Photochemistry

T₁ Population as the Driver of Excited-State Proton-Transfer in 2-Thiopyridine

Sebastian Eckert,[a] Jesper Norell,[b] Raphael M. Jay,[a] Mattis Fondell,[c] Rolf Mitzner,[c] Michael Odelius,[b] and Alexander Föhlisch[a,c]

Abstract: Excited-state proton transfer (ESPT) is a fundamental process in biomolecular photochemistry, but its underlying mediators often evade direct observation. We identify a distinct pathway for ESPT in aqueous 2-thiopyridine, by employing transient N1s X-ray absorption spectroscopy and multi-configurational spectrum simulations. Photoexcitations to the singlet S₂ and S₄ states both relax promptly through intersystem crossing to the triplet T₁ state. The T₁ state, through its rapid population and near nanosecond lifetime, mediates nitrogen site deprotonation by ESPT in a secondary intersystem crossing to the S₀ potential energy surface. This conclusively establishes a dominant ESPT pathway for the system in aqueous solution, which is also compatible with previous measurements in acetonitrile. Thereby, the hitherto open questions of the pathway for ESPT in the compound, including its possible dependence on excitation wavelength and choice of solvent, are resolved.

Introduction

Protonation and deprotonation of active atomic sites in biomolecular systems crucially define the function and activity of individual building blocks in the hydrogen-bonding network of aqueous environments. Such intermolecular coordination can be altered through photoexcitations of the system, dependent on the properties of the involved valence excited states along the decay pathways. Solute–solvent interactions can consequently affect the dominant reaction mechanisms in photochemical processes. In particular, excited-state proton transfer (ESPT), an ubiquitous process in biomolecular systems,[1–3] can be strongly dependent on, and in some cases also mediated by, the surrounding solvent molecules.[4] Here, we study the excited-state dynamics of 2-thiopyridine (2-TP) in an aqueous environment. The compound exhibits optically accessible valence excited states, which have been reported to trigger proton transfer between its nitrogen (N) to its sulfur (S) site.[5] A strong solute–solvent interaction is indicated, for example, by a chemical environment dependent N protonation.[6–8]

Three molecular species previously associated with intra- and inter-molecular ESPT in 2-TP are shown in Figure 1: 2-TP, its N to S site proton transferred tautomer 2-mercaptopryridine (2-MP), and the N site deprotonated anion (2-TP⁻). The dynamics induced by excitation with 266 nm photons at the S₀ resonance of 2-TP (aq) have been investigated by Du et al.[9] with transient Raman vibrational spectroscopy. They identified transient modes of the 2-MP tautomer in its S₅ state on nanosecond timescales and by that categorize 2-TP as an ESPT compound.[9] Recently, we observed signatures of ultrafast deprotonation of the N site of 2-TP (aq) on sub-picosecond timescales upon 400 nm excitation,[10] supporting this categorization. Additionally, Van Kuiken et al.[11] studied the excited-state dynamics of 2-TP in acetonitrile upon S₁ photoexcitation with 400 nm photons using transient S1s (tender X-ray) NEXAFS, a method

![Figure 1. Presently identified molecular species of relevance for ESPT in 2-TP: the thione–thiol tautomers 2-TP and 2-MP, and the deprotonated anion 2-TP⁻.](https://doi.org/10.1002/chem.201804166)
proven to yield detailed information on photoinduced dynamics of sulfur-containing molecules.\textsuperscript{16,17} Van Kuiken et al.\textsuperscript{15} found indications of a transition between the T\textsubscript{1} states of the 2-TP and 2-MP species on sub-ns timescales, but a distinct signature of a N site deprotonated species in its S\textsubscript{3} state was not observed. The mentioned studies differ in terms of excitation energies and solvent environment and have so far not provided a mechanistic understanding of the ESPT in 2-TP.

