Resolving the Ultrafast Dynamics of the Anionic Green Fluorescent Protein Chromophore in Water

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The chromophore of the green fluorescent protein (GFP) is critical for probing environmental influences on fluorescent protein behavior. Using the aqueous system as a bridge between the unconfined vacuum system and a constricting protein scaffold, we investigate the steric and electronic effects of the environment on the photodynamical behavior of the chromophore. Specifically, we perform ab initio multiple spawning to simulate five picoseconds of nonadiabatic dynamics after photoexcitation, resolving the excited-state pathways responsible for internal conversion in the aqueous chromophore. We identify an ultrafast pathway that proceeds through a short-lived (sub-picosecond) imidazolinone-twisted (I-twisted) species and a slower (several picoseconds) channel that proceeds through a long-lived phenolate-twisted (P-twisted) intermediate. The molecule navigates the non-equilibrium energy landscape via an aborted hula-twist-like motion towards the one-bond-flip dominated conical intersection seams, as opposed to following the pure one-bond-flip paths proposed by the excited-state equilibrium picture. We interpret our simulations in the context of time-resolved fluorescence experiments, which use short and long time components to describe the fluorescence decay of the aqueous GFP chromophore. Our results suggest that the longer time component is caused by an energetically uphill approach to the P-twisted intersection seam rather than an excited-state barrier to reach the twisted intramolecular charge transfer species. Irrespective of the location of the non-adiabatic population events, the twisted intersection seams are inefficient at facilitating isomerization in aqueous solution. The disordered and homogeneous nature of the aqueous solvent environment facilitates non-selective stabilization with respect to I- and P-twisted species, offering an important foundation for understanding the consequences of selective stabilization in heterogeneous and rigid protein environments.

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Resolving the ultrafast dynamics of the anionic green fluorescent protein chromophore in water

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Abstract:
The chromophore of the green fluorescent protein (GFP) is critical for probing environmental influences on fluorescent protein behavior. Using the aqueous system as a bridge between the unconfined vacuum system and a constricting protein scaffold, we investigate the steric and electronic effects of the environment on the photodynamical behavior of the chromophore. Specifically, we perform ab initio multiple spawning to simulate five picoseconds of nonadiabatic dynamics after photoexcitation, resolving the excited-state pathways responsible for internal conversion in the aqueous chromophore. We identify an ultrafast pathway that proceeds through a short-lived (sub-picosecond) imidazolinone-twisted (I-twisted) species and a slower (several picoseconds) channel that proceeds through a long-lived phenolate-twisted (P-twisted) intermediate. The molecule navigates the non-equilibrium energy landscape via an aborted hula-twist-like motion towards the one-bond-flip dominated conical intersection seams, as opposed to following the pure one-bond-flip paths proposed by the excited-state equilibrium picture. We interpret our simulations in the context of time-resolved fluorescence experiments, which use short and long time components to describe the fluorescence decay of the aqueous GFP chromophore. Our results suggest that the longer time component is caused by an energetically uphill approach to the P-twisted intersection seam rather than an excited-state barrier to reach the twisted intramolecular charge transfer species. Irrespective of the location of the non-adiabatic population events, the twisted intersection seams are inefficient at facilitating isomerization in aqueous solution. The disordered and homogeneous nature of the aqueous solvent environment facilitates non-selective stabilization with respect to I- and P-twisted species, offering an important foundation for understanding the consequences of selective stabilization in heterogeneous and rigid protein environments.
Introduction

Since their initial characterization in *Aequorea victoria*,1-5 fluorescent proteins (FPs) have emerged as powerful probes for *in vivo* biological function.6-11 The green fluorescent protein8 (GFP) is the monarch of fluorescent proteins, and its utility and versatility keep the green fluorophore at the forefront of science. Seminal experiments on the color and protonation state tuning of GFP12-17 inspired novel FP biomarkers, with reversible photoswitchable fluorescent proteins propelling the resolution of FPs beyond the diffraction limit.16-18 Located at the heart of GFP, a tunable chromophore is responsible for GFP’s light emission. Because the native characteristics of GFP can be altered significantly by applying point mutations near the site of the chromophore,8 we need a detailed understanding of the chromophore’s local steric and electronic environment to optimize the versatility of GFP. Frequently, *p*-hydroxybenzylidene-2,3-dimethylimidazolinone (HBDI), first synthesized by Kojima *et al.*,19 serves as a model GFP chromophore. In particular, the anionic form of HBDI (HBDI⁻) mimics the protonation state of the chromophore responsible for fluorescence in the protein.20-21

The overwhelming number of variables in a complex protein environment creates a fundamental roadblock to the rational design of FPs. Due to the rigidity of its 11-stranded β-barrel,22-23 GFP envelops its chromophore in a highly ordered pocket of amino acids. Stiff protein backbones foster heterogeneous electronic and steric fingerprints that can significantly impact the chromophore’s behavior.8 Because amino acids in GFP form complex networks of covalent and non-covalent interactions, it is difficult to predict the net effect of various residues acting on the chromophore. To filter out the effects that dictate chromophore behavior, we use HBDI⁻ to monitor the molecule’s response to incremental perturbations of the environment. Using the gas-phase dynamics of HBDI⁻ as a baseline,24 we identify the influence of specific effects on chromophore behavior. Here, we focus on a key system that unravels the role of environmental influences on GFP chromophore photodynamics – the homogeneous and disordered case of aqueous HBDI⁻.

Solvated HBDI⁻ straddles the line between the freedom permitted by the gas-phase and the confinement imposed by a protein. In water, HBDI⁻ experiences steric and electronic effects from the solvent, which are absent *in vacuo*, while in an environment that is less constricting than a protein scaffold. Time-resolved spectroscopies reveal that GFP has an excited-state lifetime on the order of nanoseconds20, 25-26 and a fluorescence quantum yield near unity.8 Yet, HBDI⁻ experiences ultrafast internal conversion in vacuum and in solution at ambient temperature.27-31 Twisting around the bridge bonds connecting the imidazolinone (I) and phenolate (P) moieties of the isolated chromophore leads to
ultrafast deactivation that decreases fluorescence quantum yield.\textsuperscript{31-33} As highlighted in previous computational studies,\textsuperscript{34-35} non-methylated HBDI\textsuperscript{−} (pHBI\textsuperscript{−}) undergoes twisted intramolecular charge transfer (TICT) as rotations of the I- and P-moieties become sufficiently large. The direction of charge transfer (CT) on the excited state depends on whether the I-moiety or the P-moiety rotates (about the $\phi_I/\phi_P$ dihedrals indicated in Figure 1, respectively). Whereas rotation of $\phi_I$ induces CT from the I-ring to the P-ring in the gas-phase, rotation of $\phi_P$ induces CT from the P-ring to the I-ring.

Using ultrafast fluorescence and polarization spectroscopy experiments,\textsuperscript{27, 36-37} Meech and co-workers identified similar short (hundreds of femtoseconds) and long (picoseconds) decay components for HBDI\textsuperscript{−} across a variety of solvents. In water, time constants of 210 fs and 1.1 ps were reported for fluorescence decay — significantly faster than the 1.3/11.5 ps excited-state lifetimes reported \textit{in vacuo}.\textsuperscript{21, 30} Early quantum chemical calculations of gas-phase pHBI\textsuperscript{−}\textsuperscript{38} suggested that a volume-conserving “hula-twist” deactivation mechanism\textsuperscript{39} is responsible for the limited sensitivity of the chromophore’s excited-state lifetime on solvent viscosity. However, more recent \textit{in silico} experiments implicate drastic torsional motion along either of the methine bridge bonds as responsible for the ultrafast deactivation of the anionic chromophore.\textsuperscript{34, 40} This behavior is corroborated by high-level quantum mechanics/molecular mechanics (QM/MM) simulations in water using extended multistate multireference second-order perturbation theory (XMS-CASPT2)\textsuperscript{41} and spin-flip time-dependent density functional theory.\textsuperscript{42} These studies found that the dominant decay pathway of pHBI\textsuperscript{−} is through the I-twisted channel. However, the limited simulation time considered (1 ps) did not allow identification of the sources responsible for the multi-timescale decay in solution. To resolve this, we simulate multiple picoseconds of the aqueous HBDI\textsuperscript{−} photodynamics while monitoring important geometric and electronic signatures of the system.

Ultrafast dispersed pump-dump-probe experiments of HBDI\textsuperscript{−} in water present excited-state decay lifetimes in the range 620-940 fs for twisted excited-state intermediates.\textsuperscript{43} Although the imperfect selectivity of a dump pulse may make it difficult to attribute time constants to specific processes, the Z/E isomerization quantum yield derived from these experiments was below the detection threshold of ~5%. While torsion along $\phi_I$ can result in the $E$-isomer ($Z/E$ isomerization), torsion along $\phi_P$ will regenerate the $Z$-isomer due to the symmetry of the P-ring ($Z/Z$ isomerization). Thus, $Z/Z$ isomerization products are experimentally indistinguishable from decay pathways that do not involve isomerization. For this reason, it remains unclear what ratio of excited-state species proceeds toward photoproduct formation ($Z/E$ isomerization), undergoes P-ring flip ($Z/Z$ isomerization), or re-populates the original ground state.
without torsion (Z-isomer). A key ingredient of QM/MM (or fully QM) simulations of photochemistry is the proper description of the potential energy surface topography around conical intersections (CIs). Many electronic structure methods are not capable of correctly describing $S_0/S_1$ CI regions\(^ {44-46}\) (e.g. time-dependent density functional theory and other single-reference methods). Following our recent gas-phase study,\(^ {24}\) we use the $\alpha$-CASSCF method,\(^ {47}\) which provides a correct and efficient description of the exact degeneracy and relative energetics along key deactivation coordinates. This enables us to extend our simulations to multiple picoseconds. Consequently, we are able to monitor the chromophore’s approach toward the CI seams and provide predictions for the photoproduct branching. The shape of conical intersections (e.g. peaked vs. sloped) can also be critically important in determining the photoproducts.\(^ {48-51}\) Although it appears that the CIs become more peaked upon solvation for the neutral GFP chromophore\(^ {52-53}\) (suggesting a shortening of the excited-state lifetime and a larger isomerization quantum yield), the shape of the CIs that govern nonadiabatic transitions for HBDI\(^ { - }\) in water has been largely unexplored.

