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Probing binding-site preferences in a propiolic acid complex with water at 0.4 K

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Carboxylic acid··water complexes serve as model systems for understanding molecular interactions that are fundamental to bio- and atmospheric chemistry. Utilizing the helium nanodroplet (HND) technique, which enables the kinetic trapping of otherwise inaccessible isomeric structures, we investigated the hydrogen bonding site preferences in a 1:1 complex of propiolic acid (HC≡C-COOH, PA) with D₂O. Mass-selective infrared (IR) spectra recorded in the C=O and C≡C stretching regions confirmed the exclusive isolation of the *cis*-PA conformer under single-molecule doping conditions of the helium droplets. Further complexation of PA with D₂O inside the droplets yielded three distinct isomers of *cis*-PA··D₂O dimer, with the dominant spectral features corresponding to two kinetically trapped structures stabilized by either a non-classical (≡C-H··OD₂) or a classical (C=O··DOD) hydrogen bond. A weak IR band corresponding to the global minimum isomer, characterized by a six-membered ring involving D₂O and the COOH moiety, is also observed. The structural assignments are aided by harmonic IR spectra of the lowest-energy isomers of PA··D₂O complexes computed at the MP2/aug-cc-pVDZ level. These structural findings demonstrate the balance of the dipole-dipole and higher-order interactions in steering aggregation dynamics in HNDs. Notably, the polar D₂O ($\mu = 1.85$ D) and PA ($\mu = 1.59$ D) promote directional association leading to the formation of local minimum structures, which are lying >20 kJ mol⁻¹ above that of the global minimum isomer. The calculated interconversion energy barriers for *cis*-PA··D₂O isomers are in accordance with kinetic trapping inside HNDs at 0.4 K.

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1. Introduction

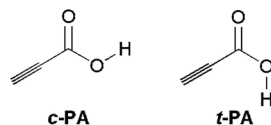
Hydrogen bonds (HBs) have been a topic of discussion since their discovery,¹ as they play pivotal roles in various chemical and physical processes across the liquid, solid and gas-phases.²⁻⁴ Beyond their well-established significance in general chemistry or biological systems,^{3,5,6} a molecular understanding of hydrogen bonding is pivotal for supramolecular chemistry,⁷ atmospheric chemistry,⁸ mineralogy,⁹ and materials science.¹⁰⁻¹² HBs were considered classically to be formed between two substantially electronegative atoms with a hydrogen atom in between (X-H··Y) where one atom acts as the HB donor (X) and the other as the acceptor (Y).¹³ In the last few decades, low-temperature experiments supported by theoretical studies have greatly reshaped the understanding of HB interaction.¹⁴⁻²⁰ For instance, C-H or π -electrons have been recognized as actively participating in HB formation.^{3,21,22} Although weaker in strength, these non-classical hydrogen bonds play a crucial role in stabilizing the three dimensional structures of biomolecules,⁵ crystal packing of chemical

compounds,²³ and the nucleation and growth of atmospheric aerosol particles.⁸

In this context, helium nanodroplets (HNDs) offer a unique and highly versatile medium for exploring unconventional intermolecular interactions, particularly through the stabilization of binary complexes. In contrast to gas-phase techniques (*e.g.* molecular beam experiments)²⁴⁻²⁶ and solid-phase methods (*e.g.* matrix isolation),^{27,28} which predominantly favor the stabilization of global minimum structures, HNDs often enable the stabilization of local minimum configurations that are otherwise challenging to access.²⁹ This distinctive capability arises primarily from the intrinsic properties of HNDs, the ultracold temperature (~ 0.4 K) and the superfluid nature.³⁰ The formation mechanism of molecular complexes within HNDs is governed by the sequential pickup of molecules with each undergoing rapid thermalization to its vibrational ground state prior to complexation. In the case of polar molecules, the long-range dipole-dipole force guides the orientation and approach of the molecules during aggregation.³⁰ It is consistently supported by experimental studies involving a broad range of systems—including homodimeric, trimeric and oligomeric clusters such as dimers of formic acid,³¹ acetic acid,³² pyruvic acid,³³ (OCS)₃,³⁴ cyclic water hexamers [cyclic-(H₂O)₆],³⁵

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Scheme 1 Schematic diagram of the *cis* and *trans* isomers of propiolic acid (PA). The acronyms **c-PA** and **t-PA** are elaborated in Section 2.3.

and hydrogen cyanide $[(\text{HCN})_n]$ chains.²⁹ Similar trends have been observed in heterodimeric systems, which can be exemplified by the propargyl alcohol $\cdot\cdot\text{D}_2\text{O}$ complex. Here, two local minimum structures located approximately 5 and 13 kJ mol^{-1} above the global minimum have been identified.³⁶

Carboxylic acids bearing additional functional groups present a rich landscape of hydrogen-bonding motifs due to the presence of multiple HB donors and acceptors. Propiolic acid ($\text{HC}\equiv\text{C}-\text{COOH}$, PA), which contains a carboxylic ($-\text{COOH}$) and an acetylenic ($-\text{C}\equiv\text{CH}$) moiety, exemplifies such a multifunctional system. Precisely, in PA the $\text{C}\equiv\text{C}$ π -electrons, the carbonyl oxygen ($\text{C}=\text{O}$), and the hydroxyl oxygen ($\text{O}-\text{H}$) serve as HB acceptors, while the acetylenic hydrogen ($\text{H}-\text{C}\equiv$) and hydroxyl hydrogen ($-\text{OH}$) act as HB donors. Scheme 1 presents two planar configurations of PA: *cis* (**c-PA**) and *trans* (**t-PA**). The global minimum species **c-PA** characterized by a 0° dihedral angle ($\text{O}=\text{C}-\text{O}-\text{H}$, along $\text{C}-\text{O}$ bond) is energetically favored by $\sim 12 \text{ kJ mol}^{-1}$ over **t-PA** (180° dihedral angle).^{37,38} The *cis*-to-*trans* interconversion barrier is calculated to be $\sim 40 \text{ kJ mol}^{-1}$,^{37,39,40} suggesting a high degree of conformational rigidity, particularly under cryogenic conditions. **c-PA** has been detected across gas, liquid and solid matrices.^{29,38,40–42} Notably, the *cis*-to-*trans* isomerization has been reported exclusively through photoinduced processes in confined environments (N_2 matrix).³⁸ The dipole moments (μ) of **c-PA** and **t-PA** are 1.59 D and 4.8 D, respectively.⁴³

