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New Synthetic Routes towards Soluble and Dissymmetric Triphenodioxazine Dyes Designed for Dye-Sensitized Solar Cells

Yohann Nicolas,*[a] Fouzia Allama,[a] Marc Lepeltier,[a] Julien Massin,[a] Frédéric Castet,[a] Laurent Ducasse,[a] Lionel Hirsch,[b] Zahia Boubegtiten,[c] Gediminas Jonusauskas,[c] Céline Olivier,[a] and Thierry Toupance* [a]

Abstract: New π-conjugated structures are constantly the subject of research in dyes and pigments industry and electronic organic field. In this context, the triphenodioxazine (TPDO) core has often been used as efficient photostable pigments and once integrated in air stable n-type organic field-effect transistor (OFET). However, little attention has been paid to the TPDO core as soluble materials for optoelectronic devices, possibly due to the harsh synthetic conditions and the insolubility of many compounds. To benefit from the photostability of TPDO in dye-sensitized solar cells (DSCs), an original synthetic pathway has been established to provide soluble and dissymmetric molecules applied to a suitable design for the sensitizers of DSC. The study has been pursued by the theoretical modeling of opto-electronic properties, the optical and electronic characterizations of dyes and elaboration of efficient devices. The discovery of new synthetic pathways opens the way to innovative designs of TPDO for materials used in organic electronics.

Introduction

Stimulated by the ever-ongoing efforts to fabricate cost-effective, lightweight, and efficient functional devices, organic alternatives to inorganic counterparts with tunable optoelectronic properties have attracted worldwide attention over the past two decades.[1] Thus, the research of original π-conjugated structures is constantly evolving to address the main issues in organic electronics such as the fine tuning of energy band gaps[2] to improve light absorption, the understanding of intermolecular arrangement by molecular design[3] to modulate the transfer integral and the charge carrier mobility in organic field-effect transistors (OFETs), or to improve the stability of the device.[4]

In this context, few new and efficient π-conjugated skeletons were directly inspired by the synthetic dyes and industrial organic pigments.[5] For instance, the diketopyrrolopyrrole core included in Ferrari-red pigment and isoidigo core has been recently and widely used as monomer for p-type polymers in organic photovoltaics.[6] Perylenes, which are also well-known pigments, were employed to increase the air-stability of n-type materials for OFETs.[7] The same cores have been successfully integrated in the design of sensitizers for dye-sensitized solar cells (DSCs).[8] On the other hand, motivated by the limited metal (as ruthenium or zinc) resource and their good flexibility in molecular tailoring, metal-free organic dyes have been developed with success for DSCs.[9] The main efforts have been devoted to the enhancement of the light-harvesting properties of the organic dyes, the optimization of the first oxidation potential level, the control of the internal charge-transfer processes and the improvement of the dye photostability.[10] So far, the donor-π-bridge-acceptor (D-π-A) design, which includes electron donor and acceptor groups connected through a π-conjugated linker, appeared to be one of the most promising arrangement to get efficient organic dyes for DSCs.

On the other hand, triphenodioxazines (TPDOs), which are well-known industrial molecules,[11] have been used as photostable pigments (i.e., C.I. PV23) and air-stable semiconducting materials in OFET.[12] Recently, TPDO derivatives have also shown high dichroic luminescence in host liquid crystals,[13] which opens the window for their integration into advanced optical devices. In addition, for the first time, a TPDO derivative has been integrated as electro-active material in organic solar cells to enhance the photoconversion efficiency for MEH-PPV:PCBM bulk heterojunctions.[14] Related compounds as phenoxazine derivatives have already been grafted on titanium oxide for efficient DSSCs due to the facile modification of their structures, low dye aggregations, and electron-donating prop-
Another π-conjugated pentacycle containing two sp² nitrogen atoms and two oxygen atoms as TPDO has newly been synthesized to increase the stability of acene derivatives and to modulate the aromaticity.\(^{[16]}\)

TPDO derivatives can therefore be envisioned as a π-conjugated platform to be integrated into D-π-A dye architectures. Nevertheless, the low solubility in common solvents and harsh synthetic conditions has limited their integration into organic electronic devices. Moreover, only one example of dissymmetrical TPDO has been reported so far.\(^{[17]}\) To implement stable TPDO structures for DSCs, we herein describe new synthetic routes extending the possible design towards soluble and unsymmetrical TPDOs substituted by numerous chemical functions such as tert-butyl solubilizing groups, electron-withdrawing or electron-releasing groups on well-defined positions (Scheme 1). Optical and electronic properties along with photobleaching of these new dyes were determined and further rationalized by the means of theoretical modeling. Finally, TPDO dyes yielded very promising power conversion efficiencies (PCE up to 6.3%) in DSCs.

### Results and Discussion

**Synthesis of dyes**

The conventional synthetic pathway towards TPDOs\(^{[11c, 18]}\) (Scheme 2) based on the nucleophilic substitution of tetrahalogeno-1,4-benzoquinone by aniline derivatives followed by an oxidation/cyclization process under harsh conditions has been unsuccessfully tested to synthesize the soluble di-alkylated TPDO 10 from 2,5-di-tert-butyl-3,6-dichlorobenzo-1,4-quinone 12.

Consequently, a new synthetic strategy involving two main steps (Scheme 3) has been established and initially applied to the simplest design of the soluble TPDO 10, by including tert-butyl groups as a solubilizing moiety. Firstly, the nucleophilic substitution of halogenobenzoquinone 12 by potassium 2-nitrophenoxide 11 resulted in the bond formation between the oxygen and the internal cycle.

Then, compound 13 was reduced either by sodium dithionite in water/ethanol or iron/acetic acid mixture to give the amine functions while leading to the hydroquinone. Subsequent oxidation of the hydroquinone moiety by air or chloranil was followed by immediate intramolecular condensation between the amino groups and the carbonyl functions of the

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**Scheme 1.** TPDO dyes designed for DSC.

