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Abstract

Attached to electron-rich aromatic systems, sulfides are very weak acceptors; however, attached to electron-poor aromatics, they turn into quite strong donors. Here, we show that this underappreciated dual nature of sulfides deserves full consideration for the design of functional systems. Tested with newly designed and synthesized planarizable push–pull mechanophores, sulfide acceptors in the twisted ground state are shown to prevent oxidative degradation and promote blue-shifting deplanarization. Turned on in the planar excited state, sulfide donors promote red-shifting polarization. Impressive Stokes shifts are the result. Demonstrating the usefulness of time-resolved broadband emission spectra to address significant questions, direct experimental evidence for the ultrafast (3.5 ps), polarity-independent and viscosity-dependent planarization from the twisted Franck–Condon S1 state to the relaxed S1 state could be secured.
**Turn-On Sulfide π Donors: An Ultrafast Push for Twisted Mechanophores**

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**ABSTRACT.** Attached to electron-rich aromatic systems, sulfides are very weak acceptors; attached to electron-poor aromatics, they turn into quite strong donors. Here, we show that this underappreciated dual nature of sulfides deserves full consideration for the design of functional systems. Focusing on twisted push-pull systems, we thought that sulfides in the position of “pushing” substituents should act as weak acceptors in the ground state and convert into strong donors only upon coupling with the “pulling” acceptor in the planarized excited state. Applied to twisted dithienothiophene-dithienothiophene S,S-dioxide probes, turn-on sulfide donors are shown to prevent oxidative degradation and to promote blue-shifting deplanarization in the twisted ground state and red-shifting polarization in the planar excited state. As a result, Stokes shifts increase up to 9400 cm⁻¹. Femtosecond time-resolved broadband fluorescence experiments reveal that the excited-state dynamics in the presence of conventional methoxy donors are mostly controlled by solvent relaxation. With turn-on sulfide donors, ultrafast (3.5 ps), polarity-independent, viscosity-dependent planarization from the twisted Franck-Condon S₃ state to the relaxed S₄ state constitutes the rate-limiting step. Supported by pertinent controls, these findings validate the concept of turn-on sulfide donors for general use.

Sulfide substituents on electron-rich aromatic systems are weak electron acceptors, whereas on electron-poor aromatic systems, they are quite strong electron donors. The very weakly accepting nature of sulfides with electron-rich aromatics is best appreciated with the Hammett σₚ = +0.03 of ethyl sulfides, a value referring to benzoic acids. This compares to σₚ = -0.83 for dimethylamino, σₚ = -0.27 for methoxy and σₚ = -0.15 for ethyl donors, and to σₚ = +0.66 for cyano, σₚ = +0.77 for ethylsulfone and σₚ = +0.78 for nitro acceptors. The strongly donating nature of sulfides to electron-poor aromatics is best appreciated by the deep red color of naphthalenedimides (NDIs) with two sulfides in the core, halfway between the yellow NDIs with two alkoxy and the blue NDIs with two alkylamino substituents in the core. The conversion of sulfides into strong donors for electron-poor aromatics is also well reflected in the σₚ = -0.60 of methyl sulfides (for cyano acceptors, for example, σₚ = σₚₑ = +0.66).

In this report, the switch of sulfide substituents from very weak acceptors to strong donors with decreasing electron deficiency of the aromatic system is used to tackle an intriguing challenge with planarizable push-pull probes. The design of these probes was inspired by the color change of lobsters during cooking and the origin of color vision. Namely, the combination of planarization and polarization in the ground state was envisioned to afford mechanosensitive fluorescent membrane probes that operate with changes in excitation rather than emission. Evolving from twisted π push-pull oligothiophenes (Figure 1A), the currently best probes operate with fluorescent sulfides. Fluorescent flippers are monomers with large surface area to better feel the environment and to keep on shining also when twisted out of conjugation (Figure 1B). In flipper probe 1, one bithiophene was bridged with a “sulfide donor” and the other with a “sulfone acceptor.” The highly fluorescent dithienothiophene S,S-dioxide acceptor is further strengthened with an aldehyde. Repulsion between methyl substituents and the chalcogenic σ holes of the endocyclic sulfur atoms next to the connecting bond is used to twist the two aromatic planes out of co-planarity. In mixed membranes, disordered domains could be imaged by excitation of deplanarized flippers 1 at shorter wavelength, whereas more ordered domains were emitting upon excitation of planarized probes at longer wavelength. However, considering the concept of planarizable push-pull probes, flipper probe 1 contains a strong aldehyde acceptor but fails to integrate a strong donor. This failure is not accidental. In conjugated push-pull systems, the electron density injected by the donor is delocalized toward the acceptor. However, when twisted out of conjugation, the electron density injected by the donor accumulates on the first aromatic system, a situation that inevitably leads to oxidative degradation. Ideally, strong donors in planarizable push-pull probes would turn on only upon planarization of the aromatic system (Figure 1C). Turn-on sulfides appeared just perfect to meet this subtle challenge.

