temperature propyne (Aldrich, ≥99% as-received) was introduced into the cryostat

through a separate needle-valve set to achieve acceptable working concentrations that favor isolated monomers and disfavor propyne clusters. Exclusively as-deposited samples (hexagonal close packed (hcp) fractions between ≈ 50 -80%, estimated using the $U_1(0)$ feature²⁹) were studied in this work; annealing was conducted post-NSC to check for reversible and irreversible changes in measured IR spectra to confirm rovibrational assignments. Sample temperatures were measured using a pair of Si-diodes (attached to the OFHC Cu optical mount making thermal contact with the BaF₂ window) connected to a digital temperature controller. One diode (T_A) is located near to the cold finger of the cryostat, with the second (T_B) at the end of the optical holder; T_A is typically too cold to register temperatures (<1.0 K), so sample temperatures are reported using $T_B \geq 1.5$ K. Temperatures are recorded autonomously (sampling rate of 1 Hz) using free chart recording software installed on the same computer used for spectroscopic measurements.

Total propyne monomer concentrations are measured via Beer's law, stated as²⁹

$$[\text{CH}_3\text{CCH}](\text{ppm}) = \frac{\ln(10) \int A_i(\tilde{\nu}) d\tilde{\nu}}{\epsilon_i^{pH_2} d} V_0 (1 \times 10^6), \quad (1)$$

where $A_i(\tilde{\nu}) = \log_{10} [I_0(\tilde{\nu})/I(\tilde{\nu})]$ is the decadic absorbance of the i^{th} vibrational mode of propyne integrated over the entire band area ($\tilde{\nu}$ to $\tilde{\nu} + d\tilde{\nu}$), $\epsilon_i^{pH_2} = 1.06 \epsilon_i^{\text{gas}}$ is the gas-phase integrated absorption coefficient corrected for the refractive index of solid $p\text{H}_2$,³² d is the thickness of the solid $p\text{H}_2$ slab,³² and $V_0 = 23.16(6) \text{ cm}^3 \text{ mol}^{-1}$ is the molar volume of solid $p\text{H}_2$ at /He temperatures.³³ Concentrations and $p\text{H}_2$ samples thicknesses are reported as weighted averages weighted by uncertainties. Integrated intensities from the literature and integration protocols are provided in the [Supporting Information](#), Table S2. Similarly, $p\text{H}_2$ sample thicknesses are reported as the weighted average of three independent measurements, integrated intensities of the $Q_1(0)+S_0(0)$, $S_1(0)+S_0(0)$ double transitions and $Q_R(0)$ IR absorptions bands³²

weighted by their absorption coefficients. Error propagation is performed in the usual way, in which experimental as well as reported uncertainties in the literature are considered; typically, the largest contribution to the stated errors stem from uncertainties in the integrated absorption coefficients. A high-resolution FTIR spectrometer (Bruker IFS 120/5; $\Delta\tilde{\nu}=0.02\text{-}0.03\text{ cm}^{-1}$) optimized for normal incidence transmission optical measurements was employed using multiple optical setups: 1) a globar (GB; MIR) source, Ge coated KBr beamsplitter and ^4He cooled InSb detector, 2) a GB source, Ge coated KBr beamsplitter and ^4He cooled HgCdTe (MCT) detector, 3) a tungsten filament (W; NIR) with a CaF_2 beamsplitter and 4) identical to setup #2 with the addition of a long pass filter (LPF) with a 3861 cm^{-1} cutoff in the IR beam path to eliminate absorptions by the $p\text{H}_2$ host. These optical setups were designed to record high-fidelity IR spectra of specific vibrational modes of propyne located within the allowed wavenumber ranges ($\Delta\tilde{\nu}=0.03\text{ cm}^{-1}$ over $600\text{-}4800\text{ cm}^{-1}$ for the MCT and $\Delta\tilde{\nu}=0.02\text{ cm}^{-1}$ over $1800\text{-}5000\text{ cm}^{-1}$ or $1800\text{-}10,000\text{ cm}^{-1}$ for InSb with the Ge coated KBr or CaF_2 beamsplitters, respectively). Furthermore, propyne concentrations deliberately span two orders of magnitude (roughly $20\text{-}300\text{ ppm}$) in order to enhance the sensitivity of target mode(s) in each experiment.

In order to study the NSC of propyne molecules isolated in solid $p\text{H}_2$ below ^4He temperatures, room temperature populations of excited rotational states of propyne must be deposited into the solid much faster than they can relax, as in $\text{NH}_3/p\text{H}_2$ studies.¹⁴ Thus, the RVD procedure was typically performed even more rapidly by setting $\Phi_{\text{H}_2}\approx 400\text{ mmol hr}^{-1}$ to enhance observations of metastable propyne. Rapid-scan FTIR spectra recorded with modest time resolution (3-5 min depending on averaging) were collected during the deposition step while calibrating the dopant flowrate with Φ_{H_2} in order to enhance S/N and maximize contributions from freshly-deposited rotating propyne molecules, and, of course, immediately post deposition

in order to monitor propyne NSC in as-deposited solid pH₂ samples near ~1.7 K until equilibration. Equilibrium was achieved when intensity changes between difference spectra were deemed undetectable, which was optimally on the order of the effective time constant of propyne NSC, that is to say, relaxation periods of $\delta t \approx 300$ min or greater.

Important experimental parameters for each experiment in this study are summarized in the [Supporting Information](#), Table S1, including optical setups, instrument resolution, H₂ flow rates, pH₂ sample thicknesses, hcp fractions, propyne concentrations and the IR bands used to determine them, and the duration of the relaxation period for each NSC kinetic run.

3. RESULTS AND DISCUSSION

Propyne (CH₃C≡CH) is the simplest linear hydrocarbon containing a methyl rotor. The gas-phase spectroscopy of this prolate symmetric top ($A''=5.31$ cm⁻¹, $B''=0.29$ cm⁻¹)³⁴ has been extensively studied as a model system to study vibrational energy flow in molecules.³⁴⁻³⁷ Belonging to the C_{3v} point group, propyne has ten fundamental vibrational modes with half of the modes, $\nu_1 - \nu_5$, having A_1 symmetry, while the other half, $\nu_6 - \nu_{10}$, are degenerate with E symmetry.^{38,39} Accordingly, parallel ($\Delta K=0$) and perpendicular transitions ($\Delta K=\pm 1$) of propyne are expected for the A_1 and E symmetry vibrations, respectively. As discussed previously, the end-over-end tumbling motion of propyne will likely be quenched similar to acetylene¹⁹ when isolated in solid pH₂, but the spinning rotation of the methyl group, corresponding to the A rotational constant, should continue to freely rotate similar to CH₃F.²¹ Based on van der Waals radii and the structure of propyne,³⁴ propyne occupies a double substitution site in solid pH₂ with cylindrical symmetry. Therefore, for propyne isolated in solid pH₂, we expect that the J and M_J rotational quantum numbers appropriate for the gas-phase molecule in free space are lost, and only the K rotational quantum number survives. Basically, the three-dimensional rotational

motion of propyne in free space transforms into one-dimensional rotation like a particle-on-a-ring, and two angular librational degrees of freedom, upon suspension in the double substitution site. Therefore, we break from using conventional gas-phase spectroscopic notation and instead use a simplified notation, namely, $\Delta K_{\nu'}(K'')$, where $\Delta K = -1, 0, +1$ are designated by P, Q, R and ν' and K'' are the vibrational quanta in the upper state and the lower state K quantum number.

Each K rotational state is entangled with a single nuclear spin wavefunction of a specific symmetry to satisfy the symmetrization postulate of quantum mechanics. For CH_3CCH with three identical hydrogen nuclei (fermions), the *ortho* ($\Gamma_{ns}=A_1, I=3/2$) nuclear spin wavefunctions combine with $K = 0, 3, 6, \dots$ rotational states in the ground vibrational state and *para* ($\Gamma_{ns}=E, I=1/2$) nuclear spin wavefunctions combine with $K=1, 2, 4, 5, \dots$ rotational states. Thus, at the high temperature limit ($T > 50$ K) a gaseous sample of propyne consists of roughly 50:50 mixtures of the *ortho* and *para* spin isomers because the factor of two greater nuclear spin weight for the *ortho* spin isomer cancels the factor of two more $K \neq 3n$ rotational states of the *para* isomer.