**Scheme 2.** Usual synthetic pathway toward TPDOs. Reaction conditions: i) AcOEt; ii) PhNO₂, benzoyl chloride, heat.

**Scheme 3.** New synthetic pathway toward soluble TPDO 10. i) DMF, 100 °C, 10 h; ii) Fe, THF, AcOH, reflux, 1.5 h; iii) Chloranil or air, overnight.
quinone to yield the new TPDO 10. As requested for further integration in DSC, compound 10 was soluble in common organic solvents as tetrahydrofuran, chloroform and pentane.

To reach the target disymmetric push–pull molecules built from the soluble TPDO core, three different synthetic routes can be considered. First of all, two different ortho-nitrophenol derivatives could be sequentially added to the quinone 12 followed by the cyclisation step described for 10. Unfortunately, this method led to low yield when the nitrophenol is substituted by a deactivated group in the para position such as methyl 4-hydroxy-3-nitrobenzoate. The second choice might consist in synthesizing a dibrominated TPDO and, then, introducing various functional groups through catalytic couplings. For example, the diphenylamine substituent has been introduced by a Buchwald–Hartwig reaction with 1:1 molar stoichiometric ratio but led in equal quantity to the mono- and bis-adduct. As the unsymmetrical TPDOs were not straightforwardly obtained by this second pathway, a third methodology has been developed. Thus, the most efficient synthetic way appeared to be the formation of the first oxazine cycle by the reaction between methyl 4-amino-3-hydroxybenzoate 14 and quinone 12 (Scheme 4). As the phenoxazine 15 was less reactive than the quinone 12, the reaction stopped after the first addition of aminophenol 14. Then, nucleophilic substitution of the remaining chlorine atom in the tricyclic intermediate 15 by 5-bromo-2-nitrophenol followed by a reduction/oxidation step as described for 10 led to the pre-functionalized disymmetric TPDO 1E. During the two first steps, the amount of byproducts was limited and the unreacted substrates could be easily recycled. Then, the substitution of the bromine atom in 1E allowed us to introduce many donor groups as diethylamine, benzamide, diphenylamine, and di(4-methoxyphenyl)amine,[19] and led to introduce many donor groups as diethylamine, benzamide, diphenylamine, and di(4-methoxyphenyl)amine,[19] and led to many donor groups as diethylamine, benzamide, diphenylamine, and di(4-methoxyphenyl)amine,[19] and led to many donor groups as diethylamine, benzamide, diphenylamine, and di(4-methoxyphenyl)amine,[19] and led to many donor groups as diethylamine, benzamide, diphenylamine, and di(4-methoxyphenyl)amine,[19] and led to many donor groups as diethylamine, benzamide, diphenylamine, and di(4-methoxyphenyl)amine,[19] and led to many donor groups as diethylamine, benzamide, diphenylamine, and di(4-methoxyphenyl)amine,[19] and led to many donor groups as diethylamine, benzamide, diphenylamine, and di(4-methoxyphenyl)amine,[19] and led to many donor groups as diethylamine, benzamide, diphenylamine, and di(4-methoxyphenyl)amine,[19], respectively). It is also worth mentioning that, during the Buchwald–Hartwig coupling with aroylamine substrates, transesterification occurred with sodium tert-butoxide. Consequently, the released sodium methoxide reacted on the brominated position to provide the methoxytriphenodioxazine ester derivative 2E. On the other hand, the use of the 4-substituted aminophenol and nitrophenol instead of the 5-substituted isomers led to new TPDO isomers. Thus, meta derivatives analogous to TPDO para-substituted by benzamide and diphenyl amine groups have been synthesized (8E, 9E). Alternatively, the following electron-releasing group (morpholine) was directly introduced by the use of the substituted nitrophenol instead of the brominated derivative to lead to the TPDO 4E. Finally, saponification of the TPDO ester derivatives followed by the acidification of reaction mixture yielded the corresponding carboxylic acids (1A–9A).

Therefore, the new synthetic strategies open the way for original soluble and disymmetric TPDOs substituted by new chemical functions with a fine control of the position isomerism and led to nine suitable dyes for DSCs (Scheme 1).

Crystallography

Although solution NMR spectroscopy and HRMS data were consistent with the expected structures for the triphenodioxazines described above, and after growing suitable crystals by slow evaporation of solutions in acetonitrile, the crystal structure of the intermediates 15, 16 (see the Supporting Information), and the ester derivative 6E (Figure 1) were solved by X-ray diffraction analysis to prove the isomerism and the structure of central rings. Each structure was consistent with that expected from the synthetic pathway. Moreover, the positions of the three internal rings and the tert-butyl substituents have been established and revealed the distortion of the fused rings due to steric hindrance caused by the bulky alkyl groups (bending angle $\approx 3^\circ$, twist angle $\approx 20^\circ$). Overall, the TPDO core was almost planar and should be highly conjugated, as further confirmed by quantum chemical calculations. Finally, the structures corroborated the invariability of isomerism throughout the synthesis.
Electronic properties

The combination of UV/Visible absorption and time-resolved emission spectroscopic investigations along with electrochemistry studies allowed an investigation into whether these new dyes showed the expected electronic properties to be efficient in DSCs (Table 1). As the acidic derivatives were only sparingly soluble in polar solvents, the study was focused on the ester analogues. Moreover, the absorption spectra of ester and acid derivatives were similar (see the Supporting Information). First of all, the absorption properties of each TPDO ester dyes were examined to better understand the effect of the nature and position of the electron-donating group in the TPDO core.

