Visible and near-infrared organic photosensitizers comprising isoindigo derivatives as chromophores: synthesis, optoelectronic properties and factors limiting their efficiency in dye solar cells

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The development of ruthenium-free organic photosensitizers showing a panchromatic absorption of the solar spectrum up to the near-infrared (IR) region is still scarce in dye-sensitized solar cells (DSSCs). Among the sensitizers with absorption beyond 700nm and developed specifically for DSSCs, only Zinc-phthalocyanine and Boron-dibenzopryromethene-based dyes have been able to reach efficiencies as high as 6%. Here we report six metal-free organic dyes based on isoindigo, thieno-isoindigo or benzo-thieno-isoindigo chromophores that strongly absorb in the UV-visible and near-IR spectral range up to 900nm. These molecules, that exhibit purple, blue or green hue, were used to sensitize TiO₂ mesoporous electrodes in order to fabricate DSSCs with iodide/triiodide-based electrolyte. A power conversion efficiency as high as 5.84% associated to a current density close to 15 mA/cm² was obtained, which is comparable to the best panchromatic organic dyes published so far. Advanced photophysical characterizations, including charge extraction, transient photovoltage, and laser transient absorption spectroscopy experiments, combined to Density Functional Theory (DFT) modeling and computational investigations allow us to fully unravel the interfacial processes at the origin of the solar cell performances and to identify the factors limiting the J₅₆. We also demonstrate in this work that the V₅₆ of the solar cells are linearly correlated to the dipolar moments of the oxidized dyes, the molecules showing the larger dipoles leading to the highest V₅₆ values. This work unravel fundamental processes that could drive new strategies of molecular engineering for organic dyes useful in DSSCs.

Introduction.

Among the third generation photovoltaic technologies, dye-sensitized solar cells (DSSCs) show some of the required features for everyday life applications and a rapid development at the industrial level.¹ Indeed they can combine a high efficiency² along with a good stability,³ and be semi-transparent and colorful, which are substantial advantages for building integrated photovoltaics (BIPV).⁴ For these reasons, they can be seen as a cost-effective alternative to conventional solar cells based on inorganic semi-conductors. For many years ruthenium complexes have been developed and employed as sensitizers in DSSCs and photoconversion efficiencies (PCE) up to 12% have been reported using them.⁵ However, the scarcity and high price of ruthenium and the toxicity of some of its derivatives have pushed the scientific community to investigate other classes of sensitizers. Highly efficient molecules have emerged from this work such as zinc porphyrin dyes, which demonstrate efficiencies in solar cells that can reach 13%.⁶ Nevertheless, other organic dyes based on various chromophores including quinoxaline derivatives,⁷ benzothiadiazole,⁸ or diketopyrrolo-pyrrole,⁹ to name a few, have also shown impressive performances. In general, organic dyes show a D-π-A structure where the electron-donor unit (D) is, in most cases, a triarylmethine group and the electron acceptor (A) unit is usually a cyano-acrylic acid that also acts as a tether group. The choice of the π-conjugated linker in these molecules is critical since it influences the energy levels of the orbitals and governs the position of the intramolecular charge transfer absorption band (ICT), which in turn determines the range of absorption and the color of the dye. It is still considered nowadays that some improvements in the efficiency of dye solar cells could come from the use of new photosensitizers with the condition that they possess a good absorption spectral match with the solar emission. Therefore, development of organic dyes showing a wide absorption up to the near-IR range (so-called panchromatic dyes) is important to explore. Pioneering work in phthalocyanines for DSSC, by Torres et al., paved the way for solar cells with a near-IR response.¹⁰ Yet, to date more than one thousand organic dyes have probably been synthesized and tested as sensitizers for TiO₂ with variable performances in solar cells, but only a small fraction of them show panchromatic absorption or have the ability to absorb photons up to the near-IR region.¹¹ Despite of
advantageous optical properties, only a few reach efficiencies close to 6%.\textsuperscript{11} One popular strategy to obtain organic dyes exhibiting a broad absorption in the visible up to the near-IR relies on the use of motives with a strong electron-withdrawing character in the backbone. The combination of such units with the electron-rich triarylamines gives rise to absorption bands (ICT) in the visible range whose position can be tuned depending on the strength of the electron-withdrawing blocks. Following this approach, in this work we have synthesized and used several isoindigo derivatives for the preparation of panchromatic sensitizers. Isoindigo is an isomer of the well-known natural and robust blue dye indigo. Isoindigo is a particularly interesting building block that shows a strong electron-withdrawing character due the presence of two lactam rings in the structure. Another interesting feature of this molecule is that it can be synthesized starting from natural synths i.e. isatin. The isoindigo building block has already been investigated for the preparation of semiconducting polymers for bulk-heterojunction solar cells\textsuperscript{13} or photosensitizers for DSSCs.\textsuperscript{14} These dyes exhibit usually a broad absorption domain up to 750nm but, in spite of this interesting characteristic, their photovoltaic efficiencies are comprised between 3 and 7.5%.\textsuperscript{15}