For A_1 symmetry vibrations, parallel transitions should lead to single peaks which correspond to overlapping $Q_1(0)$ and $Q_1(1)$ transitions out of the lowest rotational states for the *ortho* and *para* nuclear spin isomers, respectively. The fact that we observe $Q_1(0)$ transitions (vide infra) is another observable in support of the fact that end-over-end rotation of propyne is quenched; in the gas-phase no Q -branch is observed for the $K=0$ rotational sub-band.^{38,39} To the extent that the A rotational constant is the same in the upper and lower vibrational states, these transitions will overlap perfectly. However, if $A' - A'' \neq 0$, then the $Q_1(0)$ and $Q_1(1)$ transitions will occur at slightly different wavenumbers ($Q_{\nu'}(1) - Q_{\nu'}(0) = A' - A''$) and there is a possibility of resolving this K fine structure. In contrast, E type vibrations result in perpendicular bands

with well-resolved K sub-bands that correspond to the $R_1(0)$ transition for the *ortho* nuclear spin modification and $P_1(1)$ and $R_1(1)$ transitions for the *para* nuclear spin isomer. For degenerate E vibrations the vibrational angular momentum $l = \nu_i, \nu_i-2, \dots, -\nu_i$ in the upper state can align or oppose the angular momentum of the K rotation.^{38,39} This gives rise to first-order Coriolis splittings that can be modeled by the energy term $\pm 2(A\zeta)_v K l$, where ζ is the Coriolis coupling constant. Given that the ground state A'' rotational constant is $\sim 5.31 \text{ cm}^{-1}$ for propyne in the gas-phase,³⁴ at 1.7 K, transitions out of the $K'' = 2$ rotational state at approximately 21.2 cm^{-1} ought to be extremely weak due to lack of population of this level. However, if the $K=2$ level is significantly shifted to lower energies, then transitions out of this level may also be observed.

3A. Infrared Spectroscopy of Propyne in Solid $p\text{H}_2$. One of the main difficulties in making detailed assignments of the infrared spectrum of propyne isolated in solid $p\text{H}_2$ is the fine structure observed for some of the vibrational modes. This fine structure can be caused by single-axis K rotation, but can also be due to a variety of causes, such as propyne-(oH_2)_{*n*} clusters, and one wants a way to selectively identify rovibrational transitions of freely rotating propyne monomers. If the goodness of the K rotational quantum number is lost due to clustering or trapping in defect sites, then we cannot differentiate between different nuclear spin isomers using rovibrational spectroscopy. With this in mind, we begin with the assignment of the A_1 vibrational modes of propyne.

As discussed above, transitions involving A_1 vibrational modes should be observed as single peaks (or two overlapping peaks) close to the gas-phase vibrational origins. Typically, $p\text{H}_2$ matrix shifts¹⁻⁴ are less than 1% and, thus, for well-resolved single peaks, vibrational assignments are made simply by comparing the measured peak wavenumber with a potential gas-phase origin. Shown in Figure 1(a) are absorption spectra from 750 to 1500 cm^{-1} for a 2.5(2)

mm thick, 200(39) ppm propyne/pH₂ sample recorded at 1.70(1) K at 2.63 (red) and 491 min (blue) after deposition. The spectra are intentionally offset from zero to show the difference spectrum. All the A_1 vibrational modes in this region are easily assigned via comparison with the vibrational origins reported in the literature and these A_1 vibrational assignments are presented in Table 1. However, by constructing difference spectra recorded at short and long time delays after deposition, we can try to resolve these two bands based on NSC. RVD of room temperature propyne into solid pH₂ at 1.7 K kinetically traps the higher energy *para* nuclear spin isomer in the $K = 1$ rotational state due to slow NSC relative to vibrational and rotational relaxation. Assuming that NSC is slow albeit measurable, then by constructing difference spectra⁴⁰ as $(A_t - A_\infty)$ as a function of time ($t = 0$ defined as the end of deposition) allows us to identify absorptions due to the *ortho* and *para* nuclear spin isomers of propyne. On the basis of the +/- phase of the peak in the difference spectrum, positive peaks are due to the *para* spin isomer and will be maximum at $t=0$ and must decrease with time. Negative peaks are minimum at $t=0$ and increase to a maximum at $t = \infty$ (*i.e.*, ≈ 4 h) as population is transferred to the lower energy $K=0$ rotational state of the *ortho* nuclear spin isomer. Furthermore, peaks in the difference spectra can be used to selectively identify absorptions due to propyne monomers that are undergoing unhindered K rotation. As we will show, some peaks in the absorption spectra do not appear in the difference spectra and this obviously inhibits the *para* and *ortho* assignments of these peaks. Nevertheless, peaks with sufficient intensity in the difference spectra, that exhibit quantitatively the correct NSC kinetics, allow the nuclear spin isomer responsible for that transition to be rigorously assigned.

Careful inspection of the difference spectrum shown in Figure 1(b) illustrates our strategy for making detailed assignments. For A_1 symmetry parallel bands ($\Delta K=0$), the wavenumber

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3 difference in the $Q_1(0)$ and $Q_1(1)$ sub-bands must be comparable to the FWHM of the individual
4 absorptions to detect a difference signal. All the A_1 vibrational bands shown in Figure 1(a) do
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6 not show strong difference signals. Based on gas phase measurements,^{41,42} detecting difference
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8 signals for both the ν_5 and ν_4 fundamentals should be difficult because the $A' - A''$ differences are
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10 small, -0.0073 cm^{-1} and 0.017 cm^{-1} , respectively, and both peaks have modest integrated
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12 intensities (see [Supporting Information](#), Table S2). Obviously, small population differences are
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14 detected with higher sensitivity using absorptions with large integrated intensities. The $2\nu_9^0$
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16 overtone peak at 1254.47 cm^{-1} has a large integrated intensity⁴³ ($16.81(29) \text{ km mol}^{-1}$), but again
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18 based on gas phase measurements⁴⁴ the $A' - A'' = -0.00292 \text{ cm}^{-1}$ value is quite small. The small
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20 difference signal for the $2\nu_9^0$ overtone peak shown in Figure 1(b) therefore shows that the $Q_2(0)$
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22 and $Q_2(1)$ absorption features are strongly overlapped making it difficult to determine the
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24 splitting. Thus, all the A_1 vibrational modes displayed in Figure 1 do not show a clear difference
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26 spectrum and are assigned to overlapping $Q_1(0)/Q_1(1)$ peaks. However, as we will show for ν_2 ,
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28 when the $A' - A''$ difference becomes a little larger, a clear difference signal can be measured for
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30 A_1 type vibrations that provide information on the vibrational origin and the value of $A' - A''$ for a
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32 particular vibrational mode.
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41 For E -type vibrational symmetry modes that produce perpendicular bands ($\Delta K = \pm 1$), the
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43 rovibrational transitions for the *para* and *ortho* spin isomers are spectrally resolved making
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45 assignments rather straightforward. For example, also shown in Figure 1 is the ν_7 (see
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47 [Supporting Information](#), Figure S5) absorption of propyne near 1450 cm^{-1} . This mode shows a
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49 number of peaks in absorption, but the difference spectrum reveals a positive peak at 1443.54
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51 cm^{-1} and a negative peak at 1455.12 cm^{-1} for the *para* and *ortho* nuclear spin isomers,
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53 respectively. These nuclear spin assignments are rigorous, and we can therefore tentatively
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assign these two peaks to the $P_1(1)$ and $R_1(0)$ transitions of the *para* and *ortho* nuclear spin isomers, respectively. The energy difference between these two transitions predicted using the gas-phase rotational constants⁴¹ (neglecting centrifugal distortion) is 14.45 cm^{-1} , which compares favorably to the measured value of 11.58 cm^{-1} and is also consistent with assigning the higher wavenumber transition to the *ortho* nuclear spin isomer, as indicated by the difference spectrum. Similarly, the ν_8 mode shows (see inset in Figure 1b) positive and negative difference spectra peaks that are assigned to the *para* $P_1(1)$ and *ortho* $R_1(0)$ peaks, respectively. These two rovibrational transitions are typically the strongest for *E*-type vibrational modes and the $R_1(1)$ transition is also observed sometimes, but is usually much broader and harder to detect. The rovibrational assignments of *E* type vibrational bands are also listed in Table 1. In what follows, we examine various propyne bands in greater detail in order to articulate some of the challenges associated with making detailed rovibrational assignments.

After assigning *ortho* and *para* nuclear spin symmetries to the various peaks present in the spectrum of propyne using the difference spectra, we can attempt K rotational assignments using conventional energy levels expressions developed for gas-phase prolate symmetric tops.^{38,39} However, since we expect that J is no longer a good quantum number, the energy level expressions are modified to consider only terms associated with single-axis K rotation (neglecting centrifugal distortion). In the [Supporting Information](#), Section 2 various expressions are discussed for Q , R , and P transitions with $K'' = 0, 1$, and 2 for both parallel (A_1) and perpendicular (E) rovibrational bands. This information is used to compare the measured transition wavenumbers of propyne with gas-phase results to attempt detailed assignments.