Since PA has pronounced dipole moment and multifunctional hydrogen bonding capabilities, it will be insightful, particularly, for secondary organic aerosol formation and prebiotic chemistry in extraterrestrial environments to explore the binding motifs of PA with molecules like H_2O , H_2S , CO_2 , and NH_3 , which differ in polarity and bonding behaviour.⁴⁴ Here, formation of unconventional HBs involving $\equiv\text{C}-\text{H}$ and $\text{C}\equiv\text{C}$ functional groups can be expected in line with previous low temperature studies on monohydrated propyne and acetylene.¹⁶

In the current work, we present the isolation of the PA monomer and its 1:1 binary complex with D_2O in HNDs. The substantial dipole moment of D_2O ($\mu = 1.85 \text{ D}$) together with PA is expected to influence the geometry and binding orientation of the resulting binary complexes in HNDs. Mass-selective infrared spectra were recorded in the $\text{C}=\text{O}$ and $\text{C}\equiv\text{C}$ stretching regions for the PA monomer and the $\text{PA}\cdot\cdot\text{D}_2\text{O}$ dimer. A controlled formation of 1:1 complexes was achieved through sequential doping of the helium droplets using two spatially separated pickup chambers. Structural assignments were based on the comparison of the experimentally observed IR band positions and intensities with harmonic spectra calculated for the most stable isomers at the MP2 level. Furthermore, the relative stabilities of these identified structures were assessed

using computed interaction energies, offering insights into the preferred binding motifs and the influence of dipole-dipole interaction in molecular aggregation under such confined cryogenic conditions.

2. Experiment

2.1. Experimental set-up

PA and D_2O were purchased from Sigma-Aldrich and employed without further purification except degassing. The experiments were performed using the infrared helium nanodroplet spectrometer at Ruhr University Bochum.^{33,45} This instrument consists of four differentially pumped vacuum chambers – namely, expansion, pick-up, spectroscopy and quadrupole mass spectrometry (QMS) chamber – corresponding to vacuum levels ranging from moderate (10^{-5} mbar) to ultrahigh vacuum (10^{-9} mbar), respectively. These given pressure ranges (in mbar) are in the presence of a HND beam. The HNDs are generated *via* a continuous supersonic expansion of highly pressurized (40–50 bar) ultrapure (99.9999%) helium gas through a pre-cooled nozzle of 5 μm diameter into the expansion chamber. A closed-cycle helium cryostat combined with temperature controller maintains the temperature of the nozzle within 13–22 K for droplet production. The droplet size is log-normally distributed and is controlled by varying the backing He pressure and temperature of the nozzle. After expansion, droplets are collimated using a skimmer of 0.5 mm in diameter and introduced into the following pick-up chamber, which consists of two individually pumped and spatially well separated (distance $\sim 20 \text{ cm}$) pick-up cells. They are traversed by the droplet beam *via* 5-mm-diameter openings in the cell walls. Here, dopant molecules are injected into the droplet beam using these pick-up lines, namely pick-up-lines 1 and 2. After passing through the pick-up and the spectroscopy chamber, the droplets are detected using an Extrel quadrupole mass spectrometer (model 5221) in the QMS chamber.

In the present study, expansion conditions were set to ~ 45 bars of He and 19.9 K of nozzle temperature, which led to the formation of droplets containing roughly 10^4 He atoms. The droplets were doped by PA using pick-up-line 1 with the pick-up pressure set to 3.0×10^{-5} mbar. D_2O for the generation of the $\text{PA}\cdot\cdot\text{D}_2\text{O}$ complex is subsequently introduced *via* pick-up-line 2 with the pick-up pressures maintained at $\sim 3 \times 10^{-5}$ mbar.

To record the vibrational spectra of PA monomer and the $\text{PA}\cdot\cdot\text{D}_2\text{O}$ dimer, infrared light of a quantum cascade laser from DRS Daylight Solution (model number: MIRCAt-QT-Z-2400) has been employed, where the beam path was continuously purged by N_2 gas. The laser comprised of four lasing heads encompasses these ranges: 1460–1635, 1620–1860, 1960–2220 and 2290–2520 cm^{-1} . The laser light overlaps with the droplet beam in an antiparallel configuration. Pure helium droplets are transparent to the IR radiation. However, absorption of the IR photons by the embedded molecules/clusters followed by vibrational relaxation causes evaporation of several hundred helium atoms from the droplet surface (He–He binding energy $\sim 5 \text{ cm}^{-1}$). It leads to



a shrinkage of the droplet ionization cross-section and to a concomitant decrease in the ion current measured by the QMS. Now, recording the dip in the ion current for a specific mass channel (m/z) as a function of IR radiation frequency yields the mass-selective vibrational spectrum. The signal is collected using phase-sensitive-detection with a lock-in amplifier.

Pick-up curve. For the determination of the cluster size of PA, we have recorded the amplitude of the depletion signal as a function of partial pressures, the so-called pick-up curve, as shown in Fig. S1 in the SI. These pick-up curves follow Poisson statistics⁴⁶ and were used to deduce the molecular cluster size corresponding to the observed bands. According to Poisson statistics, the probability of picking up k dopant molecule(s) is given by the following equation:

$$P_k = [(\sigma\rho L)^k/k!]\exp(-\sigma\rho L)$$

where σ represents the pick-up cross-section of the droplets, ρ is the number density of the molecules in the pick-up chamber and L is the length of the pick-up region. Here, L is a constant for a given experimental setup, and σ can also be considered as a constant for a well-defined droplet size distribution. ρ is directly proportional to the partial pressure of the dopant in the pick-up chamber.

The IR features originating from the PA $\cdot\cdot$ D₂O complex are relatively weak. In addition to pick-up curve measurements (see Fig. S2 in the SI), we plotted pressure-dependent IR spectra for the PA $\cdot\cdot$ D₂O complex. In the latter case, the PA pick-up pressure (p_{PA}) was fixed at the monomer value, as determined from Fig. S1, while the D₂O pick-up pressure ($p_{\text{D}_2\text{O}}$) was systematically varied (see Fig. S3 in the SI). These measurements together with mass spectral analysis (Section 2.3) confirm the cluster size of the PA $\cdot\cdot$ D₂O complex.