In this study, we investigate the potentially different, excitation wavelength and solvent dependent, preferential decay pathways of photoexcited 2-TP in solution and identify the relevant transient states of 2-TP mediating the ESPT process. We use N\textsubscript{1}s near edge (soft) X-ray absorption fine structure (NEXAFS) spectroscopy, which has proven to be sensitive to N site protonation states\textsuperscript{13,14,18–20} and to the valence excited states of 2-TP\textsuperscript{14} in an optical pump X-ray probe experimental scheme. Together with spectrum simulations and energy levels from high-level multi-configurational quantum chemistry, this allows us to identify the molecular species and electronic states involved in the photoinduced decay cascade on pico- to nanosecond timescales. We investigate the response of 2-TP (aq) to both direct S\textsubscript{3} and S\textsubscript{1} photoexcitation on the timescale of sub-nanoseconds to tens of nanoseconds and reevaluate the partially contradictory results in the previous studies. From careful theoretical analysis of available time-resolved spectroscopic data, we put forward a fully consistent interpretation of the pico- to nanosecond decay pathway following photoexcitation of 2-TP involving a transient proton transfer process, which is independent both of excitation wavelength and of solvent.

Results and Discussion

The N\textsubscript{1}s NEXAFS spectrum of 2-TP (aq) shown in Figure 2a exhibits a distinct \( \pi^* \)-resonance at a photon energy of 400.5 eV, assigned to excitation into the 5\( \pi^* \) LUMO. For the transient spectra, we focus on intensity changes for photon energies below and on this resonance, to avoid spectral overlap with continuum contributions in the analysis. The marked excitation wavelengths in the UV/Vis spectrum of the sample, presented in the inset in Figure 2a, confirm that the two distinct optical absorption resonances are driven dominantly under the experimental conditions. In accordance with Du et al.\textsuperscript{9} our electronic structure calculations show that the lowest valence excited states of 2-TP can be grouped into singlet/triplet—pairs of identical n→\( \pi^* \) or \( \pi→\pi^* \) excitation character, as shown in Table 1. In detail, these correspond to electronic excitations from the 4\( \pi \) HOMO (S\textsubscript{3}/T\textsubscript{2} and S\textsubscript{4}) and the S site lone-pair 14\( \alpha \) HOMO-1 (S\textsubscript{3}/T\textsubscript{2} and S\textsubscript{4}) into the 5\( \pi^* \) LUMO (S\textsubscript{5}/T\textsubscript{1} and S\textsubscript{6}/T\textsubscript{1}) and 6\( \alpha^\prime \) LUMO+1 (S\textsubscript{5} and S\textsubscript{6}). Hence, the two optical absorption resonances in the UV/Vis spectrum correspond to \( \pi→\pi^* \) excitations to the S\textsubscript{3} and S\textsubscript{4} states, whereas excitation out of the lone-pair orbital and to triplet states are optically dark.

The transient NEXAFS spectra in an energy range between 394 eV and 402 eV at different pump-probe delays are presented in Figure 2b. The displayed spectra for the two different excitation wavelengths exhibit the same transient signatures in the investigated range of pump-probe delays from 0.1 ns to 40 ns. This indicates that the decay pathways from the S\textsubscript{2} and the S\textsubscript{4} valence excited states of 2-TP (aq) merge at a common intermediate (excited state) on or below a picosecond timescale, with indistinguishable subsequent kinetics. The de-
creased 2-TP $S_0$ population for positive pump-probe delays causes a bleach (vertical dark red line) of the absorption centered at 400.5 eV. The species present at a delay of 0.1 ns contribute to the first detected transient absorption feature (vertical red line) peaked at 397.8 eV. At a delay of 2.5 ns, a second feature (vertical green line) is picked up at 398.9 eV whereas the first feature at 397.8 eV is no longer present. A third transient feature (vertical gray line) at 399.8 eV (close to the bleach), as well as the bleach itself, are additionally detected at all pump-probe delays up to 40 ns.