In the last decade other isoindigo derivatives such as thieno-isoindigo or benzothieno-isoindigo have been successfully prepared to serve as building units in the preparation of low-band gap semiconducting copolymers,\textsuperscript{16} showing a better absorption in the long wavelengths.\textsuperscript{17} But surprisingly none of these synthons have been employed for the synthesis of dyes applicable in DSSCs. Although similar to isoindigo thanks to their two lactam rings that ensure a strong electron-withdrawing character, these chromophores have terminations that differ in the nature of the side aromatic rings; in benzothieno-isoindigo one phenyl is replaced by a thiophene and in thieno-isoindigo two thiophene rings are present. In this work we report the synthesis of six original organic dyes based on these three synthons. The molecules have been fully characterized by UV-Vis-NIR spectroscopy and electrochemistry prior to their use as photosensitizers in DSSCs. We show that swaping the isoindigo unit by a benzothieno-isoindigo or a thieno-isoindigo unit leads to a large shift of the absorption range toward the NIR part of the spectrum (up to 900nm) with a simultaneous increase of the molar absorption coefficient (up to 44000 M\textsuperscript{-1} cm\textsuperscript{-1}).

\textbf{Scheme 1}: Synthetic routes of the dyes TPAT8-Isol, TPAT8-Altiso, energy levels of the frontier orbitals have been determined by cyclic voltammetry (CV), differential pulse voltammetry (DPV) and estimated by Density Functional Theory (DFT) calculations. The consistent results indicate that the dyes can effectively be employed in a DSSC device to sensitize TiO\textsubscript{2} mesoporous electrodes with I/\textsubscript{3} as redox mediator in the electrolyte. Photovoltaic properties of the DSSCs based on the six dyes have been investigated under simulated 1 Sun AM 1.5 illumination and PCE ranging from 0.45% to 5.84% have been measured. Charge extraction, transient photovoltage and transient absorption spectroscopy experiments have been carried out on these devices, and DFT modeling have been used to complete the study. This has allowed us to fully unravel the origin of the low J\textsubscript{sc} and V\textsubscript{oc} obtained with some of these compounds and their limited efficiencies in spite of their strong absorption. In this work we fully identify what are the detrimental phenomena that should be suppressed to improve their performances. Our observation might be extended to other classes of near-infrared absorbers.

\textbf{Results and discussion.}

\textbf{Design and synthesis of the dyes.}

In the last decade organic dyes with a D-π-A chemical structure have been widely developed due to their ability to foster the electron transfer within the dye towards the wide band gap semiconductor. Following this approach, the dyes of this study were synthesized by embedding the isoindigo, thieno-isoindigo of benzothieno-isoindigo units in between a donating segment based on a triarylamine (D) and an electron accepting unit constituted by a phenyl-cyanoacrylic acid (A). Two different triarylamines showing different electron donating strength were used. The first one is a triphenylamine connected to an octyl-thiophene (TPAT8). This unit was initially developed in our laboratory to synthesize RK1, a dye that combines a high efficiency (over 10%) and an outstanding stability (T\textsubscript{80} over 6000h under ISOS-L2 ageing test).\textsuperscript{3a} The second triphenylamine unit that we employed consists of a fused triphenylamine with a thiophene through the formation of an indene ring (TPAF).

