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Multiconfigurational photodynamics simulations reveal the mechanism of photodecarbonylations of cyclopropenones in explicit aqueous environments†

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Gas-evolving photochemical reactions use light and mild conditions to access strained organic compounds irreversibly. Cyclopropenones are a class of light-responsive molecules used in bioorthogonal photoclick reactions; their excited-state decarbonylation reaction mechanisms are misunderstood due to their ultrafast (<100 femtosecond) lifetimes. We have combined multiconfigurational quantum mechanical (QM) calculations and non-adiabatic molecular dynamics (NAMD) simulations to uncover the excited-state mechanism of cyclopropenone and a photoprotected cyclooctyne-(COT)-precursor in gaseous and explicit aqueous environments. We explore the role of H-bonding with fully quantum mechanical explicitly solvated NAMD simulations for the decarbonylation reaction. The cyclopropenones pass through asynchronous conical intersections and have dynamically concerted photodecarbonylation mechanisms. The COT-precursor has a higher quantum yield of 55% than cyclopropenone (28%) because these trajectories prefer to break a σ_{CC} bond to avoid the strained *trans*-cyclooctene geometries. Our solvated simulations show an increased quantum yield (58%) for the systems studied here.

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Introduction

Photochemistry is an effective means by which strained, energy-dense compounds can be accessed while facilitating green chemistry through mild reaction conditions. Gas-evolving reactions produce gases such as oxygen or carbon dioxide (CO₂) and are utilized for processes such as the oxygen reduction reaction (ORR) and CO₂ reduction.^{1–4} Photochemical decarbonylation has been utilized to promote cross-coupling and form biaryl compounds,⁵ promote synthetic pathways towards strained compounds such as tetrahedrane,^{6,7} and facilitate transition-metal-catalyzed C–C bond cleavage.^{1,2,8} Small concentrations of CO have been shown to have

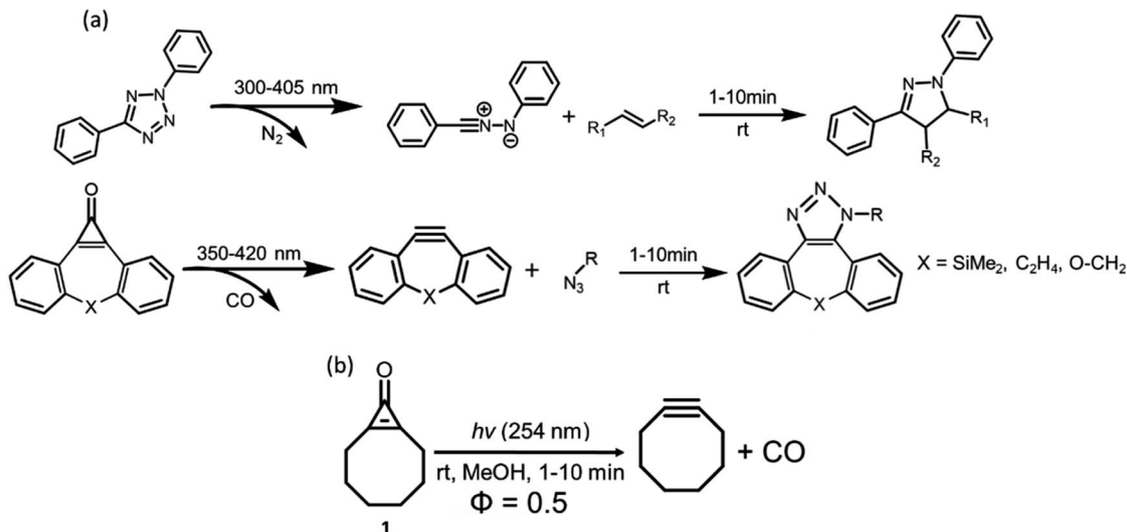
therapeutic effects, including it being used as an anti-inflammatory agent,⁹ and used to treat cancer and bacterial infections.^{9–12} CO delivery can be activated through irradiation of a photoactive CO-releasing molecule (CORM).^{13,14} Light enables spatiotemporal activation of different chemical processes, including energy storage in materials chemistry^{15–26} and pro-drugs in drug delivery.^{27–29} Reactive cycloaddends (*e.g.*, cyclooctynes and 1,3-dipoles) can be accessed using light through a class of bioorthogonal reactions known as ‘photoclick chemistry.’ Photoclick reactions require light-promoted activation to access the reactive species and a thermal strain-promoted cycloaddition reaction in the biochemical milieu. Strain-promoted and distortion-accelerated³⁰ cycloadditions can occur without Cu(I) catalysts and benefit from high reaction rates.³¹ The first photoclick reaction was reported by Lin and co-workers in 2008 and involved the tetrazole-ene cycloaddition reaction.^{32–34} Upon irradiation, N₂ is photochemically extruded from tetrazole and the remaining 1,3-dipole, nitrile imine can undergo reactions with 2 π -cycloaddends. Other photoclick reactants include photoprotected cyclooctynes,^{35–42} syndones,⁴³ azirines,^{34,43–46} Diels Alder reactions,^{47–53} and thiol click reactions.^{54–59} Scheme 1 illustrates two examples of photoclick reactions reported by Song, Poloukhine, and Wang.^{34,37,38,44,45,60}

Functionalizing the vinyl positions of **1** with aryl groups redshifts the absorption wavelength to activate decarbonylation at 350 nm. The quantum yield (QY) of 4-dibenzocyclooctynol after irradiation with 350 nm light was measured to be 0.33 by

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† Electronic supplementary information (ESI) available: The ESI contains the (10,9) active space for **1** and an analysis and visualization of the branching plane for an MECI structure of **2** in the FC-region of the S₁/S₀ hopping region. We also present two concerted hopping point structures for **1**, a bond length analysis of the hopping points for the NAMD simulations of **1** and **2** in the gas-phase, and an analysis of the S₁ lifetimes for **1** and **2** for the gas-phase and solvated dynamics trajectories. We also tested two different active spaces, presented a conformational analysis of **1**, and compared SA(2) NAMD simulations of **2** to SA(4) NAMD simulations. Finally, we present QM/MM NAMD simulations for solvated systems of **1** and **2**. All optimized structures and NAMD simulations are available online (DOI: <https://doi.org/10.6084/m9.figshare.21667808>). See DOI: <https://doi.org/10.1039/d3sc03805j>





Scheme 1 Examples of different photoclick reactions including tetrazole denitrogenation and cyclopropanone photodecarbonylation toward a strained alkyne (a). Photochemical decarbonylation reaction towards carbon monoxide and cyclooctyne from a cyclooctyne pre-cursor (**1**), (b).

Popik and co-workers.⁶⁰ The substituent effects on both cyclopropanone (**2**) and cyclooctynol were also reported by Popik and co-workers, who used multi-photon absorption with a near-IR laser to generate reactive intermediates to undergo a strain-promoted alkyne-azide cycloaddition (SPAAC) reaction (*i.e.*, photoclick reaction).^{61–63} Recently, Kunishima and co-workers have used a visible-light responsive photocatalyst to redshift the absorption wavelength required for the decarbonylation of substituted cyclopropanones.^{64,65}

While the thermal cycloadditions involved in bio-orthogonal chemistry have been intensely researched by experimentalists^{36,66–68} and theorists,^{69–72} the photochemical steps of the reactions have received less attention. We hypothesize this is due to the challenge of experimentally studying molecular excited states and reactive intermediates in the typically short timescale of photochemical reactions (10^{-13} to 10^{-8} seconds). This timescale is too short for the atomistic resolution of excited-state geometries. Fortunately, multi-configurational NAMD simulations offer high-fidelity structural information about the rapidly interconverting excited-state structures and mechanisms of organic photochemical reactions.