We now introduce the framework in which the detected transient NEXAFS signatures will be assigned to electronic states of the three possible molecular species 2-TP, 2-MP, and 2-TP$^-$. Both the simulated spectral contributions and the calculated state energies (for all possible combinations of electronic states and molecular species presented in Figure 3 and Table 1) are considered to identify the observed species through a conclusive process of elimination. As the system relaxing from the initial $S_0$ and $S_1$ excitations appears to have reached a common pathway on the investigated timescales, we employ a strict energetic argument: No excited states energetically above 2-TP $S_1$ are considered as possible detected transient species. The energies of all 2-TP and 2-MP states can be directly related. The energies of 2-TP$^-$ in its different states, however, can neither be directly related to 2-TP nor to 2-MP. The energy of the deprotonation cannot be accurately determined from the simulations, as even if a bare dissociated proton had been included, its covalent bonding with the surrounding solvent molecules would have been neglected by the polarizable continuum model (PCM). The presented energies of 2-TP$^-$ have therefore instead been shifted to align the $S_0$ state energies of 2-TP and 2-TP$^-$. This shift is well motivated to define a lower bound for the state energies of 2-TP$^-$, as 2-TP is known to be the dominant species over 2-TP$^-$ in a neutral aqueous environment$^{[10–13]}$ (i.e., for the experimental conditions, the energy of 2-TP$^-$ is higher than the energy of 2-TP in the $S_0$ state).

The simulations show that all excited states in 2-MP and 2-TP$^-$ can be excluded as possible detected transient species. Firstly, energetic limitations in the reaction kinetics render them non-accessible from the 2-TP $S_1$ state (as shown at the top of Figure 3 and in Table 1). Secondly, the associated opening of resonances at photon energies below 397 eV (see Figure 3b,c) is in contradiction with the transient experimental spectra, even if we allow for some uncertainty in the peak positions associated with the implicit solvation model. The 2-TP $S_0$ and $T_2$ states exhibit $\pi-\pi^*$ excitations from an $S$ site lone-pair orbital and consequently do not open any additional $N_1s$ absorption resonances in the region of the detected transients. Thus, the only two electronically excited species that can be responsible for the first observed transient feature (397.8 eV, red) are the 2-TP $S_2$ and 2-TP $T_1$ states, which are associated with $\sigma-\pi^*$ excitations. The spectrum simulations for these states show that in this system our probe is not directly sensitive to spin degrees of freedom; the two states exhibit virtually identical spectral signatures and must be considered as equally likely candidates from a spectroscopic point of view. Yet, with a lifetime of the excited state in the hundreds of picoseconds, we still assign this transient feature confidently to a dominant (or exclusive) 2-TP $T_1$ population. This assignment is based on the fact that the $T_1$ state is the lowest electronically excited state of 2-TP. Such intersystem crossing (ISC) to the triplet state manifold upon photoexcitation has been previously shown to occur on sub-picosecond timescales in similar thione compounds from spectroscopic measurements and simulations of excited-state dynamics$^{[21–23]}$.

The $S_0$ ground states of the 2-MP and 2-TP$^-$ species both exhibit a single $N_1s$ $\pi^*$ resonance at approximately 399 eV in the simulations. This agrees well with the previously determined experimental resonance of 2-TP$^-$,$^{[13, 14]}$ and it coincides with the second detected transient feature (398.9 eV, green). As all other combinations of electronic states and molecular species have previously either been ruled out or assigned, the associat-
ed transient species can be conclusively assigned to result from deprotonation of the N site. Owing to the spectral similarities of 2-MP and 2-TP in the S state, however, the protonation state of the S site cannot be unambiguously determined, leaving the 2-MP and 2-TP molecular species as equally viable candidates.