Each TPDO ester dye exhibited a strong and broad absorption band in the visible range with maximum molar absorptivity ($\varepsilon$) higher than 40 000 L mol$^{-1}$ cm$^{-1}$ in dichloromethane. Furthermore, the strongest electron-releasing groups (ERG) as diarylamine and dialkylamine induced a significant redshift of the maximum absorption wavelength for the dyes substituted in the para position (Figure 2). Thus, compared to that of the brominated derivative 1E, the maximum absorption band of 7E was redshifted by more than 60 nm whereas the meta-substituted analogue 9E led to a blueshift of 71 nm. These behaviors were consistent with the expected push–pull effect and the results of theoretical modeling.

Moreover, the vibrational fine structure of absorption bands linked to the rigidity of the TPDO core disappeared in the case of the push–pull structures due to the extension of frontier orbitals through free rotational bond. In the case of TPDOs meta-substituted by a strong ERG (i.e., compound 9E), a second band in the red range appeared extending the absorption from 350 to 620 nm (Figure 2). In contrast to the benzamide substituent, the absorption band redshifted from the meta to

![Figure 1. X-ray crystal structure of 6E.](image)

![Figure 2. Absorption spectrum of a) para isomers, b) meta isomers and their corresponding para isomers in dichloromethane.](image)

### Table 1. Photoelectronic properties of ester dyes.

| Dye | $\lambda_{\text{max}}$ [nm] | $\lambda_{\text{max}}$ [nm] | $E_{\text{F}}$ vs NHE | $E_{\text{F*$}}$ vs NHE | Photodestr. yield ($10^6$) |
|-----|---------------------------|---------------------------|---------------------|---------------------|-----------------------------|
| 1E  | 521 (4.7)                 | 547 (9.4) (440)           | 1.50                | 0.91                | 5.3                         |
| 2E  | 528 (4.5)                 | 633 (3.1) (810)           | 1.20                | 1.06                | 13                          |
| 3E  | 581 (4.5)                 | N/A                       | 0.80                | 1.16                | 21                          |
| 4E  | 551 (4.4)                 | 702 (6.6) (330)           | 0.91                | 1.17                | 21                          |
| 5E  | 536 (4.7)                 | 624 (23.6) (980)          | 1.20                | 1.07                | 9.1                         |
| 6E  | 571 (4.7)                 | 687 (2.5) (123, 623)      | 0.97                | 1.01                | 5.0                         |
| 7E  | 585 (4.6)                 | 752 (0.87) (65)           | 0.87                | 1.10                | 4.5                         |
| 8E  | 512 (4.6)                 | 568 (18.5) (870)          | 1.34                | 1.20                | 4.2                         |
| 9E  | 450 (4.4)                 | 546 (0.16) (900, 3100)    | 0.87                | N/A                 | 27                          |

(a) Absorption maximum wavelength in visible range of the dye solution in CH$_2$Cl$_2$ and molar absorption coefficient. (b) Emission maximum wavelength in CH$_2$Cl$_2$, quantum yield of fluorescence, and fluorescence lifetime. (c) Cyclic voltammetry measured in a solution of 0.1 M Bu$_4$NPF$_6$ in CH$_2$Cl$_2$, versus Fe$^+/Fe$, corrected by adding 0.7 V. (d) Estimated by subtracting the onset of the absorption band in solution to $E_{\text{F}}$. (e) Measured in CH$_2$Cl$_2$ solution.
the para isomer. Finally, the absorption band of dyes was centered on the strongest emissive part of the solar spectrum.

Fluorescent quantum yields and lifetimes were also determined. As expected, the fluorescent efficiency of push–pull dyes or molecules substituted by heaviest atoms were weak compared to the other dyes. We should note that a strong discrepancy in the correlation between fluorescence quantum yield and fluorescence lifetime appears for the meta-substituted compound 9E indicating an implication of multiple relaxation pathways from the excited state. Furthermore, some of these dyes appeared to be highly photostable with a photodestruction yield of 5.3 × 10^{-4}, 5.0 × 10^{-6}, and 4.5 × 10^{-6} for 1E, 6E, and 7E respectively, which is better than fluorescein isocyanate (75 × 10^{-6}) used as a reference for photodestruction measurements. By contrast, meta-substituted or alkyl-substituted derivatives were much more unstable with, for instance, a photodestruction yield of 27 × 10^{-4} for 9E.

The ground-state oxidation potential (E_{S0} in V) and the highest occupied molecular orbital/lowest unoccupied molecular orbital (HOMO–LUMO) transition energy values were evaluated by cyclic voltammetry and absorption/emission spectroscopy (Table 1). Each compound could be reversibly oxidized, consistent with the formation of stable radical-cations, which is a key requirement for efficient electronic processes in DSCs. The (E_{S0} in V) values fall within a wide range, that is, 0.8–1.5 V versus the normal hydrogen electrode (NHE), but are each more positive than that of the iodide/triiodide redox shuttle revealing that the ground-state regeneration is energetically favorable for DSC applications. On the other hand, the LUMO levels, deduced from the values of half-wave oxidation potential and the onset of the optical gaps were between 0.9 and 1.2 V versus NHE, which yields the free energy necessary to transfer an electron from the first excited state of the chromophore to the TiO_2 conduction band. On the basis of the (E_{S0} in V) and LUMO energy values, a functional DSC elaborated from the TPDO-based sensitizers, TiO_2 semi-conductor, and the iodide/triiodide redox shuttle should be energetically favorable.

Quantum chemical calculations
All calculated structures show a slight distortion of the TPDO moiety (≈ 5°), due to the steric hindrance induced by the tert-butyl groups, in close agreement with the X-ray diffraction data. Moreover, all compounds exhibit a bending in their aromatic core of about 20° (Figure 3), which is consistent with an almost planar structure as observed in their crystalline state.