The first band investigated in detail was the ν_2 (symmetric methyl CH stretch) fundamental near 2941 cm^{-1} . This vibrational mode has a large integrated band strength (see

Supporting Information, Table S2) and therefore is sensitive to small population differences. As we will show, this band can be used to effectively measure the NSC kinetics of propyne in solid pH_2 . The ν_2 spectrum recorded at 0.02 cm^{-1} resolution is shown in Figure 2. Spectra recorded 5.08 and 291.8 min after deposition are shown in Figure 2(a) as red and blue traces, respectively. Both spectra are intentionally offset from zero to better show the resulting difference spectrum. Shown in Figure 2(b), therefore, is the difference spectrum. The difference spectrum displays the signature lineshape for overlapping $Q_1(1)$ and $Q_1(0)$ transitions for the two nuclear spin isomers, namely, the positive $Q_1(1)$ peak is assigned to the *para* isomer and the negative $Q_1(0)$ peak to the *ortho* isomer. Based on the gas-phase rotational constants,^{35,45} these two transitions are predicted³⁵ to be split by 0.055 cm^{-1} (or 0.090 based on Ref. 45) with the $Q_1(0)$ transition of the *ortho* nuclear spin isomer at higher wavenumbers. For propyne isolated in solid pH_2 , we observe these two bands with the same energetic ordering as measured in the gas-phase, and with a splitting of around 0.11 cm^{-1} . These measurements demonstrate that the change in the A rotational constant upon ν_2 excitation is very small, but does not provide information on the value of A in either the ground or excited vibrational states. It also demonstrates how the difference spectrum of a parallel band can be used to measure the growth and decay in the populations of the two nuclear spin isomers of propyne. By definition, the difference spectrum is only sensitive to propyne populations that undergo K -rotation and NSC and therefore rejects overlapping propyne features that do not contribute to NSC. For example, careful examination of Figure 2 reveals that the total ν_2 lineshape is not well represented by the sum of two overlapping pseudo-Voigt peaks, but rather this absorption has a shoulder to the low wavenumber side ($\sim 2934.4\text{ cm}^{-1}$) that does not change in intensity with time. In addition, integration of the entire difference spectrum results in a value of zero indicating that the line

strengths for the two transitions are approximately equal. This is consistent with both these transitions being pure vibrational transitions (not rovibrational) and the $Q_1(0)$ transition has the same integrated band strength as the $Q_1(1)$ peak. This is a direct ramification of the loss of J as a good quantum number such that the $Q_1(0)$ transition is possible for the matrix isolated propyne.

In Figure 3, we present the full spectrum of ν_6 (antisymmetric methyl CH stretch) which is representative of a perpendicular band.^{35,46} Here we see more deviations from gas-phase behavior, signaling perturbations in the K rotational motion. For the absorption spectra recorded at short and long times after deposition, we measure approximately 7 peaks excluding the two very weak peaks at the extremes of the spectral range displayed in Figure 3. However, in the difference spectrum, three peaks are absent (marked with asterisks in Fig. 3) and we only have clear difference signals for four peaks, indicating that the three central peaks shown in Figure 3 do not show the same NSC kinetics as the others. It is possible that these peaks are due to propyne-(oH₂)_n clusters that form irreversibly due to the stronger attractive intermolecular forces between propyne and oH₂ compared to pH₂; however, assignment of these cluster peaks extends beyond the scope of this work. Based on the idea that the strongest transitions should be out of the $K = 0$ and 1 rotational states, we should observe one $R_1(0)$ peak for the *ortho* nuclear spin isomer, and two transitions for the *para* isomer, namely, $P_1(1)$ and $R_1(1)$. We do observe two peaks for the *para* spin isomer, but we also observe two $R_1(0)$ peaks for the *ortho* spin isomer.

As first demonstrated by Lee, Wu and Hougen,²¹ we can attempt to make K rotational assignments of the ν_6 band based on a simple energy level expression for single-axis K rotation and the molecular constants measured in the gas-phase (see [Supporting Information](#), Section 2). To begin, we make the simplifying assumption that we can use the average of the two observed $R_1(0)$ transitions for the $R_1(0)$ transition wavenumber ($R_1(0)_{\text{avg}} \approx 2979.7 \pm 0.7 \text{ cm}^{-1}$). As we will

show, it doesn't matter how we treat this obvious deviation from gas-phase behavior, we cannot extract a consistent set of molecular constants. We first assigned the three rovibrational transitions shown in Fig. 3 corresponding to $K'' = 0, 1$ and assuming the gas-phase Coriolis constant ($\zeta_6=0.07250$)⁴⁶ is preserved, we determined the three molecular constants presented in Table 2. This gave reasonable results consistent with reduced A constants for propyne K -rotation in solid pH_2 . However, next, we used all 5 rovibrational transitions labeled in Figure 3 corresponding to $K'' = 0, 1, 2$. Whereas the assignment of $K''=2$ transitions seems beyond a doubt, by comparison with gas phase values (Table 1), the relation $R_1(2) - P_1(2) = 2[R_1(1) - P_1(1)]$, deduced from the simplified model, is not fulfilled, independent of the value of the Coriolis constant. All variations of the fitting procedure given in the [Supporting Information](#), Section 2, fixed and floated ζ_6 , and including a D_K distortion constant resulted in fits that could not reproduce the transition wavenumbers to within experimental precision (e.g., $rms = 1.2 \text{ cm}^{-1}$). This is clearly a statement that the measured spectra, assigned via the methods described, do not readily conform to expressions determined for single-axis K rotation. Thus, comparison of the determined molecular constants with gas-phase values to make detailed K rotational assignments is likely misleading, especially when assumptions must be made given a limited number of transitions. This is an intrinsic problem with conducting these types of rovibrational assignments that only involve a few transitions and therefore lack built-in redundancies in the spectrum (such as four line combination differences), there is no way to rigorously test potential assignments. Surely, qualitative insights can be ascertained, but true predictive understanding of the energy patterns can only be advanced by comparing with fully quantum mechanical simulations that include interactions of the dopant with the surrounding pH_2 solid.

The *E* type ν_7 (CH_3 skeletal deformation) vibrational mode at roughly 1450 cm^{-1} , shown previously in Figure 1, was also studied in detail (see [Supporting Information](#), Figure S5 for the expanded spectrum). For this band, we only observed the $P_1(1)$, $R_1(0)$, and $R_1(1)$ transitions and, thus, to extract molecular constants from this mode we had to assume that the gas-phase Coriolis coupling constant ($\zeta_7 = -0.31424$)⁴¹ is largely unchanged upon solvation in solid pH_2 . Similar to the ν_6 band, the $R_1(0)$ peak consists of two partially resolved peaks and therefore we used the average peak position ($R_1(0)_{\text{AVG}} = 1454.9\text{ cm}^{-1}$). There is some doubt in the assignment of the $R_1(1)$ transition, either to a feature at 1458.4 or 1473.88 cm^{-1} . However, using either $R_1(1)$ assignment, the simplified model with the gas phase Coriolis coupling constant leads to unphysical A rotational constants, and importantly, to a different A'' constant than the ν_6 analysis. The three molecular constants determined from this analysis with $R_1(1)$ at 1473.88 cm^{-1} are presented in Table 2. Once again, this analysis demonstrates that quantitative analysis of the extracted molecular constants from K -rotational assignments of only three transitions is potentially misleading. The only other *E* type vibration where this analysis is possible is the $\nu_3+\nu_8$ combination band and the results are also presented in Table 2. Clearly, the analysis of each band leads to a different A'' constant each time. The fact that this type of analysis does not produce a consistent ground state A'' constant implies that either the assumption that the Coriolis constant does not change, or the assumed energy level expression, is incorrect. Applying a similar type of analysis to the three *E*-type rovibrational bands reported in the literature²¹ for CH_3F in solid pH_2 leads to a similar conclusion; each band produces a different ground state A'' rotational constant. This suggests that the K -rotational energy levels of both methyl rotor molecules, CH_3CCH and CH_3F , are not well described by simple single-axis K rotational energy level expression.