To elaborate on the concept of turn-on sulfide donors comprehensively, we decided to prepare a series of push-pull flippers 2-8 (Table 1, Scheme 1). “Turn-on” probes 2 and 4 contain sulfide substituents that should “turn on” as donors only in response to planarization and conjugation with sulfone and cyano acceptors (Figure 1C). In control probes 3 and 5, the turn-on sulfides are replaced by conventional methoxy donors. Turn-on probe 2, the key target, was readily accessible from dithienothiophene 9. The critical sulfide substituent could be introduced with disulfide 10 after deprotonation of the substrate 9 with LDA. Simultaneous oxidation of dithienothiophene and the turn-on sulfide donor of 11 with mCPBA afforded dithienothiophene S,S-oxide 12 with a...
ble with a broad variety of functional groups. Flippers sulfide turn-on donors from their respective disulfides is compatible with probe development, it is important to note that the introduction of amination with NBS, and S-ethylene coupling with sulfide donors.

Table 1. Structure and Spectroscopic Properties of Twisted Push-Pull Probes in Solution.

<table>
<thead>
<tr>
<th>Cpd</th>
<th>donor</th>
<th>acceptor</th>
<th>λabs (nm)</th>
<th>λem (nm)</th>
<th>Δλ (nm)</th>
<th>Δν (cm⁻¹)</th>
<th>ΔΔν (cm⁻¹)</th>
<th>QY (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>-CH₂OR</td>
<td>-CHO</td>
<td>422</td>
<td>621</td>
<td>199</td>
<td>7800</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>2</td>
<td>-SEt</td>
<td>-SO₂Et</td>
<td>403</td>
<td>635</td>
<td>232</td>
<td>9300</td>
<td>+1500</td>
<td>29</td>
</tr>
<tr>
<td>3</td>
<td>-OMe</td>
<td>-SO₂Et</td>
<td>418</td>
<td>648</td>
<td>230</td>
<td>8700</td>
<td>+900</td>
<td>22</td>
</tr>
<tr>
<td>4</td>
<td>-SEt</td>
<td>-CN</td>
<td>411</td>
<td>650</td>
<td>239</td>
<td>9400</td>
<td>+1600</td>
<td>33</td>
</tr>
<tr>
<td>5</td>
<td>-OMe</td>
<td>-CN</td>
<td>428</td>
<td>681</td>
<td>253</td>
<td>9000</td>
<td>+1200</td>
<td>23</td>
</tr>
<tr>
<td>6</td>
<td>-H</td>
<td>-CN</td>
<td>396</td>
<td>543</td>
<td>147</td>
<td>7000</td>
<td>-800</td>
<td>-</td>
</tr>
</tbody>
</table>

1  Reported for EtOAc because of the similarity to emission in membranes (Figure 3A), other solvents gave the same trends. 2  For structures, see Figure 1, Scheme 1 and Scheme S2 (7: A = -CN, D = -(n-Pr)₂; 8: A = -COR, D = -(n-Pr)₂). 3  Wavelength λabs of absorption maximum (2: ε = 19.7 mM⁻¹cm⁻¹, 3: ε = 15.9 mM⁻¹cm⁻¹). 4  Stokes shift in wavelength (nm, λem - λabs). 5  Stokes shift in frequency (wavenumbers, νem - νabs). 6  Change of Stokes shift compared to original 1. 7  Fluorescence quantum yield in CHCl₃ relative to rhodamine G (94% in EtOH). QYs are unrelated to planarization kinetics from FC to the relaxed S₁ excited state.