The third transient feature (399.8 eV, gray) is, as previously noted, present in all spectra at the different pump-probe delays. For the 2-TP T₁ state, the LUMO absorption resonance is slightly shifted with respect to 2-TP S₀ which could yield a low energy shoulder to the ground state bleach. However, as neither 2-MP nor 2-TP in their S₀ states exhibit any features in the relevant region, the transient cannot be described by the considered combinations of electronic excitations and molecular species at nanosecond timescales. Owing to its apparent formation upon decay from 2-TP Tᵰ, the transient most likely arises from the release of energy into the surrounding solvent environment upon reformation of 2-TP S₀ (and similarly also for 2-MP/2-TP S₀). Such thermalization is known to occur on a timescale of ten picoseconds. The feature is present for tens of nanoseconds after the optical excitation, which is clearly too long for the energy to remain contained within vibrational excitations of the molecule itself. The dissipation of energy into the solvent can thus be expected to appreciably affect the solution as a whole. Laser pump X-ray probe measurements of aqueous solution have shown that the initial isochoric heating leads to significant spectroscopic signatures on nanosecond timescales. The N₁s peak positions are in our case sensitive to solute–solvent interactions. Therefore, it is likely that the relaxation effects on the solvation shell can shift or broaden the spectrum of the 2-TP S₀ state sufficiently to give rise to the transient feature. Such rearrangement of solvent molecules around a photoexcited solute has previously been reported for hard X-ray NEXAFS spectra at the solute absorption edge.

This picture is also supported by the increase of the transient intensity at 399.8 eV upon decay of the first transient species within the first nanosecond (see also Figure 4a) as 2-TP in this temperature affected S₀ state can be expected to exhibit a higher absorption cross section close to the ground state π⁻ resonance than in the T₁ state. Consequently, we assign the transient to the 2-TP S₀ species in a solvent environment with an increased temperature ("hot" 2-TP S₀), which forms in competition with and subsequently to the N site deprotonated species.

The temporal evolution of the discussed transient spectral signatures is presented in Figure 4. A rate model consisting of the states discussed above and illustrated in Figure 4c was numerically solved and fitted to the data. The contribution of each state at the different energies is given in the Supporting Information. The model yields a lifetime of the 2-TP T₁ state of (0.76 ± 0.05) ns, which decays in a forked pathway to either N site deprotonated 2-MP/TP⁻ S₀ with a yield of (14±4)% or the "hot" 2-TP S₀ species with a yield of (86±4)%. The 2-MP/TP⁻ S₀ species has a lifetime of (7.3±0.3) ns and the "hot" 2-TP S₀ cools down with a time constant of (35±1) ns.

Based on comparison of transient vibrational Raman spectroscopy and quantum chemical simulations, Du et al. identified the presence of 2-TP S₀ at nanosecond scales following S₂ excitation in aqueous solution. No intermediate species were identified; the authors therefore hypothesized three different possible unresolved reaction pathways. This observation is fully consistent with our data, which also supports a deprotonated nitrogen site between roughly 1 to 7 ns. However, as Du et al. did not consider the 2-TP⁻ species, and we cannot separately separate it from 2-MP in the S₀ state in our N₁s edge measurements, it remains undetermined whether an intra-molecular proton transfer occurs directly to the S site or if the proton is fully removed from the molecule into the solvent. Still, our data clearly discriminates between the three proposed pathways, and agrees with the third option proposed by Du et al. of ESPT through ISC processes through the 2-TP T₁ state. Thus, in extension of their work, we conclusively determine the dominant reaction pathway for proton dynamics in

![Figure 4. Time traces of selected transients (photon energies marked in Figure 2) for pump-probe delay dependent transient NEXAFS of 2-TP (aq) upon S₂ excitation, with results shown on sub-nanosecond (a) to tens of nanosecond (b) timescales. A rate model described in (c) was fitted to the data set, assuming a decay of the optically accessible singlet states on timescales below our temporal resolution, yielding an initial population of 2-TP T₁ in our experiment, which decays with a lifetime t = (0.76 ± 0.05) ns through a forking into both an N site deprotonated molecular species (2-MP or 2-TP⁻), or the S₀ ground state and a temperature affected "hot" 2-TP S₀ (t = (35 ± 1) ns), with subsequent decay into the 2-TP S₀ state.](image-url)
2-TP upon S₀ excitation in aqueous solution, and can additionally confirm it to be the same also for S₁ excitation.