We repeatedly observed two $R(0)$ lines for many of the E -type propyne bands (ν_6 , ν_7 , ν_8 , $2\nu_6$). This is most likely due to the selection rules for C_{3v} symmetric tops, the fact that Coriolis perturbations are possible in upper degenerate vibrational states, and the loss of the J quantum number for propyne isolated in solid pH_2 . Neither of the rotational levels connected by the $P_1(1)$ rovibrational transition are effected by Coriolis perturbations and the $R_1(1)$ line connects E symmetry rotational levels.⁴⁷ But the $R_1(0)$ transition accesses the $K=1$ rotational state in the upper vibrational state and thus in the gas-phase accesses alternating A_1 and A_2 rotational states as a function of J . In the present case where the J quantum number is lost, Coriolis perturbations of the upper rotational state could produce two spectrally resolved $R_1(0)$ transitions. Furthermore, observation of two $R_1(0)$ transitions also contributes to the difficulty of extracting consistent molecular constants from the analysis of the $P_1(1)$, $R_1(0)$, and $R_1(1)$ transitions. Thus, it appears that the vibrational angular momentum generated in the upper degenerate vibrational state that is coupled to the K -rotation produces shifts in the transition wavenumbers on the order of the K sub-band spacing. Two $R_1(0)$ transitions could also appear when considering a removal of the degeneracy of the upper E vibrational state due to the solid environment. If the perturbation due to the solid environment is large enough to induce torsional barriers, the $K'' = 0$ and 1 levels are better represented by A and E levels, respectively, in the periodic torsional potential (splitting due to H atom tunneling) as was described in Ref. 23 in the case of the acetylacetone molecule. The analysis of the NSC process remains the same, but the structure presented in the spectrum by the $P_1(1)$ and especially the $R_1(1)$ transitions is difficult to assign.

These same types of discrepancies are not found in the interpretation of rovibrational spectra of molecules isolated in superfluid He nanodroplets.⁵⁻⁹ Typically, the measured rotational constants for molecules dissolved in He nanodroplets are reduced (39% and 95% for

heavy and light rotors) from gas-phase values, and distortion constants can be anomalously large, but the standard spectroscopic signatures are preserved along with the selection rules based on the symmetry of the gas-phase transition dipole moment.⁵⁻⁹ For example, the parallel ($\Delta K=0$) ν_1 spectrum (acetylenic C-H stretch) of propyne was recently revisited for propyne isolated in He nanodroplets.^{48,49} The spectrum displays *P*, *Q*, and *R* branches appropriate for a C_{3v} symmetric top molecule. Comparison between the fitted constants for gas-phase propyne and propyne solvated in He nanodroplets shows that the *B* rotational constant is reduced to 26.0(1)% and 25.6(1)% of its gas-phase value for the ground and $\nu=1$ excited vibrational states, respectively, and the Δ_J distortion constant increases by a factor of 5100 compared to the gas-phase. However, using these effective rotational constants, the spectrum is relatively well reproduced demonstrating that the selection rules are preserved.⁴⁹ Note that the ν_1 band is a parallel band so that the spectrum is not sensitive to the *A* constant directly, but rather the difference in *A* constants with vibrational excitation. Unfortunately, the corresponding ν_1 spectrum for propyne isolated in solid pH_2 is not very informative because the difference spectrum shows multiple positive and negative peaks (see [Supporting Information](#), Figure S6). Nonetheless, it is clear that isolation of propyne in solid pH_2 perturbs the rotational motion in a qualitatively different way; there is no end-over-end rotation in solid pH_2 while in liquid 4He the end-over-end rotational motion is preserved (although greatly hindered) and so are the selection rules. This means that the rovibrational dynamics of molecules in solid pH_2 are more difficult to assign rigorously and therefore also harder to interpret. While both He and pH_2 are quantum hosts, one is a liquid and the other a solid and this leads to very different rotational dynamics of guest molecules. In solid pH_2 , the rotational dynamics are significantly perturbed, end-over-end rotational is quenched, and the selection rules are altered (no *Q*-branch for $K=0$ in the gas-phase), such that the observed

infrared spectra of molecular dopants isolated in solid pH_2 exhibiting K -axis rotation are not so easily assigned via comparison to gas-phase spectra. However, the spectra assigned via the methods used here provide new data sets that can be modeled using quantum mechanical dynamic calculations to learn more about the pH_2 cage dynamics.

3B. Nuclear Spin Conversion Kinetics. We initiate our analysis of the NSC kinetics of propyne in solid pH_2 using the ν_2 vibrational band near 2935 cm^{-1} . As described above, since ν_2 is a parallel band, peaks associated with separate *ortho* and *para* nuclear spin isomers are not observed directly. Therefore, difference spectra recorded at various times after deposition are utilized to measure the effective rate constant for *para-to-ortho* NSC of propyne. As described in detail in the [Supporting Information](#), Section 1, each difference spectrum is fit to a sum of two normalized symmetric pseudo-Voigt profiles⁵⁰ in order to quantify time-dependent intensity changes in the two overlapping absorptions that are resolved in the difference spectrum. The two difference peaks in Figure 2 can be assigned to the *para* ($K=1, I=1/2$) and *ortho* ($K=0, I=3/2$) nuclear spin isomers. A representative difference spectrum for ν_2 fit to the sum of two pseudo-Voigt lineshape functions is shown in Figure 4; the difference spectrum is well modeled by this functional form allowing a number of parameters to be extracted from each spectral trace (see [Supporting Information](#), Section 1 for more details). The strength of this procedure is that the peak areas of components in an overlapping band, which are conserved quantities, are extracted directly from the analysis in order to study their temporal behavior. It turns out that the time dependence of the two components in the ν_2 difference spectra are well-fit to solutions of first-order kinetics equations, where the effective lifetime of propyne NSC, τ_{eff} , is statistically equivalent for both components to within the standard errors of the fitted parameters (see [Supporting Information](#), Table S4).

Representative kinetic traces for the ν_2 (A_1) vibrational mode of propyne are shown in Figure 5a. As described earlier, the red and blue data points represent the *para* and *ortho* peak areas determined from fits to the difference spectra. The lines represent the results of least-squares fits of the data to first-order kinetics equations (see below). Note that the way we calculate the difference spectra means that both peak areas should decay to zero as the propyne nuclear spin populations come into equilibrium at long times after deposition. For the data shown in Figure 5a, we did not wait long enough to achieve equilibrium as evidenced by the crossing of the fits above and below zero. However, this does not affect the determined τ_{eff} time constant due to the nature of first-order kinetics. Individual time constants are determined for each peak and are well matched. Repetition of this analysis in four separate experiments yields highly reproducible results (see [Supporting Information](#), Table S4). Examination of Figure 5a also shows that the two peak areas are symmetrically displaced around zero indicating that the line strengths associated with these two peaks are comparable, although the peak FWHM's differ (see Fig. 2).

The ν_6 and ν_7 perpendicular bands were also analyzed to determine effective NSC time constants for propyne. To quantify the NSC kinetics, the ν_6 and ν_7 bands are broken into segments; fortunately, there is enough separation between the various peaks to integrate the peaks corresponding to *ortho* ($R_1(0)$) and *para* ($P_1(1)$ and $R_1(1)$) nuclear spin isomer populations. The integrated intensities are well-fit to first-order kinetic equations given by

$$I_{para}(t) = I_{para}(\infty) + (I_{para}(0) - I_{para}(\infty))\exp(-t/\tau) \quad (2)$$

$$I_{ortho}(t) = I_{ortho}(0) + (I_{ortho}(\infty) - I_{ortho}(0))[1 - \exp(-t/\tau)] \quad (3)$$

in which the fitted lifetimes agree statistically with the ν_2 data to within error ([Supporting Information](#), Table S4). Representative kinetic traces for ν_6 and ν_7 are shown in Figure 5b and

5c, respectively. The kinetic traces in Fig. 5b and 5c are different from 5a because they were constructed from the integration of absorption spectra, not difference spectra. We did check that the total integrated intensity of each band remains constant over the time window of the kinetic analysis. Eleven separate kinetic analyses of the ν_2 , ν_6 and ν_7 bands of propyne all agree with each other and produce a weighted average time constant of $\tau_{eff} = 287(7)$ min.

As a further test of this kinetic analysis, we also fitted the difference spectrum of the ν_2 mode of $^{13}\text{CH}_3^{12}\text{C}_2\text{H}$ present in natural abundance ($\sim 1.1\%$) for the experiment with the highest propyne concentration. The other ^{13}C isotopologs likely overlap with the ν_2 mode of the most abundant isotopolog.^{35,51} As expected, this peak was also well-fit by expressions (Eq. 2 and 3) and gave a time constant of $\tau_{eff} = 170(20)$ min, which is approximately 1.7(3) times faster than the value determined for the $^{12}\text{C}_3\text{H}_4$ isotopomer. The reason for the faster NSC in $^{13}\text{CH}_3^{12}\text{C}_2\text{H}$ is likely because the $^{13}\text{C}(I=1/2)$ nucleus interacts with the H-atoms on the methyl group via magnetic dipole-dipole coupling which introduces an additional relaxation channel not open to the normal isotopolog.⁵²⁻⁵⁴ It would be interesting to put the ^{13}C on the acetylenic carbons to see if the time constant increases for the more remote atom with non-zero nuclear spin.

