

Toward Fluorescent Memories with Nondestructive Readout: Photoswitching of Fluorescence by Intramolecular Electron Transfer in a Diaryl Ethene-Perylene Bisimide Photochromic System**

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Photochromic compounds change their optical and electronic properties by reversible photochemical reactions upon irradiation and are therefore suitable for many tasks.^[1] Very recently, the application of photochromic compounds to the imaging of living cells by single-molecule photoswitching has been demonstrated.^[2] In the last few years diaryl ethenes (DAEs) have evolved as highly promising photochromic molecules for optical data storage because of their durable persistency and the thermal irreversibility of their closed and open forms.^[3]

Fluorescence photoswitching of DAEs linked to fluorescent dyes has been demonstrated even at the single-molecule level.^[4] In these systems, the readout is based on the observed fluorescence of the open form of diaryl ethene (DAE_O), whilst fluorescence quenching from the excited dye to the closed form of the diaryl ethene (DAE_C) takes place by an intramolecular Förster resonance energy transfer (FRET). The drawback of such energy-transfer-based memory systems is that FRET induces photochromic cycloreversion during the readout step, and thus destroys the memory.

Such destructive readout may be circumvented by fluorescence switching through intramolecular photoinduced electron transfer (PET)^[5] if the open and closed forms of the photochromic moieties were to possess different redox properties.^[6] The optimal case would be if the electron transfer involves oxidation of the dye and reduction of the DAE^[7,8] and the solvent-dependent driving force for PET is exergonic for the DAE_C[−]-dye⁺ but not for the DAE_O[−]-dye⁺ charge transfer (CT) state (Figure 1). Furthermore, the emission spectrum of the dye should not overlap with the absorption spectrum of any isomer of the DAE, as in FRET-based systems, but needs to be at a higher wavelength.^[9] This would facilitate writing, erasing, and reading data with light

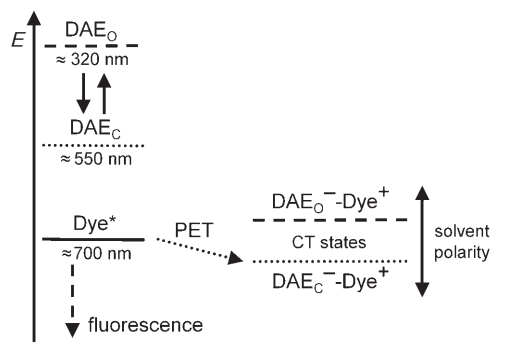
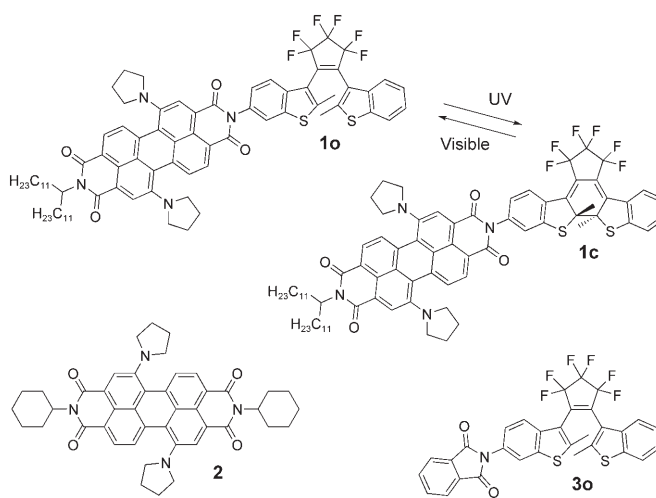


Figure 1. Schematic energy diagram for the fluorescent open form (dashed line) and the nonfluorescent closed form (dotted line) of a DAE and CT states of the DAE-dye conjugate.

sources of three different wavelengths without destructive readout.

A photochromic DAE system based on the above-mentioned optimized concept is, to the best of our knowledge, unknown.^[10] Thus, we have designed the photochromic system **1o** based on a perfluorinated DAE derivative and a bay-substituted perylene bisimide (PBI) dye (Scheme 1). The fluorescent 1,7-dipyrrolidinylperylene bisimide was chosen because of its low oxidation potential and its absorption maximum in the NIR region (around 700 nm).^[11] The photochromic compound **1o** was synthesized by imidization of the



Scheme 1. Photochromic compounds **1o** and **1c**, and reference compounds **2** (perylene bisimide dye) and **3o** (DAE).