Recent femtosecond stimulated Raman spectroscopy (FSRS) and transient absorption experiments of aqueous HBDI\(^ {31}\) fill in some of the gaps left by ultrafast fluorescence experiments. While time-resolved fluorescence experiments elucidate the temporal signatures of highly fluorescent HBDI\(^ { - }\) species, i.e., closer to the Franck–Condon (FC) point, FSRS is able to track the progression of HBDI\(^ { - }\) along vibrational modes of dark, non-fluorescent species. FSRS studies yield three time constants for the excited-state dynamics of HBDI\(^ { - }\): $\sim$350 fs, 2-3 ps, and $\sim$250 ps.\(^ {31}\) These were tentatively assigned to the formation of a charge-separated (CS) species, the transition from the CS state to a TICT state, and the ground-state recovery of the Z-isomer, respectively.

In this work, we explore the internal conversion dynamics following photoexcitation of explicitly-solvated HBDI\(^ { - }\). Using \textit{ab initio} multiple spawning\(^ {54-58}\) (AIMS), we provide insight into the primary deactivation pathways in aqueous solution and each pathway’s relative accessibility to CIs. In particular, we investigate the characteristics of HBDI\(^ { - }\) (i) at the FC region, (ii) out of the FC region as the molecule approaches CI seams, (iii) upon encountering the CI seam, and (iv) away from the CI seam as it returns to the ground state. This fully non-equilibrium behavior (chromophore and solvent response) is compared to the corresponding excited-state equilibrium solvation picture, defined by the free energy surface of aqueous HBDI\(^ { - }\). Our results are interpreted in the context of previous experiments performed \textit{in silico}, \textit{in vacuo}, and in solution. Using this approach, we discuss the impact of solvation on HBDI\(^ { - }\) and develop a more complete picture of GFP chromophore photophysics.
Computational Details and Methods

Because of its efficiency and ability to describe CIs, state-averaged59 complete active space self-consistent field (SA-CASSCF)60-62 was chosen as the electronic structure method for HBDI\(^{-}\) in water. To increase the accuracy of SA-CASSCF, we correct for the lack of dynamic electron correlation in SA-CASSCF calculations by using \(\alpha\)-CASSCF47 (Figure S1). The \(\alpha\)-CASSCF method43 uses energy scaling to approximate the effects of dynamic electron correlation at the cost of traditional SA-CASSCF, similar to previous scaling methods introduced by Olivucci and coworkers.64 Orbitals are optimized to minimize the average energy of the three lowest singlet electronic states in SA-CASSCF, as this has been shown to provide an accurate excited-state description of I- and P-torsional pathways.65 In particular, inclusion of three states captures the P-twisted minimum energy conical intersection (MECI) of HBDI\(^{-}\), which can be elusive when using a two-state averaging method with CASSCF.66 An active space of four electrons in three orbitals, i.e., SA3-CASSCF(4,3), incorporates the bonding, non-bonding and antibonding \(\pi\)-orbitals relevant to describe the allylic anionic character of the methine bridge (Figure 2). Unless stated otherwise, the 6-31G* basis set67 is used throughout.

Figure 1. Structure of the HBDI\(^{-}\) model chromophore illustrating defined geometric parameters. Cyan atoms represent carbon, red atoms represent oxygen, blue atoms represent nitrogen, and white atoms represent hydrogen. Blue, red, and orange bonds connect atoms belonging to the P-ring, I-ring, and methine bridge, respectively. \(\phi\) is defined as the dihedral angle formed by \(N_i\) and the carbon atoms of the methine bridge. \(\phi_p\) is defined as the dihedral angle formed by the carbon atoms of the methine bridge and \(C_P\). The pyramidalization of the central methine carbon atom, \(\theta_{pyr}\), is calculated as \(\cos^{-1}\left(\left(\vec{r}_p \times \vec{r}_I\right) \cdot \vec{r}_H\right) - \pi/2\) and uses the bridge bond vectors in a method similar to Radhakrishnan et al.68 For reference, the pyramidalization angle of ideal sp\(^2\) and sp\(^3\) hybridized carbon atoms using this definition are 0° and 55°, respectively.
Figure 2. Active space orbitals of HBDI from α(0.67)-SA3-CASSCF(4,3) at the FC point: the anionic, allylic (a) bonding, (b) non-bonding, and (c) anti-bonding orbitals.

The α parameter was chosen to reproduce the $S_0 \rightarrow S_1$ vertical excitation energy of the planar chromophore in solution, computed with the SA3-XMS-CASPT2(4,3) method\textsuperscript{69-71} ($1s$ orbitals of heavy atoms were excluded from the perturbation treatment and a level shift of 0.3 a.u. was used). This led to an α value of 0.67 for hydrated HBDI (Figure S1). This is very similar to the α value of 0.64 which we found for isolated HBDI\textsuperscript{-} (used in the gas-phase calculations reported herein).\textsuperscript{24} As shown in Figure S1, this α-CASSCF method reproduces XMS-CASPT2 potential energy profiles along the two bridge torsional coordinates ($\phi_I$ and $\phi_P$). Unless otherwise stated, all calculations were performed with α(0.67)-SA3-CASSCF(4,3)/6-31G* in explicit solvent. Relative energies and geometric parameters of representative optimized geometries are provided in the Supplementary Information (Figures S1-S3, Tables S1-S3). Fractional occupation number Hartree–Fock (FON–HF) was used to determine initial guess orbitals for the α-CASSCF calculations.\textsuperscript{72-75} In the FON-HF implementation, energy levels were Gaussian-broadened using temperatures of 0.2 and 0.3 a.u. for gas-phase and solution-phase systems, respectively.

Critical points were obtained using the DL-FIND geometry optimization library.\textsuperscript{76} Electronic structure calculations were performed with TeraChem,\textsuperscript{77-81} exploiting graphic processing units.

**Solvated initial conditions.** A Boltzmann sampling procedure provided initial conditions (ICs) for the solvated chromophore. Using PACKMOL,\textsuperscript{78} HBDI\textsuperscript{-} was immersed in a solvent sphere comprised of 2000 water molecules and a sodium counterion. The radius of the solvation sphere was 24.3 Å.
consistent with the density of liquid water (0.997 g/mL). Classical dynamics were run with OpenMM, and a restraining spherical force of 10 kcal/mol/Å² was applied to the water sphere during the dynamics. The solvent was equilibrated by performing a 5 ns NVT classical dynamics simulation, using a Langevin thermostat at 300K, with the geometry of the chromophore fixed. Water molecules were treated with the flexible SPC/Fw model, and the sodium ion was modelled using van der Waals parameters previously determined for use in SPC/E water. Atomic charges for the chromophore were fit using the restrained electrostatic potential (RESP) procedure with HF/6-31G*, based on the B3LYP-D3/6-31G** optimized geometry of the chromophore in the gas-phase. Van der Waals parameters for the chromophore were taken from the general Amber force field (GAFF). Other force field parameters (such as equilibrium bond lengths/angles and associated force constants) are irrelevant because the geometry of the chromophore was kept fixed during the MM simulations used to relax the solvent environment.

Following the initial solvent equilibration, the geometric constraints on the chromophore were lifted and ground-state QM/MM molecular dynamics (QM/MM-MD) within the NVT ensemble were run for 10 ps to obtain ICs for the nonadiabatic dynamics simulations. The chromophore was treated at the QM level (described above), whereas the waters and sodium ion were treated at the MM level. A D3 dispersion correction was included for the QM region, using parameters corresponding to Hartree–Fock. Spherical boundary conditions were again employed to avoid evaporation of water molecules. QM/MM-MD simulations were performed with a Langevin thermostat at 300K, using the time-reversible Niklasson integrator and a time step of 0.5 fs. After an equilibration period of 3 ps, a total of 500 temporally-equidistant samples were extracted from the remaining 7 ps of QM/MM-MD to generate an absorption spectrum. The stick spectra were convolved with a Gaussian lineshape function, using a full width at half maximum (FWHM) of 0.40 eV based on solution-phase experiments.

Nonadiabatic dynamics. A total of 50 ICs were extracted from the 500 QM/MM-MD samples (every tenth IC) and used to initiate AIMS simulations within the independent first generation approximation. Using this sampling procedure, we minimize the correlation between ICs. Moreover, this sampling procedure enables us to partition ICs based on their location in the aqueous HBDI absorption spectrum, which is necessary to investigate the wavelength dependence of excited-state deactivation. ICs are classified as either red-shifted or blue-shifted based on their excitation energy relative to the computed absorption maximum (computed as 2.84 eV, compared to the measured value of 2.91 eV). When comparing our simulations to experiment, it should be noted that the wavelength
of the pump used in previous time-resolved spectroscopy experiments was on the blue-edge of the spectrum (400 nm, i.e., 0.2 eV blue-shifted from the absorption maximum). The AIMS simulations were continued for 5 ps or until at least 99% of the excited-state population for a given initial trajectory basis function (TBF) was transferred to the ground state. If a TBF on $S_0$ went at least 5 fs without coupling with another TBF (as measured by the TBF overlap, with a threshold of 0.6), the TBF on $S_0$ was removed from the set of coupled TBFs and run independently for the remainder of the simulation. The threshold for spawning (based on the dot product of the nonadiabatic coupling vector and the velocity) was defined as 0.005 a.u. The spherical boundary conditions applied for QM/MM-MD were employed during the AIMS dynamics as well. Error bars for exponential fitting of the $S_1$ population were obtained using a bootstrapping method that included 1000 bootstrapping samples. To gauge the branching ratio of photoproducts from AIMS dynamics, each spawned TBF on $S_0$ was propagated independently for an additional 500 fs. By the end of the extended ground-state dynamics, the TBFs were classified based on the $\phi_I$ and $\phi_P$ values of their $S_0$ nuclear configuration. If the magnitude of either dihedral was above 120°, the product was classified as isomerized with respect to that angle. The resulting photoproducts were used to quantify photoisomerization quantum yield. Geometric analysis of AIMS trajectories was aided by MDTraj.