3C. Measuring the *para-to-ortho* ratio for propyne in solid pH_2 at low temperature.

The thermodynamic equilibrium concentrations for the *para* and *ortho* nuclear spin isomers of propyne are given by the respective rotational partition functions as a function of temperature. This *para-to-ortho* ratio (*POR*) is defined here as the equilibrium constant for the reversible, first-order nuclear spin conversion process between the two spin isomers of propyne, namely,

$$\textit{ortho}\text{-propyne} \leftrightarrow \textit{para}\text{-propyne}; \textit{POR} = \frac{[\textit{para}]}{[\textit{ortho}]} \quad (4)$$

As discussed earlier, in the high temperature limit the *POR* approaches unity, whereas this ratio goes to zero at $T \rightarrow 0$ K. We were therefore interested in calculating the experimental *POR*

measured in these studies after the system has come into equilibrium as another way to determine the energy splitting between the $K=1$ and $K=0$ rotational states. However, in order to do this necessitates that we transform our integrated intensities into relative populations, which means the line strengths of individual rovibrational transitions must be determined. In calculating propyne concentrations using Eq. (1) we simply integrate over the entire band and use the integrated IR band intensities given in the [Supporting Information](#), Table S2, but if we want rotational populations we must deal with rovibrational line strengths. To do this rigorously is difficult, thus, we used an approximation that has been used by others,⁵⁵ where we assume that the populations in the $K=0$ and $K=1$ follow this equation, $n_{K=1} + n_{K=0} = 1$, that is, that the system is well described as a two component system, then the relative line strengths can be determined from intensity-intensity correlation plots.⁵⁵ Correlation plots of the intensities for various transitions in ν_6 can be generated as shown in the [Supporting Information](#), Figure S9 and Table S5. Furthermore, this approach allows us to estimate the *POR* as a function of time during an experiment and also to estimate the energy of the $K=1$ excited rotational state of the *para* nuclear spin isomer.

We analyzed the ν_6 spectra for six different experiments in terms of the *POR* to see if general trends could be quantified. A representative *POR* trace is shown in Figure 6. As stated earlier, the high temperature statistical limit is $POR = 1.0$, where there is an equal mixture of *ortho* and *para* nuclear spin isomers. Upon RVD of room temperature propyne gas into solid pH_2 , the initial $POR \approx 0.9$ at ~ 2.5 K during deposition. By the end of deposition ($t=0$), the *POR* drops to a value of ~ 0.75 and after deposition it continues to decay at 1.7 K. From a fit to the experimental data, the *POR* decays to an equilibrium value of ~ 0.17 . This is an order of magnitude greater than the theoretical limit predicted using the gas-phase rotational energies at

equilibrium at 1.7 K ($POR = 0.0117$) and therefore signals a lowering of the $K=1$ level. We note that this analysis does suffer from potential systematic errors due to contributions of the non-rotating propyne absorption at 2978.29 cm^{-1} to the integrated intensity of the $R_1(0)$ peak, the fact that there are two $R_1(0)$ peaks for ν_6 , and errors in the measured temperature. However, given the difficulties in determining A'' spectroscopically, this might be the best way to measure the ground state $K=1$ and $K=0$ energy splitting.

Using this approach, we estimate the energy difference between the lowest *para* ($K=1$) and *ortho* ($K=0$) rotational states, or in other words the effective A'' constant. This analysis predicts a value of $2.2(6)\text{ cm}^{-1}$ and is compared with the spectroscopic constants in Table 2. This value suggests that propyne K -rotation is strongly perturbed. This analysis predicts that the A'' rotational constant is about 42% of the gas-phase value and therefore the population of the *para* nuclear spin isomer at equilibrium at 1.7 K is significantly greater than what is predicted using the gas-phase rotational constants. Given that we measure an equilibrium POR value of ~ 0.15 at 1.7 K, the approximate temperature where all the experiments were conducted, this implies that the elementary rate constants for the reversible first-order NSC kinetics are significantly different from the measured effective rate constant.⁵⁶ We can use the measured k_{eff} and $POR(1.7(1)\text{ K}, t=\infty)$ for five experiments (Expt. 3 was discarded) to determine $k_{p \rightarrow o} = 3.00(9) \times 10^{-3}\text{ min}^{-1}$ and $k_{o \rightarrow p} = 4.4(11) \times 10^{-4}\text{ min}^{-1}$.

Another way therefore to refine the POR determination would be to perform kinetic experiments at different temperatures. We should be able to measure different asymptotic values of POR (at equilibrium) as a function of temperature. All the experiments in the Wyoming laboratory were conducted at the deposition temperature because we did not want to complicate the analysis of the difference spectra by including changes due to annealing or oH_2 diffusion,

which can occur over comparable timescales relative to NSC at elevated temperatures.¹⁻⁴ However, the group in Orsay conducted experiments for samples that were equilibrated for longer times at 2.8 K, and were annealed at 4.0 K for brief periods of time. The group in Orsay uses a closed-cycle cryostat and thus is capable of monitoring the same sample for days. The group in Wyoming uses a bath cryostat and thus can only hold samples for approximately 10 hours without the need to refill with liquid helium. Spectra from the Orsay laboratory were analyzed using the same procedure and line strengths determined in Wyoming. The measured values from the two laboratories are presented in Figure 7. The lines presented in Figure 7 represent *PORs* calculated for single-axis *K*-rotation ($E = A''K^2$) with effective A'' constants equal to 1.00 cm⁻¹, 2.22 cm⁻¹, and 5.31 cm⁻¹, respectively. Error bars on each measurement are calculated using propagation of errors. The data from the two labs are somewhat consistent with greater *PORs* calculated at higher temperatures in Orsay. All the data lies between the two limiting values of $A'' = 1.00$ and 5.31 cm⁻¹. As can be seen in Figure 7, there is significant scatter in the measured *PORs*, but the level of agreement is promising given the difficulties in making intensity based measurements. We hope this work motivates similar measurements for different methyl rotor molecules and we want to continue to refine these measurements for propyne by conducting further experiments in both laboratories.

It is interesting to compare the NSC rate constant determined here for propyne with the one measured at 3.3 K for CH₃F isolated in solid pH₂.²¹ At 3.3 K, based on our analysis of propyne with $A''=2.2(6)$ cm⁻¹, we would predict a *POR* of approximately 0.3 which would imply $k_{eff}(3.3 \text{ K}) \approx 3.8 \times 10^{-3} \text{ min}^{-1}$ as compared to $2.2(5) \times 10^{-3} \text{ min}^{-1}$ for the fast NSC rate constant measured over the first 5 hours for CH₃F solvated in solid pH₂ at 3.3 K.²¹ Previous measurements of CH₃F NSC found bi-exponential behavior that the authors ascribed to fast oH₂

assisted NSC of CH_3F , followed by slower ($k = 3.7(8) \times 10^{-4} \text{ min}^{-1}$) NSC when the oH_2 concentration drops below a critical value.²¹ There is no evidence for bi-exponential behavior in the current NSC kinetics, but measurements over larger time windows would be helpful. Interestingly, the opposite behavior was found for NSC of ClCH_2 trapped in solid pH_2 where the conversion rate decreased by a factor of 1.5 for samples with 10% oH_2 .²⁷ All the measurements conducted in Wyoming were only performed for up to 8 hours. Preliminary results on longer timescales performed in Orsay show the time evolution of the three peaks marked with an asterisk in Figure 3 (and Figures S4 and S5), and at much longer timescales, two peaks decrease with time and one increases, meaning that the behavior at long times is more complex. An analysis of these peaks in terms of oH_2 clustering or different matrix sites must be performed to go further in this analysis.