![Scheme 1](image)

sulfone acceptor. The activated sulfone 13 was obtained by bromination with NBS, and Stille coupling with sulfide 11 afforded flipper 2 in overall four straightforward steps only. For future probe development, it is important to note that the introduction of sulfide turn-on donors from their respective disulfides is compatible with a broad variety of functional groups. Flippers 3-8 were prepared analogously, details on their synthesis can be found in the Supporting Information (Schemes S1-S2).

In the ¹H NMR spectrum of 5, the signals of the methyl protons next to the cyano acceptor and the methoxy donor appeared downfield and upfield of the methyl protons in the middle of the fluorophore, respectively (Figure 2A), and the spectrum changed on irradiation at 435 nm for long time, the decrease of emission intensity of probe 2 with turn-on sulfide donors with time was 8.2 times slower than that of carboxyfluorescein and 37.8 times slower than that of the unstable probe 3 with conventional methoxy donors.

The emission maximum of the original probe 1 in membranes is close to that in EtOAc (Figure 3A, solid). The large difference between the absorption maxima of 1 in EtOAc (and all other solvents) and solid-ordered membranes is thought to originate from ground-state planarization (Figure 3A, dashed). 6 The absorption maximum of turn-on probe 2 in EtOAc was found at λabs = 403 nm (Figure 3B, dashed, Table 1). This compared to 1 blue-shifted absorption was in agreement with an increased ground-state deplanarization in the absence of a strong push-pull dipole, i.e., with a weakly withdrawing sulfide acceptor at the partially decoupled, electron-rich dithienothiophene. The emission of turn-on probe 2 at λem = 635 nm was red-shifted compared to 1 (Figure 3B, solid, Table 1). This red shift was in agreement with a strengthened push-pull dipole in the planar S₁ state, i.e., the presence of turned-on sulfide donors. Blue shift in absorption and red shift in emission added up to a large Stokes shift, increasing by ΔΔν = +1500 cm⁻¹ from original 1 to Δν = 9300 cm⁻¹ for turn-on probe 2 (Table 1, entry 2). 10

Compared to original 1, the presence of conventional methoxy donors in 3 shifted the emission to the red but failed to shift the absorption significantly to the blue (Figure 3C, Table 1). An increased Stokes shift by only ΔΔν = +900 cm⁻¹ was the result, that is not much more than half the ΔΔν = +1500 cm⁻¹ obtained with turn-on donors in 2 (Figure 3C, Table 1). Cyanine instead of sulfone acceptors caused global red shifts (Table 1, entries 2-5). These uniform shifts imply slightly increased ground-state planarization and excited-state polarization. Despite their smaller σp, cyanine groups are thus slightly stronger acceptors than sulfones in these systems. Partially preserved blue-shifted absorption in 4 compared to 1 showed that the stronger cyanine acceptors are better suited for probe development.
significant reduced Stokes shift (the expected large blue shifts that the spectral-shift dynamics in hydrophobic solvents report on their slowing-down with increasing solvent viscosity supported group as donor increased from entry 6). Amino donors combined with cyano and ketone acceptors in sulfide donors was more pronounced in combination with sulfones in (+400 cm⁻¹) than with cyano acceptors in (±600 cm⁻¹, Table 1).

Controls 6 without any substituent in the donating position gave the expected large blue shifts in absorption and emission with significantly reduced Stokes shift (Δν = -800 cm⁻¹, Table 1, entry 6). Amino donors combined with cyano and ketone acceptors in 7 and 8, respectively, removed essentially all fluorescence in EtOAc. Appreciable emission only in the least polar solvents such as hexane (7: λabs = 445 nm, λem = 635 nm, Δν = 6900 cm⁻¹; 8: λabs = 420 nm, λem = 642 nm, Δν = 8400 cm⁻¹) implied that fluorescence quenching occurs by photoinduced intramolecular electron transfer from the amino donor, independent of the nature of the acceptor.

A novel, extremely versatile broadband fluorescence up-conversion technique, unrivaled in photometric precision, was applied to gain direct insight into the conformational and energetic relaxation processes in real-time. Contrary to the monomeric control 12 (Figure 4A), the time-resolved emission spectra of 2 and 3 in EtOAc showed a large red shift with time (using times larger 0.2 ps, Figures 4B and C). The time-resolved emission spectra of 2 with turn-on and 3 with conventional donors were quite similar. In EtOAc, the characteristic instability of sulfides, which showed basically no dependence on the solvent polarity and were distinctly slower than for 3 in apolar and polar solvents (t1e ≈ 3.5 ps, Figure 4E, O, ●).