**Umbrella sampling.** To provide insight regarding the free energy landscape and entropic effects of HBDF in water, QM/MM umbrella sampling at 300K was performed on $S_1$. A two-dimensional mapping was performed along the $\phi_I$ and $\phi_P$ twisting coordinates (ranging from -120° to 120°, using 10° intervals for a total of 625 sampling windows). To compensate for incomplete sampling of solvent configurations, we augmented the data to incorporate the symmetry with respect to twisting of the $\phi_I$ and $\phi_P$ dihedral angles (Figure S4). Specifically, each data point (i.e., energy, oscillator strength) for an observed geometry contributed to two data points: one for the observed geometry and one for its enantiomer, where $\phi_{\text{enantiomer}} = -\phi_{\text{observed}}$ for both $\phi_I$ and $\phi_P$. Convergence analysis of the two-dimensional free energy surface indicated that an equilibration time of 1.5 ps for each window provided a sufficient compromise between sampling and computational cost (Figure S5). After the equilibration, 2 ps of data was collected for each window to generate the potential of mean force (PMF). In total, over 2 ns of excited-state QM/MM-MD simulations were run to obtain the free energy surface. A harmonic biasing potential with a force constant of 100 kcal/mol/Å² was applied to both dihedrals. Contour plots were generated using a cubic interpolation between points on the free energy grid obtained from unbiasing a set of biased sampling windows via the weighted histogram analysis method.
Time-resolved fluorescence. Two-dimensional fluorescence spectra of HBDI\(^{-}\) were generated based on the AIMS simulations. Spectra were obtained as:

\[
F(t,\nu) \propto \sum_{I=S_1}^{N_{TBF}} n_I(t) \left| \mu_{S_0/S_1}(\vec{R}_I(t)) \right|^2 \left( \frac{\Delta E_{S_0/S_1}(\vec{R}_I(t))}{h} \right)^3 \delta \left( \Delta E_{S_0/S_1}(\vec{R}_I(t)) - \hbar\nu \right) \tag{1}
\]

where the sum is restricted to TBFs that are propagating on S\(_1\), \(h\) is Planck’s constant, \(n_I(t)\) is the \(S_1\) population of the \(I\)th TBF, \(\mu_{S_0/S_1}(R)\) is the transition dipole moment between \(S_0\) and \(S_1\) at molecular geometry \(R\), \(\Delta E_{S_0/S_1}(R)\) represents the \(S_0/S_1\) energy gap, and \(\vec{R}_I(t)\) is the position centroid of the \(I\)th TBF at time \(t\).\(^{98}\) The time-resolved fluorescence spectra were convolved with a Gaussian having FWHM of 0.15 eV in the energy domain (\(\nu\)) and 100 fs in the temporal domain (\(t\)) to simulate the instrument response function of previous\(^{27,\,99}\) fluorescence up-conversion experiments. For each two-dimensional fluorescence spectrum, the energy was shifted by 0.13 eV to align the fluorescence intensity maxima obtained from theory and experiment (\textit{vide infra}).\(^{31}\) Once shifted, the decay signal corresponding to an intensity slice at 500 nm was extracted and analyzed to elucidate the fluorescence behavior of HBDI\(^{-}\) over the course of the dynamics.

Results and Discussion

To begin describing the effects of solvation, we observe how the absorption spectrum of HBDI\(^{-}\) changes when the molecule is immersed in water. Figure 3a compares the simulated and experimental absorption spectra of HBDI\(^{-}\) in vacuum\(^{24}\) and aqueous solution. Solvation of HBDI\(^{-}\) blue-shifts the absorption maximum (0.41 eV), as is also observed in experiment (0.32 eV).\(^{31,\,91,\,100}\) Radial distribution functions (RDFs), based on the ground-state QM/MM-MD, between water oxygen atoms and the heteroatoms in HBDI\(^{-}\) provide a description of local solvation. As shown in Figure 3b, the RDF for the O\(_P\) atom indicates that it is more tightly coordinated to water than the O\(_I\) atom. This is consistent with the enhanced charge localization on O\(_P\), relative to O\(_I\), in the ground state (see Figure S6 for fragment-accumulated Mulliken charges).
Pinpointing the origin of this blue shift enables a more in-depth understanding of the changes that take place once the chromophore is solvated. To isolate the geometric and electronic perturbations induced by explicit solvation, we observe the shift of the absorption spectrum based on whether HBDI⁻ is isolated or surrounded by water. In particular, a comparison of the QM/MM-MD absorption spectrum of aqueous HBDI⁻ to that generated by the same HBDI⁻ configurations but in isolation allows us to directly gauge the electronic effects of solvation. Similarly, the spectral shift obtained by comparing the absorption spectrum of the solvent-removed configurations to that from a gas-phase QM/MM-MD sampling (Figure S7) enables us to isolate geometric effects of the solvent. As summarized in Table 1, the electronic properties of water are primarily responsible for blue-shifting the absorption maximum (ΔE_{abs}) of HBDI⁻ upon solvation (0.19 eV), whereas the geometric changes have a minimal effect (0.03 eV).
Table 1: Change in the absorption maximum of HBDI\textsuperscript{-} as a function of the chromophore’s environment. The calculated $\Delta E_{\text{abs}}$ from ground-state QM/MM-MD sampling, either in the gas-phase or in water, is compared to the $\Delta E_{\text{abs}}$ calculated for configurations of aqueous HBDI\textsuperscript{-} following the removal of the solvent. The same solvent-removed samples are used for comparison to gas-phase and water QM/MM-MD samples. However, since $\alpha$ is a scaling parameter, choosing an $\alpha$ value consistent with the QM/MM-MD sampling procedure is necessary to avoid spurious energy shifts upon comparison.

|            | $\Delta E_{\text{abs}}$ maximum: QM/MM-MD (eV) | $\Delta E_{\text{abs}}$ maximum: Solvent removed (eV) | $\Delta \Delta E_{\text{abs}}$ maximum:       |
|------------|-----------------------------------------------|-----------------------------------------------------|-----------------------------------------------|
| Gas ($\alpha=0.64$) | 2.56 $\pm$ 0.07                              | 2.53 $\pm$ 0.07                                      | 0.03                                          |
| Water ($\alpha=0.67$) | 2.84 $\pm$ 0.09                              | 2.65 $\pm$ 0.08                                      | 0.19                                          |

To understand the limited impact of geometric perturbations, we analyze the initial distribution of key geometric parameters to observe how the sampled configuration space changes upon solvation. Figure 4a shows the bond length distributions for the bridge bonds, $r_I$ and $r_P$, from the ground-state QM/MM-MD. In water, the bridge bonds of HBDI\textsuperscript{-} sample a narrower region of configuration space when compared to vacuum. This difference is related to whether a quantum or classical sampling technique is used, as indicated by the similar configuration space sampled by gas-phase and solution-phase HBDI\textsuperscript{-} using QM/MM-MD sampling in both instances (Figure S7). Distributions for the solvated ICs used for the AIMS dynamics are shown in Figure S8. The average $r_I$-to-$r_P$ bond length ratio in water is consistent with a stabilization of the phenolate resonance structure, in line with the RDFs in Figure 3b. Therefore, we verify that the phenolate resonance structure of HBDI\textsuperscript{-} is favored throughout the ground-state dynamics in water, as postulated by calculations in a polarizable continuum water model.\textsuperscript{39} While the phenolate and quinoid resonance structures contribute almost equally in gas-phase,\textsuperscript{24, 65} the decrease in the $r_I$-to-$r_P$ ratio upon solvation indicates a preference toward the phenolate resonance structure in solution. Interestingly, the distributions along the $\phi_I$ and $\phi_P$ dihedrals (Figure 4b) remain almost identical between systems. Both systems have broad dihedral angle distributions, with values ranging from approximately -30° to +30°, centered about zero degrees. Disrotatory motion, i.e., the concerted rotation of $\phi_I$ and $\phi_P$ in opposite directions, conserves angular momentum in both ground-state systems. The striking similarity between these angular distributions indicates that the presence of mobile water molecules does not restrict the chromophore’s motion along its twisting dihedrals on the ground state.
Electronic effects of equilibrium solvation. Before investigating the photodynamics of HBDI• out of the FC region, we start by considering the excited-state landscape in the equilibrium solvation regime. Although the difference in time scales of HBDI• intramolecular vibrations and solvent relaxation (e.g. time constants of 0.16 and 1.2 ps were reported for solvent relaxation following photoexcitation of a coumarin dye) means that an equilibrium picture is unlikely to pertain to the dynamics induced by photoexcitation, an excited-state PMF identifies the energetically favored configurations of aqueous HBDI• following excitation. In other words, the PMF indicates the stability of twisted HBDI• structures that are surrounded by (partially or fully) relaxed solvent configurations. Such insight is useful for categorizing stable excited-state intermediates and interpreting the photodynamics of aqueous HBDI•.

From two-dimensional umbrella sampling on $S_1$, along the bridge dihedrals, Figure 5 shows that single-bond twisting about either of the bridge bonds is favored. The resulting PMF highlights four energetic wells located near 90° I- and P-twisted structures on $S_1$. Interestingly, the excited-state dynamics do not follow the minimum energy path to reach these energetic wells (vide infra). Instead, the chromophore reaches highly-twisted configurations by following either the I- or P-channels illustrated in Figure 5.