The chemical structures and synthetic routes of the six dyes named TPAT8-Isol, TPAT8-Altiso, TPAT8-Tiso and TPAF-Isol, TPAF-Altiso and TPAF-Tiso are presented in Scheme 1.
These dyes have been prepared following a common synthetic strategy. First, we synthesized the isoidingo, benzothieno-isoidingo or thieno-isoidingo cores via acid-promoted condensation starting from 6-bromoisatine, 6-bromoxindole, thieno-isatin, or thieno-oxindole. The insertion of branched alkyl chains on the nitrogen atoms of the lactam rings has been either performed directly on these precursors or on the formed cores. In the substitution process, 2-ethylhexyl side chains have been employed since they ensure a good solubility of the dyes in common organic solvents and they can help reducing the dye aggregation once grafted onto TiO₂ surface.⁷ Then the isoidingo, benzothieno-isoidingo or thieno-isoidingo cores have been coupled at one extremity to benzaldehyde via a Suzuki coupling reaction. The TPAT8 or TPAF donating units were coupled at the other extremity through a Stille coupling reaction. The final dyes were then simply obtained from these aldehyde precursors by Knoevenagel condensation with cyano-acrylic acid. The preparation of all the precursors and all the dyes is fully described in ESI.

Optoelectronic and electrochemical properties.

The absorption spectra of the isoidingo, benzothieno-isoidingo, or thieno-isoidingo based dyes were recorded in CH₂Cl₂ and as adsorbed onto 2 µm-thick TiO₂ films with chenodeoxycholic acid (CDCA). This co-adsorbent was used to reproduce the real conditions of use (see Figure 1). The absorption wavelengths, molar extinction coefficients, and HOMO–LUMO energy levels of the six dyes are summarized in Table 1. The UV-vis spectra in solution exhibit two major bands, one appearing at 355-410 nm and the other prominent one at 575-740 nm. The former can be attributed to the π-π* transitions from the different aromatic rings being in conjugation. The absorption band at 575-740 nm may be assigned to the ICT transition from the triarylamine donor group to the isoidingo derivative cores connected to the cyano-acrylic acceptor unit. As expected replacing the TPAT8 donor group by a fused unit, such as TPAF, results in a bathochromic spectral shift of the ICT band (by 30-40nm depending on the core). More strikingly the modification of the core itself, i.e. replacing the phenyl groups in the isoidingo structure by thiophene rings, has a huge impact on the position and the absorption coefficient of this band. Indeed the maximum wavelength of absorption shifts toward the near-infrared leading to dyes revealing a purple, blue or green hue (see Fig. 1). For instance the λₘₐₓ of the ICT band of TPAF-Tiso is red-shifted by 129nm compared to that of TPAF-Iso. Interestingly the molar extinction coefficient of this band also rises to attain 44 000 M⁻¹ cm⁻¹. To summarize, swapping one or two benzene rings in the isoidingo core by one or two thiophene rings induces a bathochromic and hyperchromic effect on the ICT band and, consequently, the percentage of absorption of the dyes in the near-infrared region is increased to reach a maximum rate of 35%. This value corresponds to the ratio of absorption above 700nm divided by the overall absorption. Compared with the solution spectra, the maximum absorption peaks of the dyes, registered on the 2 µm-thick transparent TiO₂ films using CDCA as co-adsorbent, shift to the blue by 25, 55, 81, 28, 55 and 65 nm for TPAF-Iso, TPAF-Altiso, TPAF-Tiso, TPAT8-Iso, TPAT8-Altiso and TPAT8-Tiso, respectively (Table 1).

![Figure 1](image-url)  
**Figure 1:** (a) UV-Vis spectra of the dyes in dichloromethane, (b) pictures of the solutions and (c) UV-Vis spectra grafted on TiO₂.