The low-energy transition energies and wavelengths computed at the MPW1K/6-311G(d) level are reported in Table 2, along with the corresponding oscillator strengths and associated CI weights, charge transferred (q_CT), CT distance (d_CT, Å), and dipole moment variation (Δμ, D), calculated at the TD MPW1K/6 311G(d) level for the lowest energy excitation states of the dyes.

| Mol. | ΔE_{ex} (eV) | λ (nm) | f_{trans} | Dominant transitions | q_{CT} | d_{CT} (Å) | Δμ (D) |
|------|--------------|--------|------------|----------------------|-------|----------|-------|
| 2E   | 2.289        | 542    | 1.636      | H→L (0.70236)        | 0.434 | 1.387    | 2.886 |
| 3E   | 2.072        | 598    | 1.760      | H→L (0.70189)        | 0.468 | 2.246    | 5.291 |
| 4E   | 2.149        | 577    | 1.747      | H→L (0.70174)        | 0.480 | 2.148    | 4.941 |
| 5E   | 2.264        | 548    | 1.788      | H→L (0.70175)        | 0.434 | 2.187    | 2.664 |
| 6E   | 2.109        | 588    | 1.816      | H→L (0.69746)        | 0.533 | 2.814    | 7.221 |
| 7E   | 2.057        | 603    | 1.842      | H→L (0.69537)        | 0.553 | 3.114    | 8.242 |
| 8E   | 2.421        | 512    | 1.526      | H→L (0.70176)        | 0.443 | 0.866    | 1.869 |
| 9E   | 2.245        | 552    | 1.070      | H→L (0.67518)        | 0.680 | 3.716    | 12.137|
|      | H→L (0.17906) |       |            |                      |       |          |       |
|      | 2.780        | 446    | 0.587      | H→L (0.17871)        |       |          |       |
|      | H→L (0.65880) |       |            |                      |       |          |       |
|      | H→L (0.15533) |       |            |                      |       |          |       |

Figure 3. Front and side views of 6E, as optimized at the B3LYP/631+G(d,p) level.
mien. In accordance with the push–pull effects revealed by the spatial distribution of the MOs, and with the redox potential measured by electrochemistry, these compounds have the smallest HOMO–LUMO gaps among the series. Consistent with the UV/Vis measurements, meta-substituted compounds 8E and 9E display smaller maximum absorption wavelengths than their para-substituted analogues 5E and 7E. The significant blueshift observed when going from para to meta substitution is a consequence of a decrease in the efficiency of the electron π-conjugation in the ground state, as illustrated Figure S27 (the Supporting Information). The larger electron π-conjugation of para-substituted compounds is further confirmed by their higher stability (5E and 7E are more stable than 8E and 9E by 2.7 and 0.8 kcal mol⁻¹, respectively), as well as by the detailed analysis of the bond distances along the TPDO moiety (Table S5, the Supporting Information), which reveals a more pronounced contribution of the zwitterionic form (∆D) in the ground-state wave function. To gain more quantitative insights on the charge-transfer phenomena occurring upon photoexcitation of the dyes, the amount of charge transferred (∆q), as well as the CT distance (dCT, that is, the spatial extent of the charge delocalization) were evaluated from the difference ∆ρ between the ground- and excited-state electron densities, using the procedure described in Ref. [22]. In this procedure, ∆ρ is split into two contributions, ρ+ and ρ−, which correspond respectively to the increase (∆ρ>0) and decrease (∆ρ<0) of the density. The amount of charge transferred, qCT, is defined as the integral over space of the ρ+ (or ρ−) function, whereas dCT is the distance separating the barycenters of these two functions. The qCT and dCT values computed at the MPW1K/6-311G(d) level, as well as the light-induced dipole moment variation, are gathered in Table 2. Variation density plots are reported in the Supporting Information for all dyes.

As a general trend, it is found that the transition wavelengths within the series globally increase with ∆u and the CT distance. As illustrated in Figures S25 and S26 (the Supporting Information), the four dyes exhibiting the larger CT distances (3E, 4E, 6E, and 7E, in which dCT>2 Å) give rise to larger wavelengths, whereas compounds 2E, 5E, and 8E, in which the charge transfer occurs over smaller distances, display smaller transition wavelengths. A noticeable exception to this trend is observed for the meta-substituted compound 9E, for which the wavelength is unexpectedly low owing to its dipole moment variation and CT distance. The large values of ∆u and dCT calculated for this compound arise from a small spatial overlap between the electron-donating and electron-accepting regions, as confirmed by the small value of the Tozer’s parameter [22] (see Table S2, the Supporting Information). As discussed above, the small maximum absorption wavelength obtained for 9E is a consequence of a poor electron conjugation in the ground state. One also notes that, contrary to the corresponding para compounds, the light-induced density change is not localized on carboxyl group in meta derivatives, which is expected to reduce the efficiency of the photoexcited electron injection into the TiO2 conduction band.

**Figure 4.** Difference between excited and ground state densities (∆ρ) for 7E (top) and 9E (bottom), calculated at the MPW1K/6 311G(d) level. Blue and violet lobes are associated to positive and negative differences, respectively.