2.2. Computation

Geometry optimization and harmonic vibrational frequency calculations were performed for the PA monomer as well as its 1:1 complexes with D₂O using MP2/aug-cc-pVDZ and B3LYP-D3/cc-pVTZ methods.^{47,48} From our earlier observations,³³ we found that the helium droplet experimental spectra of organic acids and their complexes are well reproduced by harmonic IR spectra calculated at the MP2/aug-cc-pVDZ level. Therefore, these MP2-computed frequencies are employed to assist in the structural assignment of the experimental spectra and to derive zero-point-energy (ZPE) corrected relative energies (ΔE_0). A scaling factor of 1.00 (no scaling) determined by comparing the computed C=O and C≡C stretching frequencies of the PA monomer with their corresponding observed values inside HNDs is used. The BSSE-corrected [BSSE: basis set superposition error] interaction energies (D_0) for 1:1 PA $\cdot\cdot$ D₂O complexes are also calculated using the counterpoise method, where the energies of the complexes are calculated alongside the energies of the individual monomers, both with the same basis set.^{49,50} Note that relative stabilities of the isomers are discussed based on MP2-obtained ΔE_0 and D_0 values. The geometry optimization at the

DFT level were performed additionally to validate and verify real minima obtained at the MP2 level. Note that an empirical dispersion correction (D3) was included in the DFT geometry optimizations to account for dispersion interactions.⁵¹ All computations are performed using Gaussian 16 software.⁵² Optimized geometry and cartesian coordinates of all relevant structures are provided in Table S1a and b in the SI.

2.3. Structure nomenclature scheme

The labelling of the two monomer structures of PA as *cis*-PA (*c*-PA) and *trans*-PA (*t*-PA) is based on *cis* and *trans* orientations of C=O and O-H groups along the C-O single bond axis.³⁸ Structures corresponding to the 1:1 complex of PA with D₂O are systematically labelled following the scheme *c/t*-PAm-*n*, where *m* denotes the interacting molecule with PA (*m* = *w* for D₂O). The integer *n* is assigned in the ascending order of computed relative energies (ΔE_0) of the isomers where *n* = 1 is for the global minimum. Water molecules are labelled according to their role as HB donors (D) or acceptors (A). For example, a D₂O molecule simultaneously acting as a donor and an acceptor is designated as AD-D₂O.

3. Results and discussion

3.1. Mass spectra

Fig. 1 shows a comparison of the mass spectra of the pure helium droplet beam (trace a, black), doped with PA at $p_{\text{PA}} = 3.0 \times 10^{-5}$ mbar (trace b, red), droplets doped with D₂O at $p_{\text{D}_2\text{O}} = 3.0 \times 10^{-5}$ mbar (trace c, green), and droplets doped subsequently with PA and D₂O from pick-up lines 1 and 2 both set at p_{PA} and $p_{\text{D}_2\text{O}} = 3.0 \times 10^{-5}$ mbar (trace d, blue). The peaks at a multiple of $m/z = 4$ amu correspond to (⁴He)_{*n*}⁺ fragments

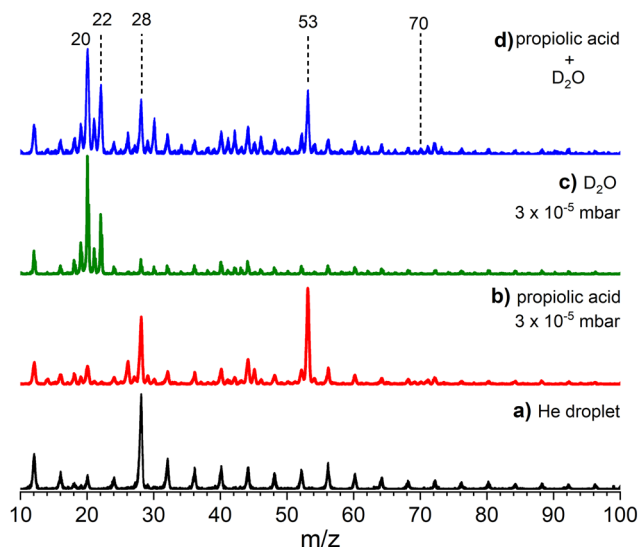


Fig. 1 Mass spectra of droplet beam of (a) pure helium and doped droplets with (b) propionic acid (PA) at a pick-up pressure of 3.0×10^{-5} mbar, (c) D₂O at a pick-up pressure of 3.0×10^{-5} mbar, and (d) PA and D₂O both set to the pick-up pressure of 3.0×10^{-5} mbar from two different pick-up lines.



(see trace 1a); the very small peak at $m/z = 18$, is due to residual water vapor in the instrument (see trace 1a). As the intensity of the $m/z = 18$ peak is five times smaller than that of $m/z = 16$ corresponding to the $(^4\text{He})_4^+$, it has no impact on the spectral measurements. The peak at $m/z = 28$, attributed to the $(^4\text{He})_7^+$ and trapped N_2 , is relatively strong. Note, the absence of a 3 : 1 intensity ratio between the $m/z = 28$ and $m/z = 32$ peaks, corresponding to the natural abundances of N_2 and O_2 , respectively, rules out any contribution from atmospheric sources. The mass spectrum of the droplets doped at $p_{\text{PA}} = 3.0 \times 10^{-5}$ mbar yields additional peaks at $m/z = 25, 28, 45, 53$ and 70 with $m/z = 53$ being the dominant one (see trace 1b). Here, $m/z = 70$ corresponds to the PA molecular ion $(\text{HC}\equiv\text{C}-\text{COOH})^+$, while the mass fragments at $m/z = 25, 28, 45, 53$ represent the species $(\text{C}_2\text{H})^+, (\text{CO})^+, (\text{COOH})^+, (\text{HC}\equiv\text{CCO})^+$, respectively. D_2O doping at $p_{\text{D}_2\text{O}} = 3.0 \times 10^{-5}$ mbar yielded the most prominent molecular mass peak $(\text{D}_2\text{O})^+$ at $m/z = 20$ as well as a smaller peak at $m/z = 22$ $(\text{D}_3\text{O})^+$ (trace 1c). This mass spectral pattern indicates the condition which corresponds to the single D_2O molecule doping. Trace 1d shows the mass spectrum obtained after sequential doping of helium droplets with PA and D_2O , both at the pick-up pressure of 3×10^{-5} mbar.