Computational approaches provide valuable structural information of **2** on the 1–10 picosecond timescale. Fang and co-workers investigated the photochemical decarbonylation of **2** in 2014 using gas phase multiconfigurational calculations (CASSCF and multistate-CASPT2) to mechanistic critical points on the S₀ and S₁ potential energy surfaces (PESs).⁷³ *Ab initio* multiple spawning (AIMS)⁷⁴ non-adiabatic molecular dynamics (NAMD) simulations were performed to investigate the photochemical reaction dynamics along the S₁ surface. The authors located one S₁/S₀ conical intersection, leading to the dissociation mechanism in 72% of trajectories. The authors also located a second S₁/S₀ conical intersection that reverts to the reactant (cyclopropanone) in 93% of trajectories. Fang reports an S₁ lifetime of 125 fs for **2**. This past study only simulated

cyclopropanone in the gas phase. To best represent biological environments, the simulation conditions should be as close to the experimental conditions as possible (*i.e.*, replicating solvent and temperature). We now compute the vertical excitation energies, absorption spectra, photochemical mechanism, and quantum yields for **1** and cyclopropanone (**2**). The NAMD simulations are performed with multiconfigurational quantum chemical calculations in the gas phase and explicitly solvated aqueous environments. We compare our results with those presented in the 2003 work by Poloukhine and Popik.⁷⁵

Results and discussion

Cyclopropanone is the prototypical pseudo-aromatic molecule on the ground state. We first computed the ground state geometries of **1**, **2**, and cyclopropanone (**3**) with density functional theory (DFT) calculations (Fig. 1).

We first compare the π_{CO} bondlengths of the pseudo-aromatic cyclopropanones (**1-S₀**, **2-S₀**); they are nearly identical (1.20 and 1.21 Å, respectively). The **1-S₀** and **2-S₀** σ_{CC} bonds are 1.44 Å in **1-S₀** and **2-S₀**. The relatively short σ_{CC} bonds in **1-S₀** and **2-S₀** is consistent with conjugated π_{CC} and π_{CO} . **3-S₀** is non-aromatic and has notably longer σ_{CC} bonds (1.47 and 1.56 Å); the π_{CO} bondlength is 1.20 Å. Next, we computed the excitation energies and electronic transitions of **1** and **2** with time-dependent density functional theory (TD-DFT) and multi-configurational complete active space self-consistent field (CASSCF). We used extended multi-state complete active space second-order perturbation theory (XMS-CASPT2) to account for dynamic correlation of the electronic states.⁷⁶ Our TD-DFT calculations included the lowest 10 singlet excited states and used the range-separated density functional CAM-B3LYP⁷⁷ with the triple- ζ aug-cc-pVTZ^{78,79} basis set. We included 10 electrons and 9 orbitals in the CASSCF active space for **1** and **2**; Fig. 2 shows the orbitals and average orbital occupancies for **2**. The same active space was used for **1**.



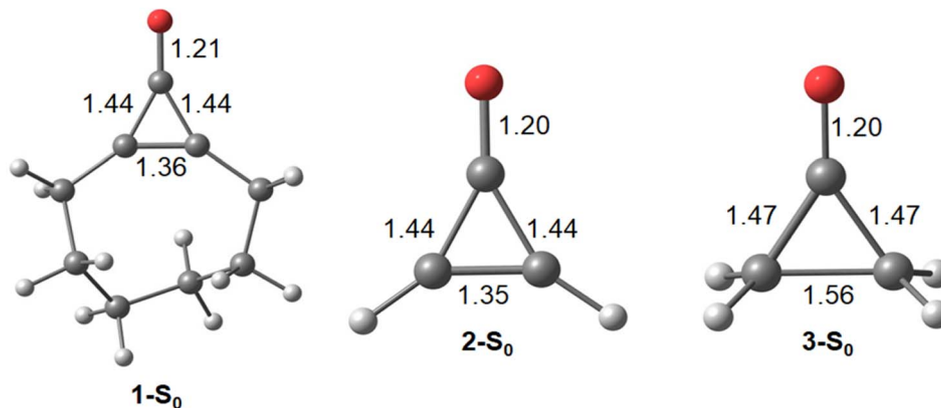


Fig. 1 Optimized S_0 global minimum geometries for 1- S_0 , 2- S_0 , and 3- S_0 . Bond lengths are presented in Angstroms (Å). All structures were optimized using PBE0-D3BJ/cc-pVDZ.

The $S_0 \rightarrow S_1$ excitation energies computed with XMS-CASPT2(10,9)/ANO-S-VDZP for **1** and **2** are 4.42 and 4.52 eV, respectively. Both transitions are $n \rightarrow \pi^*$ and have oscillator strengths of 0.0094 (**1**) and 0.0069 (**2**). The $S_0 \rightarrow S_2$ excitations have energies of 5.00 eV (**1**) and 5.68 eV (**2**); both are $\pi \rightarrow \pi^*$ transitions. The oscillator strengths of these transitions are 0 for **1** and **2**. The $S_0 \rightarrow S_3$ electronic excitations of **1** and **2** are $\pi \rightarrow \pi^*$ transitions, with excitation energies and oscillator strengths of 7.42 eV and 7.14 eV and 0.07 and 0.05, respectively. We generated an ensemble of 500 Wigner-sampled non-equilibrium geometries, computed the $S_0 \rightarrow S_n$ ($n = 1,2,3$) vertical excitation energies, and oscillator strengths to generate an absorption spectrum for **1** and **2**. Fig. 3 shows an overlay of select non-equilibrium geometries for **1** and **2** and the absorption spectrum; the intensities are normalized for the $S_0 \rightarrow S_3$ transition.

The absorption spectra for **1** and **2** show three λ^{\max} peaks centered at 190, 220, and 260 nm. The peak centered at 190 nm corresponds to a $\pi \rightarrow \pi^*$ transition; the λ^{\max} peaks centered at 220 and 260 nm correspond to $n \rightarrow \pi^*$ transitions. The λ^{\max} peaks near 190 nm correspond to $S_0 \rightarrow S_3$ excitations and have the highest intensity because they are 'allowed' transitions. The $n \rightarrow \pi^*$ transition at 220 nm corresponds to an $S_0 \rightarrow S_2$ excitation and has a relatively low intensity. The $S_0 \rightarrow S_1$ excitation for **1** and **2** ($n \rightarrow \pi^*$) has a peak centered near 260 nm. While the intensity of this peak (shown in red) is substantially lower than that of the $\pi \rightarrow \pi^*$ transition, it is sufficiently low in energy to undergo photoexcitation with the experimental light source (254 nm). The peaks corresponding to the second- and third-lowest excited states (shown in blue and yellow, respectively) correspond to excitation energies higher than the experimental light source. As such, we decided to focus on the S_1 -state to