Unlike Du et al.,[21,22] Van Kuiken et al.[15] studied the S₂ photoexcited dynamics of 2-TP in acetonitrile solution. As in the current work, the dynamics were probed with time-resolved NEXAFS spectroscopy, but from the alternative perspective of the S₁s edge. For direct comparison, we have therefore also performed simulations of the transient S₁s NEXAFS spectra of all our considered species as shown in the Supporting Information, which in large part agree with the measurements and DFT simulations by Van Kuiken et al.[13] Similarly to us, Van Kuiken et al.[13] assigned the 2-TP T_T decay to the formation of the 2-MP molecular species, followed by eventual reformation of 2-TP S₀. However, the authors proposed population of the 2-MP T₁ state as an intermediate from 2-TP T₁ to 2-MP S₀, based on the presence of a transient feature that seemingly overlaps the ground state bleach. As their measurements were performed in acetonitrile, instead of water, the reaction kinetics could in principle differ even qualitatively. Nevertheless, comparison of the state energies in acetonitrile listed in Table S2 in the Supporting Information shows that 2-MP is still 0.6 eV higher in energy than 2-TP in their T₁ states as the relative energy is here not strongly affected by the choice of solvent. An additional activation energy barrier must also be passed in a transition state structure for the transfer to occur on the T₁ potential energy surface. This makes the transition highly unlikely to occur owing to kinetic constraints. As the authors also assign the transient a 15 ns lifetime, it likely corresponds to the signature we assign to “hot” 2-TP S₀. Similarly to our data, the associated S₁s transient feature could in fact be well explained by a temperature-induced broadening and/or shift of the 2-TP S₀ resonance. As further shown in the Supporting Information, S₁s NEXAFS can, analogous to N₁s NEXAFS, hardly distinguish both T₁ from S₁ in 2-TP, and 2-MP from 2-TP in the S₀ state, thereby also leaving the initial electronic dynamics and the S site protonation states undetermined. By reasonably reassigning one of the transient species in the interpretation of Van Kuiken et al.[13] it is thereby fully consistent also with our data and rate model, indicating that the choice of solvent between water and acetonitrile only has a quantitative effect on the timescales of the reaction kinetics.

The proposed pathway for proton transfer in 2-TP results from two steps of ISC, both to and from the T₁ state of 2-TP. The probability for ISC is enhanced in such systems by the presence of the S atom generating an internal heavy atom effect, that is, increased spin-orbit coupling that mixes the spin multiplicities. Originally understood through El-Sayed’s rule,[27] ISC in N-heterocyclics is considered more efficient when occurring between states of different excitation character (i.e. (n, π*) state to (π, π*) state, or vice-versa), which leaves both Sᵢ → Sᵢ → T₁ and S₁ → T₁ → T₄ as probable decay sequences, further facilitated by the small S₁ → S₀ and T₁ → T₄ energy separations. Alternate and even forked pathways for T₁ population have also been reported.[21,22] Definitive determination of the sub-picosecond dynamics that precedes 2-TP T₁ relaxation will therefore require further investigation with femtosecond spectroscopy methods and simulations of excited-state molecular dynamics.