To separate out the electronic effects of equilibrium solvation, Figure 6 displays the average $S_0/S_1$ energy gap across all windows in solvent and solvent-stripped configurations. In water (Figure 6a),

Figure 4: Geometric characterization of ICs using the ground-state distributions of (a) carbon-carbon methine bridge bonds adjacent to the imidazolinone ($r_I$) and phenolate ($r_P$) rings and (b) dihedral angles associated with imidazolinone ($\phi_I$) and phenolate ($\phi_P$) twisting. Red indicates data for ICs sampled from gas-phase harmonic Wigner distribution at 300K. Blue indicates data for ICs sampled from aqueous QM/MM-MD simulations at 300K (via a Boltzmann sampling procedure).
the observed minima along $\phi_I$ lie close to the CI seam, whereas rotation along $\phi_P$ leads to minima on $S_1$ that are $\sim 1$ eV removed from $S_0$. This suggests that the I-twisted CI seam is more peaked than the P-twisted CI seam. The absence of electronic degeneracy associated with P-twisting hints that energetically and/or entropically unfavorable geometric changes are required to reach degeneracy along this pathway. Figure 6b shows that I-twisted intermediates have lower $S_0/S_1$ energy gaps than those of P-twisted intermediates in the gas-phase, as observed in solution. This energetic asymmetry originates from the differences in electron affinity between the I- and P-rings.\(^{24}\) However, to conclude whether or not water stabilizes one twisted structure over the other, it is necessary to look at the energetic differences caused by solvation. The difference between water and gas-phase energy gaps (Figure S9) reveals that water introduces non-selective gap reduction ($\sim 0.3$ eV) for highly-twisted ($\sim 90^\circ$) structures — the I- and P-twisted energy gaps decrease by approximately the same amount upon solvation. Yet, as previously discussed for Figure 6a, the consequences of this gap reduction are significant. This equilibrium solvation picture suggests that (i) aqueous HBDI$^-$ proceeds along one-bond-flip-dominated coordinates toward the twisted CI seam, as in gas-phase,\(^{24}\) (ii) the I-twisted CI seam is more accessible than its P-twisted counterpart, and (iii) solvation promotes internal conversion through the I-twisted CI seam, similar to its neutral counterpart.$^{52-53}$

Figure 5: Relative free energy profile on $S_1$ obtained from two-dimensional umbrella sampling along $\phi_I$ and $\phi_P$. The zero point (marked with an “X”) corresponds to the planar structure. The observed deactivation pathways through I-twisting (yellow) and P-twisting (red) are depicted schematically.
**Figure 6:** Contour plot of the average $S_0/S_1$ energy gap with respect to $\phi_I$ and $\phi_D$ dihedral angles (a) in water and (b) in gas-phase. Gas-phase energies correspond to solvated structures (5000 points total, 8 per window) from (a) after removing the solvent around the chromophore. The average dihedral angles and energy gaps were plotted for each $10^\circ$ window from the umbrella sampling, and cubic interpolation was performed between these points. The zero point (marked with an “X”) corresponds to the planar structure. All energies were computed with $\alpha(0.67)$-SA3-CASSCF(4,3)/6-31G*.

**Nonadiabatic dynamics of HBDI$^-$ in water.** Shifting into the non-equilibrium regime (i.e., the solvent does not have enough time to relax around the chromophore), we analyze the behavior of aqueous HBDI$^-$ following photoexcitation. As shown by the $S_1$ population decay in Figure 7, over half of the excited-state population is transferred to the ground state within the first 1 ps. Such a rapid decay is consistent with the ultrafast deactivation reported in experiments.$^{27, 31, 43}$ Yet, despite the sub-ps deactivation, a fraction of the population remains on the excited state for $>5$ ps. This biphasic pattern inspired a delayed bi-exponential fit to the decay profile:

$$P(t) = A \exp\left(-\frac{t - \tau_0}{\tau_1}\right) + (1 - A) \exp\left(-\frac{t - \tau_0}{\tau_2}\right)$$

where $t$ represents time, $A$ is a weighting coefficient, $\tau_0$ represents the time at which population transfer begins (lagtime), and $\tau_1$ and $\tau_2$ are characteristic lifetimes associated with the decay of the $S_1$ population over time, $P(t)$. Constants obtained from this fitting procedure, through the use of bootstrapping, are: $A = 0.50 \pm 0.19$, $\tau_0 = 204 \pm 43$ fs, $\tau_1 = 609 \pm 181$ fs, and $\tau_2 = 3.1 \pm 1.2$ ps.
Figure 7: Ground- and excited-state populations during AIMS simulations. Fifty ICs were used in the explicitly-solvated system. The average state population is represented with solid lines. The vertical, gray dotted line represents the time at which population transfer begins. The black dotted line is the delayed bi-exponential fit to the $S_1$ population decay. Error bars are illustrated using transparent lines.

Viewing the dynamics within defined time intervals reveals novel information regarding the time scales of each deactivation pathway available to aqueous HBDI — two distinct sub-populations exist with different excited-state lifetimes (Figure 8). In accordance with the decreased electronic density along the methine bridge upon excitation (as illustrated in Figure 2) facile rotation along $\phi_I$ and $\phi_P$ is permitted on the excited state. Monitoring the $S_1$ population over time, with respect to the $\phi_I$ and $\phi_P$ dihedrals, illustrates (i) the accessibility and (ii) the population transfer efficiency of each internal conversion avenue taken by the ICs. During the first 100 fs, the $S_1$ wavepacket remains localized near the FC region. Rather than following the minimum energy path along one-bond-flip coordinates (Figure 5), the disrotatory motion exhibited during the ground-state dynamics (Figure 4b) is preserved. This indicates a non-equilibrium solvation regime where the response of the surrounding waters is too slow to impede disrotatory motion out of the FC region, which is otherwise associated with a small free energy barrier (less than 0.2 eV). Once the free energy along the disrotatory path becomes too large to overcome, the molecule reaches a fork in the road. Here, the wavepacket bifurcates via a process that involves a one-bond-flip-like rotation along either $\phi_I$ or $\phi_P$ — with one path leading to highly I-twisted structures (I-channel) and the other path leading to highly P-twisted structures (P-channel). Because this drastic dihedral torsion is preceded by disrotatory motion out of the FC region, the internal conversion process resembles an aborted hula-twist mechanism. The branching associated with this mechanism becomes apparent by 500 fs as exploration into the I- and P-channels enables population transfer to the
ground state. By 1 ps, motion along the torsional angles is more prominent as the I- and P-channels arise as the exclusive avenues for population transfer. Essentially no TBFs occupy near-planar configurations after 1.75 ps of dynamics, as four distinct regions in configuration space emerge as dihedral basins. These basins coincide with the energetic wells identified in Figure 5. Interestingly, 250 fs later, no evidence of the I-twisted species is present while the P-twisted species remains on S1 for several picoseconds. The disappearance of the I-twisted species on S1 is explained by an ultrafast population transfer to the ground state, permitted by the electronic degeneracy highlighted in Figure 6a. For this part of the wavepacket, twisting along $\phi_I$ is enough to induce population transfer to the ground state. The persistence of the P-twisted species for several picoseconds on the excited state signifies that dihedral torsion around $\phi_P$ alone is not sufficient to bring the S0 and S1 electronic states close enough together to facilitate efficient population transfer. Rather, additional geometric requirements are necessary for the P-species to participate in internal conversion.

![Figure 8: Time evolution of the reduced S1 density along the bridge dihedrals during the first 2.5 ps of solution-phase AIMS dynamics.](image)

The percentages displayed in each panel represent the total S1 population at the end of the specified time window, relative to their maximum at time=0. Gray dots indicate the dihedral angles of the fifty initial conditions.

The time scale differences established in Figure 8 were refined using the S1 population profiles for the I- and P-deactivating species (Figure 9). In the case of I-twisted deactivation, the entire excited-state population is transferred to the ground state within ~1.5 ps, with decay initiating at ~175 fs. This rapid initiation and conclusion of population transfer for the I-species is due to an easily accessible I-twisted CI seam that efficiently transfers population to the ground state, in accordance with Figure 6. On
the other hand, the P-deactivation mechanism is associated with a species that persists on S₁ for ~540 fs before population transfer begins. Furthermore, this mode of population transfer is much slower and less efficient, as ~10% of the population remains on the excited state after 5 ps. A similar analysis compares the deactivation of red-shifted and blue-shifted ICs (Figure S10, Table S4).

![Figure 9: Excited-state population of HBDI in water for I- and P-deactivating ICs. Solid lines represent the average populations for each species. Transparent lines represent the error bars obtained from bootstrapping analysis. We indicate the delay time before population transfer \( \left( \tau_{I} / \tau_{P} \right) \) for each set with vertical dotted lines, emphasizing that deactivation is faster for I-twisting compared to P-twisting.](image)

After identifying the characteristic twisting pathways, the dynamics out of the FC region were probed using various electronic and geometric parameters (Figure 10, Figure S11). Here, the parent TBFs are partitioned into I- and P-twisted species based on whether the I- or P-channel was taken, respectively. Our results corroborate that the dominant mode of deactivation (66%) is through torsion along \( \phi_{I} \), in agreement with nonadiabatic dynamics simulations at the XMS-CASPT2⁴¹ and spin-flip time-dependent density functional theory⁴² levels. The violin plots in Figure 10 present the distribution of key electronic and geometric observables sampled across all parent TBFs of each species during the first 500 fs. Characteristic activity along the \( \phi_{I} \) and \( \phi_{P} \) twisting dihedrals is observed within the first 500 fs. As expected, the I-species samples along \( \phi_{I} \) more aggressively than it does along \( \phi_{P} \), and the opposite behavior is true for the P-species. Worth noting, however, is the time scale at which both species reach ~90° rotations of their respective dihedrals. Between 200 and 400 fs, both species begin to explore and populate these highly twisted configurations. As in the gas-phase,³⁵ we observe that twisting along \( \phi_{I} \) facilitates CT from the I-ring to the P-ring while twisting along \( \phi_{P} \) facilitates CT from the P-ring to the I-
ring. Furthermore, the molecule’s rapid approach toward highly I- and P-twisted structures indicates that access to their respective TICT states are downhill in solution.

