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Direct Observation of a Dark State in the Photocycle of a Light-Driven Molecular Motor

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Supporting Information

ABSTRACT: Controlling the excited-state properties of light driven molecular machines is crucial to achieving high efficiency and directed functionality. A key challenge in achieving control lies in unravelling the complex photodynamics and especially in identifying the role played by dark states. Here we use the structure sensitivity and high time resolution of UV-pump/IR-probe spectroscopy to build a detailed and comprehensive model of the structural evolution of light driven molecular rotors. The photodynamics of these chiral overcrowded alkene derivatives are determined by two close-lying excited electronic states. The potential energy landscape of these “bright” and “dark” states gives rise to a broad excited-state electronic absorption band over the entire mid-IR range that is probed for the first time and modeled by quantum mechanical calculations. The transient IR vibrational fingerprints observed in our studies allow for an unambiguous identification of the identity of the “dark” electronic excited state from which the photon’s energy is converted into motion, and thereby pave the way for tuning the quantum yield of future molecular rotors based on this structural motif.

INTRODUCTION

One of the key challenges in the development of novel materials is the ability to tune and control their macroscopic physical and chemical properties on a molecular level by external stimuli.1−5 Such a degree of control is challenging but crucial for a wide range of applications ranging from catalysis to pharmacological and medical applications.6−31 In photoactive materials light absorbed by chromophores can be converted into directed functionality, typically by means of photoinduced isomerization of nitrogen or carbon containing double bonds and pericyclic reactions.12−27 Indeed, E−Z isomerization results in large amplitude changes in structure as well as chemical and physical properties.28−32

Molecular motors based on overcrowded alkenes are particularly attractive due to the synthetically tunable properties, such as thermal stability, absorption spectra, etc., and therefore are readily applied to a wide range of functional systems. However, although the thermally activated step in the unidirectional rotary cycle is well understood and is now largely predictable, the photochemical properties and especially photochemical quantum yields are less well understood. Hence both from a fundamental perspective and in achieving the goal of complete molecular design, gaining insight into the excited-state properties of light driven molecular-sized machines is essential.33

The overcrowded alkene-based molecular rotary motors are composed of a central carbon—carbon double bond (the axle) embedded in an intrinsically chiral environment. The unidirectional rotational cycle starts with a photoinduced E−Z isomerization around the central C=C bond, generating a metastable isomer P* (Figure 1), which relaxes thermally to the stable isomer P, which in the case of a symmetric stator unit is identical to I. This isomer marks half of the rotation cycle and is the starting point for a second photoinduced isomerization step that leads again to a metastable isomer. Thermally activated conformational isomerization of this isomer brings the system back to its initial state. Overall, the rotor part has then performed a complete rotation with respect to the stator unit of the molecular motor.

Although the rotation includes both photochemical and thermal steps that involve different parts of the motor, the structural evolution is largely governed by the excited-state properties of the central C=C group, which, in turn, are
dictated by the potential energy surfaces of the excited states that are accessed upon electronic excitation. Despite the large number of theoretical\textsuperscript{34−41} and experimental studies\textsuperscript{42−44} that have focused on characterizing this process, there are several key questions outstanding including identification of the main structural coordinates and the role of various electronic states involved in the excited-state dynamics. Time-resolved studies are essential to elucidate these dynamics but have thus far focused on probing the change in the electronic structure after photoexcitation through UV/vis absorption\textsuperscript{42,43} and fluorescence\textsuperscript{43,44} studies. These techniques offer a high time resolution but provide only indirect information as to how the structure evolves spatially over time. In the present contribution we show that this information can be obtained through a combination of picosecond time-resolved vibrational spectroscopy of the electronically excited states together with quantum-mechanical calculations.

**METHODS**

The setup for the time-resolved infrared measurements has been described in detail previously.\textsuperscript{45} Mid-IR probe pulses were generated using an amplified Ti:sapphire system (Spectra-Physics Hurricane, 600 μJ, repetition rate 1 kHz) pumping a commercial BBO-based OPA (Spectra-Physics OPA-800C), the signal and idler output of which were difference-frequency-mixed in AgGaS\textsubscript{2}. UV pump pulses (400 nm, ~1 μJ) were generated by doubling part of the output of the Ti:sapphire laser. The diameter of the UV pump focus was ~180–200 μm; that of the IR probe focus, ~150 μm. The sample cell with CaF\textsubscript{2} windows spaced by 500 μm was placed in the IR focus. To avoid accumulation of the thermally unstable conformer P\textsuperscript{*} during the measurements, a flow cell was used. With a Newport ESP300 translation stage, the pump−probe delay time was scanned by mechanically adjusting the beam path of the UV pump. The temporal resolution of 200 fs has been determined from the full width at half-maximum of the pump probe cross-correlation function. The pump pulse was chopped at 500 Hz, and transient spectra were obtained by subtracting nonpumped absorption spectra from the pumped absorption spectra, both of which were recorded by spectrally dispersing the probe pulses using an Oriel MS260i spectrograph and measuring the frequency-dependent IR intensity using a 2×32 HgCdTe detector array (Infrared Associates). Compound I was available from earlier studies (see ref\textsuperscript{44}). In all experiments we investigate a 10mM solution of compound I in deuterated cyclohexane (Aldrich, 99.6 atom% D).

