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Probing the Microsolvation Environment of the
Green Fluorescent Protein Chromophore in Vacuo

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ABSTRACT. We present vibrational and electronic photodissociation spectra of a model chromophore of the green fluorescent protein in complexes with up to two water molecules, prepared in a cryogenic ion trap at 160 – 180 K. We find the band origin of the singly hydrated chromophore at 20985 cm\(^{-1}\) (476.5 nm), and we observe partially resolved vibrational signatures. While a single water molecule induces only a small shift of the \(S_1\) electronic band of the chromophore, without significant change of the Franck-Condon envelope, the spectrum of the dihydrate shows significant broadening and a greater blue shift of the band edge. Comparison of the vibrational spectra with predicted infrared spectra from density functional theory indicate that water molecules can interact with the oxygen atom on the phenolate group or on the imidazole moiety, respectively.

TOC GRAPHICS
Many proteins incorporate water molecules as a part of their functional structure.\textsuperscript{1-2} As a result, microsolvation is an important structural aspect that influences the properties of functional regions in proteins. For example, in many fluorescent proteins, such as those based on the green fluorescent protein (GFP), a water molecule binds to the anionic phenolate group of the chromophore that is situated within the double-barrel protein structure.\textsuperscript{3} The influence of the water molecule on the photophysics of the chromophore is therefore of great interest. One way of probing water-chromophore interactions is to use a model chromophore, which in GFP is a small anion consisting of a phenolate group connected by a methine bridge at the para-position to an imidazole group. This chromophore is represented very well by deprotonated \textit{p}-hydroxybenzylidene-2,3-dimethylimidazolinone (HBDI\textsuperscript{−}, see Supporting Information, Scheme S1), where the two methyl groups on the imidazole moiety stand in for the connection to the protein. The addition of water molecules to this chromophore can alter its electronic structure and such changes should be reflected in the absorption spectrum of the chromophore. The $S_1 \leftarrow S_0$ absorption spectrum of HBDI\textsuperscript{−} in aqueous solution is strongly blue-shifted from that of the protein, as the peak absorption changes from ca. 480 nm to ca. 426 nm.\textsuperscript{4} In contrast, the photodissociation spectrum of HBDI in \textit{vacuo} bears a striking resemblance to the absorption spectrum of GFP, which can be observed in even more detail for cryogenically prepared HBDI\textsuperscript{−} and GFP at 77 K, with a blue shift in the protein by just 300 cm\textsuperscript{-1}.\textsuperscript{4,6} The comparison suggests that the interactions with the protein environment, including neighboring charged residues and the water in the chromophore pocket, bring the electronic structure of the chromophore closer to vacuum than to aqueous solution. Consequently, there has been much interest in the photophysics of HBDI in \textit{vacuo} over the past two decades.\textsuperscript{4,6-32} However, the influence of water-HBDI\textsuperscript{−} interaction on the electronic spectrum of the chromophore has so far only been investigated computationally.\textsuperscript{13} One of the questions in this
context is whether the water molecule in the chromophore pocket of GFP has a strong influence on the electronic spectrum of the chromophore, which would then have to be countered by the rest of the interactions in the pocket.

In this work, we present vibrational and electronic spectroscopy on mass selected clusters of HBDI\(^-\) with up to two water molecules that have been cryogenically prepared in an ion trap. We show that in the absence of the constraints imposed by the chromophore pocket, water molecules can bind to the phenolate or to the oxygen on the imidazole moiety, and we can clearly distinguish between the two binding sites in the vibrational spectrum.