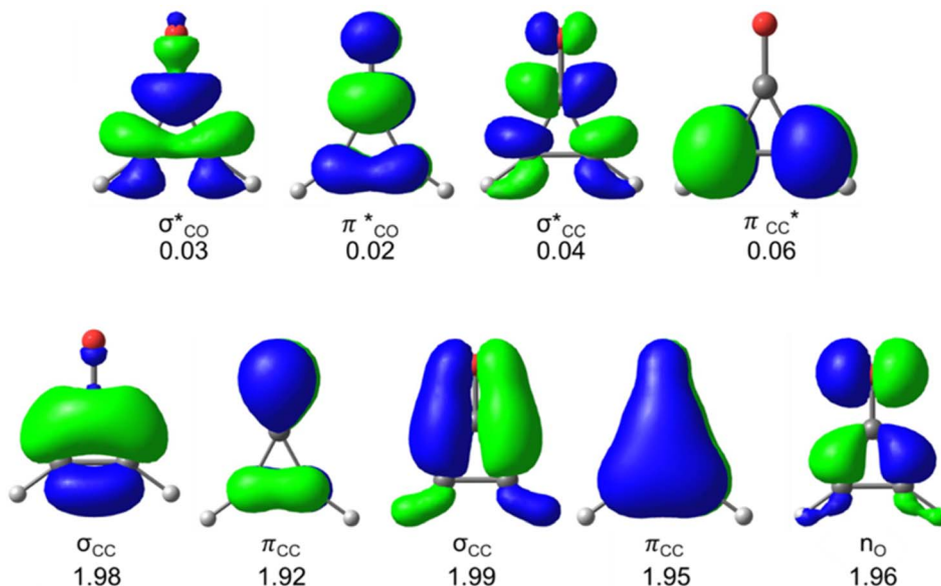


Fig. 2 CASSCF(10,9) active space of **2** with average electron occupancies. Orbitals are shown with an isosurface value of 0.06 and computed with SA(4)-CASSCF(10,9)/ANO-S-VDZP. The excitations, oscillator strengths, and electronic transition characters for **1** and **2** were computed with SA(4)-CASSCF(10,9)/ANO-VDZP, SA(4)-XMS-CASPT2(10,9)/ANO-S-VDZP, and TD-DFT (Tables 1 and 2).



Table 1 Vertical excitation energies, oscillator strengths, and electronic transition characters for the first three excited states of **1** with TD-DFT, CASSCF, and XMS-CASPT2. All energies given in eV. All calculations were run in the gas-phase on the global minimum

Method	State	Energy (eV)	Wavelength (nm)	Oscillator strength	Nature
CAM-B3LYP/aug-cc-pVTZ	S ₁	4.80	258	0.0008	n _O → π* (S ₁)
	S ₂	5.42	229	0.0001	n _O → π* (S ₂)
	S ₃	5.58	222	0.0002	π → Ryd (S ₃)
XMS-CASPT2(10,9)/ANO-S-VDZP	S ₁	4.93	281		n _O → π* (S ₁)
	S ₂	5.66	248		n _O → π* (S ₂)
	S ₃	7.21	167		π → π* (S ₃)
SA(4)-CASSCF(10,9)/ANO-S-VDZP	S ₁	5.13	242	0.0086	n _O → π* (S ₁)
	S ₂	5.75	216	0.0000	n _O → π* (S ₂)
	S ₃	7.38	168	0.0690	π → π* (S ₃)

explore the static and dynamic components of the photo-decarbonylation reaction mechanism of **1** and **2**.

Cycloproponone photodissociation mechanism

We first performed a minimum energy path (MEP) calculation along the S₁-state to determine the dominant mechanistic pathway from the FC-point for **1** and **2**. Fig. 4a and b show the MEP, S₀- and S₁-minimum geometries, and the final structure of the MEP for **1** and **2**.

Fig. 4a shows the MEP along S₁-surface and the energies for the corresponding S_{0–3} states; it contains 19 geometries. This MEP demonstrates that the S₀ and S₁ states approach a degeneracy by the 19th step (S₁–S₀ energy gap of 0.33 eV) for **1**. The steps after the FC-point reflect the geometrical changes resulting from the *n* → π* electronic transition. We defined a torsional angle parameter, θ , across C₁–C₂–C₃–C₄ for **1** and **2** (atoms labels are given in Fig. 4). **1**-S₀ and **2**-S₀ are nearly planar ($\theta = 3$ and 0°, respectively). θ increases from $\theta_0 = 3^\circ$ to $\theta_{25} = 80^\circ$. The π_{C₂–C₃} bond length increases from 1.35 Å to 1.43 Å, and C₅ is visibly pyramidalized in **1**-MEP-**19** (O–C₅–C₂ angle of 169°). The pyramidalization of C₅ is accompanied by a shortening of the C₅=O bond (1.18 to 1.16 Å), which resembles a C₅≡O triple bond. The C₅–C₃ bond is nearly broken in **1**-MEP-**19** (2.08 Å), suggesting different mechanisms for **1** and **2**.

The S₁–S₀ energy gap at the end of MEP calculations can suggest the dominant reaction mechanism in photochemical reactions. Large energy gaps suggest an S₁-minimum and radiative decay to S₀; small energy gaps suggest a degeneracy

region, indicating a possible productive pathway to photo-products. Unlike the large S₁–S₀ energy gap observed in **2**-MEP-**6**, **1**-MEP-**19** has a relatively small S₁–S₀ energy gap (0.33 eV). Prompted us to locate an S₁/S₀ minimum energy conical intersection (**1**-MECI). **1**-MECI is 2.43 eV above **2**-S₀ and has C₃–C₅ and C₂–C₅ bond lengths of 2.24 and 1.49 Å, respectively. Although there is a broken C–C bond in the cycloproponone ring, the alkene carbons (C₂ and C₃) are slightly pyramidalized ($\theta = 10^\circ$).

Fig. 4b shows the MEP along the S_{1–3} surfaces for **2**. The MEP includes 6 steps and leads to a final structure (**2**-MEP-**6**) with an S₁–S₀ energy gap of 1.46 eV, substantially larger than that of **1**-MEP-**19** (S₁–S₀ energy gap of 0.33 eV). The large S₁–S₀ energy gap of **2**-MEP-**6** suggests that the pseudo-dominant mechanistic pathway for **2** is towards an S₁-minimum. We optimized an S₁ minimum using the final MEP geometry as an input (shown in Fig. 4b). **2**-S₁ is 3.22 eV above **2**-S₀ and nearly degenerate with **2**-MEP-**6**. The π_{CC} of **2**-S₁ is pyramidalized, and the vinyl-hydrogens adopt a *trans*-relationship.