Usually the blue-shift of the λₘₐₓ going from the solution to thin films may be partially attributed to the deprotonation of the carboxylic acid.¹⁸ However this shift is far too large in our case to be solely explained by this phenomenon. Broader blue-shifts can be generally ascrib to the formation of H-aggregates.¹⁹ Despite of the fact that we employed CDCA in the dyeing solution and that we used branched alkyl chains to prevent such event, this is probably the case here. This phenomenon was already observed for isoidingo-based dyes,²⁰ nonetheless it should be noted that this effect is enhanced when one thiophene is introduced in the isoidingo core and even more pronounced when two thiophene rings are present. This can be rationally explained by the conformation of the molecules found with DFT calculations, in which the core of the molecule becomes more planar whenever a thiophene motif is embedded (see ESI). In spite of the aggregation, the dyes spectra in the solid state is broader and all the dyes show...
a panchromatic absorption from the UV to the Visible-NIR region with absorption edges ranging from 700 nm to circa 875 nm.

Table 1: Optical and electrochemical properties of isoindigo, benzothieno-isoindigo and thieno-isoindigo based dyes

<table>
<thead>
<tr>
<th>Optical properties</th>
<th>Electronic properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>λ_{max UV}^{ab} [nm]</td>
<td>λ_{max Vis-NIR}^{ab} [nm]</td>
</tr>
<tr>
<td>TPAF-iso</td>
<td>376 (355)</td>
</tr>
<tr>
<td>TPAF-Altiso</td>
<td>389 (391)</td>
</tr>
<tr>
<td>TPAF-Tiso</td>
<td>408 (408)</td>
</tr>
<tr>
<td>TPAF-Tiso</td>
<td>358 (355)</td>
</tr>
<tr>
<td>TPAF-Altiso</td>
<td>361 (369)</td>
</tr>
<tr>
<td>TPAF-Tiso</td>
<td>405 (405)</td>
</tr>
</tbody>
</table>

* Measured in CH2Cl2 at room temperature at a concentration of 10^{-3}M. Parenthesis, measured after dyeing of 2µm thick TiO2 films on glass substrate, in the presence of CDCA (dyeing solution 0.2mM Dye/ 2.0mM CDCA). Calculated from optical absorption onset. Mean values calculated from oxidative potential for the HOMO and reduction potential for the LUMO, measured by DPV and CV in in CH2Cl2 (2.10^{-3}M) at room temperature, ferrocene/ferrocenium was used as the internal standard and measured at -0.48 V (-4.8 eV). Determined by DFT calculation. Using mean values obtained from electrochemical measurements with the following equation ∆E= E_{LUMO}-E_{HOMO}.

To accurately investigate the chemical modification effects on the molecular energy levels, the electrochemical properties of these dyes were analyzed by cyclic voltammetry and differential pulse voltammetry in CH2Cl2 solution with 0.2M tetraethylammonium hexafluorophosphate (TBAPF6). All potentials reported are referenced to Fc/Fc redox couple as an internal standard. For comparison purpose, DFT calculations were performed in a CH2Cl2 solvent phase model to obtain the derived HOMO and LUMO values and their respective spatial localizations (ESI). All these data are summarized in Table 1 and Figure 2. Consistent data were found for each dye from CV and DPV measurements. Noteworthy, the mean values gathered from electrochemistry for the HOMO energy levels are in excellent agreement with the values determined by the DFT calculations. The LUMO energy levels measured using electrochemistry are almost identical for all compounds (-3.7 to -3.8eV) and a discrepancy generally more difficult to describe theoretically than the occupied ones. It appears that the chemical variations either on the TPA unit or on the core of the molecules do not change the LUMO energy level positions. To the contrary some significant variations are observed for the HOMO energy levels which are all lying between -4.9 eV and -5.2 eV. Two effects can be commented; first, the replacement of the TPAF unit by a TPAF unit, which has a stronger electron donating character, induces a positive shift of 0.1eV of the HOMO energy level position. Second, the HOMO energy level rises by 0.1eV each time that a thiophene is introduced in the core. Our results are totally consistent with the work of Estrada et al. who demonstrated that the HOMO energy levels of compounds based on isoindigo derivatives are mainly influenced by the stilbene motive whereas the LUMO energy levels are mainly related to the fumaraldehyde motive present in the core structure. As a consequence the use of electron rich aromatic of circa 0.3eV is observed with the DFT computed values. This difference is not surprising since the vacant orbitals are rings such as thiophene to replace benzene rings in isoindigo predominantly leads to the rise of the HOMO energy level.
The optimized molecular structures and isoelectronic densities are shown Figure 2. DFT calculations show that the HOMOs are localized at the TPA8T or TPAF donor moieties and they extend to the core of the molecules. The delocalization of the HOMOs is not the same in all compounds. In isoindigo-based dyes the HOMOs poorly overlap with the central core whereas in benzothieno-isoindigo and thieno-isoindigo dyes the HOMOs fully extend on these units.