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respective amino-substituted diaryl ethene with 1,7-dipyrro-
lidinylperylene monoanhydride monoimide (see the Support-
ing Information for details). Compound **1o** can be switched
between two photostationary states. Upon irradiation with
UV light ($\lambda_{\text{max}} = 350$ nm) a photostationary state containing
66% of the closed form of the PBI-DAE conjugate **1c**
(Scheme 1) was obtained,^[12] which could be isolated by
column chromatography. The open form can be regenerated
almost quantitatively (99%) by irradiation of a solution of the
closed form with visible light ($\lambda > 400$ nm).

The fluorescence quantum yields for **1o** and **1c** were
determined in a series of solvents of different polarity, with
dielectric constants (ϵ_r) ranging from 2.24 (tetrachlorome-
thane) to 36.71 (dimethylformamide). These values, divided
by the quantum yield of the reference PBI **2** (without a DAE
moiety), are shown as a function of the ϵ_r values of the
solvents in Figure 2 (see also Table S1 in the Supporting

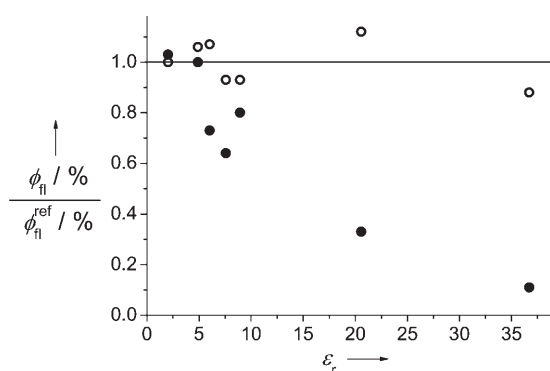


Figure 2. Fluorescence quantum yields of **1o** and **1c** (ϕ_f) divided by
the quantum yield of the reference perylene bisimide **2** (ϕ_f^{ref}) without
the DAE moiety as a function of dielectric constants of solvents; open
form ($\circ$), closed form ($\bullet$).

Information). In low-polarity solvents such as chloroform
($\epsilon_r = 4.89$), the quantum yields of **1o** and **1c** are nearly
identical to that of the reference dye **2**, which indicates that no
fluorescence quenching by the DAE_C or DAE_O moieties
occurs. Interestingly, the quantum yields of **1o** and **1c** differ
significantly at higher solvent polarity, and in dimethylforma-
mide (highly polar) almost complete quenching of the closed
form **1c** is observed. Fluorescence quenching by FRET from
the excited dye to the closed form of the DAE can be
excluded as a deactivation path. Thus, the increase in the
fluorescence quenching of **1c** as the solvent polarity increases
is highly indicative of a preferential PET from the dye to the
closed form of DAE.

The Gibbs free energy (ΔG°) for an intramolecular
charge-separated state of a covalently bound donor–acceptor
system in a given solvent can be estimated using the Rehm–
Weller Equation (1).

$$\Delta G^\circ = e[E_{\text{ox}}(\text{D}) - E_{\text{red}}(\text{A})] - E_{00} - \frac{e^2}{4\pi\epsilon_0\epsilon_s R_{\text{CC}}} - \frac{e^2}{8\pi\epsilon_0} \left(\frac{1}{r^+} + \frac{1}{r^-} \right) \left(\frac{1}{\epsilon_{\text{ref}}} - \frac{1}{\epsilon_s} \right) \quad (1)$$

Rehm – Weller equation

$E_{\text{ox}}(\text{D})$ and $E_{\text{red}}(\text{A})$ are the first oxidation potential of the
donor perylene bisimide and first reduction potential of the
acceptor DAE, respectively. E_{00} signifies the spectroscopic
excited state energy, while R_{CC} is the distance between the
centers of the donor and acceptor moieties. The effective
ionic radii of the donor radical cation and acceptor radical
anion are labeled as r^+ and r^- , respectively. The dielectric
constant of the reference solvent used in electrochemistry is
denoted as ϵ_{ref} , while ϵ_s is the dielectric constant of the given
solvent.^[13]

Compounds **1o** and **1c** exhibit the characteristic UV/Vis
spectra of 1,7-dipyrro-
lidinylperylene bisimides, with an
absorption maximum at around 700 nm. The spectrum of **1c**
displays some additional bands (in particular, a broadened
absorption band between 450 and 550 nm, see Figure 3) that
are attributed to the closed form of DAE. From the UV/Vis
and fluorescence spectra recorded in dichloromethane an
 S_0 – S_1 excitation energy of 1.71 eV was estimated for **1o** and
1c.