In contrast to the ultrafast T₁ population, ISC from the T₁ state back to S₀ is clearly a less efficient process, which here occurs on the pico- to nanosecond timescale. This is not surprising considering the S₀–T₁ energy gap of 2.66 eV even in the relaxed T₁ structure, which greatly weakens non-adiabatic coupling to the S₀ surface. The ISC is therefore likely mediated by strong fluctuations in structure and/or solvation. These fluctuations then momentarily bridge the energy gap, and mediate the proton transfer in the non-equilibrium dynamics together with the subsequent release of kinetic energy, before energy dissipation to the solvent can fully return the system to the 2-TP S₀ state in its relaxed structure. It also offers a feasible explanation of our recent measurement with time-resolved resonant inelastic X-ray scattering wherein a signature of N-H dynamics was detected on femtosecond timescales.[14] Within the framework of the dominant relaxation pathway established in this work, a fractional deprotonation can still occur within the timeframe of the relaxation from the photoexcited 2-TP S₀ or S₁ states to 2-TP T₁. As proton transfer is clearly a minority channel, even in the currently proposed pathway, it is thus reasonable that the N–H bond can be affected by the 1.23 eV and 2.47 eV respectively released in the decay from the S₀ and S₁ states in the Franck-Condon region to the relaxed 2-TP T₁. This could activate molecular vibrational modes, for example, an N–H stretch, and yield a forked relaxation pathway with partial N deprotonation as the result. The crucial role of solute–solvent interactions in the proposed proton transfer mechanism would also explain the different reaction rates observed in the two solvents: protic water and aprotic acetonitrile.

Conclusion

The relaxation pathway for excited-state dynamics following S₀ and S₁ excitation of aqueous 2-TP was determined from transient N₁s NEXAFS spectroscopy and multi-configurational spectrum simulations. The pathways populated by the two excitations merge to a common species assigned as 2-TP T₁ on sub-nanosecond timescales. Decay of the T₁ state within roughly a nanosecond results in a forked pathway. This, as majority channel, leads back to the 2-TP S₀ state or, as a minority channel, induces a deprotonation of the N site, that is, an ESPT process. The ESPT yields either the thiol tautomer 2-MP or the N site deprotonated anion 2-TP⁻, with a lifetime of a few ns. The release of kinetic energy upon electronic de-excitation results in a “hot” solution environment from which the energy fully dissipates on a scale of tens to a hundred ns.

The assigned dominant decay pathway via the 2-TP T₁ state establishes one of three previously proposed, but unresolved, pathways for ESPT. Comparison with S₁s NEXAFS measurements indicate that the same pathway is populated upon 2-TP S₁ excitation also in acetonitrile solution. The assigned photoinduced proton dynamics thus are independent of the choice of excitation wavelength and protic (aqueous) or aprotic (acetonitrile) solvent. The proposed pathway highlights dissipation of kinetic energy in non-equilibrium dynamics as the prime driver of the ESPT in 2-TP, a finding of general applicability in studies of ESPT processes as photo-protection mechanisms in biomol-
ecules. Our results provide the framework for investigations of the initial ultrafast (sub-picosecond) relaxation and further characterization of the observed ESPT as an intra- or inter-molecular process.

Experimental and Computational Section

The aqueous 2-TP solution, prepared at a concentration of 150 mM from 2-mercaptopyridine 99% supplied by Sigma–Aldrich and dissolved in deionized water, was sprayed into the experimental vacuum chamber in a liquid flat jet sample environment. The transmitted average intensity of synchrotron radiation from the beamline UE52-SGM at the synchrotron BESSY II with a bandwidth of 35 meV was monitored using a gallium arsenide photodiode to record the static N1s NEXAFS of 2-TP. A background generated from the tabulated attenuation length of the chemical composition of the solution without the N contribution was subtracted from the spectrum. The transient changes of the absorption were induced using approximately 300 fs long pulses of the third and fourth harmonic centered at 343 nm and 258 nm of a fiber laser system with a fundamental wavelength of 1030 nm. It was operated at a repetition rate of 208 kHz and the laser beam was focused to a dimension of (80×80) μm². The laser system was synchronized with the arrival time of the hybrid bunch in the gap of BESSY II fill pattern and the arrival time was shifted with respect to the arrival time of the X-ray pulses originating from the hybrid bunch to adjust the pump-probe delay. The hybrid bunch provides X-ray pulse pairs with a bandwidth of 120 meV at a rate of 1.25 MHz. The intensity of each X-ray pulse accompanied by a laser pulse as well as the pulse intensity 4 μs later, transmitting the replenished sample in its electronic ground state, were recorded from the voltage pulse of a capped (200 nm Al film) silicon avalanche photodiode. The signal was amplified by 20 dB and accumulated using a digital boxcar averager from Žürich Instrument AG. This scheme yields a differential measurement of the transient absorption changes. Details of the measurement scheme have been summarized by Fondell et al.[30] The sample thickness was estimated from the attenuation of the synchrotron radiation by the jet in comparison to tabulated attenuation values[30] to be approximately 6.5 μm for the static spectrum in Figure 2a, 6.3 μm for the transient spectra in Figure 2b, and 3.9 μm for the delay dependent absorption changes in Figure 4 for 343 nm excitation. The spectra for 258 nm excitation were accumulated at sample thicknesses in the few μm range. The pulse energies used for the excitation were 12.4 μJ for the transient spectra at 343 nm, 3.3 μJ for the delay traces at 343 nm, and 2.7 μJ for the spectra at 258 nm. The UV/Vis spectrum of 2-TP (aq) was recorded using a Shimadzu UV-2700 spectrometer at a sample concentration of 0.1 mM in a quartz cuvette, providing a 10 mm sample thickness.

