Internal Rotation in Propionic Acid: Near-Infrared-Induced Isomerization in Solid Argon

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The conformational system of propionic acid (CH₃CH₂COOH) is studied in solid argon. It is predicted by the ab initio calculations that this molecule has four stable conformers. These four structures are denoted Tₜ, Tₕ, Cₜ, and Cₕ, and they differ by the arrangement around the C=O and Cₐ–C bonds. The ground-state Tₜ conformer is the only form present at 8 K after deposition of an argon matrix containing propionic acid. For the CH₃CH₂COOH and CH₃CH₂COOD isotopologues, narrow-band excitation of the first hydroxyl stretching overtone of the conformational ground state promotes the Cₐ–C and C=O internal rotations producing the Tₕ and Cₜ conformers, respectively. A subsequent vibrational excitation of the produced Tₕ conformer by rotation around the C=O bond. In the dark, all of the produced conformers decay to the conformational ground state at different rates. The decay kinetics and its temperature dependence allow the identification of the conformers by IR absorption spectroscopy, which is supported by ab initio calculations of their vibrational spectra. For the CH₃CH₂COOD isotopologue, the excitation of molecules isolated in different matrix sites results in site-dependent photoisomerization rates for the Cₐ–C and C=O internal rotations, which also confirm the identification of the photoproducts.

Introduction

In a recent series of papers, we studied the IR-induced rotational isomerization of formic and acetic acids isolated in low-temperature inert matrices.¹⁻⁷ These studies have been mainly motivated by our interest in the conformational dynamics of the carboxylic group as a way to understand the process of intramolecular vibrational energy relaxation, which plays an important role in molecular reactivity. Interestingly, the photolysis of formic acid was shown to be conformationally dependent, being acknowledged as one of the first clear cases of optical control of chemical reactivity.⁴⁻⁸ The study of the conformational properties of simple carboxylic acids can improve our knowledge of more complex molecular systems with biological interest, where the conformation of this functional group plays an important role in determining their biological activity.⁹⁻¹¹ Furthermore, an understanding of the conformational properties of small molecules and their response to photochemical stimuli is essential to the evaluation of the potential use of more complex molecules with biconformational structure as molecular switches controlled by light.¹²,¹³

Compared to acetic acid, propionic acid (propanoic acid, CH₃CH₂COOH) has one additional internal rotational degree of freedom. Besides the methyl and hydroxyl rotations occurring in acetic acid, the internal rotation around the Cₐ–C bond should be taken into account in the case of propionic acid. Experimentally, only one conformer of monomeric propionic acid has been unequivocally characterized.¹⁴⁻¹⁶ Both gas-phase electron diffraction and microwave spectroscopy studies have shown that this conformer exhibits a planar heavy-atom backbone, with a staggered arrangement around the Cₐ–C bond and trans arrangements around the Cₐ–C and C=O bonds (C=O–C–O and C–C–O–H dihedral angles of 180°).¹⁴⁻¹⁶ Moreover, the electron diffraction results suggested that, in addition to the identified conformer, structures bearing a nonplanar C–C–O–H skeleton should also contribute to the gas-phase equilibrium conformational distribution with an estimated population of ca. 40% (T = 488 K).¹⁴

The earlier computational studies on the conformational isomerism in propionic acid, undertaken at the ab initio Hartree–Fock (HF) level of theory with the relatively modest 6-31G* basis set¹⁷ and using the molecular mechanics (MM) approach,¹⁸ predicted two stable arrangements with respect to the Cₐ–C bond, the trans and gauche (C–C–O–H dihedral angle of ca. ±60°) arrangements. The doubly degenerate gauche arrangement has been estimated by MM to be 490 cm⁻¹ higher in energy than the trans arrangement,¹⁸ whereas the HF/6-31G* calculations predicted a slightly smaller gauche–trans energy difference (~340 cm⁻¹).¹⁷ The trans–gauche barrier estimated by the HF/6-31G* calculations was ~420 cm⁻¹,¹⁷ which was in good agreement with a more recent value of 490 cm⁻¹ obtained using the density functional theory (DFT) method with the B3LYP functional and the 6-311+G(2d,2p) basis set.¹⁹ The relative stability of these two Cₐ–C arrangements can be compared with other carbonyl compounds structurally related to propionic acid, such as propionyl fluoride and propionyl chloride (CH₃CH₂COX, with X = F and Cl), which have been studied previously.²⁰,²¹ These molecules were shown to be stable in the trans and gauche conformations with respect to the Cₐ–C bond (C–C–C–X dihedrals of 180 and ca. ±60°, respectively).²⁰,²¹ In solutions of liquid rare gases, the energy differences between