Figure 1 shows the photodissociation spectra of the S\(_1\) \(\leftrightarrow\) S\(_0\) absorption band of HBDI\(^-\)·(H\(_2\)O)\(_n\) \((n = 1, 2)\) at 160 – 180 K, compared to the spectrum of bare HBDI\(^-\) taken at 30 K trap temperature\(^6\) (see Supporting Information for experimental and computational methods). The overall shapes of the spectra for HBDI\(^-\) and HBDI\(^-\)·H\(_2\)O are nearly identical. The vibrational features in the Franck-Condon profile of the spectrum of the monohydrate are less well resolved than for bare HBDI\(^-\), but we assign the band origin to a partially resolved peak at \((20985 \pm 30)\) cm\(^{-1}\), representing a 55 cm\(^{-1}\) blue shift from bare HBDI\(^-\). Additional vibrational features are a partially resolved peak at ca. 21120 cm\(^{-1}\), and a shoulder at 21195 cm\(^{-1}\). These features are likely the remnants of the Franck-Condon progression in a low-frequency bending vibration of the methine bridge.\(^6\) We assign the set of peaks around 22200 cm\(^{-1}\) to an imidazole ring breathing/in-plane CH bending mode and its combination bands with the aforementioned methine bridge bending mode.\(^6, 9, 12, 33\) While the addition of a single water molecule does not seem to cause much change in the electronic spectrum of HBDI\(^-\), the spectrum of the dihydrate shows a significantly altered envelope, with a broad feature around 21400 cm\(^{-1}\), followed by another around 22130 cm\(^{-1}\). We cannot extract a clean band origin from the spectrum, but we find that the half-maximum point on the rising edge of the
first feature is blue-shifted by ca. 180 cm$^{-1}$ from the analogous point in the spectrum of the monohydrate. A better understanding of the effects of hydration on the properties of HBDI$^-$ requires a detailed structural characterization of the clusters studied here.

**Figure 1.** From top: Absorption spectrum of wild-type GFP at 77 K (green); photodissociation spectra of HBDF at 30K (black) and HBDI$^-$·(H$_2$O)$_n$ ($n = 1, 2$, magenta and blue) in the visible spectral region; UV/vis absorption spectrum of aqueous HBDI$^-$ at pH 11. All traces were normalized to the same peak height. Photofragment signals were the loss of a methyl group for bare HBDI$^-$, and of all H$_2$O molecules for HBDI$^-$·(H$_2$O)$_n$, respectively. The vertical dashed line marks the band origin of bare HBDI$^-$ at 20930 cm$^{-1}$. The data for bare HBDI$^-$ were taken from ref. 6, the absorption spectrum of wild-type GFP was digitized from ref. 5.
We begin the structural characterization of HBDI·(H₂O)ₙ (n = 1, 2) clusters by determining which isomer of HBDI⁻ is present, as the chromophore can exist in two different isomeric forms denoted E and Z (see Scheme S1 in Supporting Information), which are connected by torsion around the methine bridge. Both of these isomers are encountered in proteins, but only the Z isomer is fluorescent (present in GFP), while the E isomer is not.⁶ We have recently used IR spectroscopy to establish that the Z isomer is the dominant species if prepared at low temperatures (30 K), in complexes with weakly bound N₂ messenger tags.⁶ In the present experiment, we formed HBDI⁻·(H₂O)ₙ clusters by condensing water molecules from the background gas of the vacuum apparatus onto HBDI⁻ at trap temperatures of 160-180 K. Figure 2 shows the infrared fingerprint region of the HBDI⁻ vibrations for HBDI⁻·(H₂O)ₙ (n = 1, 2) compared to that of HBDI⁻·N₂, which we previously reported and assigned to the Z isomer.⁶ Our assignment of the infrared spectrum to the Z isomer relies primarily on the spacing of the three intense features around 1600 cm⁻¹. Based on the comparison in Figure 2, we determine that the Z isomer is dominant in the water complexes, consistent with ion mobility results reported by Bieske and coworkers³⁰ where solvent complexes (with methanol in their work) only existed for the Z isomer.
Figure 2. Infrared photodissociation spectra of HBDI·N₂ and HBDI·(H₂O)ₙ (n = 1, 2) in the fingerprint region of HBDI⁻ vibrations. The photofragment signals were due to the loss of N₂ for HBDI·N₂ and of a single H₂O molecule for HBDI·(H₂O)ₙ, respectively. The spectrum of HBDI·N₂ was taken from ref. 6.