Multi-configurational quantum chemical calculations were performed with the complete active space self-consistent field (CASSCF)[32] method, restricted active space self-consistent field (RASSCF)[31] method, and the second-order perturbation theory restricted active space (RASPPT2)[32] multi-state[31] method, with the 6-31G basis set, as implemented in the Molcas 8.2 program package.[33] No explicit use of symmetry was employed for geometry optimization nor for the electronic structure calculations.

Geometry optimization of all molecular species (i.e., molecular structures and electronic states) was performed at the CASSCF(10,8) level (10 electrons in 8 orbitals), with an active space that included the initially occupied 4×π orbitals, the initially occupied 1×σ lone-pair orbital (S centered in 2-TP and 2-TP⁻, N centered in 2-MP), and the initially unoccupied 3×π' orbitals.

Spectra simulations were performed with RASSCF calculations and subsequent application of multi-state RASPT2, the latter utilizing an imaginary shift of 0.05 Hartree to accelerate convergence. Interactions with the aqueous surrounding were approximately included through a polarizable continuum model (PCM) equilibrated to the initial state of NEXAFS process.[30] The active space included the same orbitals in the RAS2 space (no restrictions, equivalent to the complete active space method) as in the corresponding geometry optimization and, additionally, the N1s core orbital in the RAS1 space (restricted to a maximum of one electron-hole). The N1s orbital was frozen for the RASSCF orbital optimization, to avoid relaxation from core- to valence-excitations in the final state calculations. For each spectrum, five (three for 2-MP owing to convergence problems of the higher states) non-core-excited states (ground state and valence-excited) and ten core-excited states, were separately obtained within the state averaging formalism. Core-excited states were obtained through a separate orbital constraint on the N1s orbital (to avoid rotation with N2s) and the highly excited states (HEXS) method[36] (which eliminates contributions for all electronic configurations with doubly occupied N1s). Transition electronic dipole moments for NEXAFS intensities were obtained with the RAS state interaction method (RASSI).[37,38] For comparison with the measured NEXAFS spectra, the simulated discrete transitions were convoluted with a Gaussian broadening of 0.7 eV accounting for the bandwidth of incident radiation in combination with inhomogeneity and dynamical effects in the sample through solute–solvent interactions. The Lorentzian lifetime broadening with full-width at half-maximum (FWHM) was neglected (estimated as 0.11 eV[39] from N2 and N2O). An absolute shift of −3.78 eV (to account for, for example, the frozen N1s orbital and limitations in dynamical correlation) was applied to the energies of all core-excited states, which aligns the absorption resonance of 2-TP S0 to the experimental value of 400.5 eV.

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Conflict of interest

The authors declare no conflict of interest.