To better explain this observation, the bond length alternation (BLA) of the chemical bonds located at each side of the cores was calculated. The BLA are comprised between 0.059 and 0.076 Å in isoindigo dyes and they are characteristic of aromatic systems. In benzothieno-isoindigo dyes these distances decrease to attain values ranging from 0.037 to 0.064 Å and in thieno-isoindigo-based molecules the BLA are comprised between 0.035 and 0.040 Å; these values are typical from quinoidal compounds, which is in good agreement with the electrochemically and optically derived data. On the other hand, the LUMOs of the six molecules are primarily localized on the cores and the phenyl-cyanoacrylic acid unit. Such a directional electronic distribution is preferred for good electron injection into TiO2 from dye anchoring sites.

To summarize this section, the six dyes present broad absorptions covering the solar spectrum from the UV to the NIR that could give rise to large photocurrents. Their LUMO energy levels positioned around -3.8 eV are located 0.25 eV above the energy level of the conducting band of the oxide (around -4.05 eV) which is the minimum driving force required to achieve an efficient injection of electrons in the oxide. However, the experimentally estimated HOMO levels of the dyes, are comprised between -5.2 eV and -4.9 eV. Some of them are very close to the Nernst potential of the triiodide/iodide-based redox electrolyte which is around -4.9 eV (i.e. 0.4 V versus NHE). This means that for TPA8T-Tiso, TPAF-Tiso, TPA8-Altiso and TPAF-Altiso the $\Delta E_{\text{red}}$ might not be sufficient to regenerate the oxidized species of the dyes. Finally the larger overlap between HOMO and LUMO, deduced from the computed structures of these four dyes, might also be detrimental for the photocurrent generation.

Photovoltaic properties.

The performances of the DSSCs with these dyes were measured with a mask under the irradiation of AM 1.5 (1000 W/m²) simulated solar light after calibration. For a direct comparison, we fabricated solar cells with the same photoelectrode composition consisting of a double layer TiO2 (12µm-thick transparent layer and 4µm-thick scattering layer) from Solaronix. This thickness was chosen taking into account a previous work that showed that thicker electrodes give higher photocurrent densities and performances. This hypothesis was further confirmed through tests with some of our dyes (see ESI). Then, we started to optimize some of the fabrication parameters of the solar cells and we focused our preliminary investigations of the isoindigo dyes TPAF-Iso and TPA8-Iso.

By analyzing the conclusion of several papers from literature we noticed that most of the devices comprising isoindigo-based sensitizers did not contain tert-Butyl-Pyridine (tBP) in the electrolyte. For other class of dyes, classical electrolytes contain 0.5M of tBP. The reason behind this statement is the following: the LUMO energy level of the isoindigo dyes is usually close to the energy level of the CB of the TiO2, and tBP is known to shift positively the CB band of the oxide through a dipole effect. By suppressing this dipole effect, the CB of TiO2 is located at a lower energy and the injection process is favored. On the other hand, the removal of tBP has a detrimental effect on the $V_{oc}$ of the solar cells, which becomes lower because the energy difference between this energy level and the redox potential of the electrolyte decreases. Consequently, an optimum concentration has to be found to obtain a good trade-off between $J_{sc}$ and $V_{oc}$.