**TPDO-Sensitization of porous TiO2 films and performances of DSC**

Dyes 2A–9A were first chemisorbed onto 3 μm-thick nanoparticulate anatase TiO2 films in the presence or absence of cheno-deoxycholic acid (CDCA) as a co-adsorbent. [24] Regardless of the dye nature, the shape of absorption spectrum of the dye-modified TiO2 layers showed strong and broad absorption bands in the visible range (Figure 5). Moreover, a significant hypsochromic shift of the maximum absorption band compared to the spectrum in solution was clearly evidenced (Figure 5), this effect being even more pronounced without co-adsorbent. This blueshift could be attributed to the dye aggregation at the oxide surface but also to the coupling between the dyes and TiO2, as it was recently proposed in the case of triarylaniline dyes. [25]

Then, two types of devices were elaborated involving TPDO-sensitized nanoporous TiO2 films of two different thicknesses coated over transparent conductive electrode as anode, a commercial iodide/triiodide based liquid electrolyte and a platinum counter electrode (see the Experimental Section). For thick porous TiO2 films (9 μm-thick TiO2, 20 nm nanoparticles + 6 μm light-scattering TiO2 particles), the para-substituted dyes 2A–7A showed good photovoltaic responses with power conversion efficiency (PCE, η) ranging from 1.91 to 6.30% under AM1.5G illumination (100 mA/cm²), the best performances being reached with the arylamine donor groups (Table 3 and the Supporting Information). Thus, the diphenylamine-substi-
tuted TPDO 6A led to η of 6.30% with a short-circuit current density ($J_{sc}$) of 12.44 mA cm$^{-2}$, an open-circuit photovoltage ($V_{oc}$) of 685 mV and a fill factor (ff) of 74.1%. This performance constitutes one the best ever reported for such a simple design. Furthermore, the incident photon-to-electron conversion efficiency (IPCE) measured under light bias for the best cells (see the Supporting Information) exhibited a maximum efficiency centered on the maximum absorption band of thin film to exceed 60% in the case of 6A (Figure 6). The good agreement between the photocurrent action spectrum and the UV/Visible absorption spectrum of the 6A-modified TiO$_2$ electrodes definitely shows that the sensitization process actually takes place. For the 9A dye, the photoconversion was equally observed at 450 and 550 nm, independently to the nature of absorption bands. By contrast, the meta isomers 8A–9A led to a drastic collapse of the power conversion efficiencies mainly related to a collapse in $J_{sc}$. For instance, the photocurrent density for the 9A dropped by 62% from para to meta substitution, which evidences the key role of the para substitution to reach good photoconversion efficiencies. As discussed above, the lack of frontier orbital electron density and light-induced density changes on the anchoring group could explain lower injection rates leading to the downfall of $J_{sc}$.

Recent development in DSCs have shown that the replacement of the corrosive and volatile iodide/triiodide based liquid electrolyte by other redox shuttles (as Co$^{2+}$/Co$^{3+}$) or solid hole-transporters requires the use of thinner TiO$_2$ photoanodes. To evaluate the performance of TPDO-based devices built with thinner TiO$_2$ film (3 μm-thick TiO$_2$ 20 nm nanoparticles + 6 μm light-scattering TiO$_2$ particles), three soaking methods have been compared: 1) Dipping in a pure solution of the dyes; 2) Dipping in a solution containing both dye and CDCA; 3) Successive soaking in the dye solution and, then, in a CDCA bath. The photovoltaic performances of the corresponding devices were still high for diphenylamino and amido donor substituents in the para position. Thus, for para-benzamide TPDO 5A and the same soaking method, the overall energy conversion yield remained unchanged (η$_{oc}$ = 8.44 mA cm$^{-2}$, $V_{oc}$ = 0.699 V, ff = 0.716, η = 4.22%). The weaker current density, which could be due to the decrease of light absorption, was compensated by a higher open-circuit voltage. On the other hand, the photoconversion efficiency of our best dye 6A obtained for thin film reached 4.61, 4.54, and 5.06%, respectively, for soaking in the solution of the pure dyes, in a solution of both dye and CDCA, and in two baths as described above (Figure 7 and the

![Figure 5](image_url). a) Normalized spectrum of thin nanoporous TiO$_2$ film sensitized by dyes 2A–9A. b) Absorption spectrum of a 6E solution ( ), a thin film sensitized by 6A only ( ) and with co-adsorbent CDCA ( ).

![Figure 6](image_url). IPCE of 6A sensitized DSC ( ) compared to the absorption spectrum of 6A sensitized TiO$_2$ layer. ( ).

| Dyes | $J_{sc}$ [mA cm$^{-2}$] | $V_{oc}$ [mV] | ff [%] | η [%] |
|------|-------------------|--------------|--------|-------|
| 2A   | 6.36              | 649          | 67.6   | 2.79  |
| 3A   | 4.57              | 578          | 72.1   | 1.91  |
| 4A   | 6.77              | 640          | 68.2   | 2.96  |
| 5A   | 8.70              | 679          | 72.8   | 4.30  |
| 6A   | 12.44             | 685          | 74.1   | 6.30  |
| 7A   | 9.93              | 644          | 72.4   | 4.62  |
| 8A   | 1.18              | 553          | 70.7   | 0.46  |
| 9A   | 3.79              | 634          | 71.5   | 1.72  |

[a] Short circuit current density. [b] Open circuit photovoltage. [c] Fill factor. [d] Power conversion efficiency.
Supporting Information). Although the part of aggregated dyes was greater without CDCA and contrary to what we observed for other dyes, the performances remained high.

Conclusion

An original and versatile route has been successfully established to synthesize nine new TPDOs possessing high solubility and unsymmetrical structures. This synthetic pathway allowed for controlling precisely the position isomerism as evidenced and unsymmetrical structures. This synthetic pathway allowed for controlling precisely the position isomerism as evidenced by X-ray diffraction of single crystals of TPDOs. Moreover, the molecular structure revealed the bending of the π-conjugated system, which was confirmed by modeling, due to the steric hindrance of two tert-butyl substituents. On the other hand, the TPDO derivatives covered a large part of the solar spectrum with strong molar extension coefficients while preserving the high photo-stability, commonly observed for TPDO pigments. The visible absorption band has been attributed to the charge transfer between the two external substituents, consistently with the push–pull structure of the dyestuffs. Moreover, the HOMO and LUMO energy levels calculated at the DFT level and estimated from cyclic voltammetry were suitable for the integration of TPDO dye in DSC containing titanium oxide as transparent semiconductor and iodide/triiodide as redox shuttles. As a consequence, all these properties allowed us to integrate the TPDO core into DSC by soaking of porous TiO₂ films, which are currently in progress in our laboratories. These latter should provide additional information for optimizing the performances of the next generation of TPDO dyes.