For the investigation of the ground-state conformers (including vertical excitation energies and forces acting at the Franck–Condon point) quantum mechanical calculations were performed using the TURBOMOLE suite of programs.\textsuperscript{46} Geometry optimization and normal-mode analysis were performed by (time-dependent) density functional theory (\textit{(TD)-DFT}) using the hybrid functional B3LYP.\textsuperscript{47} For the geometry optimization the D3 correction method of Grimme et al.\textsuperscript{48} was used to include dispersion effects. For all elements the cc-pVDZ or aug-cc-pVDZ basis sets were employed, respectively.\textsuperscript{49,50} Second derivatives were computed analytically.
with the AOFORCE module. The vibrational frequencies have been scaled by 0.98. Post-Hartree–Fock calculations were performed on the basis of spin-component scaled second-order Møller–Plesset perturbation theory\(^5\) and the approximate coupled-cluster singles-and-doubles model CC2. All perturbation method calculations were performed using the resolution of identity (RI) approximation\(^6\) and the efficient RICC2 module\(^7\) implemented in the TURBOMOLE package together with the standard auxiliary basis sets.\(^8\) The Gaussian09 program package\(^9\) was employed to study the excited-state properties in more detail and in particular to optimize the structure of the dark-state species. Geometry optimization and normal-mode analysis were performed in the framework of TD-DFT at the B3LYP/cc-pVDZ level.\(^{45,53–59}\)

**RESULT AND DISCUSSION**

The structural evolution from the initial state I to isomer P* (Figure 1) was considered first in terms of the spectral properties of the initial state and the first metastable product (P*). The steady-state UV/vis absorption spectra of the two isomers are distinct, reflecting a change in the conjugated π system (Figure 1a). The FTIR spectra of the two isomers (Figure 1c) show that the carbon–carbon stretch and ring vibrations couple to a large degree as a consequence of the extended π system; however, DFT calculations enable us to assign the bands in the 1550–1650 cm\(^{-1}\) range as being mainly associated with modes involving the central C=C double bond (Supporting Information). The metastable isomer P* shows a shift of these modes to lower frequencies confirming a decrease in bond strength. Comparison of the optimized geometries of the two isomers (I (P*)), CC bond length 1.369 (1.377) Å; CC=C=CC dihedral angle 14.1° (29.5°)) evidence the corresponding lower π bond character as well.

Time-resolved vibrational spectroscopy\(^{51–67}\) was employed to achieve the required temporal resolution as well as structural sensitivity to elucidate the structural evolution that follows photoexcitation with a short UV-pump pulse (center wavelength 400 nm). The resulting structural evolution was followed by time-resolved differential absorption spectra in cyclohexane-d\(_\text{4}\); with a time resolution of 200 fs (Figure 2). The transient IR spectra show, as expected, pronounced differences between the IR absorption spectrum of the molecule in the initial and excited states. A first point of note is that the spectrum does not evolve further after ca. 30 ps, and the difference spectrum corresponds perfectly with the difference spectrum obtained upon continuous wave excitation (Figure 1); i.e., the metastable isomer P* has formed fully within the 30 ps after excitation.

The time-resolved IR spectra obtained within the first 30 ps exhibit several remarkable features. The first and most striking is a broad and unstructured absorption over the entire mid-IR range (1200–3000 cm\(^{-1}\)). This absorption, which decays rapidly, is particularly prominent at early time delays (<5 ps). Individual absorption bands with remarkably high integrated intensities in the 1400–1600 cm\(^{-1}\) range are superimposed on the broad spectral feature. Notably, these bands are significantly broader than the bleached bands in the transient spectra and bands in the FTIR spectra. The solvent response and the dependence of the UV pump intensity (Supporting Information) confirm that these features are intrinsic to the compound and are not due to solvent or two-photon processes.