The positions of the most intense vibrational signatures of HBDI·N₂ shown in Figure 2 are only weakly influenced by the microsolvation environment (by up to ca. 10 cm⁻¹ per water molecule, see Supporting Information, Table S1), showing a minor influence of the water adducts on the robust HBDI⁻ frame, in line with computational predictions by Krylov and coworkers.¹³ The observed changes in vibrational frequencies are likely due to small changes in the charge distribution that results from water molecules binding to either of the oxygen atoms (see
Supporting Information, Table S2). Some of the observed changes suggest that hydration occurs on the phenolate moiety, for example a slight blue shift observed for the imidazole CO stretching vibration found in HBDI·N₂ at ca. 1680 cm⁻¹, and the enhancement of a phenolate CO stretching signature at 1396 cm⁻¹ in the dihydrate spectrum. However, a weak band around 1650 cm⁻¹ is consistent with hydration on the imidazole, so we can expect a mixture of conformers to be present. A more detailed discussion of the vibrational spectrum in the fingerprint region of HBDI·(H₂O)ₙ⁻ and comparison with predicted spectra for various hydration conformers is given in Supporting Information.

While the fingerprint region (1000 cm⁻¹ to 2000 cm⁻¹) shows the (weak) effects of microhydration on the solute, the OH stretching region encodes microhydration from the solvent point of view, and presents a clearer picture, as the OH stretching frequencies of the water molecules are highly sensitive probes of their hydrogen bonding environment. Figure 3 shows the experimental infrared spectra of HBDI·N₂, and of the mono- and dihydrate of HBDI⁻ over the range of 2700 to 3800 cm⁻¹. The spectra of HBDI·(H₂O)ₙ are dominated by broad features in the region of 3100 cm⁻¹ to 3500 cm⁻¹, which contain the signatures of the hydrogen bonded (H-bonded) OH groups of the water molecules. The free OH signature appears at 3703 cm⁻¹. In the range of 2800 cm⁻¹ to 3100 cm⁻¹, the spectrum of HBDI·N₂ carries the signatures of the CH stretching vibrational modes, which are also observed in the spectra of the hydrated ions (see Supporting Information for a more detailed discussion of the CH stretching modes).
Figure 3. Infrared photodissociation spectra of HBDI·N$_2$ (top, black) and HBDI·(H$_2$O)$_n$ ($n = 1, 2$, magenta and blue) from 2700 to 3800 cm$^{-1}$. The photofragment signals were the loss of N$_2$ and a single H$_2$O molecule for HBDI·N$_2$ and HBDI·(H$_2$O)$_n$, respectively.

Figure 4 shows a comparison of the experimental and simulated infrared spectra for several calculated conformers of the monohydrate complex, HBDI·H$_2$O. We assign the broad feature around 3180 cm$^{-1}$ to the signature of the symmetric stretching mode of a water molecule solvating the phenolate group (conformer Ia), which is largely localized on the OH group H-bonded to the ion. There is an analogous conformer (Ia’, see Supporting Information, Figure S1) anti to the methine bridge CH group, which is isoenergetic (within 5 meV) with conformer Ia and has the same infrared spectrum. The feature around 3340 cm$^{-1}$ carries the signature of a water molecule
solvating the imidazole oxygen (conformers Ib and Ic). The observation of both features in the spectrum of HBDI·H₂O shows that a mixture of the two hydration conformers is present in the experiment. The two conformers for hydration at the imidazole (Ib and Ic) are separated by a barrier of ca. 40 meV. The widths of the infrared bands reflect the fact that the complexes are formed through an evaporative ensemble, with internal energies of the order of the binding energy of a water molecule (see discussion below). As a result, the water molecule is likely to explore a significant range of OH···O⁻ angles through large amplitude motions, both in the plane of the HBDI⁻ ion and perpendicular to it, and this motion results in broadening of the corresponding OH stretching transitions (see Supporting Information).