Figure 7. I/V curves of DSCs made of 3 μm thick nanoparticulate anatase TiO₂ films sensitized with a pure solution of 6A ( ), a solution of 6A and CDCA ( ), by successive soak in a solution of 6A and solution of CDCA ( ).

Experimental Section

Materials and general methods

1H and 13C NMR spectra were recorded on a DPX-300 spectrometer (δ given in ppm relative to tetramethylsilane). Electrospray mass spectra (ESI-TOF HRMS) were carried out by the CESAMO (Bordeaux, France) on a Qstar Elite mass spectrometer (Applied Biosystems). The electrospray needle was operated at room temperature. Spectra were recorded in the positive-ion mode using the reflextron and with an accelerating voltage of 20 kV (high resolution). Elemental microanalyses were performed at the CESAMO (Bordeaux, France). UV/Visible absorption studies have been carried out with a UV-1650PC SHIMADZU and Cary 5G (Varian) spectrophotometers. Steady-state emission spectra were recorded on Fluorolog (Jobin Yvon) spectrophluorimeter. Time-resolved fluorescence was measured with a streak camera Hamamatsu 5680 equipped with a fast single sweep unit MS676 (temporal resolution 2 ps) and coupled to a spectrograph (Chromex 250). The fluorescence excitation light pulses were obtained from Ti:Sapphire femtosecond laser system (Femtopower Compact Pro) followed by an optical parametric generator with frequency mixers (Topas C). The procedure to determine photodestruction quantum yields has been described elsewhere[31] and consists in the measurement of absorption decay during irradiation of the solution at appropriate wavelength followed by a mathematical treatment of the obtained kinetic. Cyclic voltammetry studies were carried out with a potentiostat/galvanostat Autolab PGSTAT100 using a three-electrode device (working electrode: Pt disc; reference electrode: Ag/AgCl calibrated with ferrocenium/ferrocene as the internal reference; counter electrode: Pt).

Syntheses

2,5-di-tert-butyl-3,6-bis(2-nitrophenoxy)-1,4-benzoquinone (13): Under an inert atmosphere, potassium nitrophenoxide (1.51 g, 8.52 mmol) was added to a solution of 2,5-di-tert-butyl-3,6-dichloro-1,4-benzoquinone 12 (1.231 g, 4.26 mmol) in DMF (10 mL). After stirring at 100 °C for 10 h, the mixture was poured into chloroform (50 mL). Then, the organic phase was washed with HCl 1M aqueous solution (50 mL) and NaOH 1M aqueous solution (50 mL). After evaporation of the volatiles, the brown solid was successively rinsed with water, methanol, and pentane to give a yellow powder (1.48 g, 70 %).1H NMR (400 MHz, [D₆]DMSO, 25 °C, TMS): δ 8.05 (d, 3J(H,H) 2 Hz, 1H), 7.64 (t, 3J(H,H) 8.1 Hz, 2H), 7.49 (s, 1H).13C NMR (101 MHz, [D₆]DMSO, 25 °C, TMS): δ 181.64, 149.66, 149.41, 139.13, 138.79, 134.40, 125.36, 123.08, 118.18, 35.27, 29.53 ppm; HRMS (ESI): m/z calcd for C₂₆H₂₆N₂O₈+: 517.1581 [M+Na⁺]; found: 517.1598.
ethyl acetate (300 mL). The organic phase was then washed with chloroform (30 mL) to remove the excess of iron. After addition of chloranil (2.72 g, 11.1 mmol) and stirring overnight, the organic phase was washed with water and 2% Na2CO3 aqueous solution, dried over MgSO4 and concentrated. The crude material was purified by column chromatography over silica gel (petroleum ether/chloroform 9:1) and isolated as a red powder (195 mg, 44%). 1H NMR (400 MHz, CDCl3, 25 °C, TMS) δ 7.42 (dd, JHH 7.7 Hz, JHH 1.6 Hz, 2H), 7.17 (td, JHH 7.7 Hz, JHH 1.4 Hz, 2H), 7.02 (dd, JHH 7.9 Hz, JHH 1.3 Hz, 2H), 1.64 ppm (s, 18H); 13C NMR (101 MHz, CDCl3, 25 °C, TMS) δ 151.16, 144.28, 143.81, 133.33, 128.50, 129.74, 122.38, 114.60, 37.07, 39.01 ppm; HRMS (ESI): m/z calc for C22H24ClNO4 + H+: 399.2075 [M+H]+; found: 399.2075.

Methyl 1,4-di-tert-butyl-2-chloro-3-oxo-3H-phenoxazine-7-carboxylate (15): Sodium bicarbonate (143.9 g, 1.73 mol) was added to a solution of methyl 4-amino-3-hydroxybenzoate (17.3 mmol) in dimethylacetamide under nitrogen. The reaction mixture was stirred at 80 °C overnight. Then, hydrochloric acid 1H), 7.86 (d, JHH 1.6 Hz, 1H), 7.46 (d, JHH 8.0 Hz, 1H), 7.30 (m, 3H), 3.96 (s, 3H), 1.65 (s, 9H), 1.64 ppm (s, 9H); 13C NMR (75 MHz, CDCl3, 25 °C, TMS) δ 166.21, 152.77, 150.85, 144.17, 144.08, 143.88, 143.34, 136.73, 132.29, 129.61, 120.05, 128.73, 126.64, 127.57, 123.27, 123.11, 121.62, 117.94, 115.93, 52.49, 37.17, 31.89, 31.85 ppm; HRMS (ESI): m/z calc for C16H19BrNO4 + H+: 535.1252 [M+H]+; found: 535.1293.