3.2. Experimental infrared spectra

PA and $\text{PA}\cdot\cdot\text{D}_2\text{O}$. In Fig. 2 and Fig. 3, we display mass-selective [$m/z = 53, (\text{HC}\equiv\text{C}-\text{C}=\text{O})^+$] IR spectra of PA and the $\text{PA}\cdot\cdot\text{D}_2\text{O}$ complex, trapped inside HNDs, in the C=O and the C \equiv C stretching frequency range, respectively. The IR traces of PA recorded at the $p_{\text{PA}} = 3 \times 10^{-5}$ mbar are shown in black (Fig. 2a and 3a). The red traces (Fig. 2b and 3b) show the IR spectra of the $\text{PA}\cdot\cdot\text{D}_2\text{O}$ complex formed *via* sequential pickup of PA and D_2O in HNDs, with both pick-up pressures maintained at 3×10^{-5} mbar.

The IR spectrum of PA in the C=O stretching range is dominated by the strong feature located at 1762.5 cm^{-1} , designated as A_1 (see trace 2a, black). In addition, two weak bands (~ 15 times less intense than A_1) are observed within 3 cm^{-1} . These absorptions are marked with asterisks in trace 2a.



Fig. 2 Mass-selected ($m/z = 53$) IR spectra in the C=O stretching region of (a) the PA monomer ($p_{\text{PA}} = 3 \times 10^{-5}$ mbar) and (b) the $\text{PA}\cdot\cdot\text{D}_2\text{O}$ complex *via* a sequential pickup of PA ($p_{\text{PA}} = 3 \times 10^{-5}$ mbar) and D_2O ($p_{\text{D}_2\text{O}} = 3 \times 10^{-5}$ mbar) in helium nanodroplets. IR absorption bands of PA and the $\text{PA}\cdot\cdot\text{D}_2\text{O}$ complex are denoted by A_1 and a_1 – a_3 , respectively. In the inset, we plot the range 1725 – 1750 cm^{-1} displaying the bands a_2 and a_3 . The substantially weak features in trace 2a are marked with asterisks.



Fig. 3 Mass-selected ($m/z = 53$) IR spectra in the C \equiv C stretching region of (a) PA monomer ($p_{\text{PA}} = 3 \times 10^{-5}$ mbar) and (b) $\text{PA}\cdot\cdot\text{D}_2\text{O}$ complex *via* a sequential pickup of PA ($p_{\text{PA}} = 3 \times 10^{-5}$ mbar) and D_2O ($p_{\text{D}_2\text{O}} = 3 \times 10^{-5}$ mbar) in helium nanodroplets. Traces a and b are normalized to the intensity of the A_3 band. The strongest IR absorption of the monomer is denoted as A_3 and the shoulder to A_2 . The bands marked with a_4 and a_5 correspond to the $\text{PA}\cdot\cdot\text{D}_2\text{O}$ complex.

In the C \equiv C stretching region (Fig. 3a), a moderately intense feature at 2142.6 cm^{-1} (A_3) is detected with a shoulder at 2141.6 cm^{-1} (A_2). Pick-up curve analysis indicates that the dominant features at 1762.5 and 2142.6 cm^{-1} are due to the PA monomer (see Fig. S1).

Previous studies in solid matrices (N_2 , Ar and Ne) identified the C=O and C \equiv C stretching vibrations of the *cis*-PA (**c-PA**) conformer at around 1755 and 2140 cm^{-1} , respectively.^{38,43} In gas-phase FTIR measurements, two broad features centered around 1745 and 2137 cm^{-1} are attributed to **c-PA**.⁴² Therefore, the observed bands A_1 and A_3 in HNDs can be assigned to the C=O and C \equiv C stretching modes of the **c-PA** conformer, respectively (see Table 1). In line with the present helium droplet study, matrix isolation measurements also reported weak features adjacent to the dominant IR bands (A_1 and A_3). These additional bands marked by asterisks in trace 2a and the A_2 peak in trace 3a could be due to anharmonic effects—such as combination bands, overtones, or Fermi resonances (*vide infra*).

Now, formation of the $\text{PA}\cdot\cdot\text{D}_2\text{O}$ dimers inside the HNDs results in the emergence of three new IR absorption bands, located at 1758.5 cm^{-1} (a_1), 1743.3 cm^{-1} (a_2), and 1729.5 cm^{-1} (a_3), in the C=O stretching region (summarized in Table 1). The a_1 feature overlaps slightly with a tiny band from the PA monomer. Hence, this peak is delineated by a dashed line to aid visual identification (see trace a *vs.* trace b, Fig. 2). A magnified view of the 1725 – 1750 cm^{-1} range (inset) provides a clearer view of a_3 along with its relative intensity compared to a_2 (Fig. 2). It is important to note that the narrowest band, a_1 (FWHM = 0.5 cm^{-1}), exhibits only a redshift of 4 cm^{-1} relative to the C=O stretch of the PA monomer (A_1 , 1762.5 cm^{-1}). However, a substantial bathochromic shift of $\sim 34 \text{ cm}^{-1}$ is observed for the broadest band, a_3 (FWHM = 2.0 cm^{-1}). The feature a_2 displays a redshift of approximately 19 cm^{-1} with FWHM $\sim 1.0 \text{ cm}^{-1}$. This trend reveals an inverse correlation between redshift and spectral linewidth (FWHM) across the three bands suggesting that they are likely originating from structural motifs of varying hydrogen bonding strengths.