Overall, we obtain similar structures and energetic ordering as found by Krylov and coworkers. The lowest energy structure of the monohydrate corresponds to hydration on the phenolate (structure Ia). Hydration on the imidazole is ca. 100 meV higher in energy, where the oxygen lone pair of the water molecule can be weakly tethered to the nearest methyl group (structure Ib) or to the CH group on the methine bridge (Ic, the only imidazole-binding structure reported by Krylov and coworkers). Conformers Ib and Ic have different OH stretching frequencies and the geometries sampled by large amplitude motion connecting the two structures will contribute to the broadening of the OH stretching feature of the OH group H-bonded to the imidazole (see Supporting Information).
Figure 4. Calculated infrared spectra of several conformers of monohydrated HBDI (lower traces), compared to the experimental spectrum (upper trace). The spectrum contains signatures for CH stretching vibrations (CH), H-bonded OH stretching on the phenolate (red-shaded region) and imidazole (blue-shaded region) groups, and the free OH stretching mode (F). All calculations have been scaled by 0.957 to match the experimentally observed free OH stretching vibrations. The calculated energies (at 0 K) of the conformers relative to the lowest energy conformer (Ia) are given in each simulated trace.

In order to obtain a coarse estimate of relative populations for the two hydration sites, we fitted the two H-bonded OH features with Gaussian profiles and compared the peak areas scaled by the calculated infrared intensities for conformers Ia and Ib. The result suggests that in ca. 60% of the
complexes the water molecule is bound to the phenolate, and ca. 40% have a hydrated imidazole structure. Interestingly, the estimated population ratio of the two conformers is not consistent with a thermal population. In principle, the different binding energies of the water molecules to the different binding sites could lead to overestimating the imidazole population by suppressing photodissociation from the more strongly bound conformers. The binding energy at the imidazole site (calculated at 3740 cm$^{-1}$) is less than at the phenolate (calculated at 4550 cm$^{-1}$). However, there is considerable energy in the complexes, since they are formed in the ion trap from an evaporative ensemble. The energy in a complex formed this way is of the order of the binding energy of a water molecule in the dihydrate, which is calculated to be between 3300 cm$^{-1}$ and 4500 cm$^{-1}$. As a result, the internal energy in the complex assists in dissociation after photon absorption. This is also evidenced by the fact that we can record infrared photodissociation spectra in the carbonyl stretching region (see Figure 2). To explain the estimated population ratio, we assume that water migration between the two binding sites is hindered by a large barrier, leading to kinetically trapped species. We hypothesize that the formation of the two different hydration conformers can be better described by cross sections for the capture of a water molecule in the ion trap by the two different binding sites of HBDI$^-$. We note that future experiments that could employ separation of water adduct formation and messenger tagging together with hole-burning spectroscopy could allow the observation and assignment of individual hydration conformers.

The spectrum of the dihydrate in the OH stretching region is quite similar to that of the monohydrate. It is dominated by a broad feature around 3190 cm$^{-1}$ and a second broad feature around 3340 cm$^{-1}$. The two water molecules can be either H-bonded to the same oxygen, or one can each be bonded to the two binding sites, resulting in three possible binding site occupation combinations. There are several conformers for the same occupation of binding sites, since there
is some conformational flexibility within each binding site, and these conformers are close in energy. Figure 5 shows a comparison of calculated infrared spectra for several conformers with the experimental spectrum.

**Figure 5.** Calculated infrared spectra of several conformers of HBDI\( \cdot \)\((\text{H}_2\text{O})_2\) (lower traces), compared to the experimental spectrum (upper trace). The points in the upper trace are raw data, the full black line is an 11-point gliding average to guide the eye. The spectrum contains signatures for CH stretching vibrations (CH), OH stretching H-bonded to the phenolate (red-shaded region) and imidazole (blue-shaded region) groups, and the free OH stretching mode (F). All calculations have been scaled by 0.957 to match the experimentally observed free OH stretching vibrations. The calculated energies (at 0 K) of the conformers relative to the lowest energy conformer (IIa) are given in each simulated trace. Arrows in the lowest trace show how the infrared features of conformer IIa change as the H-bond between the water molecules is lost (see text). Arrows in the
trace for conformer IIId show the expected analogous behavior upon breaking the water-water H-
bond for a conformer with both water molecules on the imidazole. The full and dashed lines in the 
trace for conformers IIc and IIc’ correspond to the water molecule positions indicated by full and 
dashed circles, respectively.