In the gas-phase, bond length alternation and bridge pyramidalization are the main coordinates used by HBDI\(^-\) to reach the CI seam.\(^{24}\) As shown in Figure 10 (300 – 500 fs), displacement along the pyramidalization coordinate is attenuated for the I- and P-twisted species in solution relative to gas-phase (|θ\(\text{pyr}\)| of 30° and 48° for I- and P-twisted MECIs,\(^{24}\) respectively). Because solvation essentially closes the energy gap of the I-twisted species but not completely for the P-twisted, the bridge pyramidalization requirement for reaching the I-twisted CI seam becomes almost negligible. Compared to species along the I-channel, larger-amplitude displacements of the pyramidalization coordinate are required to trigger population transfer along the P-channel. This suggests that the path to the I-twisted CI seam is essentially energetically downhill, while this process is energetically uphill for the P-twisted case.

![Figure 10](image_url)

**Figure 10:** Violin plots that monitor density distributions of geometric and electronic observables out of the FC region for I-deactivating (red) and P-deactivating species (blue). Shown data monitors observables for the parent TBFs from each IC during the first 500 fs.

In addition to monitoring the geometric and electronic changes that occur during the first few hundred femtoseconds following photoexcitation, the response of the solvation environment during the first 1 ps was investigated. Figure 11 shows the evolution of the first solvation peak for the parts of the wavepacket following the I- and P-twisted pathways. In the case of the I-deactivating species in Figure
11a, there are no significant changes to the RDF profiles of O₁ or N₁ during the first 1 ps. However, the first solvation peak of Oᵢ experiences a slight decrease over time. This contrasts the evolution of RDFs for the same atoms in the P-deactivating species in Figure 11b. Over the first 1 ps, the maximum of the first solvation peak for Oᵢ and O₁ gradually decreases and increases, respectively, toward the RDFs associated with equilibrated ϕᵢ-twisted structures. This trend is consistent with the TICT behavior identified in Figure 10. Upon torsion along ϕᵢ, CT occurs from the P-ring to the I-ring. Consequently, a P-twisted intermediate will have (i) a more negative I-ring and (ii) a more positive P-ring than its pre-twisted counterpart. Thus, it is expected that the surrounding solvent would rearrange to compensate for this CT process by shifting increased solvent density from near the Oᵢ to near the O₁. Interestingly, the opposite trend is absent for the I-species. We attribute this difference to the disparate deactivation time scales along ϕᵢ and ϕᵦ that are presented in Figure 9. Because deactivation through the I-channel occurs so rapidly, water does not have time to relax fully around I-twisted structures before internal conversion. Such behavior explains the absence of large shifts of excited-state RDF peaks in Figure 11a. On the other hand, deactivation through the P-channel is more gradual than decay through the I-channel, due to the barrier to reach the P-twisted CI seam from the corresponding TICT intermediate. Consequently, the solvent has time to reorient itself around P-twisted structures on the excited state. As a result, we observe gradual shifts of excited-state RDF peaks in Figure 11b.

**Figure 11:** Radial distribution function (RDF) for HBDI heteroatoms with water oxygen atoms during the first 1 ps of AIMS dynamics on S₁. Each RDF shown for (a) I-deactivating species and (b) P-deactivating species begins and ends within the specified time window. Each figure focuses on the first solvation shell of the specified atom. The ϕᵢ- and ϕᵦ-twisted RDFs are obtained from umbrella sampling windows on Sᵢ, with the specified angle restrained to ±90° while the other dihedral is restrained to zero degrees. The purple line represents the average of the 90°- and -90°-twisted RDFs on Sᵢ.
A geometric analysis at the points of population transfer elucidates the difference between the deactivation profiles of the I- and P-species. For I-deactivation, the spawning events that lead to population transfer generally occur near geometries with $\phi_I$ angles near 90° and $\phi_P$ angles that lie relatively close to planarity, and vice versa for P-deactivation (Figure 12). Noting the occurrence of spawning events in relation to the key dihedral and pyramidalization angles reveals that, in addition to lasting longer on $S_1$, P-twisted TBFs demand larger pyramidalization angles than I-twisted TBFs to reach their respective CIs. To investigate the implications of this difference, the absolute population transfer was calculated for each spawning event as a function of the pyramidalization and dihedral angle values upon entering the spawning region (Figure 12a). Coupling these observations with AIMS data supports our claim that TBFs that reach highly I-twisted structures rapidly approach electronic degeneracy and transfer their $S_1$ population to the ground state. In contrast, the TBFs that proceed through P-twisting transfer their population less effectively and therefore persist on $S_1$ for several picoseconds. Movie S1 provides a time lapse of these population and angle parameters to clarify this point. For a direct comparison of the spawned geometries associated with each pathway, the pyramidalization angle was measured for each spawned geometry of I-deactivating (Figure 12b) and P-deactivating (Figure 12c) TBFs. From these distributions, it is clear that pyramidalization requirements for P-deactivating TBFs are more demanding (peaked at $\sim$30°) than the requirements for I-deactivating TBFs (peaked at $\sim$10°). The larger pyramidalization angles needed to reach the P-twisted CI gate access to the P-twisted region of the CI seam. We conclude that limited access to an uphill CI seam is responsible for the lingering $S_1$ population of the P-species.

In addition to being more accessible than the P-twisted CI seam, the I-twisted CI seam appears to be more efficient at transferring population to the ground state. On average, population transfer events mediated by the I- and P-channels transfer 56 ± 32% and 35 ± 26% of the incoming $S_1$ population to $S_0$ per spawning event, respectively (Figure S12). This is expected based on the more direct approach toward the I-twisted CI seam, than for its P-twisted counterpart, as supported by Figures 5 and 6. The uphill pyramidalization requirements and lack of efficiency explain the more gradual deactivation of the P-twisted species, when compared to the I-twisted species.

Relative to gas-phase HBDI$^-$, aqueous HBDI$^-$ generally demands less pyramidalization to reach I- and P-twisted CI seams (Figure 12b-c). This difference between systems becomes most apparent when comparing internal conversion tendencies associated with the I-channel. In gas-phase, I-twisted
configurations transfer population to $S_0$ while sampling along a larger range of $\phi_P$ values than in water. Based on our analysis regarding pyramidalization gating, we conclude that this behavior is due to the topology differences of the I-twisted CI seam across systems. Because solvation reduces the $S_0/S_1$ energy gap for I-twisted structures, aqueous HBDI$^-$ has smaller pyramidalization requirements to reach the I-twisted CI seam than does gas-phase HBDI$^-$. Thus, the solvated chromophore spends little time sampling $\phi_P$ as it undergoes internal conversion through a near-barrierless process. The I-twisted spawning geometries of aqueous HBDI$^-$, which are associated with ultrafast spawning tendencies, are consistent with rapid accessibility out of the FC region via an aborted hula-twist mechanism, as presented in Figure 8. Due to the larger pyramidalization angles required to reach the I-twisted CI seam in gas-phase, more extensive sampling along $\phi_P$ occurs for gas-phase HBDI$^-$ before the entire $S_1$ population is transferred to $S_0$. This logic extends to the comparison between I- and P-twisted spawning occurrences. Because reaching the P-twisted CI seam from a P-twisted TICT intermediate is an energetically uphill process, P-twisted intermediates significantly sample configurations above 90°. The enhanced sampling along this coordinate may explain the higher proportion of spawning geometries skewed toward the $E$-isomer, relative to this proportion for $\phi_I$ from I-twisted spawns (Figure S13).

![Figure 12:](image)

(a) Absolute population transfer and pyramidalization angle associated with the bridge carbon atom for spawned geometries of aqueous HBDI$^-$ from parent TBFs. Absolute population transfer is defined by the fraction of original $S_1$ population transferred to $S_0$. Plotted values of $\phi_I$, $\phi_P$, and $\theta_{\text{pyr}}$ characterize geometries that initiate population transfer. A histogram (using 5° bins) of pyramidalization angle magnitudes is provided for spawned TBFs that transfer population through torsion dominated by (b) $\phi_I$ or (c) $\phi_P$ dihedrals. Pyramidalization angles measured for I- and P-twisted MECIs in gas-phase$^{24}$ are represented with vertical lines in (b) and (c).
Based on vibrational signatures obtained from FSRS experiments, Taylor et al. proposed that the formation of a CS species and the transition from this CS state to a TICT state were responsible for measured short (~350 fs) and long (~2-3 ps) time components, respectively.\textsuperscript{31} Our data suggests a different origin of these time constants. As supported by the substantial torsional displacements sampled within the first ~200 fs (Figure 10), the formation of TICT states is essentially barrierless. In other words, we do not see evidence of a non-twisted CS intermediate in water. Rather, we observe two competing processes: (i) an I-twisting process that easily reaches a downhill CI seam and (ii) a P-twisting process that encounters a barrier as it approaches an uphill CI seam. As an alternative, we propose that the different accessibility to the CI seams, from the corresponding TICT states associated with competing I- and P-twisted processes, is responsible for the short (caused by mainly I-species) and long (caused by mainly P-species) time components reported from the FSRS experiments. However, to rigorously establish this interpretation of the FSRS fingerprints requires calculation of the vibrational signatures of I- and P-twisted structures and remain a task for future work.