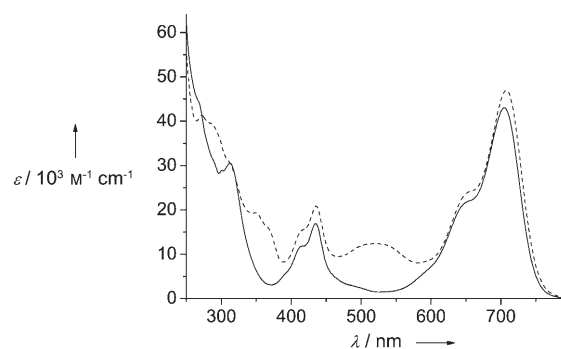


Figure 3. UV/Vis spectra of **1o** (solid line) and **1c** (dashed line) in
dichloromethane.

The redox potentials of **1o** and **1c** and the reference
compounds (see Table S2 in the Supporting Information)
were obtained by cyclic voltammetry measurements in
dichloromethane, with ferrocene used as an internal reference
(Figure 4). The cyclic voltammogram of **1o** is identical to that
of the reference dye **2** (see Figure S1 in the Supporting
Information). Thus, additional redox processes associated
with the DAE_O moiety are apparently not observable within
the available potential window. The first half-wave oxidation
potentials of the PBI moiety were observed at 0.22 V and
0.21 V for **1o** and **1c**, respectively. The closed form **1c** showed
additional waves for the DAE moiety at -1.60 V and 1.00 V,
in agreement with the reference DAE compound **3c** bearing a
phthalic imide residue (see Figure S3 in the Supporting
Information). The first reduction potential of **3o** was
observed at -1.98 V while the respective wave for **1o** was
not observable within the available potential range (up to ca.
 -2.0 V). Donor–acceptor distances of 1.32 nm and 1.24 nm
were estimated from the optimized geometries of **1o** and **1c**
conjugates, respectively.

Since all experimental values were determined in
dichloromethane, the solvent-related term in the Rehm–
Weller equation can be neglected in the present case, and by

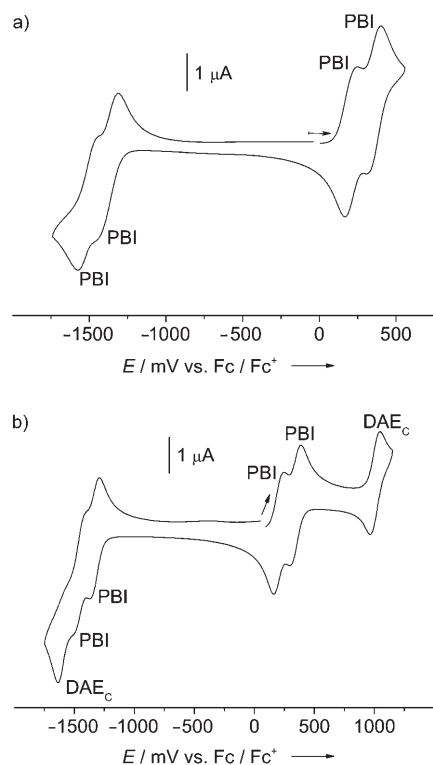


Figure 4. Cyclic voltammograms of a) **1o** (2.7×10^{-4} M) and b) **1c** (3.8×10^{-4} M) in dichloromethane with tetrabutylammonium hexafluorophosphate (0.1 M) as the supporting electrolyte, scan rate 100 mVs^{-1} . Fc/Fc^+ = ferrocene/ferrocenium couple.

applying the obtained data in this equation, a ΔG° value of 0.37 eV for **1o** and a slightly exergonic value of -0.03 eV for **1c** in dichloromethane can be calculated. These energy values imply that the electron-transfer process, clearly endergonic for the open form, is thermodynamically feasible only for the closed form in dichloromethane. In more polar solvents such as dimethyl sulfoxide ($\epsilon_r = 46.68$), the CT state for **1c** ($\text{DAEc}^- - \text{PBI}^+$) becomes more stable^[14] and PET is more likely to take place (Figure 2). Fluorescence lifetime measurements in dimethyl sulfoxide solution provided values of 1.9 ns for **1o** and 0.23 ns for **1c**, thus indicating that the excited fluorophore is subject to an additional quenching process when the photochrome is in its closed form (see Table S3 in the Supporting Information).