The two new absorptions at 2134.6 cm^{-1} (a_4) and 2141.7 cm^{-1} (a_5) in the C \equiv C stretching region are assigned to the



Table 1 Comparison of observed IR band positions of the propiolic acid (PA) monomer and 1 : 1 PA \cdots D₂O complexes isolated in helium nanodroplets with MP2/aug-cc-pVDZ computed harmonic frequencies. Very weak absorption features from PA monomer are denoted by asterisks, see trace 2a. Previously reported values of the PA monomer trapped in N₂, neon (Ne), argon (Ar) matrices and in the gas-phase are included for reference. The MP2-calculated transitions from Section 3.3 were employed to assist in the vibrational assignments

Species	Observed transition (cm ⁻¹)				Gas-phase ^d	Calculated (cm ⁻¹) ^e	Vibrational assignments
	He droplet	N ₂ ^a	Ne ^b	Ar ^c			
PA	1759.1(*)		1756				
	1761.1(*)						
	1762.5 (A ₁)	1754	1759	1754	1745	1762	C=O str. (ν_3)
	2141.6 (A ₂)		2136				
PA \cdots D ₂ O	2142.6 (A ₃)	2140	2140	2137	2137	2140	C \equiv C str. (ν_4)
	1758.5 (a ₁)					1759	C=O str. (c-PAw-3)
	1743.3 (a ₂)					1745	C=O str. (c-PAw-2)
	1729.5 (a ₃)					1731	C=O str. (c-PAw-1)
	2134.6 (a ₄)					2132	C \equiv C str. (c-PAw-3)
	2141.7 (a ₅)				2143	C \equiv C str. (c-PAw-2)	

^a Observed in the N₂ matrix at 12 K.³⁸ ^b Observed in the Ne matrix at \sim 10 K.⁴³ ^c Observed in the Ar matrix at 9 K.⁴⁰ ^d FTIR measurements in the gas-phase at room temperature.⁴² ^e MP2/aug-cc-pVDZ computed IR band frequencies in the current study.

PA \cdots D₂O complex (see trace 3b vs. trace 3a). Note that peak a₅ overlaps significantly with the features from the PA monomer. However, a normalization of traces a and b of Fig. 3 to the intensity of peak A₃ supports the existence of the band a₅. The a₄ feature exhibits a red shift of approximately 8 cm⁻¹ relative to the corresponding A₃ band of the PA monomer, whereas a₅ shows a shift of less than 1 cm⁻¹ (Table 1). This contrasting behavior suggests distinct binding motifs: for a₄, the C \equiv C unit is likely directly involved in the interaction with D₂O, while in the case of a₅, the C \equiv C moiety remains largely unaffected, indicating an indirect or more distant binding configuration.

We have measured pick up curves (see Figure S2 in the SI) and plotted pressure-dependent IR spectra (Figure S3) to confirm the assignment of a₁, a₂, a₃ and a₄ to the 1 : 1 PA \cdots D₂O dimer. A similar analysis for band a₅ is not possible due to the significant overlap with monomer features.

PA \cdots H₂O. To account for isotopic effects, IR spectra of the PA \cdots H₂O complex were recorded in both the C=O and C \equiv C stretching regions (Fig. S4 and S5, SI). In monohydrated systems, the C=O stretching frequency is generally weakly affected by isotopic (H₂O/D₂O) substitution. Consistent with this, bands a₂ and a₅ exhibit only minor shifts (within \pm 0.5 cm⁻¹), while bands a₃ and a₄ show no detectable change compared to the PA \cdots D₂O complex. Based on the observation, we propose that group (a₁, a₄) and (a₂, a₅) are associated with two different isomers. This spectral grouping is verified later in Section 3.3.

3.3 Structural assignments

Geometry optimization and harmonic frequency calculation for the plausible isomers of the PA monomer and the 1 : 1 complex of PA \cdots D₂O were carried out to aid structural assignment of the experimental spectra. The lowest-energy structures of the PA monomer and PA \cdots D₂O dimer according to their relative energies (ΔE_0) obtained at MP2/aug-cc-pVDZ level are plotted in the left and right panels of Fig. 4, respectively. Additionally, Table 2 and Table S3 in the SI provide ΔE_0 and structural parameters obtained both at MP2 and DFT levels. BSSE-

corrected interaction energies (D_0) in Table 2 are obtained at the MP2 level.

3.3.1. PA monomer. As mentioned earlier, **c-PA** is reported to be the global minimum whereas **t-PA** is 12–16 kJ mol⁻¹ higher in energy.³¹ In accordance with the previous findings, our current MP2 and DFT calculations confirm **c-PA** as the global minimum, with **t-PA** residing approximately 12 kJ mol⁻¹ above (Table 2). The optimized structures and corresponding dipole moment vectors (arrows in cyan) of these conformers are presented in Fig. 4 (left panel).



Fig. 4 ZPE-corrected relative energies ΔE_0 obtained at the MP2/aug-cc-pVDZ level for the lowest energy structures of the PA monomer and the **c-PA** \cdots D₂O dimer are presented in the left and right panels, respectively, with color coded bars. The nomenclature scheme is elaborated in Section 3.2. The name of the species identified in helium droplets are in bold. Bond distances and angle are in picometers (pm) and degrees ($^\circ$), respectively. The orientation of dipole moments for PA conformers are shown by arrows. Additional lowest energy structures for 1:1 complexes can be found in Table S1 in the SI.



Table 2 ZPE-corrected relative energies (ΔE_0) for the lowest energy isomers of the propiolic acid (PA) monomer and the PA \cdots D₂O dimer calculated at MP2 and DFT levels. The hydrogen bond distances (r_{HB}) and BSSE-corrected interaction energies (D_0) at the MP2 level are in picometers and kJ mol⁻¹, respectively

Species	Isomers	MP2/aug-cc-pVDZ				B3LYP-D3/cc-pVTZ	
		Symm.	ΔE_0 (kJ mol ⁻¹)	r_{HB} (pm)	D_0 (kJ mol ⁻¹)	ΔE_0 (kJ mol ⁻¹)	r_{HB} (pm)
PA	c-PA	C_s	0.0			0.0	
	t-PA	C_s	11.7			11.5	
PA \cdots D ₂ O	c-PAw-1	C_1	0.0	176 & 204	32.6	0.0	173 & 199
	c-PAw-2	C_s	21.0	201	12.4	29.1	202
	c-PAw-3	C_s	22.9	211	11.3	29.0	209
	c-PAw-4	C_1	26.5	207	6.0	34.5	214

The vibrational spectra of PA, recorded in both the solid state (Ne matrix) and gas phase (vapor at room temperature), display distinct bands corresponding to the C=O and C \equiv C stretching modes. Precisely, the C=O stretching vibration appears at 1759 cm⁻¹ in the Ne matrix and shifts to 1745 cm⁻¹ in the vapor phase. The C \equiv C stretching band is observed at 2140 cm⁻¹ in the solid neon and at 2137 cm⁻¹ in the vapor phase (see Table 1). Ar matrix data are also in accordance, reporting C=O and C \equiv C stretching vibrations at 1754 and 2137 cm⁻¹, respectively.⁴⁰ In addition, a recent matrix-isolation study in solid N₂ reported the C=O stretch of **c-PA** at \sim 1754 cm⁻¹ and of **t-PA** at \sim 1780 cm⁻¹. In the N₂-matrix, the higher energy **t-PA** conformer was selectively populated *via* UV irradiation.³⁸

We compare the IR spectra of the PA monomer in helium nanodroplets recorded at the C=O and C \equiv C stretching regions with the MP2-calculated spectra of its possible conformers (Fig. 5). The observed bands at 1762.5 cm⁻¹ (A₁) and 2142.6 cm⁻¹ (A₃) are in excellent agreement with the computed transitions of **c-PA** (trace 5a vs. 5b). So, the *cis* conformer of the PA monomer has been exclusively isolated inside the He droplet which is consistent with previous gas- and solid-phase findings.