Several preliminary tests in solar cells using a dye/CDCA ratio of 0.2/2mM were carried out with various amount of tBP in the electrolyte or without tBP (see ESI), and we found that 0.1M is an optimum concentration to reach the highest performances. We highlight within this first series of tests the tremendous effect of this additive on the performances and the necessity to tune the formula of the electrolyte with this class of dyes. Indeed, with a standard amount of tBP (0.5 M) the performances of TPAF-Iso and TPA8-Iso are limited to 0.88% and 1.27% with $J_{sc}$ below 2.7 mA/cm², and the $V_{oc}$ values are 0.64 and 0.61 V, respectively. When the concentration of tBP is lowered the performances are better, and for an optimum concentration of 0.1M, performances around 4% are achieved with $J_{sc}$ reaching 9.6 mA/cm² and 10.2 mA/cm². As expected the $V_{oc}$ of the resulting cells are slightly lower (around 0.55 V) but this loss is largely compensated by a more efficient injection process, which produces four times higher photocurrents. The strong influence of the tBP concentration on the performances further confirms that the LUMO energy levels of the dyes are close to the limit for an efficient electron injection.

From the UV-Vis spectra of the molecules recorded on TiO2 films (vide supra), we learned that our dyes have a large propensity to form aggregates even in the presence of CDCA. Aiming to reduce this dye aggregation higher amounts, up to 12 mM of CDCA, were used in the dyeing bath of TPA8-Iso for further tests, and the performances of the corresponding solar cells were measured. The I-V curves and photovoltaic
parameters are given in ESI. When the concentration of the co-
adsorbent is fixed between 10 to 12 mM, $J_{sc}$ as high as 15.9 mA/cm$^2$ are observed, leading to PCE of up to 5.84%. Solar cells were fabricated with the six dyes using these optimized conditions, the I-V curves are presented in Figure 3 and the corresponding photovoltaic parameters are summarized in Table 2.

Table 2: Photovoltaic parameters of the solar cells obtained using the optimized fabrication conditions. Dyeing bath: dye/CDCA 0.2/10 mM, EtOH/CHCl$_3$ (4:1), electrodes: 12 × 4 μm, size 0.36 cm$^2$, Electrolyte: BMII 0.5 M, I$_2$ 0.03 M, LiI 0.5 M, Gthio 0.5 M, tBP 0.1 M. Irradiation 1 Sun, using a mask. First lines correspond to the results of the best cells, second lines correspond to the parameters obtained from three or four devices.

<table>
<thead>
<tr>
<th>Dye/CDCA</th>
<th>$V_{oc}$ (V)</th>
<th>$J_{sc}$ (mA/cm$^2$)</th>
<th>FF</th>
<th>η (%)</th>
<th>Dye Loading 10$^4$ Mol/cm$^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>TPAF-Iso</td>
<td>0.53</td>
<td>10.97</td>
<td>0.69</td>
<td>4.01</td>
<td>3.3 x 10$^8$</td>
</tr>
<tr>
<td></td>
<td>0.54±0.02</td>
<td>10.09±0.87</td>
<td>0.72±0.03</td>
<td>3.88±0.13</td>
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</tr>
<tr>
<td>TPAF-Altiso</td>
<td>0.47</td>
<td>9.69</td>
<td>0.59</td>
<td>2.66</td>
<td>7.0 x 10$^8$</td>
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<tr>
<td></td>
<td>0.5±0.02</td>
<td>8.44±1.18</td>
<td>0.60±0.01</td>
<td>2.51±0.19</td>
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<td>TPAF-Tiso</td>
<td>0.38</td>
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<td>0.49</td>
<td>8.2 x 10$^8$</td>
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<td></td>
<td>0.37±0.02</td>
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<tr>
<td>TPAT8-Iso</td>
<td>0.59</td>
<td>14.55</td>
<td>0.68</td>
<td>5.84</td>
<td>4.3 x 10$^8$</td>
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<tr>
<td></td>
<td>0.57±0.02</td>
<td>14.91±1.23</td>
<td>0.65±0.04</td>
<td>5.54±0.49</td>
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</tr>
<tr>
<td>TPAT8-Altiso</td>
<td>0.50</td>
<td>11.28</td>
<td>0.53</td>
<td>2.99</td>
<td>9.9 x 10$^8$</td>
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<tr>
<td></td>
<td>0.48±0.03</td>
<td>9.01±1.44</td>
<td>0.59±0.04</td>
<td>2.52±0.29</td>
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</tr>
<tr>
<td>TPAT8-Tiso</td>
<td>0.38</td>
<td>3.33</td>
<td>0.55</td>
<td>0.70</td>
<td>13.0 x 10$^8$</td>
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<tr>
<td></td>
<td>0.38±0.01</td>
<td>2.89±0.53</td>
<td>0.55±0.01</td>
<td>0.61±0.12</td>
<td></td>
</tr>
</tbody>
</table>