Photochromic compounds **1o** and **1c** were also investigated by femtosecond transient absorption spectroscopy in dimethyl sulfoxide by excitation at 680 nm (the $S_0 \rightarrow S_1$ transition of the perylene bisimide dye). An intense bleaching between 650 and 770 nm that arises from the depopulation of the ground-state molecules and the stimulated emission between 770 and 800 nm is observed as a combined negative signal in the spectrum of **1o** (Figure 5a) that decays with a lifetime of approximately 1.5 ns. Compound **1c** shows the same qualitative behavior (Figure 5b), but with a greatly reduced lifetime of 230 ps. The fact that the quenching of the dye S_1 state is not accompanied by a distinctive accumulation of a transient charge-separated species suggests that charge separation is slower than charge recombination and the

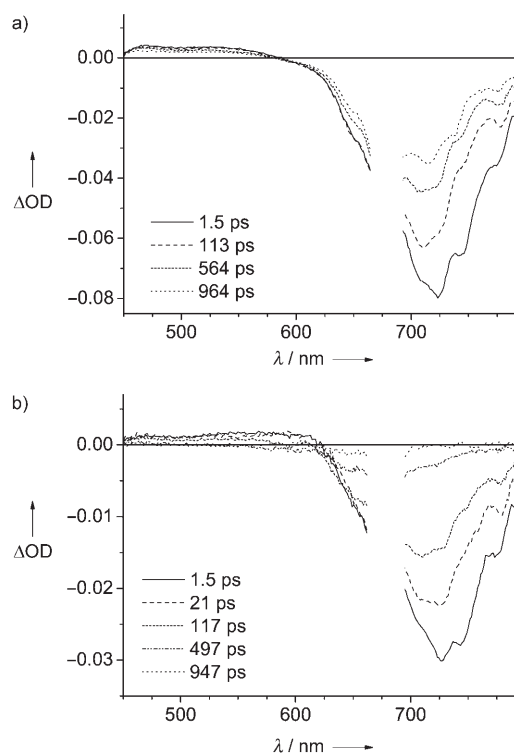


Figure 5. Ultrafast transient absorption spectra of a) **1o** and b) **1c** in dimethyl sulfoxide; $\lambda_{\text{ex}} = 680$ nm.

former is the rate-determining step in $^*\text{PBI} - \text{DAEc}$ deactivation. Clear evidence for the occurrence of photoinduced electron transfer is obtained from femtosecond transient absorption experiments by excitation at 400 nm (the $S_0 \rightarrow S_2$ transition of the dye). In this case, while the behavior of **1o** is virtually identical to that observed upon excitation at 680 nm (see Figure S4 in the Supporting Information), an additional short-lived positive absorption band with a maximum centered at 613 nm is observed for **1c** (Figure 6). The latter can be assigned to the radical cation of the 1,7-pyrrolidinylperylene bisimide moiety,^[15] which was verified by spectroelectrochemistry (see Figure S5 in the Supporting Information). In this case, population of the $\text{DAEc}^- - \text{PBI}^+$ state from the upper S_2 state of the dye, rather than from the S_1 state, is fast

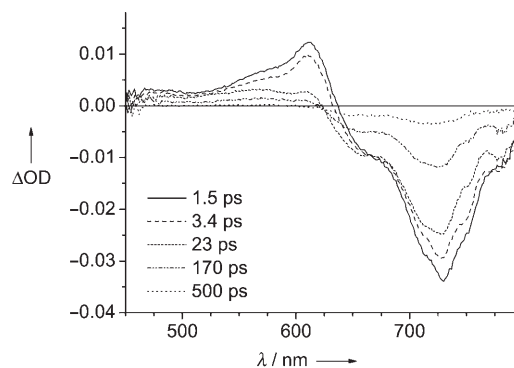


Figure 6. Ultrafast transient absorption spectra of **1c** in dimethyl sulfoxide; $\lambda_{\text{ex}} = 400$ nm.

enough to lead to appreciable transient accumulation of the charge-separated product.

In summary, this new design of photoswitch is effective in a polar environment where an intramolecular electron transfer from the excited PBI dye to the closed form, but not to the open form of the DAE takes place, leading to fluorescence quenching by a PET mechanism. Thus, a prototype photochromic switch for nondestructive readout optical memory systems has been introduced.

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