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the trans and the gauche conformers and the trans \(\rightarrow\) gauche isomerization barrier were found to be 329 and 680 cm\(^{-1}\) in propionyl fluoride and 505 and 820 cm\(^{-1}\) in propionyl chloride, respectively.\(^{22,24}\) For chloroacetic acid (CH\(_2\)ClCOOH), the gauche arrangement around the \(\text{C}^–\text{O}\) bond was estimated to be 120–400 cm\(^{-1}\) higher in energy than the trans form, with a trans \(\rightarrow\) gauche isomerization barrier of ca. 500 cm\(^{-1}\) for the monomer isolated in rare-gas matrices.\(^{22}\)

Computationally, in addition to the two stable arrangements around the \(\text{C}^–\text{C}\) bond, propionic acid has two stable conformations differing in the arrangement around the \(\text{C}–\text{O}\) bond, corresponding to the planar trans and cis configurations characterized by \(\text{C}–\text{C}–\text{O}–\text{H}^\dagger\) dihedrals of 180 and 0\(^\circ\), respectively.\(^{18,19}\) Similarly to the situation in formic and acetic acids,\(^{1,5}\) in propionic acid the trans arrangement of the carboxylic group is the lowest-energy configuration. According to MM and DFT/B3LYP calculations, the energy difference between the two configurations is ca. 1700–2100 cm\(^{-1}\).\(^{18,19}\) The trans to cis barrier associated with internal rotation around the \(\text{C}–\text{O}\) bond estimated by the same methods is 4300–4600 cm\(^{-1}\).\(^{19}\) These results are in good agreement with the data for formic and acetic acids (energy difference of \(\sim 1400–1800\) cm\(^{-1}\) and a trans to cis barrier of \(\sim 3900–4400\) cm\(^{-1}\)).\(^{25–28}\) For the matrix-isolated monomers of formic and acetic acids, the cis conformer is short-lived, even at 8 K, because of its conversion to the trans conformer by tunneling with a rate of \(\sim 10^{-2}–10^{-3} \text{s}^{-1}\), which complicates the experimental characterization of the cis conformer of these molecules. A similar phenomenon can also be expected for propionic acid.

In the present study, the rotational isomerization of monomeric propionic acid isolated in solid argon was investigated. The ground-state potential energy surface (PES) and the vibrational spectra of its different conformers were calculated at the MP2/6-311G\(^{++}\)(2d,2p) level of theory. The computational results support the analysis of the experimental data obtained using the low-temperature matrix-isolation technique combined with narrow-band pumping of individual vibrational transitions.

**Experimental and Computational Details**

The gaseous samples were prepared by mixing propionic acid (>99%), degassed by several freeze-pump-thaw cycles, with high-purity argon (99.9999%), in a 1:1000 or 1:500 ratio. The CH\(_3\)CH\(_2\)COOD species was obtained from the fully hydrogenated isotopologe by HD exchange on both the inner surface of the sample container and the deposition line saturated with D\(_2\)O. The OH and OD isotopologues of propionic acid are expected for propionic acid.

In agreement with previous theoretical predictions,\(^{15–19}\) the present ab initio calculations give four nonequivalent minima on the PES of propionic acid. The geometries of the four conformers, their relative energies, and the barriers to internal rotation around the \(\text{C}^–\text{O}\) and \(\text{C}^–\text{C}\) bonds are shown in Figure 1. The potential energy profiles for conformational interconversion are shown in Figure 2. Only one possible arrangement around the \(\text{C}^–\text{C}\) bond is stable, which minimizes steric repulsion between the methyl and methylene hydrogen atoms. In the present article, the stable arrangements

![Figure 1. Conformers of propionic acid. Shown are the ab initio [MP2/6-311++G(2d,2p)] relative conformational energies and energy barriers for the internal rotation around the \(\text{C}^–\text{C}\) and \(\text{C}–\text{O}\) bonds (in cm\(^{-1}\)).](image)
with respect to the C–O bond (trans and cis) are denoted by the capital letters T and C, respectively, and the small letters refer to the arrangements around the Cα–C bond (t for the trans and g for the two degenerate gauche arrangements).