To prove the reproducibility and reliability of our results, at least three optimized solar cells for each dye were fabricated and characterized under exactly the same conditions. The current-voltage characteristics of the best performing cells are given in Figure 3, some of them clearly show a more or less pronounced S-shape which is unusual. From the best-cell and average parameters summarized in Table 2, it appears that isoidigo-based dyes show the best performances, with a PCE of 4.01% and 5.84% for TPAF-Iso and TPAT8-Iso, respectively. This is mainly explained by the rather high $J_{oc}$ (up to circa 15 mA/cm$^2$) delivered by these solar cells and the absence of S-shape in the I-V curves that keep the FF at high values. One should note here that these compounds led, however, to the lower dye-up-takes on the electrodes. The solar cells containing benzothieno-isoindigo-based dyes show rather similar PCE but the solar cells with TPAT8-Altiso lead to higher $J_{sc}$. Notably for these dyes the $V_{oc}$ is lower and the FF is strongly deteriorated by the S-shape of the I-V curve. Finally, the dyes containing thieno-isoindigo units show the worst performances, below 1%. The $J_{sc}$ supplied by TPAT8-Tiso and TPAF-Tiso solar cells is extremely low despite that these devices exhibit the higher dye-loading and the larger absorption. As explained before, TPAT8-Tiso and TPAF-Tiso show a strong tendency to stack in the solid state. This phenomenon is known to induce a photocurrent loss because the aggregates cause the quenching of excited state of the dyes.29

The IPCE curves show a trend that is comparable with the device performances and the photocurrent action spectra are all extended beyond 700 nm. (see Figure 4).

Figure 4: IPCE spectra for the devices containing the six dyes

After measuring the I-V curves for all organic sensitizers in DSSC using iodine/iodide electrolyte, we observed that the photocurrent was strikingly low for some of the dyes. At first sight, one could think that this is due to difficulties related to the dyes excited-state energy with respect to the TiO$_2$ conduction band. However, both experimental results and theoretical experiments lead to the conclusion that the dyes excited-states are well above the TiO$_2$. In fact, Laser Transient Absorption Spectroscopy measurements (L-TAS) shown later on this manuscript (Figure 7) show efficient charge injection from the dyes excited state to the TiO$_2$ conduction band.