The S₁-MEPs of **1** and **2** lead to a conical intersection and an S₁-minimum, respectively. While both molecules are theoretically capable of undergoing *cis*–*trans* isomerization, this reaction is highly disfavored in **1** because of the resulting *trans*-cyclooctene, which is highly strained. *Trans*-cyclooctene is higher in energy than *cis*-cyclooctene by 16 kcal mol^{–1}.⁸⁰ **2**, however, avoids the additional strain energy and undergoes the *cis*–*trans* isomerization to **2**-S₁. Although MEP calculations provide mechanistic information, they omit dynamical effects,

Table 2 Vertical excitation energies, oscillator strengths, and electronic transitions for the first three excited states of **2** at TD-DFT, CASSCF, and XMS-CASPT2 levels of theory. All energies are given in eV

Method	State	Energy (eV)	Wavelength (nm)	Oscillator strength	Nature
CAM-B3LYP/aug-cc-pVTZ	S ₁	4.36	284	0.0004	n _O → π* (S ₁)
	S ₂	5.46	227	0.0000	n _O → π* (S ₂)
	S ₃	6.20	200	0.0062	π → Ryd (S ₃)
XMS-CASPT2(10,9)/ANO-S-VDZP	S ₁	4.52	274		n _O → π* (S ₁)
	S ₂	5.68	218		n _O → π* (S ₂)
	S ₃	7.14	174		π → π* (S ₃)
SA(4)-CASSCF(10,9)/ANO-S-VDZP	S ₁	4.61	269	0.0069	n _O → π* (S ₁)
	S ₂	5.70	218	0.0000	n _O → π* (S ₂)
	S ₃	6.95	178	0.0484	π → π* (S ₃)



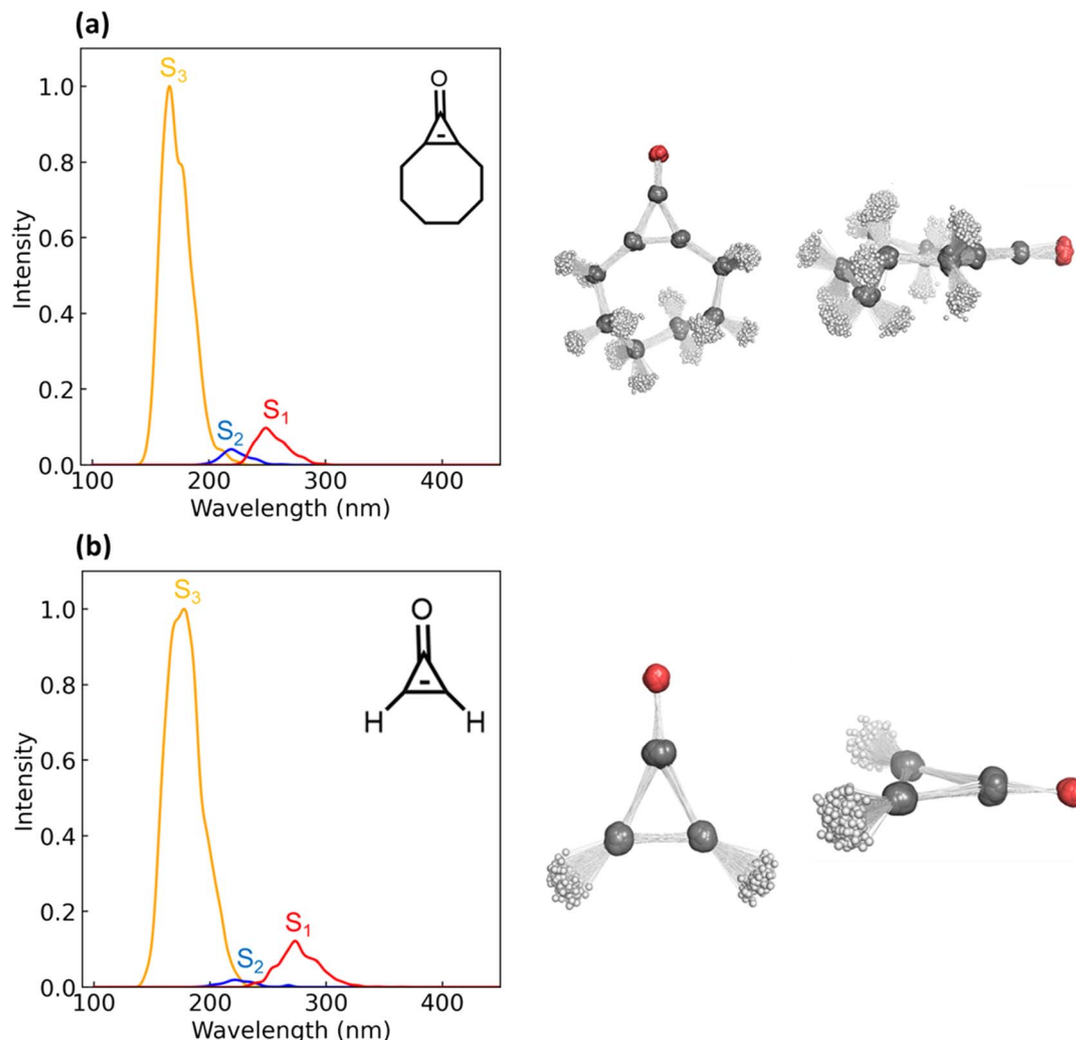


Fig. 3 Absorption spectra and geometric overlays corresponding to Wigner sampled geometries of **1** (a) and **2** (b). Both spectra were computed in the gas-phase using SA(4)-CASSCF(10,9)/ANO-S-VDZP, and the Wigner sampled geometries were generated from using the vibrational frequencies of a ground-state geometry optimized with SA(4)-CASSCF(10,9)/ANO-S-VDZP.

which control photochemical reaction mechanisms and quantum yields. As such, we performed NAMD simulations starting from the S_1 FC region for **1** and **2** to enumerate all possible mechanistic pathways.

Non-adiabatic molecular dynamics simulations

We performed 1 picosecond (ps) NAMD simulations using the FSSH algorithm^{81,82} on **1** and **2**, using Wigner-sampled initial conditions. For **1**, 497 of the trajectories crossed to the S_0 after 1 ps (99%), while 3 (1%) remained on the S_1 . Of the 497 trajectories that passed through S_1/S_0 hopping points, 274 underwent decarbonylation ($\Phi_{\text{comp}} = 55\%$). The remaining 223 trajectories (45%) reverted to **1**. For **2**, 464 trajectories crossed to the ground state after 1 ps. 130 of the 464 trajectories corresponded to the decarbonylation reaction ($\Phi_{\text{comp}} = 28\%$), while 334 trajectories (72%) reverted to cyclopropanone. We plotted the two σ_{CC} bond lengths in **1** and **2** to determine whether the reaction was concerted or stepwise. Fig. 5 shows these bond lengths

throughout the 1 ps trajectories; the black dots correspond to the S_1/S_0 hopping points.

Fig. 5 shows the σ_{CC} distances throughout each of the trajectories. Examples of hopping point structures from both seam regions are shown in Fig. 5c and d. Both the asynchronous hopping point geometries (**1-as** and **2-as**) contain one σ_{CC} bond in the cyclopropanone ring that is fully broken (2.31 Å and 2.36 Å, respectively). In contrast, the other cyclopropanone σ_{CC} bond is lengthened from **1-S₀** (1.57 and 1.41 Å, respectively). All trajectories with σ_{CC} bonds exceeding 3.3 Å were considered dissociated. The QYs for the dissociation reaction of **1** and **2** are 55% and 28%, respectively. Optimized MECI structures for **1** and **2** are shown as yellow points on both plots, and the black dots represent S_1/S_0 hopping points. The S_1/S_0 hopping points for **1** and **2** have a large range of σ_{CC} bond distances (1.36–2.77 Å). We grouped the hopping points structures based on their structural similarities. The blue region corresponds to hopping points with at least one $\sigma_{\text{CC}} > 1.80$ Å and are labeled