**Photoproducts.** To evaluate the branching ratio of HBDI\textsuperscript{−} upon deactivation to the ground state, dynamics on S\textsubscript{0} were continued for an additional 500 fs (Figure S14, Table S5). Classifying the resulting products provides an estimate for the branching ratio with respect to Z/E isomerization, Z/Z isomerization, and ground-state recovery to the Z-isomer. Most of the population (87\%) returns to the Z-isomer (Figure 13), in line with the low Z/E isomerization quantum yield reported by Vengris, et al.\textsuperscript{43} Despite I-torsion being the dominant deactivation pathway, associated CIs are highly inefficient at producing E photoproducts. Thus, HBDI\textsuperscript{−} is much less efficient at generating E photoproducts in solution (~13\% yield) than in gas-phase (~40\% yield).\textsuperscript{24} Only 20\% of the I-twisted spawns undergo Z/E isomerization. P-twisted CIs are more efficient (37\%) at producing P-flipped products; however, the symmetry of the P-ring regenerates the Z-isomer whether or not the P-ring isomerizes. Verifying the Z/Z isomerization quantum yield experimentally would require asymmetric substitution on the P-ring. The respective isomerization quantum yields are likely linked to the barrier (or lack thereof) between the TICT state and CI seam associated with each pathway. Because the I-twisted CI seam is reached without significant pyramidalization, I-twisted structures readily proceed through the CI seam as soon as twisting along \(\phi_I\) nears 90° (Figure S13a). On the other hand, the P-twisted CI seam is not reached so easily. As a consequence, P-twisted intermediates sample along \(\phi_P\) more robustly to reach significantly pyramidalized structures. This results in a majority of spawns with \(\phi_P\) greater than 90° (Figure S13b) and likely contributes to the higher quantum yield for Z/Z isomerization than that of E/Z isomerization.
Figure 13: Photoproducts of HBDI\(^{-}\) in water for spawned TBFs, following 500 fs of QM/MM-MD on $S_0$. A total of 264 spawned TBFs were analyzed. $\phi_I$ and $\phi_P$ branching percentages represent the fraction of TBFs that participated in each twisting pathway. Bar graphs illustrate the percentage of each product formed by the spawned TBFs for their respective pathway. The Z-1 product represents a Z-isomer of HBDI that is not the result of ZZ isomerization, whereas the Z-2 product corresponds to a Z-isomer of HBDI following ZZ isomerization.

**Time-resolved fluorescence.** Solvated HBDI\(^{-}\) has been the subject of many time-resolved fluorescence experiments.\(^{27,37,102-104}\) To link our simulation data to experimental findings, time-resolved fluorescence spectra were computed over the course of the AIMS simulations (Figure 14). A line out at 500 nm (based on experiment\(^{99}\)) is extracted for various sets of TBFs. Across all TBFs (Figure 14a), the two-dimensional fluorescence spectrum experiences a significant red-shift spectral evolution after a few hundred femtoseconds. The red-shift and the reduction in fluorescence signal are attributed to a decrease in the $S_0/S_1$ energy gap and emission strength, respectively, as the chromophore begins twisting its bridge dihedrals upon departure from the FC region (Table S3, Figure S16). Since twisting along $\phi_I$ and $\phi_P$ is correlated with a decrease in oscillator strength, we may expect the steeper (compared to XMS-PT2) $\alpha$-CASSCF gradients beyond 30° torsion about either the I or P bonds (Figure S1) to artificially accelerate fluorescence quenching in our AIMS simulations. The fluorescence spectrum was decomposed to obtain the contributions from the I- and P-deactivating species (Figure 14b, Figure 14c). By comparing the two-dimensional fluorescence spectra, it becomes obvious that the P-twisted channel significantly prolongs the fluorescence intensity of aqueous HBDI\(^{-}\) when compared to the signal from the I-twisted channel. From the time-resolved fluorescence spectrum for all TBFs at 500 nm (Figure 14d), AIMS simulations predict a fluorescence signal that loses roughly 50% of its intensity by 500 fs. Despite this rapid quenching, fluorescence signatures are observed beyond 1.5 ps. An I- and P-pathway decomposition of this signal shows that I-twisting species quench their fluorescence rapidly (within a
picosecond). Yet, the P-twisting species has a more pronounced fluorescence signal that survives until ~1.6 ps. Therefore, the long-lived signal in the time-resolved spectrum, across all TBFs, is attributed to the species that undergoes internal conversion through the P-channel. We attribute the faster fluorescence decay of our simulations, relative to experiment, to the slightly steeper α-CASSCF gradients for partially twisted geometries, when compared to XMS-CASPT2. The underestimated S0/S1 energy gap for I- and P-twisted structures at the α-CASSCF level (Figure S1) likely red-shifts the fluorescence signal of I- and P-twisted structures more than expected. However, this issue likely affects the fluorescence signal of P-twisted ICs, which remain on S1 for several picoseconds, more than the signal of I-twisted ICs. Another possible contributor to the faster-than-expected simulated fluorescence signal is a higher ratio of P-to-I species in experiment than sampled for our AIMS simulations, which would increase the intensity of the fluorescence tail. Despite these discrepancies, our simulations provide evidence that the deactivation pathway taken by the chromophore has a significant impact on the observed time-resolved fluorescence.

Figure 14: Two-dimensional fluorescence spectrum, probed at 500 nm, for (a) all TBFs, (b) I-deactivating TBFs, and (c) P-deactivating TBFs. (d) Time-resolved fluorescence signals of HBDI calculated from AIMS simulations in water. The maximum fluorescence intensities of I- and P-deactivating species are scaled based on the relative proportion of ICs that deactivate through the I- and P-channels. Experimental data, following 400 nm excitation, for HBDI in water was provided by Meech and co-workers.27, 99 To align the experimental and theoretical fluorescence wavelength maxima, a shift of 0.13 eV was applied to the theory spectra (Figure S15). Transparent lines represent the error obtained from bootstrapping analysis.
Conclusions

AIMS dynamics support the presence of two ground-state recovery mechanisms after the photoexcitation of the aqueous HBDI\textsuperscript{−} to S\textsubscript{1}. The dominant mode of deactivation, through I-twisting, proceeds via an ultrafast mechanism and is responsible for the rapidly-decaying signals in time-resolved fluorescence and FSRS experiments. The minor pathway, through P-twisting, remains on S\textsubscript{1} for several picoseconds and its access to the CI seam is gated by pyramidalization of the central methine bridge carbon. Our results provide evidence for the following mechanism: 1) evolution from the FC region via near-barrierless disrotatory motion (along $\phi_I$ and $\phi_P$) that is consistent with ground-state QM/MM-MD simulations, 2) formation of a TICT state through an aborted hula-twist mechanism as either $\phi_I$ or $\phi_P$ approaches 90°, 3) population transfer from S\textsubscript{1} to S\textsubscript{0} as the two states reach electronic degeneracy (through I-twisting for the I-species and through P-twisting along with pyramidalization for the P-species), and 4) re-population of the ground state with low Z/E and Z/Z isomerization quantum yields. This process is summarized in Figure 15. The ultrafast differentiation between I- and P-species indicates that both twisted species are accessible from the FC region with little/no energetic barrier, as supported by Figure 5 and Figure S1. Relative to the uphill approach needed to reach the P-twisted CI seam, the downhill approach to the I-twisted CI seam makes it easier to transfer population to the ground state through the I-channel. However, the P-twisted species has an isomerization quantum yield about two times as large as its I-twisted counterpart.

Our AIMS simulations provide novel mechanistic and electronic insight into the behavior that governs the photodynamics of the GFP chromophore. QM/MM-MD simulations indicate that surrounding HBDI\textsuperscript{−} with mobile solvent molecules has minimal impact on the configuration space sampled by the chromophore in water. Yet, the electronic effects of water blue-shift the absorption spectrum. During gas- and solution-phase dynamics on S\textsubscript{0}, preferential twisting along $\phi_I$ and $\phi_P$ in a disrotatory fashion is observed. The subsequent internal conversion mechanisms predominantly occur through rotation of $\phi_I$ in both systems.\textsuperscript{24} The absence of rigid electronic and steric constraints on the chromophore likely explains the ultrafast deactivation tendencies reported by experiments for both systems.\textsuperscript{21,27,30-31} However, the stabilization of highly-twisted structures and the promotion of electronic degeneracy, especially with respect to $\phi_I$ which leads to a downhill CI seam, are responsible for shorter lifetimes in water when compared to gas-phase. Furthermore, the fitted time constants for the S\textsubscript{1} population decay (609 ± 181 fs and 3.1 ± 1.2 ps) are associated with internal conversion through the I- and P-channels, respectively. Through a direct comparison of AIMS simulations and experimental
fluorescence spectra, we provide mechanistic insight to explain the observed fluorescence profile of HBDI− in water. We conclude that the short experimental time constant (210 fs) fluorescence of HBDI− in water is attributed to departure from the FC region (of both I- and P-deactivating species) and that the longer experimental fluorescence time constant (1.1 ps) corresponds to the longer-lived, P-deactivating species.

**Figure 15:** Summary of the photophysics of the solvated GFP chromophore. Percentages indicate the relative population that proceeds along a given pathway. \( \tau_{CT} \) represents the time range at which TICT occurs, \( \tau_0 \) represents the time at which population transfer begins, and \( \tau_{ex} \) represents the order of the excited-state lifetime. \( \theta_{pyr} \) represents the peak in the pyramidalization angle distribution for spawned geometries. Heavy atoms are colored gray, hydrogen atoms are colored white, and a reference carbon atom is colored purple to distinguish P-ring-flipped and original Z products. Percentages listed in boxes indicate the total percent of spawned TBFs that formed the associated photoproduct.