Now, to address the assignment of weakly intense features (marked with asterisk and A₂, Fig. 5), we performed

anharmonic calculations. Because the positions and intensities of the IR bands originated from anharmonicity are highly sensitive to the ground-state structure, we computed the anharmonic IR spectra of **c-PA** at multiple levels of theory, each producing noticeable variations (see the bottom trace of Fig. S6 in the SI). The anharmonic spectrum computed at the VPT2/MP2/aug-cc-pVDZ level (see Fig. S6 in the SI) does not accurately reproduce the weak experimental features, preventing an unambiguous mode assignment. However, tentatively the tiny feature A₂ can be attributed to a combination band involving the C–C–OH stretching and O–H in-plane bending vibrations. Since these bands are extremely weak in intensity compared to the dominant feature A₁ and A₃, we refrain from making further vibrational assignments for them.

3.3.2. PA \cdots D₂O dimer. PA comprised of carboxylic (–COOH) and acetylic (HC \equiv C–) functional groups provides multiple potential hydrogen bonding sites for D₂O. As **c-PA** has been exclusively identified as the isolated species in helium droplet (see Section 3.3.1), we focused only on complexes of **c-PA** with D₂O. A comprehensive potential energy surface scan was performed to identify the lowest-energy geometries of the **c-PA** \cdots D₂O complex. Four distinct HB sites were found: (i) AD-D₂O doubly hydrogen bonded to COOH group (**c-PAw-1**), (ii) D-D₂O singly hydrogen bonded to carbonyl oxygen atom (**c-PAw-2**), (iii) A-D₂O singly hydrogen bonded to H–C \equiv C moiety (**c-PAw-3**), and (iv) D-D₂O singly hydrogen bonded to hydroxyl oxygen atom (**c-PAw-4**). These structures are presented in Fig. 4 (left panel).

Briefly, the global minimum **c-PAw-1** is the lowest symmetric (C_1) structure and characterized by a six-membered ring formation where D₂O and PA moieties simultaneously serve as HB donors and acceptors. The HB length in which the water molecule functions as the acceptor, *i.e.*, –O–H \cdots OD₂, is predicted to be approximately 15% shorter than the one where D₂O serves as the donor, *i.e.*, –C=O \cdots DOD (see Table 2 and Fig. 4). The second-lowest energy structure, **c-PAw-2**, lying 21 kJ mol⁻¹ above **c-PAw-1**, adopts C_s symmetry. The corresponding HB length is calculated to be 201 pm, closely matching the weaker HB length in **c-PAw-1** (204 pm). Isomer **c-PAw-3** (C_s symm.) found 22.9 kJ mol⁻¹ above the global minimum exhibits a so-called non-classical HB. In this case, the D₂O molecule acts as a HB acceptor, while the acetylenic hydrogen (\equiv C–H) is the HB donor. The resulting HB distance is comparatively longer, 211 pm. The C₂ principal axis of D₂O (aligned with its dipole



Fig. 5 Comparison of (a) the experimental infrared spectrum of the propiolic acid (PA) with the MP2/aug-cc-pVDZ calculated harmonic IR spectra for the two lowest energy structures (b) **c-PA** and (c) **t-PA**. ZPE-corrected relative energies (ΔE_0) in parenthesis are in kJ mol⁻¹.



moment vector) is nearly colinear with the C≡C–H axis of PA, forming an angle of $\sim 179^\circ$ (Fig. 4). The 4th lowest energy isomer **c-PAw-4** ($\Delta E_0 = +26.5$ kJ mol $^{-1}$, C_1 symm.) exhibits a HB of 207 pm distance. Note, a similar trend in ΔE_0 value is also found at DFT level as listed in Table 2.

Nevertheless, stability of these binding motifs is further substantiated by calculating interaction energies (D_0) for each isomer at the MP2 level. The D_0 value for **c-PAw-1** is 33 kJ mol $^{-1}$, which is more than twice as stabilizing as that of the **c-PAw-2** (12 kJ mol $^{-1}$) as listed in Table 2. This substantial D_0 value for **c-PAw-1** directly reflects the cooperative effect of its dual hydrogen-bonded motif. Notably, D_0 values for **c-PAw-2** and **c-PAw-3** are nearly similar, whereas **c-PAw-4** exhibits a marked difference (see Table 2).

For completeness, geometry optimization seeking for lowest energy isomers of **t-PA** ··· **D₂O** complex is also performed. The structures and their corresponding ΔE_0 values relative to the global minimum **c-PAw-1** are provided in Table S1 in the SI.

The IR spectrum of the **c-PA** ··· **D₂O** 1:1 complex (trace a, Fig. 6) displays five prominent vibrational bands: a_1 (1758.5 cm $^{-1}$), a_2 (1743.3 cm $^{-1}$), a_3 (1729.5 cm $^{-1}$), a_4 (2134.6 cm $^{-1}$) and a_5 (2141.7 cm $^{-1}$). According to the analysis at Section 3.2 above, (a_1 , a_4) and (a_2 , a_5) are attributed to two different isomers. Anyways, MP2-computed harmonic vibrational frequencies in