The width and intensity of the broad excited-state absorption bands excludes that they arise from vibrational transitions. Indeed, the absorbance is closer to that expected for electronic transitions, albeit these typically occur at much higher energies (in the UV/vis range). The electronic excited-state manifold of the compound was therefore investigated computationally. The initial state of the compound (I) shows an absorption band at 390 nm (Figure 1), which is reproduced well by B3LYP/cc-pVDZ calculations in the gas phase (403 nm; \(f = 0.428\)). The S\(_1\) ← S\(_0\) transition has predominantly HOMO → LUMO character (Figure 1). In contrast to previous assignments,\(^{12}\) however, we find that this electronic transition is not localized on the stator but involves to a significant extent excitation from the bonding π orbital of the central double bond to the antibonding π* orbital (Figure 1). As a consequence, the double bond character of this bond is reduced as well as the barrier to rotation around this bond. In line with calculations on ethylene and its derivatives, we find a dark state (S\(_2\)) associated with the HOMO→1 → LUMO excitation at slightly higher excitation energies that is not directly accessible from the ground state (Figure 1, Table 1). Because the HOMO→1 orbital is localized on the aromatic stator unit, S\(_2\) has a significantly larger dipole moment than S\(_1\) (4.3 vs 2.2 D at the SCS-CC2/cc-pVDZ level). It should be emphasized that the description of the lower-lying electronic states in the compound studied here is thus entirely different from ethylene and its derivatives where, as discussed by Salem and others,\(^{68–75}\) for torsion angles of around 90° the two low-lying electronic states have either a biradical or zwitterionic character depending on the occupation of the two central 2p-orbitals and require multireference methods for a proper description. In the present study, in contrast, we investigate the dynamics occurring at
small torsional angles (vide infra) for which the employed single-reference methods not only have been validated. In addition we employ the results of multireference calculations on smaller motor molecules as reported by Liu and Morokuma to make a further comparison with our experimental results.

For vertical excitation our calculations predict that S₁ and S₂ are nearly degenerate (Table 1). They thus strongly suggest that the broad and structureless absorption feature observed in the time-resolved IR spectra is associated with the electronic S₁ → S₂ transition. The presence of two close-lying states also explains why the induced absorption bands have such high intensities. Our calculations show that the energy gap between the two states strongly depends on structural parameters such as the C═C bond length that are involved in the 1400–1600 cm⁻¹ normal modes. For these modes one therefore expects that their transition moments will be enhanced by vibronic coupling as is indeed observed. The conclusion that the excited-state dynamics of the molecular rotor should be described in a multistate model is further supported by recent multireference calculations on smaller rotor molecules that also find very small energy differences between the S₁ and S₂ states.

The excited-state dynamics were elucidated further using a global analysis of the transient spectra, which shows that the spectra can be described well with two exponential decay terms with time constants of 1.53 ± 0.03 and 12 ± 1 ps. We note that our experiments have a time resolution of 150–200 fs and therefore do not allow observation and confirmation of the ultrafast dynamics reported previously in fluorescence-upconversion studies. Species-associated spectra (SAS) (Supplementary Figure 2) were derived by using a two-step sequential kinetic model.

The SAS of the first species shows an overall background signal with broad and intense features superimposed on it that indicate vibronic coupling. Furthermore, the increase (decrease) of absorbance in the 1550 (1600) cm⁻¹ region indicates a substantial change in bond character of the central double bond (Figure 2c). This spectrum was related to the structure of this excited-state species using excited-state geometry optimizations. Full optimization of the “dark” state without any structural constrains associated with the HOMO–LUMO excitation leads to a stable minimum that can be reached from the Franck–Condon excited geometry by relatively small structural changes (Supplementary Figure 4), and an excited-state IR absorption spectrum that is in good agreement with the SAS of species 1 (Figure 2). It is notable that for a proper comparison with the experimental spectrum, the theoretically predicted stick spectrum needs to be convolved with 20 cm⁻¹ Gaussian line shapes, suggesting that in the “dark” state a wide range of structures is sampled. Our experiments and calculations thus demonstrate that excitation and excited-state decay involve two different electronic states. In earlier studies, time-resolved fluorescence and visible absorption data were interpreted in terms of a decay from a single, isolated electronic state decaying to the ground state after structural relaxation in the excited state. The present data show that the S₁ state actually decays to an intermediate, dark electronic state, and from there to the ground state.

In the SAS of the second species (Supplementary Figure 2) the broad background is absent, and bands have widths comparable to those in the S₀ state. The absence of the broad background indicates that this species is no longer in an electronically excited state but back in the (vibrationally hot) electronic ground state. Indeed, the vibrational activity observed in the 1550 cm⁻¹ region follows closely the IR absorption spectrum of P*. In addition, repopulation of the original state (I) is observed also (i.e. the quantum yield for isomerization is less than unity), and the SAS of the second species contains contributions from both vibrationally hot ground-state I and metastable P* isomers. Vibrational cooling of these isomers leads to the SAS of the final species, which is in excellent agreement with the steady-state FTIR difference spectrum of room-temperature I and P*.

