Bridging the Gap between the Gas and Solution Phase: Solvent Specific Photochemistry in 4-tert-Butylcatechol

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ABSTRACT: Eumelanin is a naturally synthesized ultraviolet light absorbing biomolecule, possessing both photoprotective and phototoxic properties. We infer insight into these properties of eumelanin using a bottom-up approach, by investigating a subunit analogue, 4-tert-butylcatechol. Utilizing a combination of femtosecond transient electronic absorption spectroscopy and time-resolved velocity map ion imaging, our results suggest an environmental-dependent relaxation pathway, following irradiation at 267 nm to populate the $S_1$ ($\pi\pi^*$) state. Gas-phase and non-polar solution-phase measurements reveal that the $S_1$ state decays through coupling onto the $S_2$ ($\pi\pi^*$) state that is dissociative along the non-intramolecular hydrogen bonded ‘free’ O–H bond. This process is mediated by tunneling beneath an $S_1$/S2 conical intersection and occurs in $4.9 \pm 0.6$ ps in the gas-phase and $27 \pm 7$ ps in the non-polar cyclohexane solution. Comparative studies on the deuterated isotopologue of 4-tert-butylcatechol in both the gas- and solution-phase (cyclohexane) reveals an average kinetic isotope effect of $\sim 19$ and $\sim 7$, respectively, supportive of O–H dissociation mediated by a quantum tunneling mechanism. In contrast, in the polar acetonitrile, the $S_1$ state decays on a much longer timescale of $1.7 \pm 0.1$ ns. We propose that the $S_1$ decay is now multicomponent, likely driven by internal conversion, intersystem crossing and fluorescence, as well as O–H dissociation. The attribution of conformer driven excited state dynamics to explain how the $S_1$ state decays in the gas and non-polar solution-phase versus the polar solution-phase, elegantly demonstrates the influence the environment has on the ensuing excited state dynamics.

1. INTRODUCTION

The photoexcited state dynamics of biomolecules have borne extensive interest in the field of photochemistry due, in part, to an apparent relative photostability present in key biochemical systems, despite the fact that they contain ultraviolet (UV) chromophores. The absorption of UV radiation has the ability to cause bond dissociation within molecules, which can lead to undesired chemical changes; mutagenesis in DNA being a prime example. In order for biomolecules to be photostable after UV absorption, they must be able to efficiently redistribute any excess energy away from such harmful pathways. Typically, these non-dissociative processes must occur on an ultrafast (femtosecond (fs) to picosecond (ps)) timescale in order to kinetically out-compete the destructive pathways; however, notall exceptions of longer-lived excited states do exist, e.g. DNA excimers.

Employing a bottom-up approach, in which the UV chromophore subunit of a larger biomolecule is isolated and studied, can lend valuable insights from which we may begin to understand the photochemistry of the larger biomolecule as a whole. Much of the previous work in this field has been performed in the isolated gas-phase, which proves successful in recovering information about the molecular dynamics of these UV chromophores without the presence of environmental effects. Whilst this provides a good starting point for the interpretation of excited state dynamics in larger systems, interactions with surrounding solvent and solute molecules need to be taken into consideration in order to develop a more complete picture of photostability in nature. Placing a chromophore in solution allows for environmental perturbations on the chromophore to be studied, more closely matching the native environment of such biomolecules. A comparison of gas- and solution-phase dynamics has been previously performed on an amino acid subunit, the UV chromophore phenol. Subtle differences between the two phases were observed, supporting the postulate that the local environment of biomolecules can play an important role in the molecule’s photodynamics, whilst also highlighting that knowledge of the dynamics in the gas-phase is highly transferable and complementary to solution-phase studies.

Many other phenol analogues have also been studied in the gas-phase, one of which is catechol (1,2-dihydroxybenzene) - a UV chromophore in a range of biomolecules, one of which is eumelanin. As one of three types of melanin, eumelanin is largely responsible for skin’s frontline defense to UV exposure. However, gaining a complete understanding of its photodynamics is ongoing, with much progress to be made, evidenced by recent work implicating the long-term (after many hours of UVA exposure) phototoxicity of eumelanins.

Previous gas-phase studies of catechol have shown that after excitation to the first excited $\pi\pi^*$ state ($S_1$), H-atom elimination is observed from dissociation of the non-intramolecular hydrogen bonded ‘free’ O–H bond, leading to the formation of the catechol radical in approximately 10 ps. Immediately, this suggests that catechol is not photostable following UV exposure, and yet it is a subunit of a photoprotective biomolecule. In nature, eumelanin resides within an organelle as large hetero-polymer fibrils, thus gas-phase studies are unable to fully capture the photodynamics of catechol in this native environment. Therefore, we ask: do the natural surroundings of catechol influence its photodynamics?