4. CONCLUSIONS

We present here a detailed spectroscopic analysis of propyne isolated in solid pH_2 . This analysis shows how NSC can be used to rigorously assign the nuclear spin isomer responsible for a given rovibrational transition, provided that the molecules exhibit single-axis K rotational motion. Further, through the analysis of three separate perpendicular rovibrational bands, a detailed K rotational assignment is not possible that delivers a consistent set of ground state rotational constants. This demonstrates that the rotational levels of propyne solvated in solid pH_2 are not well described by simple one-dimensional free rotor expressions, and that quantum calculations that include intermolecular interactions with the quantum host are likely necessary for acceptable agreement with experiment. This data therefore contains detailed experimental information on how the quantum cavity influences the large amplitude rotational motion of a dopant, but the

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3 complication of how best to extract this information still remains. It is hoped that the procedures
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5 used in this study are applicable to a wide variety of molecules containing methyl rotors to
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7 permit systematic studies.
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10 The NSC kinetics of propyne in solid pH₂ show single-exponential decay indicative of a
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12 reversible first-order process. Careful analysis of three separate bands all lead to an average
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14 effective time constant of $\tau_{eff} = 287(7)$ min for NSC at 1.7 K. This time constant is comparable
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16 to the fast time constant extracted for CH₃F suspended in solid pH₂, but as discussed in the
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18 previous section, the NSC of propyne should be measured over longer time periods to see if it
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20 too shows bi-exponential behavior similar to CH₃F. Using intensity-intensity correlation plots,
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22 we are able to determine the relative line strengths of different propyne rovibrational transitions
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24 that allow the *POR* to be calculated. This data suggests that using the RVD technique we are
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26 able to trap ~75% of the higher rotational energy *para* nuclear spin isomer immediately post
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28 deposition. Further, by measuring the *POR* after full thermal equilibration, analysis from both
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30 labs suggests that the *A''* rotational constant is significantly reduced from its gas phase value. A
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32 reduction in the *A''* rotational constant would be suspected for a hindered rotor in solid pH₂ and
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34 thus further experiments could better determine this value and hopefully bring the predicted *A''*
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36 values estimated from spectroscopic measurements in the two laboratories into better agreement.
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38 We plan to study NSC in a series of molecules containing methyl groups in order to see if trends
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40 emerge and to further develop these analytical tools and procedures by testing them on a variety
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Table 1. Peak Positions and Widths (FWHM) in cm⁻¹ of Propyne (¹²C₃H₄) Trapped in Solid pH₂ at 1.7 K.

Mode	NSI ^a	$\Delta K_v(K'')$ ^b	gas ^c	pH ₂ (FWHM)	Shift	Ref.
2v ₁₀ ⁰ (A ₁)	<i>o/p</i>	Q ₂ (0)/Q ₂ (1)	650.352	665.44(0.82)	15.09	42
v ₅ (A ₁)	<i>o/p</i>	Q ₁ (0)/Q ₁ (1)	930.277	929.81(0.13)	-0.46	42
2v ₉ ⁰ (A ₁)	<i>o/p</i>	Q ₂ (0)/Q ₂ (1)	1254.347	1254.47(0.99)	0.12	44
v ₄ (A ₁)	<i>o/p</i>	Q ₁ (0)/Q ₁ (1)	1385.579	1384.09(1.04)	-1.49	41
2v ₈ ⁰ (A ₁)	<i>o/p</i>	Q ₂ (0)/Q ₂ (1)	2066.33	2058.65(0.50)	-7.68	57
v ₃ (A ₁)	<i>o/p</i>	Q ₁ (0)/Q ₁ (1)	2137.87	2138.34(0.04), m	0.47	57
(v ₃ +2v ₉) ⁺³ (A ₁)	<i>o/p</i>	Q _{1,2} (0)/Q _{1,2} (1)	2336.95	2310.35(1.25)	-26.60	34
v ₇ +v ₈ +v ₉ (A ₁)	<i>o/p</i>	Q _{1,1,1} (0)/Q _{1,1,1} (1)	2760	2763.71(0.77)	-23.40	58
2v ₄ (A ₁)?	<i>o/p</i>	Q ₂ (0)/Q ₂ (1)	2760	2751.99(3.65)	-8.01	58
2v ₇ ⁰ (A ₁)	<i>ortho</i>	Q ₂ (0)	2879	2870.8(1.1), f	-7.8	34
2v ₇ ⁰ (A ₁)	<i>para</i>	Q ₂ (1)		2871.4(0.9), f		
v ₂ (A ₁)	<i>o/p</i>	NR		2934.4, sh		
v ₂ (A ₁)	<i>para</i>	Q ₁ (1)	2940.944	2934.49(0.15), f	-6.46	35,45
v ₂ (A ₁)	<i>ortho</i>	Q ₁ (0)	2941.000	2934.60(0.16), f	-6.40	35,45
v ₃ +v ₅ (A ₁)	<i>o/p</i>	NR		3065.7, sh		
v ₃ +v ₅ (A ₁)	<i>para</i>	Q _{1,1} (1)	3070.123	3066.01(0.04)	-4.11	59
v ₃ +v ₅ (A ₁)	<i>ortho</i>	Q _{1,1} (0)	3070.141	3066.14(0.30)	-4.00	59
v ₁ (A ₁)	<i>o/p</i>	Q ₁ (0)/Q ₁ (1)	3335.066	3328.52(0.07), m	-6.54	35
v ₃ +2v ₉ ⁰ (A ₁)	<i>o/p</i>	Q _{1,2} (0)/Q _{1,2} (1)	3381.15	3377.93(1.09)	-3.22	60
2v ₂ (A ₁)	<i>ortho</i>	Q ₂ (0)	5781	5766.77(1.20), f	-14.23	37
2v ₂ (A ₁)	<i>para</i>	Q ₂ (1)		5769.10(1.11), f		
v ₂ +2v ₇ ⁰ (A ₁)	<i>para</i>	Q _{1,2} (1)		5815.14(1.61), f		
v ₂ +2v ₇ ⁰ (A ₁)	<i>ortho</i>	Q _{1,2} (0)	5828	5814.72(0.99), f	-13.28	37
2v ₆ ⁰ (A ₁)	<i>para</i>	Q ₂ (1)		5906.14(0.89), f		
2v ₆ ⁰ (A ₁)	<i>ortho</i>	Q ₂ (0)	5917.32	5906.40(0.85), f	-10.78	61
v ₁ +v ₂ (A ₁)	<i>o/p</i>	Q _{1,1} (0)/Q _{1,1} (1)	6275.841	6262.86(0.67)	-12.98	62
v ₁ +v ₃ +v ₅ (A ₁)	<i>o/p</i>	Q _{1,1,1} (0)/Q _{1,1,1} (1)	6399.463	6361.05(0.19)	-29.18	62
2v ₁ (A ₁)	<i>o/p</i>	Q ₂ (0)/Q ₂ (1)	6567.878	6555.13(0.17), m	-12.75	62
v ₁ +v ₃ +2v ₉ ⁰ (A ₁)	<i>o/p</i>	Q _{1,1,2} (0)/Q _{1,1,2} (1)	6660.222	6652.00(2.11)	-8.22	62
v ₉ (E)	<i>o/p</i>	$\tilde{\nu}_0$ /NR	638.575	633.47(0.76)	-5.11	42
v ₈ (E)	<i>para</i>	P ₁ (1)	1030.554	1029.76(0.11)	-0.79	42
v ₈ (E)	<i>o/p</i>	NR		1029.93(0.06)		
v ₈ (E)	<i>o/p</i>	$\tilde{\nu}_0$ /NR	1036.148	1031.30(0.17)	-4.85	42
v ₈ (E)	<i>ortho</i>	R ₁ (0)	1037.384	1034.8, sh	-2.58	42
v ₈ (E)	<i>ortho</i>	R ₁ (0)	1037.384	1035.05(0.22)	-2.34	42
v ₇ (E)	<i>para</i>	P ₁ (1)	1444.678	1443.54(0.45)	-1.14	41
v ₇ (E)	<i>o/p</i>	NR		1444.93(0.46)		
v ₇ (E)	<i>o/p</i>	$\tilde{\nu}_0$ /NR	1450.271	1446.04(0.39)	-4.24	
v ₇ (E)	<i>o/p</i>	NR		1451.46(0.61)		
v ₇ (E)	<i>ortho</i>	R ₁ (0)	1459.131	1454.67(0.12), sh	-4.46	41
v ₇ (E)	<i>ortho</i>	R ₁ (0)	1459.131	1455.12(1.06)	-4.01	41
v ₇ (E)	<i>para</i>	R ₁ (1)	1472.926	1473.88(4.60)	0.96	41
2v ₇ ⁺² (E)	<i>para</i>	P ₂ (1)		2887.17(0.62)		
2v ₇ ⁺² (E)	<i>ortho</i>	R ₂ (0)		2890.08(0.86)		
2v ₇ ⁺² (E)	<i>para</i>	R ₂ (1)		2898.63(0.44)		
v ₆ (E)	<i>para</i>	P ₁ (2)	2965.400	2963.15(1.49)	-2.25	46
v ₆ (E)	<i>para</i>	P ₁ (1)	2975.259	2971.48(0.60)	-3.77	46