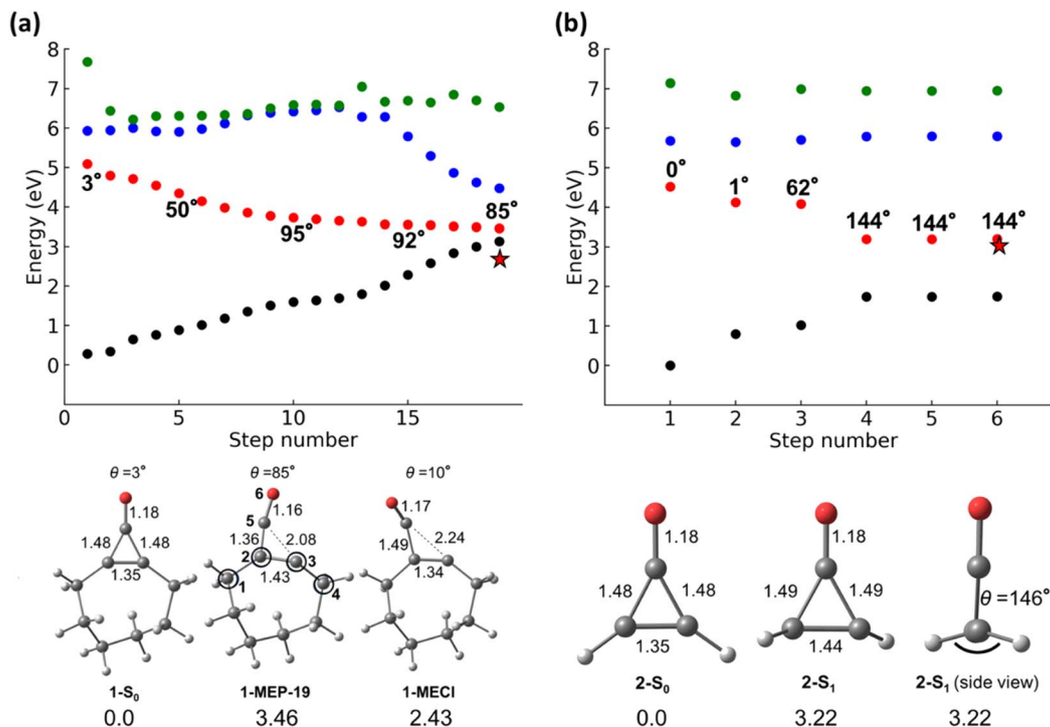


Fig. 4 Minimum energy path calculations for **1** (a) and **2** (b). The dots on the graphs correspond to the S_0 (black), S_1 (red), S_2 (blue), and S_3 (green) states. Relevant structures (S_0 and S_1 minima, MECI structures, and endpoints of the MEP calculations) are shown below the plots. **1-MECI** (left) and **2-S₁** (right) are shown on the plots with red stars. For both optimized points, the final MEP geometry from the calculation was used as the initial guess structure. The MEP calculations were run with SA(4)-CASSCF(10,9)/ANO-S-VDZP, and SA(4)-XMSCASPT2(10,9) single-point energy corrections were performed on each MEP step. All optimized structures were performed with SA(4)-CASSCF(10,9)/ANO-S-VDZP with SA(4)-XMS-CASPT2(10,9) energy corrections.

asynchronous because the breaking σ_{CC} distances are different by 0.5 Å. The hopping points in the red region correspond to structures with both σ_{CC} bonds <1.80 Å (labeled FC-region).

We computed the branching ratios for asynchronous and FC-region hopping points to determine if the hopping points' structures influenced the QYs of each reaction. Only 87 (18%) of the 497 NAMD trajectories of **1** passed through a FC-region hopping point. All 87 trajectories that passed through a hopping point in the FC-region reformed **1**. The remaining 410 (82%) NAMD trajectories passed through asynchronous hopping points. The strain energy associated with *trans*-cyclooctene geometries redirects many of the trajectories (82%) away from the FC-region conical intersection seam and towards through more productive asynchronous conical intersection seam. These hopping points in the asynchronous region 274 (67%) dissociate, while 136 (33%) reformed the reactant. 269 (58%) of the 464 trajectories of **2** passed through a FC-region hopping point, and 195 (42%) passed through asynchronous hopping points. Of those 269 FC-region hopping points, 262 hopping points revert to the reactant, and only 7 trajectories dissociate to the product. 123 of the trajectories (63%) from the asynchronous region undergo photo-decarbonylation dissociate, and 72 trajectories (37%) reform the reactant. For **1** and **2**, there is an overwhelming preference to reform the reactant after surface hopping occurs in the FC-region, and a $\sim 2 : 1$ preference to dissociate after surface hopping occurs in the asynchronous region. We hypothesize that the preference for reactant

reformation is due to the energy required to break the cyclopropanone ring structure on the ground state. We also present the branching space for an MECI optimized along the FC-region of the S_1/S_0 hopping seam in the ESI.† For trajectories that hop in the asynchronous region, there is a slight preference for breaking the second C–C bond on the S_0 state after hopping, as these bonds are already extended in the excited state.

The θ values of **1-fc** and **1-as** are 95° and 74°, respectively. This hopping point preference for **1** is nearly double that of **2** (42%). This agrees with the MEP results in Fig. 4, which suggest a direct path to an asynchronous S_1/S_0 hopping point (one broken σ_{CC} bond in the cyclopropanone ring). We find an overwhelming preference for a non-productive reaction outcome from hopping points in the FC-region (100% and 97% for **1** and **2**, respectively). Our interest in understanding **2** as a bioorthogonal photoclick reagent drove us to understand the role of solvation on the QYs and mechanisms. To that end, we generated solvated initial conditions for **1** and **2** and ran 250 fs NAMD simulations from their respective S_1 FC-regions.

Explicitly solvated NAMD simulations

We used an explicitly solvated fully quantum mechanical NAMD simulations to determine the photodecarbonylation mechanism of **1** and **2** because multiscale QM/MM simulations cannot capture the electronic effect of intermolecular hydrogen bonding between the water molecules and the cyclopropanones. Historically, dynamics simulations involving solvent utilize