In the present study, we try to address this question. Owing to catechol’s poor solubility in non-polar solvents, the functionalized catechol, 4-tert-butylcatechol (termed 4-TBC hereon), is

\[ \text{[4-tert-Butylcatechol]} \]
used instead, shown in Figure 1. In order to gauge how well 4-TBC performs as a proxy for catechol, we performed gas-phase time-resolved velocity map ion imaging (TR-VMI) and time-resolved ion yield (TR-IY) measurements. To model the role that the surrounding solvent plays in the excited state dynamics of 4-TBC after excitation to the S1 state, we employ fs transient electronic (UV-visible) absorption spectroscopy (TEAS), using the weakly perturbing, non-polar solvent, cyclohexane, and the highly perturbing polar solvent, acetonitrile.

II. METHODS

The detailed experimental procedures pertaining to TR-VMI, TR-IY and TEAS have been extensively described elsewhere. Briefly, a commercially available Ti:Sapphire oscillator and amplifier system (Spectra-Physics) produces 3 mJ laser pulses of ~40 fs duration centred at 800 nm and a repetition rate of 1 kHz. For TR-VMI and TR-IY two optical parametric amplifiers (Light Conversion TOPAS-C), each pumped with ~1 mJ/pulse at 800 nm, produce tunable UV pump and probe pulses (hv_pump and hv_probe respectively), hv_pump is centred at 267 nm (~6 µJ/pulse) and is temporally varied with respect to the probe using a hollow gold retroreflector mounted on a motorized stage, which enables a maximum temporal delay of Δτ = 1.2 ns to be achieved. hv_probe is set to 243 nm (~7 µJ/pulse) to resonantly ionize H-atoms. Both beams are then focused, near-collinearly, into the VMI spectrometer, perpendicularly intersecting a molecular beam of 4-TBC (95%, Sigma-Aldrich) seeded in 2 bar of helium. The molecular beam is formed by supersonic jet expansion from an Even-Lavie pulsed solenoid valve34 heated to 100 °C, and is passed through a 2 mm conical skimmer. The VMI spectrometer is in line with the molecular beam and follows the standard Eppink and Parker design.35 After photolysis of 4-TBC with hv_pump, resulting H-atoms are ionized with hv_probe. The VMI optics project the 3-D velocity distribution of H+ ions towards a position sensitive detector which consists of a pair of microchannel plates and a P-43 phosphor screen (Photek, VID-240), the emitted light being captured on a CCD camera (Basler, A-312f). The original 3-D velocity distribution is reconstructed from the resulting 2-D image using an image reconstruction algorithm.36 Radial pixels on the image are converted into total kinetic energy release (TKER) using the appropriate Jacobian, calibration factor and co-graft mass, generating the desired 1-D TKER spectra. For TR-IY, we simply record the appearance of the parent ion signal (4-TBC+) as a function of Δτ, by gating the detector on this signal and recording the total emitted light registered on the detector.

For TEAS, a 1 mJ/pulse 800 nm laser beam is split into two beams of: (i) 0.95 mJ/pulse and (ii) 0.05 mJ/pulse. (i) is used to generate hv_pump centred at 267 nm (1–2 mJ/cm²) through second and then third harmonic generation using two beta-barium borate crystals. (ii) is used to generate hv_probe, a white light continuum (330-675 nm). Pump-probe polarizations are held at magic angle (54.7°) relative to one another. Changes in optical density (ΔOD) of the sample were calculated from probe intensities, collected using a spectrometer (Avantes, AvaSpec-ULS1650F). The delivery system for the samples (4-TBC in either cyclohexane (100%, VWR) or acetonitrile (99.9%, VWR)) is a flow-through cell (Demountable Liquid Cell by Harrick Scientific Products, Inc.). The sample is circulated using a PTFE tubing peristaltic pump (Masterflex) recirculating sample from a 50 ml reservoir in order to provide each pump-probe pulse-pair with fresh sample.