As generally found for simple carboxylic acids,

\[ \text{cis arrangement around the C–C bond is higher in energy than the trans configuration by ca. 1800 cm}^{-1} \]

and the isomerization barrier for conversion of the trans to the cis arrangement is ca. 4000 cm\(^{-1}\) (\(E_{\text{T}t\rightarrow\text{C}t}\), \(E_{\text{T}c\rightarrow\text{C}c}\)).

The calculated energy differences between the conformers agree with the earlier computational studies, whereas our T–C barrier is somewhat lower than the results at a lower level of approximation.

In agreement with known experimental and computational results, the most stable arrangement around the Cα–C bond corresponds to the trans configuration, whereas the degenerate gauche configurations (C=C–C=O dihedral angle of approximately \(\pm 75^\circ\)) are somewhat higher in energy than the corresponding trans forms (\(\Delta E_{\text{g}t\rightarrow\text{C}t}\)).

The calculated energy difference between the gauche and trans Cα–C arrangements in propionic acid is similar to the corresponding values in lipids, saturated hydrocarbons (150–400 cm\(^{-1}\)), and other \(\alpha\)-substituted carbonyl compounds such as propionyl fluoride, propionyl chloride, and chloroacetic acid. The energy barrier predicted for the trans \(\rightarrow\) gauche Cα–C internal rotation in propionic acid is closer to those found in propionyl fluoride and propionyl chloride (680–820 cm\(^{-1}\)) than to the average values reported for saturated hydrocarbons (1000–1400 cm\(^{-1}\)).

Direct interconversion between the two degenerate gauche conformations of the C–C–C–O moiety (i.e., the \(C_g^+ \rightarrow C_g^-\) or \(C_t^+ \rightarrow C_t^-\) process) can also take place (Figure 2). The transition state for these processes corresponds to the syn configuration around the Cα–C bond. The calculated syn barriers for the \(C_g^+ \rightarrow C_g^-\) and \(C_t^+ \rightarrow C_t^-\) processes are \(\approx 180\) and 100 cm\(^{-1}\), respectively. These values are somewhat smaller than those obtained previously at a lower level of theory (240 cm\(^{-1}\)).

The direct conversion between the conformers differing by internal rotation about both C–O and Cα–C bonds (the \(C_g^+/T_t^+\) and \(C_T^+/T_g^+\) pairs of conformers, see Figure 1) can be also estimated. The second-order transition state associated with the simultaneous rotation along the two torsional coordinates (Cα–C=O) was found in the 2D potential energy curve at \(\approx 4740\) cm\(^{-1}\) above the conformational ground state. Interestingly, in this second-order transition state, the C–C–C–O and C–C–O–H dihedral angles agree with the first-order transition states found along each of the torsional coordinates (116 and 83\(^\circ\), respectively).

2. Experimental Results. The T\(_t\) conformer is the only form present in the as-deposited matrix. For both PA-OH and PA-OD, the spectrum of T\(_t\) shows the splitting of bands due to the interaction of the isolated molecule with inhomogeneous local surroundings (matrix site effect). For PA-OH, the absorption bands of the fundamental and first overtone of the hydroxyl stretching mode (\(\nu\text{OH}\) and \(2\nu\text{OH}\), respectively) are shown in Figure 3. Conformational isomerization was induced site selectively by using tunable narrow-band radiation as found for HONO\(^{22}\) and formic acid.\(^{23}\) The site selectivity upon excitation of the \(2\nu\text{OH}\) mode of PA-OH is demonstrated in Figure 3, which shows holes burned in the \(\nu\text{OH}\) absorption envelope of T\(_t\) upon pumping at four different frequencies.