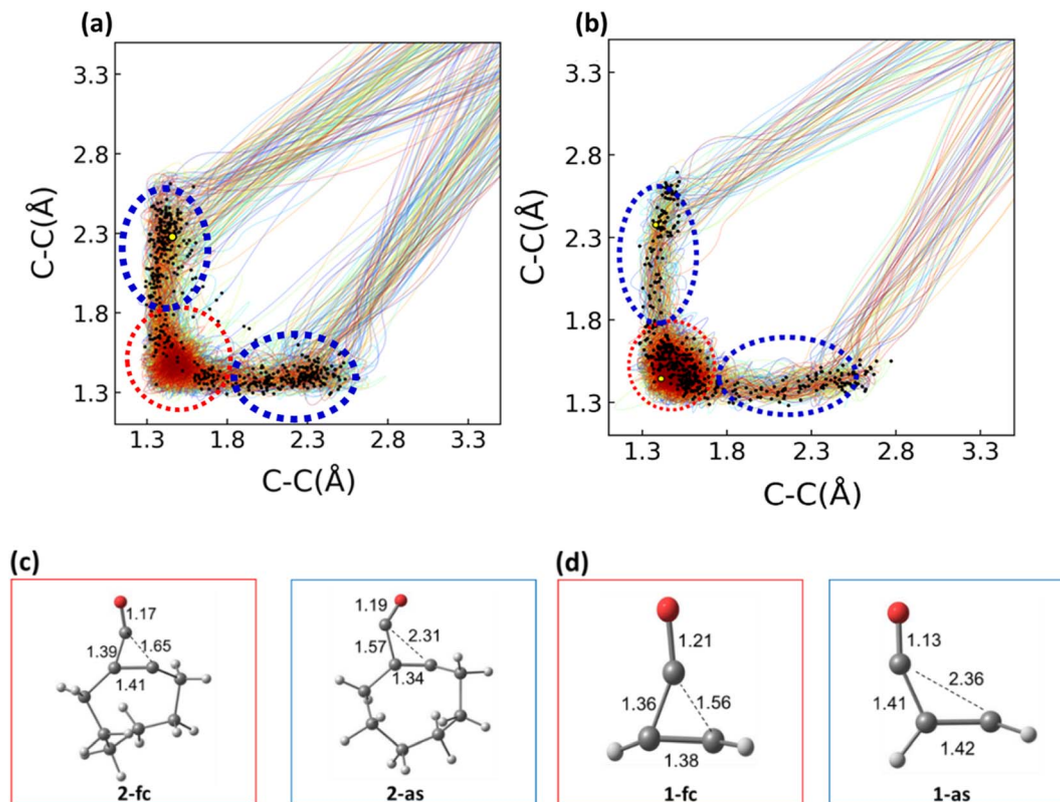


Fig. 5 C–C bonds plotted against each other for **1** (a) and **2** (b). The bond lengths we calculated are depicted by red dotted lines. The solid multicolor lines on each plot show the bond lengths over time, and the black dots represent the S_1/S_0 surface hopping points for each trajectory. The minimum energy conical intersections (MECIs) are marked on the plots with a yellow point. Representative hopping points are given for the reactant (red) and asynchronous (blue) regions for **1** (c) and **2** (d). Hopping points from the FC region have the suffix 'fc' and hopping points from the asynchronous region have the suffix 'as'. NAMD simulations were run with SA(4)-CASSCF(10,9)/ANO-S-VDZP. 500 initial conditions were sampled by an S_0 Wigner distribution. Trajectory simulations were run for 1 ps with a 0.5 fs timestep. The hopping points in (c) and (d) are snapshots obtained directly from the NAMD simulations.

a multiscale approach where the chromophore is treated with QM method, and the surrounding environment is treated using MM. QM/MM approaches are particularly useful for calculating large systems, which generally include proteins and solvent molecules. To compare to the explicitly solvated all-QM method, we used a QM/MM approach for simulating the photochemical pathways along the S_1 for **1** and **2**; the presentation and discussion of these results are given in the ESI.† Initial conditions were sampled in line with the report by Gonzalez and co-workers last year.⁸³ We observed an increased quantum yield for the decarbonylation reaction for both systems compared to the gas-phase simulations (58% for **1** and **2**).

We performed 250 fs NAMD simulations on solvated spheres (20 water molecules) of **1** and **2**, starting with initial conditions sampled using MD simulations in CP2K. The NAMD simulations utilized the FSSH surface hopping algorithm and were propagated from the S_1 state. The σ_{CC} bond lengths in the cyclopropanone ring were measured throughout the trajectories and are represented in Fig. 6.

The inclusion of an explicit solvent in our NAMD simulations resulted in the Φ_{comp} for the dissociation products increasing from 53% to 58% for **1**, and from 28% to 58% for **2**. This increased yield can be explained by the ratio of trajectories

going through the asynchronous and FC regions of the S_1/S_0 seam. For the solvated NAMD simulations of **1**, nearly all trajectories crossed to the S_0 in the asynchronous region (94%), corresponding to a 12% increase compared to the gas-phase simulations. While the decarbonylation yield from the asynchronous region is smaller than for the gas-phase simulations (48% and 64%, respectively), the overwhelming preference for the asynchronous region ultimately leads to a higher quantum yield. From the FC-region of **1**, there was a larger number of trajectories (9) that dissociated than those that reformed the reactant (**4**), which was significantly different from the gas phase. For **2**, the number of trajectories hopping in the asynchronous region increased for the solvated and gas-phase systems (70% and 52%, respectively); the quantum yield from the asynchronous region was slightly smaller for the solvated system compared to the gas-phase (59% and 63%, respectively), the higher ratio of trajectories crossing through this region leads to a higher Φ_{comp} . To understand the solvent effect on the reactivities of **1** and **2**, we hypothesized that hydrogen bonding intermolecular interactions between the solvent and the chromophores were important in the S_0 , S_1 -surface, and conical intersection seams.