Comparative TR-VMI and TEAS studies also were carried out on the deuterated isotopologue of 4-TBC, i.e., 4-TBC-d2, in which we have selectively deuterated both O–H bonds (CdH2O2D2) synthesized and characterized according to the description provided in the supporting information (SI). For the TEAS, a bespoke sample delivery system was implemented in order to keep the sample in a dry, inert atmosphere to minimize isotopic exchange. Further details are provided in the SI.

III. RESULTS AND DISCUSSION

a. Gas-phase studies

Figure 2a shows an example TKER spectrum with the image from which it was extracted shown inset; the left side corresponds to the recorded H+ image while the right half shows the reconstructed slice through the centre of the 3-D ion distribution (the black arrow indicates the electric field polarization, e, of the pump pulse). A pump wavelength of 267 nm (4.65 eV) was used to photoexcite 4-TBC to the S1 state, with the generated H-atoms, associated with O–H dissociation, subsequently ionized via a [2+1] resonance enhanced multiphoton ionization (REMPI) scheme. The pump-probe time delay (Δτ) used when recording this particular image was set at Δτ = 1.2 ns. A high kinetic energy (KE) feature is apparent in the TKER spectrum centered at ~7000 cm⁻¹, which returns to the baseline by ~10,000 cm⁻¹. This high-KE signature strongly accords with our previous work in catechol,32 and is associated with H-atoms generated through dissociation of the ‘free’ (non-hydrogen bonded) O–H bond (see Figure 1a) to yield the 4-tet-butylicateoxy radical C6H4O2X (termed 4-TBC* hereon) plus H photoprotein, via the dissociative (O–H coordinate) 1π* surface (S2).

Collecting a series of TKER spectra at varying Δτ and then integrating the high KE feature over the range ~5500-8500 cm⁻¹ results in the H+ transient shown in Figure 2b (blue circles). In order to obtain a time constant for the S2 mediated O–H dissociation, a kinetic fit to the data is applied, comprising two exponential rise functions convoluted with our instrument response (~120 fs full width half maximum), shown by the blue line. Two (gas-phase, g) time constants (τ) are returned: τMP > 30 fs, which is associated with multiphoton processes (MP),37.
results of both the D+ and 4-TBC-d2+ transients are shown in Figure 2c. Kinetic fits to the transients return time-constants of \( \tau_D = 98 \pm 2 \) ps and \( \tau_D = 102 \pm 3 \) ps and thus an average kinetic isotope effect (KIE, \( k_D/k_H \)) of \(-19\). Here the time-constant typography has the same meaning as above, but applied to the deuterated isotopologue. The large KIE obtained here is a tell-tale signifier that the ‘free’ O–H dissociation is very likely mediated through tunneling under the S1/S2 CI. For completeness, we also note that \( \tau_{IVRD} = 1.0 \pm 0.1 \) ps compares favorably with the value extracted for the non-deuterated isotopologue. Notably, the KIE obtained for 4-TBC accords well with similar measurements carried out in catechol, which returned a KIE \(-30\), validating the use of 4-TBC as a proxy for catechol (vide supra).

b. Solution-phase studies

The non-polar solvent, cyclohexane, was used as the starting point for unraveling the solution-phase dynamics of 4-TBC, as the weakly perturbing environment of cyclohexane serves as a good model for the gas-phase environment. Figure 3a shows transient absorption spectra (TAS) of 4-TBC in cyclohexane, following 267 nm (4.65 eV) excitation, having two main features at early \( \Delta t \) (\( \sim 1-5 \) ps); a peak centered around 370 nm and another broader feature centered at 495 nm. These features are attributed to the excited state absorption (ESA) from the S1 state to higher-lying S2 states. At \( \Delta t = 5 \) ps, peaks at 375 and 390 nm appear to lie on top of the S1 absorption; these peaks are the signature absorption of 4-TBC+ in concordance with literature on other phenols. The peak at 390 nm (and to a lesser extent 375 nm) appears to narrow as the spectra evolve over time (\( \Delta t = 5-40 \) ps). This may be explained by quenching of the initially produced hot radical through vibrational energy transfer (VET). Figure 3b (red circles) shows a transient acquired by integrating a 5 nm wide slice of the TAS, centered on 450 nm, as a function of \( \Delta t \). This wavelength is chosen due to the absence of radical absorption in this region, ensuring that signal at this wavelength will come primarily from S1 absorption. This transient was fit with a tweta-exponential decay function (red line); the first three exponentials account for the cyclohexane solvent response alone (see SI for further details). The single time constant obtained for (non-deuterated i.e. H) 4-TBC in cyclohexane \( \tau_{H} = 27 \pm 7 \) ps, is assigned to excited state population flux out of the S1 state. TAS of 4-TBC-d2+ (see SI for details) were also recorded to acquire a corresponding 4-TBC-d2+ transient centered at 450 nm (again using a 5 nm integration around this spectral region) (Figure 3b, blue diamonds). Our kinetic fit (blue line) returns an S1 lifetime for the deuterated (D) species of \( \tau_{D} = 190 \pm 20 \) ps and thus a KIE of \(-7\). Whilst the KIE returned from the cyclohexane measurements is less than that obtained from the gas-phase measurements (cf. KIE \(-19\)), this still suggests O–H dissociation likely occurs along a pathway that is mediated by tunneling through a potential barrier.