When the deposited PA-OH(D)/Ar matrix is irradiated at \(\approx 6960\) cm\(^{-1}\) (\(2\nu\text{OH}\)) or \(\approx 5170\) cm\(^{-1}\) (\(2\nu\text{OD}\)), the bands of the T\(_t\) conformer decrease, and new bands emerge. The light-induced spectral changes are due to conformational isomerization induced by vibrational excitation of the T\(_t\) conformer. The emerging bands can be divided into two sets assigned to the \(T_g^\pm\) and \(C_t\) forms (see later). The two sets of bands are distinguished by their behavior upon IR pumping and dark decay kinetics at 8–15 K.

In the case of PA-OH, the spectral changes induced by excitation of the T\(_t\) form are shown in trace a of Figure 4. Trace b illustrates the changes taking place after several minutes in the dark. One set of light-induced bands quickly decreases (\(k \approx 6 \times 10^{-3} \text{ s}^{-1}\)) after interrupting the pumping, whereas the other set is much more stable (\(k \approx 10^{-6} \text{ s}^{-1}\)). The quickly decaying bands (marked with dots in Figure 4) originate from the \(C_t\) conformer, and the more stable bands belong to the \(T_g^\pm\) form. In both cases, the decay of the photoproduced conformers regenerates the T\(_t\) form. Remarkably, the \(T_g^\pm \rightarrow T_t\) recovering process is strongly enhanced at higher temperatures as shown.
in Figure 5 (1 order of magnitude variation in the rate constant upon changing between 8 and 12 K).

Excitation of the $\nu_{\text{OH}}$ mode of the $T_g^\conform$ conformer (at $\sim6950$ cm$^{-1}$) promotes its conversion back to the $T_t$ form and its isomerization to a new species identified as the $C_g^\conform$ (see trace c in Figure 4). The photoinduced $T_g^\conform \rightarrow T_t$ conversion is a very efficient process, transferring ca. 70% of the molecules from the $T_g^\conform$ to the $T_t$ state in a few minutes of irradiation. The bands of the $C_g^\conform$ conformer (marked with an asterisk in Figure 4) can be observed only during the pumping of $T_g^\conform$ because they quickly decay in the dark. The decrease of the $C_g^\conform$ bands is accompanied with the growth of the $T_t$ bands, showing that this conformer decays to the most stable conformer. The fast dark decay of both $C_t$ and $C_g^\conform$ conformers is due to a phonon-assisted tunneling process, which has been shown to be common for internal rotation around the C–O bond in simple carboxylic acids.\textsuperscript{3,7}

For the deuterated isotopologue (PA-OD), the $C_t$ and $T_g^\conform$ conformers produced by excitation of the $T_t$ form are quite stable at 8 K. In fact, when compared to PA-OH, the $C_t$ to $T_t$ conversion by tunneling slows down by ca. 4 orders of magnitude. Nevertheless, the spectral signatures of the two photoproduced conformers could also be reliably separated for this isotopologue. First, the emerging bands of the two photoproduced conformers grow at different rates when pumping at different frequencies within the $2\nu_{\text{OH}}$ absorption of the $T_t$ conformer (high-frequency pumping (HF) at $5175$ cm$^{-1}$ and low-frequency pumping (LF) at $5170$ cm$^{-1}$). These results are shown in Figures 6 and 7 (trace a in both Figures) as difference spectra, where the $C_t$ and $T_g^\conform$ bands increase and the $T_t$ bands decrease. Second, the two conformers decay at different rates in the dark upon annealing. The $T_g^\conform \rightarrow T_t$ isomerization at 15 K is faster at least by 2 orders of magnitude than the $C_t \rightarrow T_t$ process (see trace c in Figure 7). Finally, it is also possible to selectively excite the $2\nu_{\text{OD}}$ mode of the $T_g^\conform$ conformer (which appears to be slightly redshifted from the $2\nu_{\text{OD}}$ absorption of $T_t$). The excitation of $T_g^\conform$ decreases its bands, a new set of bands emerges that can be assigned to the $C_g^\conform$ conformer, and the $T_t$ form is partially recovered. Without the pumping of $T_g^\conform$, the $C_g^\conform$ conformer converts to the most stable $T_t$ conformer. As for the $C_t \rightarrow T_t$ tunneling, $C_g^\conform \rightarrow T_t$ is much slower in PA-OD than in PA-OH (roughly by 2 orders of magnitude).