Conformer IIa is the lowest energy conformer (at 0 K), with both water molecules binding to the 
phenolate, and forming a water-water H-bond. Note, however, that the experimental spectrum does 
not show any feature that would indicate that the two water molecules form a water-water H-bond, 
which is predicted at ca. 3600 cm\(^{-1}\). Experiments by Johnson and coworkers on \(\Gamma \cdot (H_2O)_2\) 
demonstrated that the water-water H-bond in that cluster does not survive at the trap temperatures 
in the present experiment, easily explaining the absence of the water-water H-bond signature.\(^{40}\) 
Based on their work, we expect the two ionic H-bond signatures to shift closer together, 
concomitant with breaking the water-water H-bond, behaving like independent monomers rather 
than dimers. The corresponding conformer (IIb) in the present case is in fact a stable minimum 
energy structure, which is practically isoenergetic with conformer IIa. At the internal energies of 
the complexes, the significantly higher entropy of conformer IIb will drive breaking the water-
water H-bond, and we expect that conformer IIa is not in fact populated at this temperature. As a 
result, the lower frequency feature in the experimental spectrum is still indicative of hydration on 
the phenolate (conformers IIa, IIb), while the higher frequency feature signals hydration on the 
imidazole (conformers IIc, IIc’, IIId). We note that at 0 K, there is no minimum energy structure 
with two independent water molecules at the imidazole. However, we still expect the same entropy 
driven loss of the water-water H-bond to occur for this hydration configuration, with the 
concomitant behavior of the calculated infrared signatures of conformer IIId indicated by the arrows 
shown in Figure 5.
The observed dihydrate spectrum is consistent with a mixture of conformers corresponding to doubly and two singly hydrated binding sites, but due to the multitude of different conformers, we refrain from estimating the ratio of the different conformers here. The free OH signature in the dihydrate spectrum is broader than in the monohydrate spectrum, reflecting the composition of the spectrum from several hydration conformers with slightly varying free OH frequencies. The relative energies for the dihydrate favor hydration of the phenolate by both water molecules, followed by one water molecule on each binding site (ca. 70 meV higher in energy), with two water molecules on the imidazole being highest in energy (ca. 180 meV above dihydrated phenolate), in line with calculations by Krylov and coworkers.13

With our insight into the structures of hydrated HBDI\textsuperscript{-}, we can return to the electronic spectra of these clusters. Calculations by Krylov and coworkers\textsuperscript{13} at the SOS-CIS(D)/cc-pVTZ level predict a blue shift of 240 cm\textsuperscript{-1} for hydration at the phenolate moiety and a red shift by 160 cm\textsuperscript{-1} for hydration at the imidazole group. Our calculations, done at the CAM-B3LYP/def2-TZVPP level of theory, suggest a 9 cm\textsuperscript{-1} red shift and a 54 cm\textsuperscript{-1} blue shift for hydration at the phenolate and imidazole respectively. Based on the experimental electronic spectrum of cryogenically prepared HBDI\textsuperscript{-}, the vertical excitation energies are equal to the band origin energies in the Franck-Condon region, which represents a shallow minimum on the S\textsubscript{1} potential energy surface.\textsuperscript{6} We assume that the first feature in the monohydrate spectrum at 20985 cm\textsuperscript{-1}, which is broader than the band origin feature for bare HBDI\textsuperscript{-}, contains the band origins of both hydration conformers. The splitting between the band origins of the two hydration conformers in our calculations are quite close to the experimentally observed width of this feature, although the good agreement is likely to be fortuitous. The splitting calculated by the Krylov group\textsuperscript{13} (400 cm\textsuperscript{-1}) is too large to maintain the experimentally observed width of the Franck-Condon profile. It is worth comparing the small blue
shift upon hydration with the blue shift caused by betaine, an adduct with a much stronger dipole moment.\textsuperscript{31} The shift induced by betaine (ca 12 D dipole moment\textsuperscript{31}) is 640 cm\textsuperscript{-1}, while the shift induced by water (1.85 D dipole moment) is 55 cm\textsuperscript{-1}, i.e., the shift introduced by water is even lower (by a factor of 1.8) than expected based on the electric field of betaine.