Although our simulations do not include an explicit protein environment, our findings imply that the mobility and electronic distribution of residues surrounding the chromophore in a protein can tune the chromophore’s behavior on the excited state. From our simulations, we predict that modifications to the protein environment can lead to internal conversion pathways with notably different population transfer rates, fluorescence profiles, and isomerization quantum yields. Using Dronpa2 variants, Romei and co-workers have demonstrated that chromophore modifications can bias the preferred...
directionality of the CT process and modify observables of interest in the protein. Our direct observation of TICT processes in solution provides evidence that steering the chromophore toward a given pathway can significantly alter its properties. To characterize the photochemical processes that drive the dynamics of the chromophore in protein scaffolds, we will extend our simulations and analyses to fluorescent protein systems.

**Author Contributions**

CMJ contributed to data curation, formal analysis, investigation, methodology, visualization and writing (original draft) of the presented work. NHL contributed to data curation, investigation, methodology and project administration. TJM contributed to methodology, project administration, funding acquisition, resources, and supervision. All authors contributed to conceptualization of the project and review and editing of the manuscript.

**Conflicts of Interest**

There are no conflicts to declare.

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Jones, List, Martínez – Photodynamics of solvated HBDI anion – Page 29

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Supporting Information

Resolving the ultrafast dynamics of the anionic green fluorescent protein chromophore in water

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Table of Contents

I. Validation of α-CASSCF ................................................................................................................. 2
II. Critical points of aqueous HBDF using α-CASSCF ................................................................. 5
III. Benchmark of α-CASSCF along an AIMS trajectory ............................................................... 6
IV. Isoenergetic enantiomers of HBDF .......................................................................................... 7
V. Equilibration of free energy during umbrella sampling ............................................................ 8
VI. Fragment-accumulated Mulliken charges across solvated initial conditions ...................... 9
VII. Geometric analysis for gas-phase and solvated initial conditions ........................................ 10
VIII. Geometric analysis for solvated AIMS initial conditions .................................................... 11
IX. Electronic effects of solvation .................................................................................................. 12
X. Excited-state population decay of red-shifted and blue-shifted initial conditions .............. 13
XI. Flapping angle of HBDF during AIMS dynamics ..................................................................... 14
XII. Relative population transfer of HBDF spawning events .................................................... 15
XIII. Dihedral angle distribution for spawned HBDF geometries ................................................. 16
XIV. Photoproducts of HBDF ......................................................................................................... 17
XV. Comparison of experimental and theoretical fluorescence maxima .................................... 18
XVI. Oscillator strength as a function of twisting angles ............................................................... 19
XVII. Geometric parameters of critical points in water ................................................................. 20
XVIII. Comparison of S0/S1 energy gap across multiple complete active space methods .......... 21
XIX. Oscillator strength of geometries used to benchmark α-CASSCF ........................................ 22
XX. Photoproducts of red-shifted and blue-shifted initial conditions ........................................ 23
XXI. Classifications for TBF photoproducts ................................................................................. 24
I. Validation of $\alpha$-CASSCF

Prior to running our nonadiabatic dynamics simulations, we benchmarked the validity of $\alpha$-CASSCF based on reference XMS-CASPT2 calculations. After choosing an $\alpha$ parameter in optimal agreement with the $S_0/S_1$ energy gap of XMS-CASPT2 at the Franck-Condon (FC) point, the qualitative and quantitative aspects of $\alpha$-CASSCF were evaluated. As seen in Figure S1, using an $\alpha$ correction for the solvated system is beneficial for describing the potential energy surface (PES) topology of the ground and excited states. Using $\alpha(0.67)$-SA3-CASSCF(4,3)/6-31G* with a D3 dispersion correction, geometries along characteristic $\phi_I$ and $\phi_P$-twisted dihedrals produce a PES with the same qualitative features as XMS-CASPT2. Most notably, the energetic gradient of SA3-CASSCF(4,3) on $S_1$ appears much steeper (after 30° rotation along $\phi_I$ or $\phi_P$) than that of $\alpha(0.67)$-SA3-CASSCF(4,3) and XMS-CASPT2, which may artificially accelerate excited-state dynamics. Quantitatively, $\alpha$-CASSCF predicts electronic state energies that are much closer to XMS-CASPT2 values than SA3-CASSCF for these geometries. Critical points for the QM/MM system were obtained using the same partitioning scheme described for the QM/MM-MD, treating the chromophore at the $\alpha(0.67)$-SA3-CASSCF(4,3)/6-31G* level. Optimized geometries along $\phi_I$- and $\phi_P$-twisting pathways were reached by coordinate driving along the respective dihedral, keeping the other dihedral angle fixed at zero degrees and allowing all other degrees of freedom to relax. I- and P-twisted minimum energy conical intersections (MECIs) were reached by initiating an MECI search from the 90° I- and P-twisted minima, respectively.

Although MECIs do not necessarily reflect geometries that are encountered during the dynamics, they can provide useful information about notable geometric features of the PES. It is known that HBDI- has two primary MECIs in water: one dominated by torsion along $\phi_I$ and the other dominated by torsion along $\phi_P$.\(^1\) For our explicitly-solvated systems, these conical intersections are encountered with $\alpha$-CASSCF (Figure S2). Associated with these MECIs are characteristic geometric features (Table S1). In the case of the I-twisted MECI (MECI-I), degeneracy between the ground and excited states is achieved by rotation along $\phi_I$ with little pyramidalization of the bridge carbon. Additionally, the bridge bond closer to the I-ring ($r_I$) is considerably longer than the bridge bond closer to the P-ring ($r_P$). For the P-twisted MECI (MECI-P), this relationship between the bridge bond lengths is reversed and the pyramidalization
requirement increases by more than a factor of two. Reaching the MECI-P also requires non-negligible twisting along $\phi_I$, making the structure less dominated by a single rotation than its I-twisted counterpart. Reaching these MECIs through incremental twisting along their respective dihedrals suggests that both MECIs lie below the FC point, consistent with previous claims that solvation permits energetically accessible CIs between the excited state and the ground state.$^2$-$^3$
Figure S1: Relative energy comparison for aqueous HBDI⁻ upon twisting along the $\phi_I$ and $\phi_P$ angles. (a) Energies of HBDI⁻ geometries in explicit water. The planar geometry (labeled as “0”) was optimized on $S_0$ as a reference FC structure. The $\phi_I$- and $\phi_P$-twisted structures were optimized on $S_1$ in succession (using 15° intervals and the previous optimized geometry as a starting point) to avoid large disruptions to the local solvent, keeping the non-twisted dihedral angle fixed at 0°. SA3-CASSCF(4,3)/6-31G* and XMS-CASPT2/6-31G* energies were computed using single point energy calculations on $\alpha(0.67)$-SA3-CASSCF(4,3)/6-31G* optimized geometries. (b) Effect of solvent stabilization for non-equilibrated configurations of aqueous HBDI⁻. Gas-phase energies were derived from single point energy calculations on aqueous HBDI⁻ geometries in (a), after stripping the solvent. Optimized gas-phase energies are from optimized stripped-solvent geometries. Lower markers represent $S_0$ energies and upper markers represent $S_1$ energies.
II. Critical points of aqueous HBDI⁻ using α-CASSCF

Figure S2: Representative geometries of the ground-state ($S_0$) minimum and $S_0$/$S_1$ MECIs of HBDI⁻ in water treated with α(0.67)-SA3-CASSCF(4,3)/6-31G*.
III. Benchmark of $\alpha$-CASSCF along an AIMS trajectory

*Figure S3*: Benchmarked performance of $\alpha$-CASSCF relative to XMS-CASPT2(4,3) for HBDI$^-$ in water. Energy shown corresponds to the $S_0/S_1$ energy gap along an I-deactivating $\alpha$-CASSCF AIMS trajectory. XMS-CASPT2(4,3) energies were computed for each geometry along the AIMS trajectory. The 6-31G* basis set was used for both methods.
IV. Isoenergetic enantiomers of HBDI⁻

Figure S4: Optimized $S_1$ geometries of gas-phase HBDI⁻, fixed at specific dihedral angles, to illustrate isoenergetic enantiomers with respect to the $\phi_I$ and $\phi_P$ twisting dihedral angles. Geometries were optimized to illustrate symmetry with respect to (a) I-twisting, (b) P-twisting, and (c)-(d) simultaneous I- and P-twisting. Geometries were optimized using $\alpha(0.67)$-SA3-CASSCF(4,3)/6-31G*. 
V. Equilibration of free energy during umbrella sampling

Figure S5: (a)-(d) Potential of mean force (PMF) differences between various time periods during the umbrella sampling of aqueous HBDI$^-$ on $S_1$. In each panel, the absolute relative free energy difference of the two specified PMFs was taken to estimate the equilibration time for the solvent and chromophore across all windows. Once these fluctuations dropped below 0.05 eV, the system was considered equilibrated.
VI. Fragment-accumulated Mulliken charges across solvated initial conditions

Figure S6: (a) Labeled atoms of HBDI with positive and negative dihedral rotations illustrated. Average $S_0$ and $S_1$ Mulliken charges localized on (b) atoms and (c) moieties of HBDI in water. Average charges were obtained across 500 initial conditions using the $\alpha(0.67)$-SA3-CASSCF(4,3)/6-31G* level of theory.
VII. Geometric analysis for gas-phase and solvated initial conditions

Figure S7: (a) Distributions of carbon-carbon methine bridge bonds adjacent to the imidazolinone ($r_I$) and phenolate ($r_P$) rings. (b) Distributions of dihedral angles associated with imidazolinone ($\phi_I$) and phenolate ($\phi_P$) twisting. Blue indicates data for Boltzmann-sampled initial conditions from aqueous QM/MM-MD simulations at 300K, using $\alpha(0.67)$-SA3-CASSCF(4,3)/6-31G*. Red indicates data for Boltzmann-sampled initial conditions from a gas-phase QM/MM-MD at 300K, using $\alpha(0.64)$-SA3-CASSCF(4,3)/6-31G*. 
VIII. Geometric analysis for solvated AIMS initial conditions

Figure S8: As in Figure S7, but restricted to initial conditions that were used for the AIMS dynamics in explicit water at 300K.
IX. Electronic effects of solvation

**Figure S9**: Difference between $S_0/S_1$ energy gaps in water (Figure 6a) and in gas-phase (Figure 6b) systems. Gas-phase energies correspond to umbrella-sampled solvated structures after removing the solvent around the chromophore. The average dihedral angles and energy gaps were plotted for each $10^\circ$ window from the umbrella sampling, and cubic interpolation was performed between these points. The zero point (marked with an “X”) corresponds to the planar structure.
X. Excited-state population decay of red-shifted and blue-shifted initial conditions

Based on the ratio of I- and P-twisted pathways taken by red- and blue-shifted ICs (Table S4), the excitation wavelength may impact the subsequent deactivation path. An increased I-twisted/P-twisted ratio for blue-shifted ICs, relative to red-shifted ICs, explains the faster population transfer observed for blue-shifted ICs when compared to the population transfer of their red-shifted counterparts. However, there is only limited sampling to support this claim of wavelength-dependent branching ratios, and further investigations are necessary to establish and explain the origin of this difference.