Tables 1 and 2 present the observed frequencies for the four conformers of matrix-isolated PA-OH and PA-OD together with...
the corresponding calculated [MP2/6-311++G(2d,2p)] frequencies, intensities, and PEDs.

Discussion

1. Conformational Cooling during Sample Deposition.

Taking into consideration the calculated relative energies of the four conformers of propionic acid, the populations of the $T_t$ and $T_g^\pm$ conformers are expected to be approximately 73 and 27% at room temperature (accounting for the degeneracy of the $T_g$ forms), whereas those of the $C_t$ and $C_g^\pm$ should be below 1%. Note that at 488 K the estimated populations of the $T_t$ and $T_g^\pm$ conformers increase, and the $T_t$ bands decrease.

In the 3300–1100 cm$^{-1}$ spectral window available for observation of the $C_t$ conformer, the bands assigned to this form are observed at ~1800 cm$^{-1}$ ($\nu(C=O)$), ~1360 cm$^{-1}$ ($\omega(CH_2)$), and ~1156 cm$^{-1}$ ($\nu(C=O)$), in good agreement with the computational values of 1773, 1361, and 1134 cm$^{-1}$, respectively (Figure 4). In addition, two doublets are observed at ~1276 and ~1248 cm$^{-1}$, which are ascribed to $\delta(CO)$, predicted at 1254 cm$^{-1}$. The fine structure observed for the $\nu(C=O)$ and $\omega(CH_2)$ vibrations of $C_t$, as well as for the bands ascribed to $\delta(CO)$ Table 1), is due to Fermi resonance with the $\gamma(CH_2)$ vibrations of $C_t$. Therefore, if the low-temperature matrix could efficiently freeze the equilibrium population of the deposited gaseous conformational mixture, it should have been possible to detect the presence of both $T_t$ and $T_g^\pm$ forms in the deposited matrix. However, for both PA-OH and PA-OD, only one conformer was found to be present in the deposited matrixes, whose spectral signature closely matches that predicted for the most stable $T_t$ conformer (Figures 5 and 6 and Tables 1 and 2). The absence of the $T_g^\pm$ conformer can be attributed to its fast conversion to the lowest-energy conformer during deposition at 15 K. This conformational cooling is commonly associated with low isomerization barriers ($\sim$400 cm$^{-1}$). Indeed, the computational barrier for the $T_g^\pm \rightarrow T_t$ isomerization is $\sim$60 cm$^{-1}$, which supports the fact that the $T_g^\pm$ form, presumably present in the gaseous mixture, can easily decay to the ground conformational state. In agreement with this interpretation, we observed the decay of the photoproduced $T_g^\pm$ conformer back to $T_t$ by an over-barrier process with a strong temperature dependence, which is consistent with the calculated $T_g^\pm \rightarrow T_t$ isomerization barrier.

2. Assignment of the Spectra of PA-OH. The spectra of the $T_t$ and $T_g^\pm$ conformers are compared in Figure 5. The difference spectrum shown by trace a was obtained by subtracting the spectrum of the deposited matrix from the spectrum recorded after several minutes of irradiation of $T_t$ and 1 min in the dark. The dark period leads to a decay of the $C_t$ conformer to the $T_t$ form, hence this spectrum shows only the result of the $T_t \rightarrow T_g^\pm$ photoprocess. A comparison of trace a with the calculated spectra of $T_t$ and $T_g^\pm$ shown in trace b demonstrates good general agreement between the experiment and theory. This agreement makes the assignment of the spectrum of both the $T_t$ form present in the deposited matrix and the photoproduced $T_g^\pm$ form straightforward. However, two discrepancies between the calculated and experimental spectra of $T_g^\pm$ can be noticed. First, none of the four bands of $T_g^\pm$ predicted to appear in the 1400–1200 cm$^{-1}$ region are clearly observed experimentally, and the absence of the medium-intensity $\nu(C-O)$ band predicted at 1350 cm$^{-1}$ is especially remarkable. Second, the high-intensity band predicted at 1174 cm$^{-1} (\delta(CO))$ appears as a doublet at 1174 and 1157 cm$^{-1}$. The first discrepancy is probably due to the computational overestimation of the band intensities and/or to the broadening of the experimental bands in this spectral region, which makes them difficult to discriminate from the background. Indeed, four broad weak bands are seen in the 1400–1200 cm$^{-1}$ region that can correspond to the apparently missing $T_g^\pm$ vibrations (Table 1). However, it is highly probable that the observed doublet at 1174/1157 cm$^{-1}$ is due to a Fermi resonance between $\delta(CO)$ and the first overtone of the C–O torsion whose fundamental is observed at 584 cm$^{-1}$. Similar splitting of the $\nu(C=O)$ or $\delta(CO)$ bands due to Fermi resonance was reported for formic and acetic acids.