the C=O and C≡C stretching regions for all four isomers, **c-PAw-1** to **c-PAw-4**, are compared with the experimental IR spectrum recorded in helium droplet to carry out structural assignment (see Fig. 6). As the water binding motifs in each of these four lowest energy isomers are distinctive, characteristic differences in their predicted IR spectra are evident. Calculated spectra for **c-PAw-3**, **c-PAw-2**, **c-PAw-1**, and **c-PAw-4**, are plotted in trace b (red), c (blue), d (green), and e (violet) of Fig. 6, respectively. The experimental bands a_1 (1758.5 cm $^{-1}$) and a_4 (2134.6 cm $^{-1}$), highlighted in red, are closely reproduced by the spectrum computed for the isomer **c-PAw-3**. The predicted frequencies for the C=O and C≡C stretching modes in **c-PAw-3** are 1759 cm $^{-1}$ and 2132 cm $^{-1}$, respectively—both within $\Delta\nu = \pm 2$ cm $^{-1}$ of the experimental values, indicating good agreement between theory and experiment (traces a vs. b). Furthermore, bands a_2 (1743.3 cm $^{-1}$) and a_5 (2141.7 cm $^{-1}$), displayed in blue, are reproduced by the IR transitions computed for the species **c-PAw-2**. Precisely, the predicted frequencies corresponding to the C=O and C≡C stretching modes for **c-PAw-2** are 1745 cm $^{-1}$ and 2153 cm $^{-1}$, respectively—both within $\Delta\nu = \pm 2$ cm $^{-1}$ of the experimental values, establishing also satisfactory agreement between theory and experiment (traces a vs. c).

The broadest peak a_3 (1729.5 cm $^{-1}$, green) lies near to the computed C=O stretching mode of the global minimum isomer **c-PAw-1** at 1732 cm $^{-1}$ (trace a vs. trace d). The frequency deviation is within $\Delta\nu = \pm 3$ cm $^{-1}$, further supporting this structural assignments. However, the associated C≡C stretching bands predicted for the **c-PAw-1** was not observed experimentally. This is consistent with computational predictions, which indicate that the C≡C stretching modes exhibit infrared intensities approximately one-fifth of those associated with the C=O stretching vibrations. Besides, the feature is predicted to be in overlap with stronger absorptions of the PA monomer. In summary, we have identified the following structures: **c-PAw-3**, **c-PAw-2** and **c-PAw-1** inside HNDs based on the comparison between experiment and theory. The vibrational assignments are listed in Table 1.

3.4. Discussion

The three isomers, **c-PAw-1**, **c-PAw-2**, and **c-PAw-3**, observed upon **PA** ··· **D₂O** dimerization within HNDs, exhibit a broad range of relative stabilities and distinct hydrogen-bonding motifs (classical and non-classical) involving both C=O and C≡C moieties. The stabilization of local minimum structures (**c-PAw-2** and **c-PAw-3**) is likely governed by the kinetic trapping at ultra-low temperature (0.4 K) inside HNDs.

Low-temperature spectroscopic measurements combined with computational study on the complexes of acetylene (HC≡CH) and propyne (CH₃–C≡CH) with a single water molecule has previously revealed distinct binding motifs where non-classical HB generation is dominating.¹⁶ Particularly, the presence of α -hydrogens on the methyl group adjacent to the C≡C bond in CH₃–C≡CH enables the formation of a five-membered ring involving the C≡C bond, methyl C–H and the AD–H₂O in ≡C···H(H)O···HC– fashion. Contrarily, the

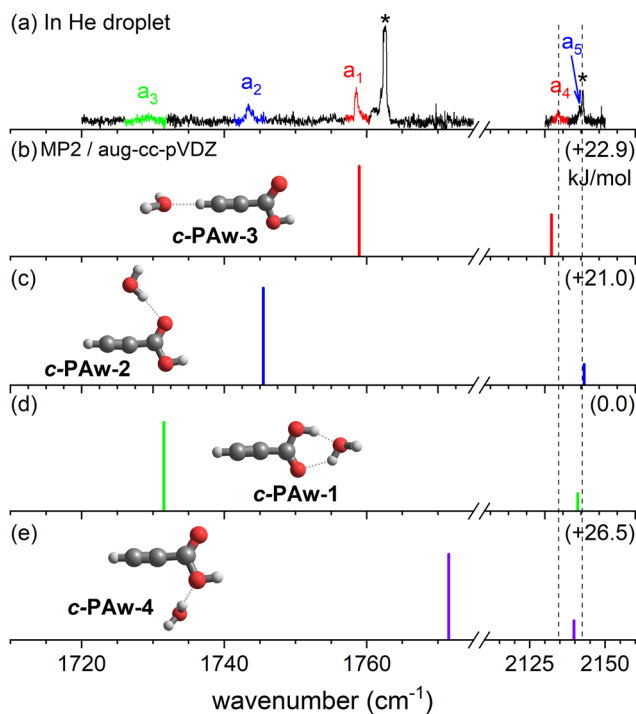


Fig. 6 Comparison of (a) the experimental IR spectrum of the **c-PA** ··· **D₂O** 1:1 complex (peaks a_1 , a_2 , a_3 , a_4 and a_5) with the MP2/aug-cc-pVDZ calculated harmonic IR spectra for the lowest energy structures (b) **c-PAw-3**, (c) **c-PAw-2**, (d) **c-PAw-1** and (e) **c-PAw-4**. Structural assignments are denoted by color coding. ΔE_0 (kJ mol $^{-1}$) are in parenthesis. Peaks belonging to the **c-PA** monomer in trace a are marked with asterisks. The peak position of the overlapped feature a_5 is shown by an arrow in blue. Dashed lines are used as a guide to the eye for the >2000 cm $^{-1}$ region.



Table 3 HB motifs for the 1 : 1 complex of **c-PA** ($\text{HC} \equiv \text{CCOOH}$) with D_2O stabilized inside helium nanodroplets as marked with \checkmark . Calculated interaction energies (D_0) are in kJ mol^{-1} . The dipole moment (μ) vector of each individual molecular units (*i.e.* PA and D_2O) are shown by arrows. The μ values for **c-PA** and D_2O are 1.59 and 1.85 D, respectively

Hydrogen Bond Motifs	$\text{HC} \equiv \text{CCOOH} + \text{D}_2\text{O}$	
	He droplet	D_0 (kJ mol^{-1})
c-PAw-3 	\checkmark	11
c-PAw-2 	\checkmark	12
c-PAw-1 	\checkmark	32