1							
2							
3	$\nu_6(E)$	<i>o/p</i>	NR		2972.95(0.50)		
4	$\nu_6(E)$	<i>o/p</i>	$\tilde{\nu}_0$ /NR	2980.852	2975.96(0.53)	-4.90	46
5	$\nu_6(E)$	<i>o/p</i>	NR		2978.29(0.24)		
6	$\nu_6(E)$	<i>ortho</i>	$R_1(0)$	2985.666	2979.00(0.46)	-6.67	46
7	$\nu_6(E)$	<i>ortho</i>	$R_1(0)$	2985.666	2980.37(0.46)	-5.30	46
8	$\nu_6(E)$	<i>para</i>	$R_1(1)$	2995.479	2984.93(1.60)	-10.55	46
9	$\nu_6(E)$	<i>para</i>	$R_1(1)$	2995.479	2985.5, sh	-9.98	46
10	$\nu_6(E)$	<i>para</i>	$R_1(2)$	3005.266	2997.19(4.32)	-8.07	46
11	$\nu_3+\nu_8(E)$	<i>para</i>	$P_{1,1}(1)$	3170.484	3165.99(0.13)	-4.49	47
12	$\nu_3+\nu_8(E)$	<i>o/p</i>	NR		3166.17(0.06)		
13	$\nu_3+\nu_8(E)$	<i>o/p</i>	$\tilde{\nu}_0$ /NR	3176.077	3167.37(0.16)	-8.70	47
14	$\nu_3+\nu_8(E)$	<i>o/p</i>	NR		3170.90(0.13)		
15	$\nu_3+\nu_8(E)$	<i>ortho</i>	$R_{1,1}(0)$	3177.373	3171.31(0.20)	-6.07	47
16	$\nu_3+\nu_8(E)$	<i>para</i>	$R_{1,1}(1)$	3183.759	3175.82(0.31)	-7.94	47
17	$\nu_1+\nu_9(E)$	<i>para</i>	$P_{1,1}(1)$		3956.55(50)		
18	$\nu_1+\nu_9(E)$	<i>o/p</i>	$\tilde{\nu}_0$ /NR	3947.60	3957.97(0.32)	10.37	63
19	$\nu_1+\nu_9(E)$	<i>ortho</i>	$R_{1,1}(0)$		3962.04(0.94)		
20	$\nu_1+\nu_9(E)$	<i>para</i>	$R_{1,1}(1)$		3965.97(0.52)		
21	$\nu_2+\nu_6^{\pm 1}(E)$	<i>o/p</i>	NR?		5780.17(1.49)		
22	$\nu_2+\nu_6^{\pm 1}(E)$	<i>para</i>	$P_{1,1}(1)$		5784.68(0.18)		
23	$\nu_2+\nu_6^{\pm 1}(E)$	<i>o/p</i>	$\tilde{\nu}_0$	5790			37
24	$\nu_2+\nu_6^{\pm 1}(E)$	<i>ortho</i>	$R_{1,1}(0)$		5789.85(7.48)		
25	$\nu_6^{\pm 1}+2\nu_7^0(E)$	<i>para</i>	$P_{1,2}(1)$		5835.37(0.07)		
26	$\nu_6^{\pm 1}+2\nu_7^0(E)$	<i>o/p</i>	$\tilde{\nu}_0$	5852			37
27	$\nu_6^{\pm 1}+2\nu_7^0(E)$	<i>ortho</i>	$R_{1,2}(0)$		5841.14(1.05)		
28	$2\nu_6^{\pm 2}(E)$	<i>para</i>	$P_2(1)$		5938.42(0.75)		
29	$2\nu_6^{\pm 2}(E)$	<i>o/p</i>	NR		5939.62(0.42)		
30	$2\nu_6^{\pm 2}(E)$	<i>o/p</i>	$\tilde{\nu}_0$ /NR	5953	5941.89(0.28)	-11.11	37
31	$2\nu_6^{\pm 2}(E)$	<i>o/p</i>	NR		5945.38(0.48)		
32	$2\nu_6^{\pm 2}(E)$	<i>ortho</i>	$R_2(0)$	5958	5947.05(0.82)	-10.95	63
33	$2\nu_6^{\pm 2}(E)$	<i>ortho</i>	$R_2(0)$	5958	5948.27(0.37)	-9.73	63
34	$2\nu_6^{\pm 2}(E)$	<i>para</i>	$R_2(1)$		5792.66(0.46)		
35	$\nu_1+\nu_6^{\pm 1}(E)$	<i>para</i>	$P_{1,1}(1)$		6284.58(0.03)		
36	$\nu_1+\nu_6^{\pm 1}(E)$	<i>o/p</i>	$\tilde{\nu}_0$ /NR	6315.786	6286.61(0.15)	-29.18	62
37	$\nu_1+\nu_6^{\pm 1}(E)$	<i>ortho</i>	$R_{1,1}(0)$		6288.82(0.14)		
38							
39							

^aNuclear Spin Isomer: *ortho* and *para* spin isomers; *o/p* indicates unresolved contributions. ^bNR stands for non-rotating; see text for details. ^cTaken from literature values or calculated using literature values and ground state spectroscopic constants of Ref. 64. f = value from fit; m = multiple peaks (principal peak listed), sh = shoulder.

Table 2. Parameters Determined from Observed Transition Frequencies and *POR*.

Parameter	$\nu_6(E)$	$\nu_7(E)$	$\nu_3+\nu_8(E)$	$\nu_6(E)$, <i>POR</i>
$\tilde{\nu}_0 / \text{cm}^{-1}$	2976.59	1445.50	3170.53	---
A' / cm^{-1}	3.62	5.77	4.13	---
ζ_i	0.07250 ^a	-0.31424 ^a	0.4052 ^a	---
A'' / cm^{-1}	5.10	1.96	4.54	2.22(55)
K'' values	0, 1	0, 1	0, 1	---

^a Fixed to gas-phase values: for ν_6 see Ref. 46, for ν_7 see Ref. 41, and for $\nu_3+\nu_8$ see Ref. 47.

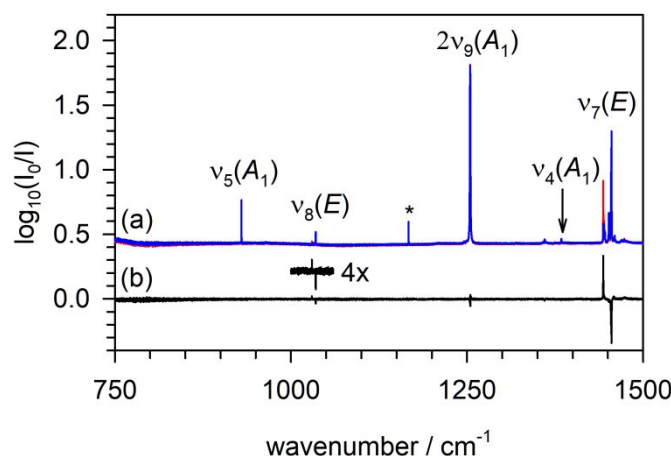


Figure 1. Infrared absorption spectra of propyne trapped in solid pH_2 at 1.70(1) K showing propyne absorptions in the 750-1500 cm^{-1} region. (a) Spectra recorded 2.63 (red) and 491 min (blue), respectively, after deposition of a 2.5(2) mm thick, 200(39) ppm propyne/ pH_2 sample. (b) Difference spectrum ($A_t - A_\infty$) calculated using the two spectra (before – after) displayed in trace (a) showing changes in peak intensities with time. The inset above (b) shows the difference signal multiplied by 4. The peak marked with * is the $U_0(0)$ transition of solid pH_2 . The $\nu_8(E)$ and $\nu_7(E)$ absorptions display peaks in the difference spectrum consistent with *para-to-ortho* NSC after deposition. See text for details.

(single column)

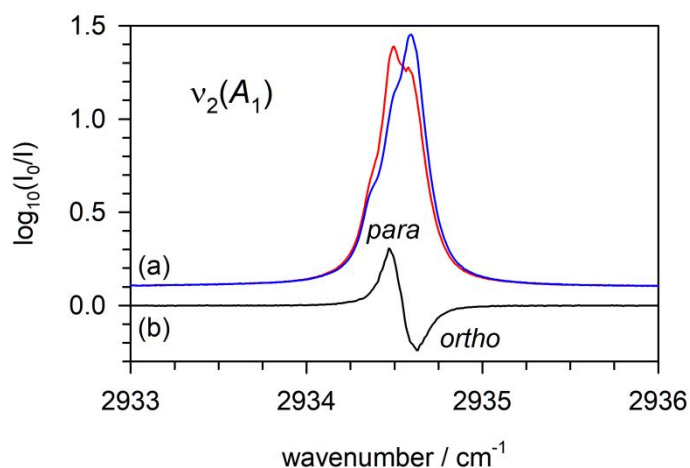


Figure 2. Infrared absorption spectra of propyne in solid pD₂ at 1.72(1) K in the region of the $\nu_2(A_1)$ fundamental. (a) Spectra recorded 5.08 (red) and 291.8 min (blue), respectively, after deposition of the sample shown in Figure 1. (b) Difference spectrum ($A_t - A_\infty$) calculated using the two spectra displayed in trace (a) showing changes in absorption intensity with time after deposition. The positive and negative peaks are labeled *para* and *ortho*, respectively, indicating the nuclear spin symmetry in the ground vibrational state associated with each peak. See text for details.