We now begin to evaluate the environmental influence on the excited state dynamics of 4-TBC by using a more perturbing, polar solvent. Acetonitrile was selected due to its high polarity and (apparent) low reactivity towards 4-TBC. Figure 3c shows the TAS of 4-TBC in acetonitrile. The main feature at early \( \Delta t \) is a broad ESA across the entire probe window, which we once again attribute to absorption from the S1 state to higher-

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Figure 2. a) TKER spectrum of TBC following excitation at 267 nm and probing H-atom photoproducts with a 243 nm probe pulse at \( \Delta t = 1.2 \) ns. Inset: H+ velocity map image, left half showing raw image, right half showing reconstructed image. b) Normalized integrated H+ signal transient (blue circles) and the corresponding parent ion signal transient (purple diamonds), solid lines show the kinetic fits. c) Normalized integrated D+ signal transient (dark brown circles) and the corresponding parent (4-TBC-d2+) ion signal transient (light brown diamonds). Solid lines show kinetic fit.
Figure 3. a) Selection of TAS of 35 mM 4-TBC in cyclohexane with an excitation wavelength of 267 nm. b) Transient slices of 35 mM 4-TBC in cyclohexane (red circles) and 35 mM 4-TBC-d2 in cyclohexane (blue diamonds) acquired by integrating over a 5 nm window centered at 450 nm; solid lines are the kinetic fits. c) Selection of TAS of 35 nm mM 4-TBC in cyclohexane (red circles) and 30 mM 4-TBC-d2 in acetonitrile (bottom trace) with an excitation wavelength of 267 nm. d) Transient slices of 35 mM 4-TBC in acetonitrile (red circles) and 30 mM 4-TBC-d2 in acetonitrile (blue diamonds) acquired by integrating over a 5 nm centered at 450 nm; solid lines are the kinetic fits. Note that optical density is plotted in the range mOD = 3-12 to magnify the extended decay.

The second extracted time-constant, τhh = 1.7 ± 0.1 ns, is assigned to excited state population flux out of the S1 state in accord with our previous studies on guaiacol.32 Intriguingly, we note there is a >50-fold increase in the lifetime of the S1 state in acetonitrile relative to cyclohexane (cf. τhh = 27 ± 7 ps), the significance of which is discussed below. TAS of 4-TBC-d2 in acetonitrile collected and then integrated with a 5 nm wide slice centered around 450 nm yielded a similar transient (Figure 3d). Our kinetic fit (blue line) returned an S1 lifetime of τhh = 3.3 ± 0.1 ns (as well as τVR(D) = 180 ± 50 fs for completeness) and thus a KIE of ~1.9.

The parent transient, which directly reports on the S1 lifetime, returns a time-constant of τhh = 4.9 ± 0.6 ps whilst the S1 lifetime in cyclohexane is τhh = 27 ± 7 ps. The almost six-fold increase in the S1 lifetime in cyclohexane may be attributed to the very modest polarity of cyclohexane (dielectric constant (εr) = 2.02), which will perturb, albeit weakly, the electronic state energies. The KIE returned from both the gas- and solution-phase (gas ~19, solution ~7) strongly suggests that, following excitation to S1, O–H dissociation proceeds along a barred pathway. Our attribution of tunneling mediated dissociation along the ‘free’ O–H bond, beneath an S1/S2 CI, also accords with previous studies on phenols in the gas- and solution-phase (see reference 2 and references therein). Studies on substituted phenols have also shown the quite dramatic effect substituents have on the topography of either the S1 or S2 (or both) potential energy surface (PES), resulting in noticeable effects on the S1/S2 tunneling probability and
hence $S_1$ lifetime.\cite{17,18,21,45,46} Similar effects on the local topography of the $S_1$ or $S_2$ PESs from a weakly perturbing solvent will undoubtedly have a similar effect on the $S_1$ lifetime as observed here.