$\text{HC} \equiv \text{CH}$ molecule forms a linear complex ($\equiv \text{C}-\text{H} \cdots \text{OH}_2$) where the water molecule acts as $\text{A}-\text{H}_2\text{O}$. The present study disclosed a similar binding pattern in the isomer **c-PAw-3**. So, the absence of α -hydrogens adjacent to the $\text{C} \equiv \text{C}$ group in PA facilitates the $\equiv \text{C}-\text{H} \cdots \text{OD}_2$ binding. However, D_2O in **c-PAw-2** and **c-PAw-1** forms HB involving the carboxylic acid group only. These findings highlight the diverse hydrogen bonding topologies accessible to PA under superfluid conditions, which is in line with earlier helium droplet experiments on prototypical systems, *i.e.* monohydrated propargyl alcohol ($\text{H}_2\text{O} \cdots \text{HC} \equiv \text{C}-\text{CH}_2\text{OH}$).³⁶ $\text{HC} \equiv \text{C}-\text{CH}_2\text{OH}$ has a dipole moment of 1.53 D and the complexation proceeded *via* a dipole-steering mechanism leading to the formation of two local minimum structures stabilized by $\equiv \text{C}-\text{H} \cdots \text{OD}_2$ and $-\text{CH}_2\text{O}(\text{H}) \cdots \text{DOD}$ HB formation. In helium droplets, the typical time interval between successive pickup events is on the order of microseconds, while the cooling of molecules inside the droplets occurs within nanoseconds.³⁵ Consequently, each molecular unit picked up from different pickup events becomes fully thermalized to the droplet temperature (~ 0.4 K) before aggregation. At this ultralow temperature, thermal energy is negligible, and molecular aggregation is initially guided by long-range electrostatic forces, *i.e.* dipole-dipole interactions, which have $1/R^3$ distance (R) dependence. Considering PA (1.59 D) and D_2O (1.85 D) with substantial permanent dipole moments, dipole-dipole steering pathway plays a fundamental role in their aggregation process.

Now, the relative orientations of the dipole moment vectors of individual molecules in the complexes stabilized inside the helium droplets, **c-PAw-3**, **c-PAw-2**, and **c-PAw-1**, are found not to be head-to-tail but rather appear to be quite random (see Table 3). This observation can be explained by considering a two-step mechanism of dimer formation: (i) initial long-range approach governed by dipole-dipole realignment, (ii) subsequent short-range stabilization driven by the most feasible hydrogen bonding interactions during the time of contact. This successive pathway leading to the formation of local minimum structure for pyruvic and formic acid dimers in HNDs are already reported.^{21,31,33,53}

It is also important to emphasize that kinetic trapping to local minima at ~ 0.4 K is possible, as the system lacks

sufficient thermal energy to overcome even modest interconversion barriers. This phenomenon accounts for the significant experimental populations observed for the local minimum structures **c-PAw-3** ($\Delta E_0 = +22.9 \text{ kJ mol}^{-1}$) and **c-PAw-2** ($\Delta E_0 = +21.0 \text{ kJ mol}^{-1}$), despite the greater thermodynamic stability of the global minimum structure **c-PAw-1**. Notably, the order of interaction energies (D_0) among these isomers does not correlate with their observed abundances, further supporting the role of kinetic and not thermodynamic control under ultracold conditions (see Tables 2 and 3). The computed barrier for interconversion between **c-PAw-2** and **c-PAw-1**, involving reorientation of the water molecule, is approximately 2 kJ mol^{-1} (Fig. S7, left panel in the SI). While relatively low on an absolute energy scale, this barrier remains prohibitively high at 0.4 K. A relaxed scan along the angle between the $\text{C}=\text{O}$ group of the acid and the O atom of the water molecule was performed, which resulted the transition from **c-PAw-1** to a structure close to **c-PAw-4**. The pathway was found to be barrierless (see Fig. S7, right panel). Since we have already confirmed **c-PAw-4** as a real minimum, this indicates that a very low barrier exists from **c-PAw-4** to **c-PAw-1**. However, this barrier is small enough to be overcome by the energy released at time of interaction between PA and D_2O , which rationalizes why **c-PAw-4** is not observed experimentally. A similar scenario was observed in acetylene-furan trimer formation inside helium droplet.⁵⁴

So, the 1 : 1 $\text{PA} \cdots \text{D}_2\text{O}$ dimer highlight the crucial interplay between long-range dipole-dipole forces and shorter-range interactions in guiding aggregation pathways at ultracold temperatures. The vibrational features of these kinetically trapped higher-energy isomers in the $\text{PA} \cdots \text{D}_2\text{O}$ system may serve as distinctive spectral markers for identifying similar binding motifs in both terrestrial and extraterrestrial environments.

4. Conclusions

We report the characterization of the propiolic acid (PA) monomer and its binary complexes with D_2O in helium droplets using mass-selective infrared spectroscopy in the $\text{C}=\text{O}$ and $\text{C} \equiv \text{C}$ stretching regions. Five IR absorption bands of 1 : 1 $\text{PA} \cdots \text{D}_2\text{O}$ species reveal the stabilization of three isomers, with D_2O accessing multiple hydrogen-bonding sites on the acid. The most intense absorption bands are assigned to the local minimum structures stabilized by a single non-classical HB formation. The calculated interaction energies along with interconversion barriers between isomers clearly depicted the kinetic trapping of species at 0.4 K, a phenomenon commonly observed in helium droplets. These findings highlight the competing roles of dipole-dipole and higher-order interactions in directing complex formation at ultracold temperatures. The substantial dipole moments of D_2O (1.85 D) and PA (1.59 D) promote long-range directional hydrogen bonding, which is subsequently complemented by short-range stabilization through the most favorable hydrogen-bonding interactions upon contact.



Author contributions

A. C. has designed and carried out the experiments and computations. M. H. has set-up a helium droplet experiment in Bochum, managed the project and provided funding. Data analysis and interpretation were carried out by A. C., G. S., and M. H. S. H. has contributed to mathematical analysis. All authors participated in the preparation and proofreading of the manuscript.

Conflicts of interest

There is no conflict to declare.

Data availability

The data generated in this study have been deposited in our open-access repository: <https://doi.org/10.17877/RESOLV-2025-MF12V240>.

Supplementary information (SI): pressure dependent pick-up curves for the IR bands of the propiolic acid (PA) monomer and PA $\cdot\cdot$ D₂O dimers, comparison of the IR spectra of PA $\cdot\cdot$ D₂O varying the pick-up pressure of D₂O, calculated anharmonic spectra of PA, and optimized structures and coordinates of possible PA $\cdot\cdot$ D₂O and PA $\cdot\cdot$ H₂S dimers are provided. See DOI: <https://doi.org/10.1039/d5cp02794b>.

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