(single column)

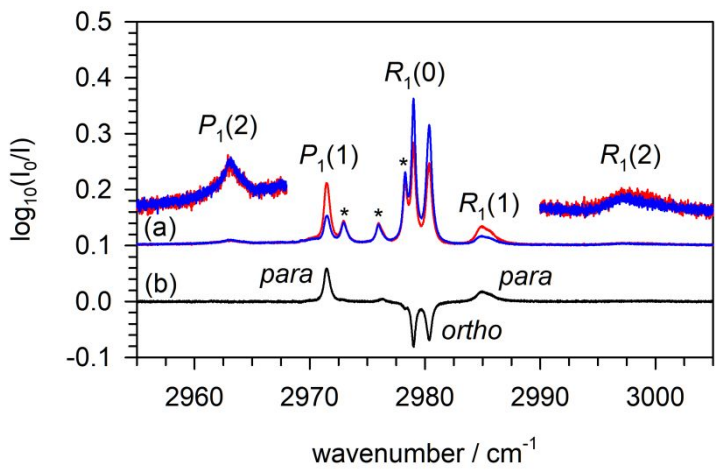


Figure 3. Infrared absorption spectra of propyne in solid pH₂ at 1.72(1) K in the region of the $\nu_6(E)$ fundamental. (a) Spectra recorded 5.08 (red) and 291.8 min (blue), respectively, after deposition of the sample shown in Figure 1. (b) Difference spectrum ($A_t - A_\infty$) calculated using the two spectra displayed in trace (a) showing how the absorption intensity changes with time after deposition. The positive and negative peaks are labeled *para* and *ortho*, respectively, indicating the nuclear spin symmetry in the ground vibrational state associated with each peak. Peaks marked with an asterisk do not show up in the difference spectrum. See text for details.

(single column)

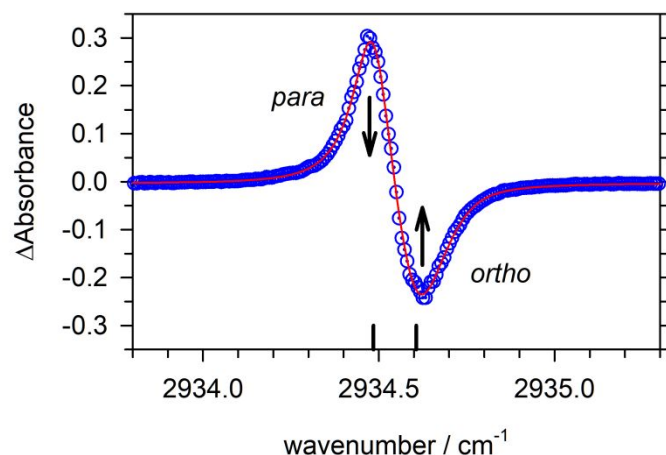


Figure 4. Fits of the ν_2 difference spectrum of propyne in solid pD_2 calculated using spectra recorded 5.08 and 291.8 min after deposition of the sample shown in Figure 1. The data are the blue circles and the results of a least squares fit to an analytic function is shown as a red line. The two black lines represent the average peak positions determined from fits to all the spectra for this kinetic run.

(single column)

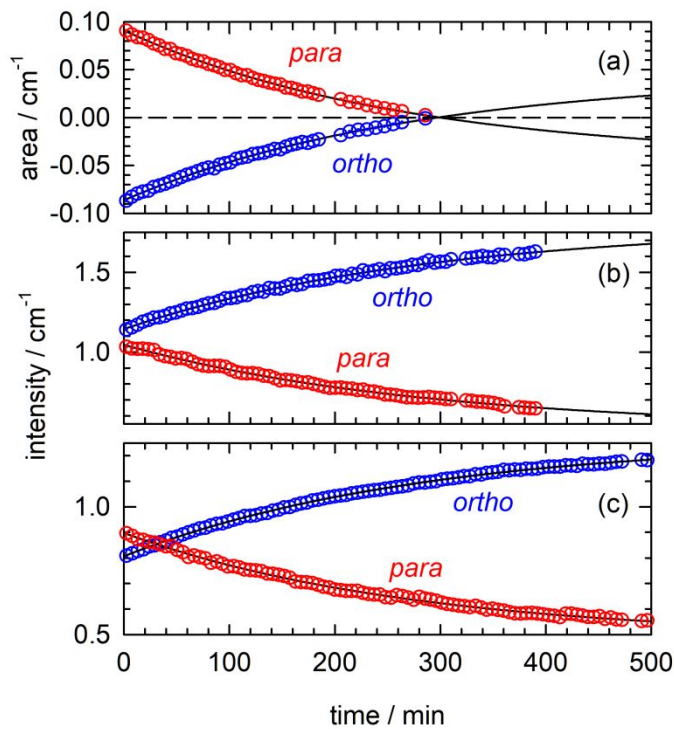


Figure 5. Representative kinetic traces of propyne NSC for propyne isolated in solid pH_2 at 1.7 K. (a) The red and blue data points correspond to the areas of the *para* and *ortho* peaks in the ν_2 difference spectrum of propyne for the sample shown in Figure 1. Integrated intensities of the *ortho* and *para* peaks assigned to (b) ν_6 and (c) ν_7 absorption bands for samples containing 289(16) and 200(39) ppm of propyne, respectively. The red and blue data points in (b) and (c) correspond to the integrated intensities of the *para* and *ortho* peaks.

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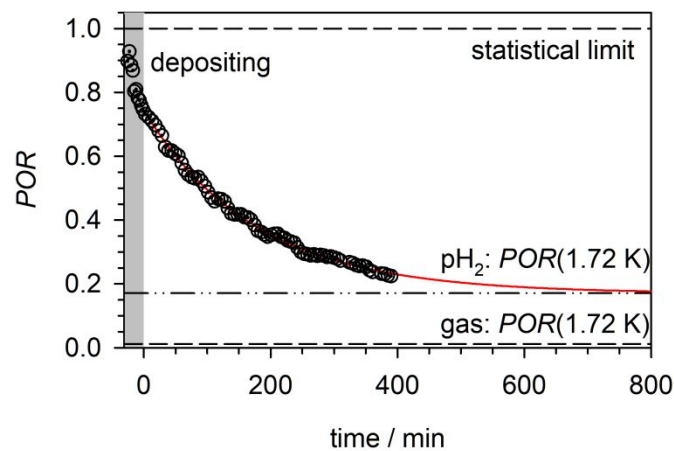


Figure 6. Representative kinetic trace (data shown in Fig. 5b) of the propyne *para*-to-*ortho* ratio (*POR*) measured as a function of time after deposition for propyne isolated in solid pH₂ at 1.7 K.

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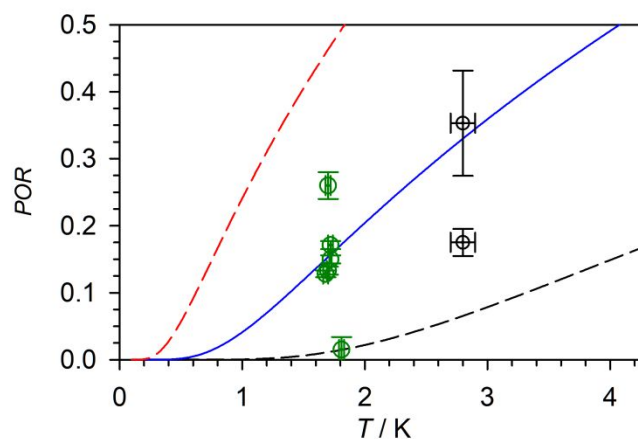


Figure 7. Plot of the equilibrium *POR* versus temperature (K) measured in this study. Data from samples grown in Wyoming and Orsay are plotted as green and black circles, respectively. The lines represent theoretical *POR* curves calculated for single-axis *K*-rotation with effective *A''* constants equal to 1.00 cm⁻¹ (red dashed line), 2.22 cm⁻¹ (blue line), and 5.31 cm⁻¹ (black dashed line). All the measured values lie between the two limiting cases and are best represented by *A''*=2.22 cm⁻¹. See text and [Supporting Information](#), Tables S6 and S7 for more details.