Methyl 6,13-di-tert-butyl-3-diphenylamino-phenoxazine-10-carboxylate (16E): Brominated derivative 1E (100 mg, 0.187 mmol) and the sodium tert-butoxide (20 mg, 0.21 mmol) was added to a solution of the diphenylamine (0.187 mmol), [Pd 2(dba)3] (6.8 mg, 7.5 mmol); dba tris(dibenzylidenecacetonato) and diphenylphosphine ferroene (8.3 mg, 15 mmol) in toluene (30 mL) under nitrogen. The reaction mixture was then stirred at reflux during 3 h; it turned dark purple. Then, hydrochloric acid 1M (50 mL) was added. The product was extracted into ethyl acetate (2x100 mL) and the organic phase was washed with brine, dried over MgSO4 and the solvent was evaporated. Then, the product was purified by column chromatography over silica gel (petroleum ether/ethyl acetate 90:10) to yield a purple powder (43%). 1H NMR (300 MHz, CDCl3, TMS) δ: 7.78 (dd, JHH 8.4 Hz, JHH 1.8 Hz, 1H), 7.66 (dd, JHH 1.8 Hz, 1H), 7.36 (m, 7H), 7.18 (m, 7H), 6.84 (dd, JHH 8.7 Hz, JHH 2.4 Hz, 1H), 6.74 (d, JHH 2.4 Hz, 1H), 3.97 (s, 3H), 1.69 (s, 9H), 1.62 ppm (s, 9H); 13C NMR (101 MHz, CDCl3, TMS): δ 166.59, 153.81, 149.52, 148.50, 147.41, 145.33, 144.90, 144.16, 144.09, 137.77, 130.08, 129.34, 128.95, 128.87, 127.40, 125.94, 124.76, 123.75, 122.53, 119.33, 115.99, 107.35, 52.65, 37.49, 37.31, 32.20, 31.96 ppm; IR (ATR): ν̃ 2954, 1723, 1625, 1596, 1590 cm−1; HRMS (ESI): m/z calc for C54H43N3O4 + H+: 826.2862 [M+H]+; found: 826.2846.

6,13-Di-tert-butyl-3-diphenylamino-phenoxazine-10-carboxylic acid (6A): KOH (10 equiv) and a solution of tetrabutylammonium fluoride 1M in THF (0.1 mL) was added to a solution of the ester derivative 6E in toluene. The reaction mixture was heated at reflux for 4 h. Then, HCl 1M (50 mL) was added and the product was extracted into ethyl acetate (3x50 mL). The organic phase was washed with saturated NaCl (aq., 100 mL), dried over MgSO4 and the solvent was evaporated. Concentrated HCl was added to the solution until complete precipitation of the acid (pH 1). Then, the pure acid was filtered, washed with distilled water and dried under vacuum to a purple powder (45%). 1H NMR (300 MHz, CD3THF, TMS): δ: 7.74 (d, JHH 8.1 Hz, 1H), 7.61 (s, 1H), 7.33 (m 6H), 7.12 (m, 6H), 6.80 (dd, JHH 8.7 Hz, JHH 2.1 Hz, 1H), 6.69 (d, JHH 8.4 Hz, JHH 1.6 Hz, 2H), 7.17 (td, JHH 7.7 Hz, JHH 1.4 Hz, 2H), 7.02 (dd, JHH 7.9 Hz, JHH 1.3 Hz, 2H), 1.64 ppm (s, 18H); 13C NMR (101 MHz, CDCl3, 25 °C, TMS) δ: 178.36, 165.62, 151.86, 150.24, 148.46, 145.56, 142.36, 138.43, 137.83, 133.82, 133.02, 129.60, 128.42, 127.35, 126.02, 125.36, 120.42, 116.84, 77.16, 52.88, 37.85, 36.10, 32.09, 30.55 ppm; HRMS (ESI): m/z elemental analysis calcd [%] for C30H20BrNO4 + H+: 583.1080 [M+H]+; found: 583.1100.
Quantum chemical calculations

Molecular geometries were optimized at the density functional theory (DFT) level using the B3LYP exchange-correlation (XC) functional and the 6-311+G(d,p) basis set. The absorption properties of the dyes were calculated using the time-dependent DFT (TD-DFT) method with the 6-311g(d) basis set and the MPW1K XC functional.[29] This hybrid functional, which contains 42.8% of Hartree-Fock (HF) exchange, has been shown to provide absorption spectra in good agreement with experiments for other push pull organic systems.[30] To obtain an accurate description of the photoinduced charge transfer (CT) phenomena along the dyes, we also performed TD-DFT calculations using the long-range corrected ωB97X-D functional, in which the amount of HF exchange increases with the inter-electron distance (22.20% of HF exchange at short-range and 77.80% at long-range, with a range separation parameter of 0.20).[31] Moreover, additional calculations performed using the 6-311g(d) basis set demonstrated that the absorption properties are not significantly affected by the addition of diffuse functions (see the Supporting Information). Solvent effects (dichloromethane) were included both in geometry optimizations and TD-DFT calculations by using the polarizable continuum model in its integral equation formalism (IEF-PCM).[32] All calculations were performed with Gaussian 09.[32]

Device fabrication and characterization

FTO-coated conducting glass substrates (NSG10, 10 ohm/square, glass thickness 3.2 mm, XOPFisica) were cleaned in ethanol followed by an ultrasonic treatment in an alkaline detergent solution. The conducting glass was then treated under UV/O3 during 20 min allowed by an ultrasonic treatment in an alkaline detergent solution. Glass thickness 3.2 mm, XOPFisica) were cleaned in ethanol followed by an ultrasonic treatment in an alkaline detergent solution. The conducting glass was then treated under UV/O3 during 20 min followed by an ultrasonic treatment in an alkaline detergent solution. The role of this co-adsorbent and de-aggregating agent. The sensitization efficiency of 0.20).[30] Moreover, additional calculations performed using the 6-311g(d) basis set demonstrated that the absorption properties are not significantly affected by the addition of diffuse functions (see the Supporting Information). Solvent effects (dichloromethane) were included both in geometry optimizations and TD-DFT calculations by using the polarizable continuum model in its integral equation formalism (IEF-PCM).[32] All calculations were performed with Gaussian 09.[32]