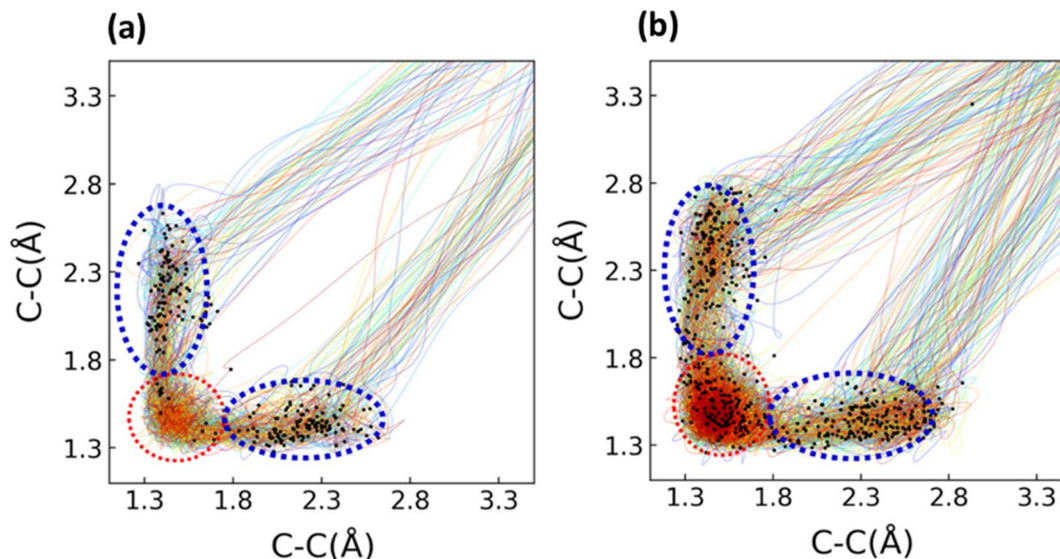


Fig. 6 Trajectory plots of **1** (a) and **2** (b) were solvated with 20 water molecules. The bonds being measured over are the breaking C–C σ bonds in the cyclopropanone ring. The NAMD simulations were run using SA(4)-CASSCF(10,9). The ANO-S-VDZP basis set was used for the chromophores **1** and **2**, and the ANO-S-MB was used for the water molecules. The S_1/S_0 crossing seams for the solvated systems of **1** and **2** are indicated with black dots. The distribution of S_1/S_0 hopping points for **1** and **2** have a large range of σ_{CC} bond distances (1.42–3.25 Å). For **1**, there were 276 initial conditions propagated from the S_1 . Of these 276, 253 crossed to the S_0 by the end of the simulation. Of these 253, there were 19 trajectories where the endpoint was an intermediate structure on the ground state (one σ_{CC} bond in the cyclopropanone ring >1.80 Å and the other σ_{CC} bond in the cyclopropanone ring <1.60 Å). These intermediate structures were not counted in the final quantum yield. An example intermediate structure is provided in the ESI.† Of the remaining 234 trajectories, we observed 135 trajectories that underwent decarbonylation ($\Phi_{\text{comp}} = 58\%$) and 99 that reformed the reactant (42%). For the solvated system of **2**, we ran simulations on 481 initial conditions. Of these 481, 20 trajectories (4%) ended at an intermediate structure on the ground state (one σ_{CC} bond in the cyclopropanone ring >1.80 Å and the other σ_{CC} bond in the cyclopropanone ring <1.60 Å). 277 trajectories underwent decarbonylation ($\Phi_{\text{comp}} = 58\%$), and 173 trajectories (36%) reformed the reactant. We also calculated the ratio of trajectories for **1** and **2** that crossed through the asynchronous (one σ_{CC} bond of 1.80 Å or longer) and FC-regions (both σ_{CC} bonds less than 1.80 Å) of the S_1/S_0 crossing seam. For the solvated system of **1**, 237 trajectories (94%) went through a hopping point in the asynchronous region, while 16 (6%) crossed at a hopping point in the FC-region. There were 114 trajectories from the asynchronous hopping region that dissociated (48%) and 88 trajectories that reformed the reactant (37%). From the FC-region of the solvated system of **1**, 9 trajectories formed the decarbonylation product (56%), and 4 reformed the reactant (25%). For the solvated system of **2**, there were 337 trajectories (70%) that crossed through hopping points in the asynchronous region (one σ_{CC} bond of 1.80 Å or longer), while 144 trajectories (30%) crossed through hopping points in the FC-region. From the asynchronous region, 198 trajectories (59%) went on to form the products, whereas 116 (34%) reformed the reactant structure. The 20 trajectories with intermediate structures (6%) exclusively went through hopping points in the asynchronous region. We located 78 (54%) trajectories from the FC region that showed decarbonylation and 57 (40%) which reformed the reactant. The hopping point structures from each region of the hopping seam resemble those from the gas phase simulations; the asynchronous hopping points feature a broken σ_{CC} bond in the cyclopropanone ring, whereas the FC region contains points where both bonds are still intact.

Hydrogen bonding analysis of solvated trajectory simulations

We analyzed the hydrogen bonding throughout all the explicitly solvated NAMD trajectories **1** and **2** with the VMD software.⁸⁴ We noted a hydrogen bond when the donor–acceptor distance (denoted as the $O_{\text{water}} - O_{\text{carbonyl}}$ distance) was 3.1 Å, and the donor–acceptor angle ($O_{\text{carbonyl}} - H_{\text{water}} - O_{\text{water}}$) was near linearity (25°). These criteria were chosen based on previous work examining and measuring intermolecular H-bonding.^{85–87} Fig. 7 shows a snapshot for the solvated systems of **1** and **2** and the H bonding interaction between the carbonyl oxygen (the hydrogen bond acceptor) and a water molecule (the hydrogen bond donor) for **2**.

Our analysis found that the initial condition geometries 64% of **1** and 54% of **2** had H-bonding interactions. To evaluate if H-bonding in the initial condition geometry affects the mechanistic outcome, we divided the initial condition geometries into productive and non-productive trajectories. For **1**, 64% of the

non-reactive initial conditions and 72% of the reactive initial conditions had H-bonding, respectively. For **2**, 54% of the non-reactive initial conditions and 57% of the reactive initial conditions featured H-bonding, respectively. While the reactive initial conditions have slightly more H-bonding for both **1** and **2**, there is no clear correlation between H-bonding in the ground state and photochemical decarbonylation.

While H-bonds are a strongly stabilizing intermolecular force on the ground state between the oxygen and water molecule(s), this interaction should become repulsive in the S_1 . To test this hypothesis, we quantified the degree of H-bonding in the productive and non-productive trajectory pathways to determine how this electronic structure affected the intermolecular H-bonding interactions. After analyzing the full trajectory pathways of **1**, we computed that 22% of productive trajectories have hydrogen bonding, while 34% of non-productive trajectories have hydrogen bonding. For the



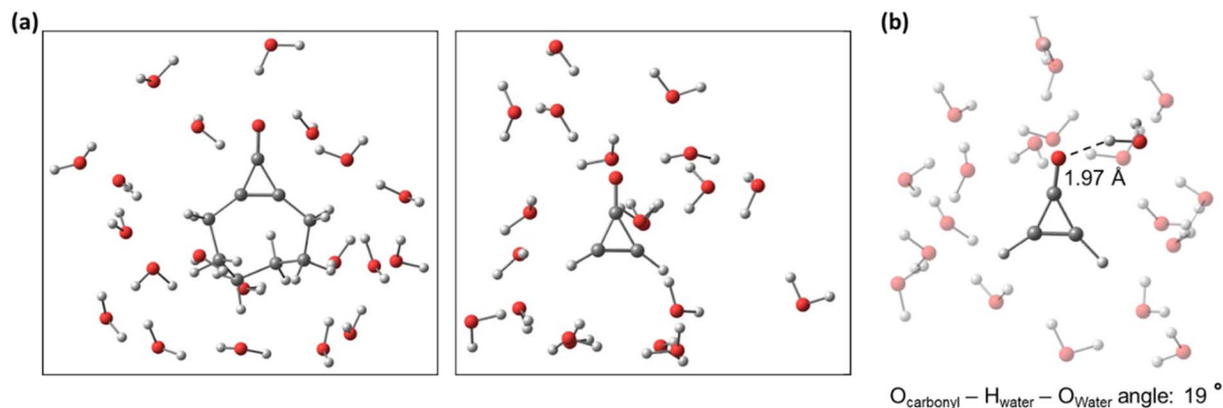


Fig. 7 Snapshots of solvated trajectories of **1** and **2** (a) and an example of a solvated structure with intermolecular hydrogen bonding present between the O_{carbonyl} and a water molecule (b).

trajectory pathways of **2**, we computed that 17% of productive trajectories have hydrogen bonding, while 43% of non-productive trajectories have hydrogen bonding. The productive trajectories overwhelmingly access the asynchronous hopping point geometries, which feature a broken σ_{CC} bond in the cyclopropenone ring. The lack of hydrogen bonding throughout the productive trajectories suggests that it is not stabilizing for the O_{carbonyl} to be interacting with the surrounding water molecules. The non-productive trajectories have much higher levels of H-bonding, we hypothesize this is because the O_{carbonyl} possesses two lone pairs to participate in intermolecular H-bonding.