The calculated vertical excitation energies for the three hydration conformer families of the dihydrate span ca. 300 cm\textsuperscript{-1}, and several of the conformers exhibit a red shift (see Supporting Information, Table S3). These calculations do not explain the blue shift of the spectrum or deviation from the characteristic band envelope found in HBDI\textsuperscript{-}\cdot N\textsubscript{2} and HBDI\textsuperscript{-}\cdot H\textsubscript{2}O. Previous calculations by Krylov and coworkers\textsuperscript{13} arrived at a difference of 960 cm\textsuperscript{-1} between the different hydration conformers for the dihydrate, which could lead to a change in the observed band shape. We hypothesize that the change observed in the envelope of dihydrate spectrum is indeed caused by a mixture of different hydration conformers. However, in light of the complications arising from the mix of hydration conformers and the lack of clear computational description, we refrain from a more definitive assignment. Clearly, the currently available calculations are not able to satisfactorily explain the dihydrate spectrum, particularly its significant blue shift compared to that of the monohydrate. We note that the larger effect of the second water molecule is unexpected, since most often the first one or two water molecules each introduce a significant shift, while additional water adducts result in less dramatic changes.\textsuperscript{41-42} This is clearly not the case here, where the second water adduct brings the spectrum much closer to bulk solution than the first (see Figure 1). Future work with larger hydrated clusters could shed light on the transition to full hydration.

In summary, clusters of HBDI\textsuperscript{-} with up to two water molecules are based on the Z isomeric form of the chromophore itself. There are two binding sites for water molecules in HBDI\textsuperscript{-}, the oxygen
atoms on the phenolate and the imidazole moieties, respectively, and we observe hydration on both. For the monohydrate, the relative population on each binding site cannot be described by thermal conformer distributions, and we assume that the populations are given by capture of a water molecule into either binding site and subsequent cooling and kinetic trapping. The dihydrate shows a mixture of hydration conformers, and at the range of trap temperatures in the present work, we do not find the signatures of water-water hydrogen bonding. In contrast to the dihydrate, whose electronic spectrum envelope deviates from that of bare HBDI\(^-\), a single water molecule interacting with HBDI\(^-\) does not significantly change the envelope of the \(S_1 \leftarrow S_0\) electronic band, and the shift of the band origin from bare HBDI\(^-\) is only +55 cm\(^{-1}\). In the context of interpreting HBDI\(^-\) as a model for the GFP chromophore, this observation implies that the ca. 300 cm\(^{-1}\) shift caused by the chromophore environment is not caused by the functional water molecule that binds to the phenolate moiety in the protein.

**ASSOCIATED CONTENT**

**Supporting Information.** Isomers of HBDI\(^-\); experimental and computational methods; vibrational assignments for HBDI\(^-\)·N\(_2\) and HBDI\(^-\)·(H\(_2\)O)\(_n\) (n = 1, 2) in the fingerprint region; calculated selected charges and bond lengths in HBDI\(^-\) and different position isomers for HBDI\(^-\)·(H\(_2\)O)\(_n\) (n = 1, 2); calculated vertical excitation energies for all structures; discussion of the influence of water adducts on the fingerprint region of HBDI\(^-\) vibrational modes; comparison of calculated and experimental infrared spectra of HBDI\(^-\)·(H\(_2\)O)\(_n\) (n = 1, 2) from 1350 cm\(^{-1}\) to 1800 cm\(^{-1}\); calculated hydrogen bonded OH stretching frequency as a function of hydration geometry in HBDI\(^-\)·H\(_2\)O in the plane of HBDI\(^-\); calculated hydrogen bonded OH stretching frequency as a
function of hydration geometry in HBDI·H$_2$O perpendicular to the plane of HBDI; discussion of CH stretching modes; all calculated structures with atomic coordinates and frequencies.

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