Device fabrication and characterization

FTO-coated conducting glass substrates (NSG10, 10 ohm/square, glass thickness 3.2 mm, XOPFisica) were cleaned in ethanol followed by an ultrasonic treatment in an alkaline detergent solution. The conducting glass was then treated under UV/O3 during 20 min to remove the remaining carbon residues. To increase photoanode adhesion while reducing recombination between TCO and I2 at this interface, the conducting glass was then treated by a 40 nm TiCl4 aqueous solution at 70 °C for 30 min. The photoanode was prepared by using the screen-printing method using commercially available titania pastes. First a transparent layer was deposited. A second layer of around 5 μm thick was printed afterwards. The role of this transparent layer. Ethyl-cellulose and terpineol contained in the titania pastes were removed by gradual thermal treatment under air flow at 325 (5 min), 375 (5 min), 450 (15 min), and 500 °C (15 min). The as-obtained films were then assembled with a platinum counter electrode using a hot-melt Surlyn polymer gasket and the iodine-based electrolyte (EL-PMe, DyeSol) was introduced in the cell by vacuum back-filling as previously described.[30] The as-obtained cells (working area 0.159 cm2 defined using a black mask) were then illuminated by a AM1.5G solar simulator calibrated with a radiometer (IL 1400BL) to provide an incident irradiance of 100 mWcm−2 and the l/I responses were measured by using a Keithley model 2400 digital source meter (Keithley). The incident photon-to-charge carrier efficiency (IPCE) data were collected from 300 to 1000 nm by using a Xe lamp associated with a monochromator (Triax 180, Jobin Yvon). A Bias light was employed to illuminate the cell to reach 10% of Jsc. The current produced was measured by steps of 1 nm after 0.2 s of radiation exposure with a Keithley 6487 picoammeter to be in steady-state conditions. The incident photon flux was measured with a 6-inch diameter calibrated integrated sphere (LabSphere) and a silicon detector.

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Keywords: dyes/pigments · donor–acceptor systems · photostability · solar cells · synthetic methods

1. a) P. M. Beaujuge, J. M. J. Fréchet, J. Am. Chem. Soc. 2011, 133, 20009 20029; b) T. M. Figueira Duarte, K. Mullén, Chem. Rev. 2011, 111, 7260 7314.
2. a) P. M. Beaujuge, C. M. Amb, J. R. Reynolds, Acc. Chem. Res. 2010, 43, 1396 1407.
3. a) C. Wang, H. Dong, W. Hu, Y. Liu, D. Zhu, Chem. Rev. 2012, 112, 2208 2267; b) M. Mas Torrent, C. Rovira, Chem. Rev. 2011, 111, 4833 4856.
4. X. Zhan, A. Facchetti, S. Barlow, T. J. Marks, M. A. Ratner, M. R. Wasielewski, S. R. Marder, Adv. Mater. 2011, 23, 268 284.
5. M. J. Robb, S. Y. Ku, F. G. Brunetti, C. J. Hawker, J. Polym. Sci. Part A 2013, 51, 1263 1271.
6. a) S. Qu, H. Tian, Chem. Commun. 2012, 48, 3039 3051; b) H. Bronstein, Z. Chen, R. S. Ashraf, W. Zhang, J. Du, J. R. Durrant, P. S. Tuladhar, K. Song, S. E. Watkins, Y. Geerts, M. M. Wiemken, R. A. J. Janssen, T. Anthopoulos, L. Searinghouse, M. Hennay, I. Mulillo, J. Am. Chem. Soc. 2011, 133, 3272 3275; c) E. Wang, Z. Ma, Z. Zhang, K. Vondewal, P. Henriksson, O. Inganás, F. Zhang, M. R. Anderson, J. Am. Chem. Soc. 2011, 133, 14244 14247.
7. a) R. Schmidt, J. H. Oh, Y. S. Sun, M. Deppisch, A. M. Krause, K. Radacki, H. Braunschweig, M. Könnemann, P. Erk, Z. Bao, F. Würthner, J. Am. Chem. Soc. 2009, 131, 6215 6228; b) C. P. Piliego, D. Jarzab, G. Gigli, Z. Chen, A. Facchetti, M. A. Loi, Adv. Mater. 2009, 21, 1573 1576; c) B. A. Jones, A. Facchetti, M. R. Wasielewski, T. J. Marks, J. Am. Chem. Soc. 2007, 129, 15259 15278; d) K. Hwang, R. R. Dasari, M. Fenoll, V. Alain Rizzo, A. Dindar, J. W. Shim, N. Deb, C. Fuentes Hernandez, S. Barlow, D. G. Bucknall, P. Audebert, S. R. Marder, B. Kippelen, Adv. Mater. 2012, 24, 4445 4450.
8. a) S. Qu, C. Qin, A. Islam, Y. Wu, W. Zhu, J. Hua, H. Tian, L. Han, Chem. Commun. 2012, 48, 6972 6974; b) T. Edvinsson, C. Li, N. Pschirer, J. Schonboem, F. Eickemeyer, R. Sens, G. Boschloo, A. Herrmann, K. Mullen, A. Hagfeldt, J. Phys. Chem. C 2007, 111, 15137 15140; c) C. Li, J. H. Yum, S. J. Moon, A. Herrmann, F. Eickemeyer, N. G. Pschirer, P. Erk, J. Schoeboem, K. Mullen, M. Grätzel, M. K. Nazeeruddin, ChemSusChem