In the 3300–1100 cm$^{-1}$ spectral window available for observation of the $C_t$ conformer, the bands assigned to this form are observed at ~1800 cm$^{-1} (\gamma(C=O))$, ~1360 cm$^{-1} (\omega(CH_2))$, and ~1156 cm$^{-1} (\nu(C=O))$, in good agreement with the computational values of 1773, 1361, and 1134 cm$^{-1}$, respectively (Figure 4). In addition, two doublets are observed at ~1276 and ~1248 cm$^{-1}$, which are ascribed to $\delta(CO)$, predicted at 1254 cm$^{-1}$. The fine structure observed for the $\nu(C=O)$ and $\omega(CH_2)$ vibrations of $C_t$, as well as for the bands ascribed to $\delta(CO)$ (Table 1) is presumably due to matrix site effects. However, the large splitting between the two $\delta(CO)$ doublets (~30 cm$^{-1}$) might have a different origin. A possible explanation for this observation is a Fermi interaction with the $\gamma(CH_2)$
TABLE 1: Experimental and Calculated [MP2/6-311++(2d,2p)]* Frequencies for the Four Conformers of the CH₂CH₂COOH Monomer

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</tbody>
</table>

*Only the bands affected by the NIR excitation experiments are here listed. The majority of the observed vibrational modes exhibit a site splitting of the absorption. The ab initio harmonic frequencies are rescaled using the 0.94 and 0.97 correction factors for the regions above and below 2000 cm⁻¹, respectively. The calculated intensities were normalized by the intensity of the strongest band (C₁ conformer at 1254.4 cm⁻¹). In general, symmetry coordinates with contributions higher than 15% to a particular vibrational mode are shown. Symbols: ν - stretching; δ - bending; γ - rocking; ω - wagging; τ - twisting; τ - torsion; and FR - involved in Fermi resonance. (See details in the text.)

3. Assignment of the Spectra of PA-OD. The assignment of the bands of the most stable conformer in the deuterated isotopeology is straightforward because of the generally good agreement between the experimental and calculated spectra for this form. The assignment of the T₂ and C₁ conformers relies to a great extent on the results of the irradiation experiments where the pumping frequency was varied. As already mentioned, two pumping frequencies were used to excite the T₁ conformer: high-frequency pumping (HF) at ~5175 cm⁻¹ and low-frequency pumping (LF) at 5170 cm⁻¹ (Figures 4 and 7).

The νOD and νC=O spectral regions are useful for the identification of both the T₂ and C₁ forms. These two modes are sensitive to the conformation of the carboxylic group. The νOD and νC=O modes of C₁ appear to be blue shifted from the T₁ bands by 46 and 30 cm⁻¹, respectively (Table 2). The calculations predicted these shifts at 47 and 30 cm⁻¹, in excellent agreement.
agreement with the experimental results. The observed shifts are also in agreement with the available data for acidic acid, where a change in conformation from the trans to the cis arrangement around the C=O bond leads to blue shifts in these modes of 44 and 30 cm⁻¹, respectively. In the case of T₉⁺, both the νOD and νC=O modes were predicted to be shifted from those of T₁ by less than 1 cm⁻¹. Accordingly, the bands assigned to this conformer are observed at 2627 (νOD) and 1780–1760 cm⁻¹ (νC=O, quartet), which are close to the corresponding bands of T₁ observed at 2630/2631 and 1770–1762 cm⁻¹, respectively.

Most of the T₈⁺ and C₆ bands appear to be split because of matrix-site effects. For each conformer, the positions of the emerging bands remain unchanged for the two pumping experiments (HF and LF), as shown in Figures 6 and 7, but their relative intensity changes with the pumping frequency. This is due to the fact that the pumping efficiency of various matrix sites depends on the pumping frequency, as discussed elsewhere. However, very interestingly, the HF excitation of the νC=O(55) features originating from molecules isolated in various sites.