![Graph showing population decay](image)

**Figure S10**: Ground-state (red) and excited-state (blue) population of HBDI in water for (a) red-shifted initial conditions ($S_0/S_1$ excitation energy < 2.84 eV) and (b) blue-shifted initial conditions ($S_0/S_1$ excitation energy > 2.84 eV). Populations are averaged across all initial conditions for each species. Solid lines represent the average populations for each state. The vertical, gray dotted line represents the time at which population transfer begins ($\tau_0$). The black dotted line is the delayed bi-exponential fit (consisting of $\tau_1$ and $\tau_2$ components) to the $S_1$ population decay. Transparent lines represent the error obtained from bootstrapping analysis.
XI. Flapping angle of HBDI\(^{-}\) during AIMS dynamics

**Figure S11**: Flapping angle ($\phi_{\text{flapping}}$) between the I- and P-rings during AIMS dynamics of HBDI\(^{-}\) in water. Blue lines represent parent TBFs, and red lines represent spawned TBFs. Atoms involved in calculating the flapping angle are highlighted in purple, and the flapping angle is defined as the dihedral angle illustrated by the inset.
XII. Relative population transfer of HBDI⁻ spawning events

Figure S12: (a) Visual of the 90° rotation of HBDI⁻ that leads to the orientation in (b), which illustrates positive and negative $\theta_{\text{pyr}}$ angles. (c) Relative population transfer and pyramidalization angle associated with the bridge carbon atom for spawned geometries of HBDI⁻ in water from parent TBFs. $\phi_{I}$ and $\phi_{P}$ dihedral angles correspond to geometries upon entering a spawning region. Relative population transfer is defined by the fraction of original $S_1$ population transferred to $S_0$ relative to the $S_1$ population upon entering the spawning region. (d) Relative population transfer distributions of I- and P-spawning events in water.
XIII. Dihedral angle distribution for spawned HBDI$^-$ geometries

![Graphs showing dihedral angle distribution](image)

**Figure S13:** Dihedral angle distribution for spawned geometries of HBDI$^-$ in water (from $S_1$ to $S_0$). (a) $\phi_I$ dihedral angle distribution for TBFs of initial conditions that undergo I-deactivation (b) $\phi_P$ dihedral angle distribution for TBFs of initial conditions that undergo P-deactivation.
XIV. Photoproducts of HBDI⁻

**Figure S14:** Photoproducts of HBDI⁻ initial conditions, following 500 fs of ground-state QM/MM-MD simulations for child TBFs spawned on S₀. Photoproducts are shown for spawned geometries reached via (a) I-twisted and (b) P-twisted deactivation.
Figure S15: Fluorescence maxima of solvated HBDI\(^-\) from theory and experiment.\(^4\) The theory spectrum consists of data from all fifty initial conditions. Based on this energy gap, the theory spectra were shifted by 0.13 eV to align the two maxima for analysis purposes.
XVI. Oscillator strength as a function of twisting angles

Figure S16: Oscillator strength of aqueous HBDI$^-$ as a function of twisting along $\phi_I$ and $\phi_P$ dihedral angles. Data is obtained from average dihedral angles and oscillator strengths recorded for 10° umbrella sampling windows on $S_1$. The “X” marker at the origin represents the point of complete planarity with respect to the $\phi_I$ and $\phi_P$ dihedral angles.
XVII. Geometric parameters of critical points in water

Table S1: Key geometric parameters for representative α(0.67)-SA3-CASSCF(4,3)/6-31G* critical points of HBDI− in water.

|                  | \( r_1 \) (Å) | \( r_P \) (Å) | \( \phi_I \) (degrees) | \( \phi_P \) (degrees) | \( \theta_{\text{pyr}} \) (degrees) |
|------------------|----------------|----------------|-------------------------|-------------------------|----------------------------------|
| \( S_0 \) minimum | 1.38           | 1.41           | 4.28                    | 11.67                   | 1.97                             |
| MECI-I           | 1.45           | 1.41           | 84.31                   | 4.31                    | 14.39                            |
| MECI-P           | 1.43           | 1.48           | 17.03                   | 72.68                   | -34.23                           |
XVIII. Comparison of $S_0/S_1$ energy gap across multiple complete active space methods

Table S2: Difference between $S_0$ and $S_1$ electronic state energies for geometries used to benchmark $\alpha$-CASSCF(4,3) in water. The FC geometry is optimized on $S_0$, and all other geometries are optimized on $S_1$. The $\alpha(0.67)$-SA3-CASSCF(4,3) method was used for optimization. XMS-CASPT2(4,3) and SA3-CASSCF(4,3) single point energy calculations were run on the optimized $\alpha$-CASSCF geometries. A 6-31G* basis set was used for all methods.

|       | $\alpha(0.67)$-SA3-CASSCF(4,3) (eV) | XMS-CASPT2(4,3) (eV) | SA3-CASSCF(4,3) (eV) |
|-------|-----------------------------------|----------------------|----------------------|
| FC    | 2.90                              | 2.90                 | 4.33                 |
| Planar| 2.92                              | 2.92                 | 4.36                 |
| I15 / P15 | 2.54 / 2.62                     | 2.55 / 2.61          | 3.80 / 3.90          |
| I30 / P30 | 2.28 / 2.45                     | 2.31 / 2.45          | 3.40 / 3.65          |
| I45 / P45 | 1.83 / 2.08                     | 1.90 / 2.13          | 2.73 / 3.11          |
| I60 / P60 | 1.25 / 1.42                     | 1.36 / 1.56          | 1.86 / 2.11          |
| I75 / P75 | 0.45 / 0.95                     | 0.63 / 1.24          | 0.66 / 1.42          |
| I90 / P90 | 0.09 / 0.86                     | 0.36 / 1.24          | 0.13 / 1.28          |
XIX. Oscillator strength of geometries used to benchmark $\alpha$-CASSCF

**Table S3:** Oscillator strength of $S_0 \rightarrow S_1$ electronic transitions for geometries used to benchmark $\alpha$-CASSCF(4,3) in water. The FC geometry is optimized on $S_0$, and all other geometries are optimized on $S_1$. The $\alpha(0.67)$-SA3-CASSCF(4,3)/6-31G* method and basis set were used for optimization.

|       | Oscillator strength, $S_0 \rightarrow S_1$ (a.u.) | Energy gap, $S_0 \rightarrow S_1$ (eV) |
|-------|-----------------------------------------------|--------------------------------------|
| FC    | 1.15                                          | 2.90                                 |
| Planar| 1.09                                          | 2.64                                 |
| I15 / P15 | 1.01 / 1.08                                    | 2.54 / 2.62                          |
| I30 / P30 | 0.80 / 0.96                                    | 2.28 / 2.45                          |
| I45 / P45 | 0.53 / 0.67                                    | 1.83 / 2.08                          |
| I60 / P60 | 0.30 / 0.28                                    | 1.24 / 1.42                          |
| I75 / P75 | 0.08 / 0.07                                    | 0.45 / 0.95                          |
| I90 / P90 | 0.00 / 0.01                                    | 0.09 / 0.86                          |
XX. Photoproducts of red-shifted and blue-shifted initial conditions

Table S4: Classifications for red-shifted ($S_0/S_1$ excitation energy < 2.84 eV) and blue-shifted ($S_0/S_1$ excitation energy > 2.84 eV) initial conditions used for AIMS simulations of HBDI$^-$ in water. Each initial condition is classified based on whether rotation around $\phi_I$ or $\phi_P$ is dominant on $S_1$. The I/P ratio average and standard deviation for red-shifted and blue-shifted initial conditions are based on a bootstrapping analysis using 1000 bootstrapping samples.

|                  | Initial conditions | I-products | P-products | I/P ratio |
|------------------|--------------------|------------|------------|-----------|
| Red-shifted      | 20                 | 12         | 8          | 1.7 ± 1.1 |
| Blue-shifted     | 30                 | 21         | 9          | 2.7 ± 1.6 |
XXI. Classifications for TBF photoproducts

Table S5: Classifications for I- and P-twisted photoproducts, based on the configuration of spawned geometries after 500 fs of ground-state QM/MM-MD. A threshold of 120° is used to determine whether a Z-1 product (less than 120°) or an isomerized product (greater than 120°) is generated. The Z-1 product represents a Z-isomer of HBDI$^-$ that is not the result of Z/Z isomerization.

|         | Total spawns | Z-1 products | Isomerized products |
|---------|--------------|--------------|---------------------|
| I-channel | 181          | 145          | 36                  |
| P-channel | 83           | 52           | 31                  |
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