Whilst the gas-phase studies compare favorably with those in the non-polar cyclohexane, as expected, this is not the case in acetonitrile ($\Delta t = 37.5$). The $S_1$ lifetime, $\tau_{S_1} = 1.74 \pm 0.1 \text{ ns}$, is >50 times that of 4-TBC in cyclohexane and >350 times that of gas-phase 4-TBC ($\tau_{S_1} = 27 \pm 7 \text{ ps}$ and $\tau_{S_0} = 4.9 \pm 0.6 \text{ ps}$ respectively). The polar acetonitrile is thought to induce the same conformer change to 4-TBC as with catechol, where the intramolecular hydrogen bond is broken, resulting in two intermolecular hydrogen bonds with the solvent.\cite{27,36} To explore the role of conformational change on the observed dynamics, time-dependent density functional theory calculations were performed with the Gaussian09\cite{97} package using both the M052X\cite{48} and CAM-B3LYP\textsuperscript{\textit{95}} functionals and a 6-311G** basis set. These calculations show that the ‘open’ and ‘closed’ conformers are planar in the $S_0$ state (see Figure 1 and SI). In contrast, in the $S_1$ state, whilst the ‘open’ conformer retains planarity, the ‘closed’ conformer becomes distorted (see SI).

In the gas-phase and in cyclohexane, the photoexcited $S_1$ state is non-planar. The non-planar ‘closed’ conformer has been proposed to explain the dramatic decrease in $S_1$ lifetime in catechol relative to phenol (due to symmetry enhanced tunneling).\cite{20,21} A similar reasoning can be applied here for 4-TBC in both non-perturbing environments. In the polar solvent acetonitrile, the photoexcited $S_1$ state likely retains a planar ‘open’ geometry. Such a conformer change (with solvent polarity) has also been observed in guaiacol,\textsuperscript{31,52} resulting in considerably altered photodynamics.\cite{32} In 4-TBC we see an analogous alteration in the observed photodynamics of the $S_1$ excited state in the form of an increased lifetime, an appearance of the triplet state absorption at large values of $\Delta t$ and a smaller radical absorption appearing at $\Delta t \sim 300 \text{ ps}$ and beyond; dynamics which are extremely similar to those seen for guaiacol in methanol\textsuperscript{12} as well as bearing resemblance to their archetype, phenol in cyclohexane.\textsuperscript{15,16} This leads to the conclusion that a solvent-induced conformer change causes the suppression of the O–H bond fission pathway (likely due to an enhanced barrier to tunneling) allowing for additional decay pathways such as internal conversion (IC), fluorescence and intersystem crossing (ISC) to effectively compete with ‘free’ O–H dissociation.\textsuperscript{15,32,33} The absence of 4-TBC-$d_2$ out to $\Delta t = 10 \text{ ns}$ in our deuterated studies supports this idea; O–D bond fission is kinetically outcompeted by other pathways, which is unsurprising given that D-atom tunneling through the barrier beneath an $S_1/S_0$ CI will be significantly suppressed. As such, we see an increase in the triplet state signal.

On the basis of our discussion above, a proposed schematic summarizing the various decay pathways in 4-TBC, both in perturbative and non-perturbative environments, is given in Figure 4. Photoexcitation with 267 nm radiation, prepares 4-TBC in the $S_0$ state. In the gas-phase, the excited state decays through dissociation of the ‘free’ O–H bond, along a pathway that has a barrier. The KIE of ~19 strongly suggests that O–H dissociation is mediated through tunneling beneath the $S_1/S_0$ CI. In the weakly perturbing cyclohexane solvent, the photoexcited $S_1$ state decay appears to follow a similar path: tunneling mediated O–H dissociation. While a milder KIE ~7 makes this supposition less conclusive, it is still strongly supportive of a barrier to O–H dissociation coordinate. The left side of Figure 4 contains a single, dominant process involved in the non-perturbative, gas-phase and weakly perturbative, (cyclohexane) solution-phase: $k_S$ signifies the rate constant of O–H dissociation. In the strongly perturbing acetonitrile solvent, we have contrasting dynamics. The ‘open’ conformer, which now dominates, displays a diminished 4-TBC$^*$ feature. Upon deuteration, the 4-TBC-$d_2$ absorption signature is almost completely extinguished. Instead, the excited state decay reveals an absorption feature in the TAS that is reminiscent of triplet state absorption. As a consequence, we propose that excited state decay in 4-TBC (acetonitrile) is multicomponent, including O–H dissociation, ISC and likely IC and fluorescence, characterized by associated rate constants $k_{\text{IC}}, k_{\text{ISC}}, k_{\text{IC}}$ and $k_F$ respectively. The right side of Figure 4 shows this process.