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For example, there are three C-S absorptions in the 1500–1100 cm⁻¹ region affected essentially by the LF excitation (Figure 6). These are the band at 1368 cm⁻¹ (ωCH₂, predicted to be a medium-intensity band at 1360 cm⁻¹), the weak doublet at ~1263 cm⁻¹ (twCH₂, predicted to be a very weak band at 1274 cm⁻¹), and the very strong doublet observed at ~1244 cm⁻¹ (νC−O, predicted at 1178 cm⁻¹ to be the most intense band of the spectrum of this conformer). In the same spectral window, the only band of C-S clearly connected with the HF excitation is the relatively strong band observed at 1194 cm⁻¹, which is assigned to the νC−O mode of the site mostly influenced by the HF excitation (Figure 7). The Tₙ bands observed at 1380/1377 and 1295/1267 cm⁻¹ are essentially associated with the sites affected by the LF excitation, which leads mainly to the Tₙ → Cₙ conversion (Figure 6). The two observed doublets for Tₙ agree reasonably well with the theoretical spectrum where two relatively intense bands are predicted at 1376 (ωCH₂) and 1200 cm⁻¹ (νC−O) (Table 2). However, the Tₙ bands that are more affected by the HF excitation are observed at 1372 (ωCH₂) and 1207/1204 cm⁻¹ (νC−O) (Figure 7), which can be correlated with the bands of Tₙ₊₋₀ observed at 1332 cm⁻¹ (ωCH₂ mode, predicted at 1337 cm⁻¹), 1290 cm⁻¹ (twCH₂ mode, predicted at 1293 cm⁻¹), and 1242 cm⁻¹ (νC−O, predicted at 1226 cm⁻¹).

Another interesting observation is the large splitting (40–90 cm⁻¹) between the two νC−O features responding differently to the HF and LF excitations, which is observed for both Tₙ and Cₙ conformers. The bands associated with the LF excitation exhibit a larger blue shift (60–100 cm⁻¹) relative to the ab initio calculated values, whereas those associated with the HF excitation almost match the calculated values. This observation points to different mode coupling in the two sites, either between νC−O and other internal modes or between νC−O and matrix modes, which seem to be more important in the case of the LF-sensitive sites. Similar deviations (30–90 cm⁻¹) between the observed and calculated νC−O frequencies were previously observed for formic and acetic acids.⁶,⁸

The assignment of the C-S bands is based on the spectral changes induced by the excitation of Tₙ₊₋₀, although the relatively low concentration of the C-S conformer limits the number of observed bands of this form. As expected from the calculations, the observed bands of C-S appear to be slightly shifted but clearly distinguishable from those of the C₁ form (Table 2), and they fit the most intense bands predicted by the calculations (νOD at 2670, νC=O doublet at 1798/1784, and νC=O doublet at 1238/1216 cm⁻¹).

4. Can the C−O and Cₕ−C Rotations Occur in a Concerted Way? Excitation of the 2νOH(D) mode introduces an energy in the molecule higher than the computed barriers for isomerization along each of the two internal rotation coordinates (Cₕ−C and C−O). All stepwise processes are energetically allowed, and for the excitation of Tₙ and Tₙ₊₋₀, they were experimentally observed as described earlier. The energy barrier associated with the direct Tₙ → Cₛ₊₋₀ and Tₙ₊₋₀ → C₂ₕ processes, via concerted C−O/Cₕ−C internal rotation, is predicted to be 4300–4800 cm⁻¹ (Figure 1), which is also smaller than the excitation energy. However, no experimental evidence that excitation of the 2νOH(D) modes of either Tₙ or Tₙ₊₋₀ conformers induces the concerted process is obtained. Indeed, for both PA-OH and PA-OD, neither the C-S conformer upon excitation of the Tₙ form nor the C₂ₕ conformer upon excitation of the Tₙ₊₋₀ form is produced.