The present experimental and theoretical data allow a complete picture of the mechanism by which the molecular rotary motor undergoes isomerization after electronic excitation (Figure 3). Photon absorption brings the motor to a “bright” state from which it decays on an ultrafast time scale and within the time resolution of the present experiments to a different electronically excited “dark” state. In line with previous calculations on similar systems, our calculations indicate that this is a barrierless process proceeding via a conical intersection. Previously, it was assumed that structural evolution on the HOMO → LUMO potential energy surface is responsible for the observed ultrafast decay of the fluorescence, and (following commonly accepted models on photoisomerization of carbon–carbon double bonds) that it primarily proceeds along torsional and pyramidalization coordinates. The present study, in contrast, leads to the conclusion that the fluorescence decay is mediated by the passage through the conical intersection of the HOMO → LUMO and HOMO−1 → LUMO potential energy surfaces. Inspection of the structural changes that occur upon geometry relaxation to the minimum on the HOMO−1 → LUMO potential energy surface show that the central C=C bond length increases from 1.381 to 1.461 Å and that only relatively small changes occur in the torsional and pyramidalization coordinates. In combination with the forces that act on the molecule at the Franck–Condon excited “bright” state (Supplementary Figure 3), this suggests that motion along the C=C stretch coordinate needs to be taken into account as well for understanding the decay of the fluorescence.

Hence, it can be concluded that internal conversion to the electronic ground state takes place from the “dark” state. The decay rate for this process indicates that it is mediated by conical intersections between the two states as has indeed been proposed in previous theoretical studies. These studies concluded, however, as well that to reach these

<table>
<thead>
<tr>
<th>geometry</th>
<th>B3LYP/aug-cc-pVDZ*</th>
<th>SCS-CC2/cc-pVDZ*</th>
</tr>
</thead>
<tbody>
<tr>
<td>S₀</td>
<td>3.020 (0.428)</td>
<td>3.642 (0.583)</td>
</tr>
<tr>
<td>S₁</td>
<td>2.228 (0.004)</td>
<td>2.353 (0.002)</td>
</tr>
<tr>
<td>S₂</td>
<td>3.065 (0.963 × 10⁻³)</td>
<td>2.632 (0.495)</td>
</tr>
<tr>
<td>S₃</td>
<td>3.662 (0.018)</td>
<td>2.788 (0.592)</td>
</tr>
</tbody>
</table>

*For S₀ the optimized structure on the B3LYP+D/aug-cc-pVDZ level was used to calculate the vertical excitation energies.

**Table 1. Calculated Vertical Excitation Energies (Evert, eV) and Corresponding Oscillator Strength in Brackets of the First Two Excited States of the Molecular Motor at the Optimized S₀ and S₁ Geometries**

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The photodynamics of a unidirectional molecular motor has been studied by time-resolved IR spectroscopy in which both vibrational and electronic structures are probed. In combination with quantum mechanical calculations, the high structural sensitivity and real-time character of the technique allows us to understand fundamental aspects of the operation mechanism. The results show that conversion of photon energy into directed motion proceeds on the potential energy surfaces of two different electronic states, which is the more important as it implies that the conical intersection between the “dark” electronically excited state and the ground state determines the efficiency of the motor. To rationally tune the performance of photoinitiated unidirectional rotation thus involves a multicoupled-state optimization of nuclear motion.

**SUMMARY AND CONCLUSION**

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpca.6b09644.

Solvent response and power dependency of the transient IR signal, transient IR global data fitting and FTIR spectra, structure showing the gradient vectors acting on the molecule at the Frank–Condon point, structure of the dark state, structure showing the force vectors associated with the normal mode, vibrational band assignments of conformers I and P* (PDF)

Movie of the normal mode, I 1625.43 cm$^{-1}$ (MOV); movie of the normal mode, I 1618.85 cm$^{-1}$ (MOV); movie of the normal mode, I 1598.05 cm$^{-1}$ (MOV); movie of the normal mode, I 1587.39 cm$^{-1}$ (MOV); movie of the normal mode, I 1567.67 cm$^{-1}$ (MOV); movie of the normal mode, P* 1611.44 cm$^{-1}$ (MOV); movie of the normal mode, P* 1593.63 cm$^{-1}$ (MOV); movie of the normal mode, P* 1576.22 cm$^{-1}$ (MOV); movie of the normal mode, P* 1573.24 cm$^{-1}$ (MOV); movie of the normal mode, P* 1573.24 cm$^{-1}$ (MOV); movie of the normal mode, dark-state 119.44 cm$^{-1}$ (MOV) (ZIP)

**AUTHOR INFORMATION**

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**Notes**

The authors declare no competing financial interest.

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