These observations allow for two conclusions. Firstly, the excited-state planarization (the exclusive dynamics being monitored in apolar solvents) is distinctly slower for 2 than for 3. The conventional strong donor in 3 thus supports planarization already in ground state, as expected for push-pull systems and demonstrated by red-shifted absorption. Thus planarization from the less twisted Franck-Condon (FC) S1 state to the planar relaxed S0 state requires little structural rearrangement. In clear contrast, turn-on donors in 2 support deplanarization while acting as acceptors in the ground state, as expected for pull-pull systems and demonstrated by blue-shifted absorption. As a result, they need correspondingly larger amplitude motion, and thus more time, to planarize from the more twisted FC state to the relaxed S0 state while transforming from weak acceptors to strong donors.

Secondly, when moving toward polar solvents, the slower diffusive solvent relaxation starts to dominate the relaxation dynamics of the S0 state of 3. In clear contrast, planarization of 2 is slower than solvent relaxation and thus the excited-state dynamics of 2 are independent of solvent polarity. This could indicate that in 2, planarization is a prerequisite for the excited-state charge transfer, which may be interpreted as a direct experimental support of the concept of turn-on sulfide donors. These insights could be secured only with ultrafast broadband fluorescence in a judicious choice of solvents, thus allowing us to assign the timescales of the different relevant relaxation processes without the complications arising from overlapping spectral contributions when using transient absorption spectroscopy or the limited access to spectral lineshapes in single wavelength fluorescence measurements.

Direct evidence for a twisted FC S1 state from time-resolved emission spectra was in agreement with strong positive solvatochromism of the emission but not the absorption maxima (Figures S1-S6). This key characteristic of planarizable push-pull probes.
contrasts clearly to positive solvatochromism in both absorption and emission found for standard push-pull fluorophores with planar ground and excited states. The dependence of emission maxima on polarity index was linear and similar for all push-pull probes (slope $v_e = 13300-17400 \text{ cm}^{-1}$, 2-4<3<5). As with the Stokes shifts, solvatochromism of 6 without additional donor was much weaker ($v_e = 7600 \text{ cm}^{-1}$, Figure S6).

In summary, the dual nature of sulfide substituents is applied to tackle a most annoying challenge with planarizable push-pull probes, i.e., rapid decomposition with strong donors. Blue shifts in absorption confirm sulfides as weak, stabilizing and deplanarizing acceptors in the twisted ground state, whereas red shifts in emission confirm their transformation into polarizing donors upon planarization in the excited state. Demonstrating the usefulness of time-resolved broadband emission spectra to address significant questions, direct experimental evidence for the ultrafast, polarity-independent and viscosity-dependent planarization from the twisted FC $S_1$ state to the equilibrium $S_1$ state could be secured. Compared to controls with conventional donors, turn-on donors provide access to twisted probes that, i) most importantly, are stable, ii) have maximal Stokes shifts, iii) reasonable quantum yields, and iv) mechanosensitive excited-state dynamics that are controlled by planarization rather than by solvation. These findings became possible because the concept of turn-on sulfide donors, i.e., the conversion from very weak acceptors for electron-rich aromatics to strong donors for electron-poor aromatics. Compared to isostructural sulfur redox switches, the more subtle concept of turn-on sulfide donors has attracted much less attention for the design of functional systems. The reported results suggest that it certainly could, maybe should.

**ASSOCIATED CONTENT**

**Supporting Information.** Detailed experimental procedures. This material is available free of charge via the Internet at http://pubs.acs.org.

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The authors declare no competing financial interest.

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(16) As an amorphous solid, turn-on probe 2 is highly fluorescent, whereas in single crystals, 2 is fully planarized and does not fluoresce. Studies on ground-state planarization in confined space are ongoing and will be reported in due course.


(20) Involvement of a twisted intramolecular charge-transfer (TICT) state can be excluded because the substantial red shift of the fluorescence observed in apolar solvents and the polarity-independent excited-state dynamics of 2 cannot be rationalized in terms of a TICT state.