However, the C-S to Tₙ decay occurs when the NIR irradiation is interrupted (at 8 K, with incident globar radiation below 3300 cm⁻¹). It is then, in principle, possible that the direct Cₛ₊₋₀ → Tₙ conversion is taking place via the concerted mechanism. Because the calculated isomerization barrier for the Cₛ₊₋₀ → Tₙ process is 2542 cm⁻¹, the globar radiation below 3300 cm⁻¹ should not play a determinant role in the concerted process. In fact, the only modes with energy above the barrier that can be excited by the globar are the C−H stretching modes (predicted within the 3000–2900 cm⁻¹ spectral range), but their cross sections are so small that they are not experimentally observed. Hence, although the concerted mechanism cannot be excluded, a stepwise process for the conversion of Cₛ₊₋₀ to Tₙ seems to be more plausible. The Cₛ₊₋₀ form can first decay by tunneling to a vibrationally excited Tₙ₊₋₀ form, and then partial dissipation of the excess vibrational energy into the Cₕ−C rotational coordinate, above the Tₙ₊₋₀ → Tₙ isomerization barrier, can lead to the occurrence of this last process. The isotopic effect in the Cₛ₊₋₀ → Tₙ decay rate (slower rate by at least 2 orders of magnitude for PA-OD) supports a conversion mechanism involving tunneling. The proposed stepwise mechanism is consistent with the model for the dark C→T tunneling decay in formic and acetic acids from the torsional ground state of the cis conformer to a vibrationally excited state of the trans form.³,⁷

Concluding Remarks

The conformers of propionic acid were studied by means of ab initio calculations, IR absorption spectroscopy, and narrow-band NIR excitation in solid Ar. The calculations predicted the existence of four conformers, differing by internal rotation around the Cₕ−C and C−O bonds. For both PA-OH and PA-OD species, internal rotations were induced by excitation of the first overtone of the hydroxyl stretching modes of different conformers, and IR absorption spectroscopy was used to probe the photoinduced conformational changes. The main results of this study can be summarized as follows:

1. The IR absorption spectra of the four conformers of propionic acid isolated in solid argon are obtained and assigned on the basis of the ab initio calculated spectra (Tables 1 and 2).
2. Because of conformational cooling, the ground conformational state (Tₙ) is the only form initially present in the deposited argon matrix. The remaining three conformers (Tₙ₊₋₀, C₁ and Cₛ₊₋₀) can be photoproduced by irradiation in the NIR region. Excitation of the 2νOH(D) mode is efficient in promoting the Cₕ−C and C−O internal rotations leading to the Tₙ → Tₙ₊₋₀, Tₙ → C₁, Tₙ₊₋₀ → Tₙ, and Tₙ₊₋₀ → Cₛ₊₋₀ isomerization processes. Although energetically allowed, no direct Tₙ → Cₛ₊₋₀ or Tₙ₊₋₀ → C₁ photoisomerization by concerted rotation around the Cₕ−C and C−O bonds was observed. The comparison of photochemical processes on propionic, acetic, and formic acids can be found elsewhere.⁴¹
3. For PA-OH, the C₁ and Cₛ₊₋₀ conformers are short-lived species due to fast C₁ → Tₙ and Cₛ₊₋₀ → Tₙ conversions in the absence of irradiation. The much slower decay of these two forms in the case of PA-OD (by ca. 2–4 orders of magnitude) proves the crucial role of tunneling in these isomerization processes.
4. The 2νOH(D) absorption band is split by the matrix-site effect. For PA-OD, excitation of Tₙ was performed at two different frequencies within the 2νOD absorption band, leading to site-dependent isomerization efficiencies that aid the discrimination between the spectra of the photoproducts (Tₙ₊₋₀ and C₁).
5. The Tₙ₊₋₀ → Tₙ isomerization takes place in the dark with an appreciable rate (k ≈ 10⁻⁷ s⁻¹), even at the lowest working temperature (8 K). The very low isomerization barrier associated...
with the $T_g^{\pm} \rightarrow T_1$ process (computationally estimated as 60 cm$^{-1}$) suggests that this isomerization could occur over the barrier. The strong temperature dependence of this process experimentally supports a low barrier for the $T_g^{\pm} \rightarrow T_1$ process.

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**Supporting Information Available:** Definition of the internal symmetry coordinates used in the normal coordinate analysis of propionic acid. This material is available free of charge via the Internet at http://pubs.acs.org.

**References and Notes**