Conclusion

We have used multiconfigurational quantum mechanical calculations and NAMD simulations to predict the photochemical decarbonylation of cyclopropenones with direct applications in photomedicine, bioorthogonal, and photoclick chemistry. We identified that the mechanistically relevant electronic transitions are $S_0 \rightarrow S_1$ ($n \rightarrow \pi^*$) transitions and have excitation energies of 5.13 eV (**1**) and 4.61 eV (**2**). Our MEP calculations suggest that **1** will prefer to proceed to an S_1/S_0 crossing region while **2** will approach a minimum on the S_1 surface. Our gas-phase NAMD simulations provide dynamical mechanistic information and predict a quantum yield for the dissociation products of 55% and 28% for **1** and **2**, respectively. The time constants of the S_1 were calculated to be 77 and 89 fs for **1** and **2** in the gas phase. We conclude from these simulations that nearly all productive trajectories involve an ultrafast dynamically asynchronous reaction mechanism with one σ_{CC} bond breaking in the S_1 -state and the second σ_{CC} bond breaking in the S_0 -state. The cyclopropenones immediately undergo *cis-trans* isomerization after excitation to S_1 , leading to the cleavage of one σ_{CC} bond. The fused cyclopropenone of **1** has higher QYs because these trajectories avoid the FC-region that lead to highly strained *trans*-cyclooctene, thus preferring to proceed through the productive asynchronous region of the S_1/S_0 seam. We also report the first explicitly solvated NAMD simulations needed to emulate the aqueous environment of the *in vivo*

reaction. In solvated environments, we observed increased quantum yields of 58% for both **1** and **2** and S_1 time constants of 56 fs (**1**) and 68 fs (**2**). We found that H-bonding between the O_{carbonyl} and explicit water molecules was energetically stabilizing hopping points in the FC-region. There were significantly higher rates of H-bonding in trajectories that revert and overwhelmingly undergo surface hopping in the FC-region (34% and 43% for **1** and **2**, respectively) compared to those that dissociate and prefer to hop in the asynchronous region (22% and 17% for **1** and **2**, respectively).

Computational methods

Single reference methods

We used density functional theory (DFT) to optimize the ground-state global minima of **1**, **2**, and **3**. Geometries were optimized using the PBE0-D3BJ/cc-pvdz.^{88–90} Time-dependent density functional theory (TDDFT)⁹¹ was used to calculate the excitation energies and wavelengths for the first 10 excited states of **1**, **2**, and **3**. The TD-DFT calculations were run using the CAM-B3LYP⁷⁷ functional with the aug-cc-pVTZ basis set.^{78,79} DFT includes dynamical correlation between electronic states, which provides accurate vertical excitations and transition natures for small organic molecules. This is an improvement over the multiconfigurational method (CASSCF) which omits dynamical correlation and tends to overestimate excitation energies. While it can accurately predict excitation energies into the Franck-Condon region where there is very little coupling between states, DFT cannot accurately predict the electronic structure on regions of a PES where there is significant coupling between two electronic states (*e.g.*, avoided crossings and conical intersections). We used multiconfigurational methods to predict molecular electronic structure more accurately near these energetic degeneracies. All DFT calculations were run using the Gaussian 16 software.⁹²

Multiconfigurational methods

The multiconfigurational calculations were performed with a state-averaged complete active space self-consistent field (SA-



CASSCF) using OpenMolcas 21.02.⁹³ The methods are described with the format SA(N)-CASSCF(m,n), where N is the number of singlet states the calculation is averaged over. The variables m and n denote the number of electrons and orbitals used in the active space, respectively. For all included molecules, we used a (10,9) active space consisting of two σ and σ^* orbitals, two π and π^* orbitals, and one lone-pair orbital, including a lone pair on the oxygen atom in the cyclopropenone moiety. The active space is shown in Fig. 2. We used SA(4)-CASSCF(10,9)/ANO-SVDZP multiconfigurational calculations to optimize the geometries for the ground and excited states and to run minimum energy path (MEP) calculations to track the steepest path along the excited state, starting from the Franck–Condon (FC) point. We used the extended multi-state complete active space perturbation theory (XMS-CASPT2)⁷⁶ single-point energy correction to account for dynamic correlation for all multiconfigurational calculations. All optimized structures had their vibrational frequencies computed to confirm that they were stationary points (zero imaginary frequencies).

Non-adiabatic molecular dynamics

We used NAMD to track the photochemical pathways of **1** and **2** along the PESs of the ground and excited states. We sampled the initial conditions (nuclear positions and velocities) from the vibrational modes using the Wigner distribution at the zero-point energy (ZPE). The electronic structure calculations used SA(4)-CASSCF(10,9)/ANO-SVDZP. The trajectories started from the S_1 (FC)-point, and the simulation time was 1 ps with a 0.5 fs time step. We computed the surface hopping probability at each step using Tully's fewest switches surface hopping method^{82,94} in OpenMolcas.⁹³

Molecular dynamics simulations

All molecular dynamics (MD) simulations of **1** and **2** in the presence of explicit water molecules were performed in CP2K version 9.1 (ref. 95 and 96) using Generalized Amber Force Field (GAFF)⁹⁷ for the chromophore and SPC/Fw⁹⁸ flexible water forcefield. First, a single geometry of the chromophore was solvated by 200 water molecules in a cubic box. Next, the systems were energy minimized and equilibrated in an isobaric-isothermal (NPT) ensemble for 0.5 ns and a canonical (NVT) ensemble for 1 ns. The last snapshot of this pre-equilibration process was taken, and respective Wigner geometries replaced the chromophore to initialize 500 parallel MD simulations for each molecule (**1** and **2**). During these MD simulations, the chromophores were kept frozen at their Wigner positions. The systems were minimized and equilibrated with a 0.1 ns NPT simulation followed by a 2 ns NVT simulation. All MD simulations used the velocity Verlet algorithm to integrate the equations of motion with a time step of 0.5 fs except during the energy minimization steps in which the conjugate gradient algorithm was used. All short-range non-bonded interactions were evaluated with a cutoff of 8 Å. To compute the coulombic interactions, Smooth Particle-Mesh Ewald (SPME)⁹⁹ summation was used. To maintain a temperature of 300 K and 1.0 bar pressure during the MD simulations, Nose–Hoover

thermostat^{100,101} and barostat were used as needed with time constants of 0.2 ps and 1.0 ps, respectively. The final snapshots of the NVT simulations were used to generate droplets of 20 water molecules surrounding the chromophore for subsequent ground and excited state CASSCF dynamics simulations.

Explicitly solvated NAMD simulations

After running the gas-phase NAMD simulations, we next prepared the solvated NAMD simulations to be run. The production-run solvated trajectories contained our chromophores of interest (**1** and **2**) centered within a droplet of 20 water molecules. For **1** and **2**, the initial geometries for the chromophore were obtained from Wigner sampling using the ground state vibrational frequencies. The initial conditions of the water molecules were taken from the final snapshots of the NVT simulations described in the MD section. The non-adiabatic coupling terms (NACTs) were computed between neighboring states at a given timestep using Tully's FSSH algorithm in OpenMolcas at the SA(4)-CASSCF(10,9) level of theory. To avoid overcoherence between states, we implemented a decoherence correction of 0.1 Hartree, as reported by Persico and Granucci.¹⁰² The velocity rescaling was handled at a constant temperature of 300 K using the Nosé–Hoover method. To leverage computational resources and accuracy, the ANO-SVDZP basis set was used only for the chromophores (**1** and **2**), and ANO-S-MB was used for all 20 water molecules throughout the NAMD simulations. 276 initial conditions were simulated for the solvated system of **1**, and 500 initial conditions were computed for the solvated system of **2**. All simulations were 250 fs in length using a timestep of 0.5 fs.

Data availability

The ESI† is available on Figshare and includes all static and dynamics calculations (<https://doi.org/10.6084/m9.figshare.21667808.v1>).

Author contributions

D. Adrion – calculations, data analysis, manuscript writing, figure preparation. W. V. Karunaratne – calculations, manuscript writing. S. A. Lopez – project conceptualization, research supervision, and manuscript editing.

Conflicts of interest

The authors declare no conflicts of interest.

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