We close by returning to the original question regarding whether the natural surroundings impact the photostabiliy of biomolecules. In 4-TBC, a close analogue to the catechol chromophore, it is clear that the photoinduced dynamics in non-perturbing environments compare favorably with those in the gas-phase; both cases involve O–H dissociation to yield the phototoxic radical species 4-TBC$^*$.\textsuperscript{54} When 4-TBC is placed into a polar solvent, this pathway is severely suppressed. This is a consequence of a change in molecular geometry through solvent interactions. This structural change reduces the excited state decay, with the phototoxic radical pathway becoming less favorable, and opens up alternative (and competitive) relaxation pathways, including potentially harmful triplet state formation.\textsuperscript{29} Interestingly, the triplet state formation accords with previous studies by Sundström and coworkers on 5,6-dihydroxynindole,\textsuperscript{28} one of the main building blocks of eumelanin, which incorporates the catechol chromophore. Thus, the present bottom-up study provides a key steppingstone between a model UV chromophore in the gas phase and a larger biomolecule within its
native environment. This emphasizes the critical role of environment-induced photodynamics.

IV. CONCLUSIONS

The excited state dynamics of 4-TBC have been investigated in both gas- and solution-phase using a combination of TR-VOI, TR-IY and TEAS, following photoexcitation of the S1 (~ππ*) state at 267 nm. Through the recorded gas-phase TKER spectra and associated H+ transient, a time constant for H-atom elimination of τH = 5.9 ± 0.3 ps is found. Complementary TR-IY of the parent transient yields a time constant of τP = 4.9 ± 0.6 ps. Comparative studies in 4-TBC d2 yield time constants of τH = 98 ± 2 ps and τDP = 102 ± 3 ps and thus an average KIE of ~19. A dominant dissociation of the non-intramolecular hydrogen bonded ‘free’ O–H bond is deduced, very likely mediated through tunneling beneath an S1/S2 CI. Solution-phase studies in the weakly perturbing cyclohexane solvent yield excited state decay time constants of τH = 27 ± 7 ps and τDP = 190 ± 20 ps for 4-TBC and 4-TBC-d2 respectively. The KIE of ~7 implies, together with the emergent 4-TBC-d2 feature, that a similar relaxation pathway in 4-TBC/cyclohexane is operative as in 4-TBC in the gas-phase. In the strongly perturbing, polar acetonitrile solvent, excited state decay occurs with time constants of τH = 1.7 ± 0.1 ns and τDP = 3.3 ± 0.1 ns for 4-TBC and 4-TBC-d2 respectively. The dominant decay channel in acetonitrile is no longer O–H dissociation, but rather multicomponent, involving ISC, IC and fluorescence together with O–H dissociation. The dramatic differences in excited state lifetime in the two solvents, is attributed to conformational change between 4-TBC (and 4-TBC-d2) in cyclohexane and acetonitrile, in which, respectively, the closed and open conformers dominate. This result thus serves to highlight the critical importance of the influence of structure on dynamics.

ASSOCIATED CONTENT

Supporting Information
Static absorption spectra, additional experimental details, fitting procedures for presented transients, deuterated TAS, solvent-only transients and further computational details.

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Notes
The authors declare no competing financial interest.

ACKNOWLEDGMENTS

The authors are grateful to Ms Faye Monk, Mr Connor Galagher and Mr Peter Brindley for experimental assistance and Dr Gareth Roberts (University of Bristol) for helpful discussions. M.D.H. and J.D.Y. thank the University of Warwick for EPSRC doctoral training awards. L.A.B., W.D.Q. thank the EPSRC for providing studentships under grant number EP/F500378/1, through the Molecular Organisation and Assembly in Cells Doctoral Training Centre. M.S. and S.E.G thank the EPSRC and the Warwick Institute of Advanced Studies, respectively, for postdoctoral funding. M.S. also thanks Prof. Martin Paterson (Heriot Watt) for use of his computational facilities. V.G.S. thanks the EPSRC for an equipment grant (EP/I007153) and the Royal Society for a University Research Fellowship.

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