Hence, we analyzed in depth the photo-induced charge present in the solar cells for each dye in complete solar cells at...
different light intensity by using the charge extraction, as reported previously.\textsuperscript{30} As shown in Figure 5, the amount of charge is similar for all dyes at a given voltage close to the $V_{oc}$ at 1 Sun (for example 0.5 V, the photo-induced charge density is $0.15 \times 10^{18}$ electrons/cm$^2$). This value is among the normal values for DSSC having dye coverages between $3-15 \times 10^{14}$ mol cm$^{-2}$. Thus, the differences seen in the IPCE and the measured photocurrent at short circuit ($V_{oc}$ at 1 Sun) must be due to a different reason but not due to the lack of efficient charge injection. Once the charge density was measured, we turned to analyze the carriers lifetime that is related to the recombination kinetics between the photo-injected electrons and the oxidized electrolyte. To do so, we have used the Transient Photo-Voltage (TPV) measurements that have been reported earlier.\textsuperscript{31} Figure 6 shows the measured lifetime vs. charge density for all the solar cells, which have been fabricated identically except for the different sensitizers used.

**Figure 5:** Charge extraction data showing electron density as a function of induced voltage for the different dyes

![Charge extraction data showing electron density as a function of induced voltage for the different dyes](image)

**Figure 6:** Transient photovoltage (TPV) data showing electron lifetimes versus electron density for DSSC devices based on the six dyes

![Transient photovoltage (TPV) data showing electron lifetimes versus electron density for DSSC devices based on the six dyes](image)

All carriers lifetimes ($\tau$) fall in the milliseconds time range (0.01-0.001 s) which is usual in DSSC. However, the slope of the linear plots is quite different for the TPAT8-Tiso, which shows the lowest $V_{oc}$ at 1 Sun and a greater slope. The slope is related to the order of recombination kinetics, which clearly is much higher for TPAT8-Tiso than for the other dyes. In fact, the trend follows quite well the $V_{oc}$: the solar cells with higher slope, as TPAT8-Tiso, show the lower $V_{oc}$ at 1 Sun.

Hence, from the TPV measurements we can correlate the observed differences in $V_{oc}$ with the recombination kinetics (the order of the recombination kinetics) between the photo-injected electrons at the TiO$_2$ and the oxidized electrolyte. Yet, we have not found a convincing explanation for the lower photocurrent measured for some dyes. In particular, the TPAT8-Tiso and the TPAT8-Tiso. As mentioned above, neither TPAT8-Tiso nor TAPT8-Tiso are limited by the nature of their excited state to inject electrons into the TiO$_2$. Thus, we decided to study the interfacial charge injection and charge recombination kinetics of complete devices using L-TAS. As per comparison purposes we also measured solar cells made with TPAT8-isos that shows photocurrent as high as 15 mA/cm$^2$ at 1 Sun.

Figure 7 shows the decay for each solar cells in the presence and absence of the electrolyte. As can be seen, for TPAT8-iso and TPAT8-tiso, and in absence of the electrolyte, the decay corresponds to the oxidized dye after efficient injection of electrons into the TiO$_2$. The measured half-lifetime of the decay and the HWHM (Half Width at the Half Maximum) is of 0.02 s. On the contrary, in the presence of the iodine/iodide, the electrolyte fully regenerates the oxidized dye, the half-lifetime is of tenths of microseconds, and a slow phase is present that has been previously assigned to the $I_3$ species.

For TPAT8-Altiso, the decay is slower. In clear contrast, for TPAT8-Tiso and TPAT8-Tiso the presence of the electrolyte does not regenerate the dye, and a slower decay is observed.

A plausible explanation for this new decay trace might be the formation of a stable complex between the dye and the oxidized species of the electrolyte but this hypothesis is not confirmed by the theoretical calculations (vide supra). Another explanation could be that the regeneration is extremely slow for these dyes which are not fully regenerated. The hypothesis of an extremely slow regeneration processes is fully consistent with the low $\Delta E_{reg}$ values deduced from cyclic voltammetry measurements. This low driving force may cause the increase of the total series resistance of the DSSC and could explain the low FF and S-shape of the (IV) curves observed for these devices. Indeed a low FF is generally ascribed to high sheet resistance and/or a high series resistance in the device.\textsuperscript{32}

However, to shed light on the results extracted from L-TAS experiments the interaction between iodide and the six dyes has been studied computationally (ESI). The dyes have been optimized in acetonitrile in their neutral and radical cationic forms and the electrostatic potential ($V_r$) of the latter have been obtained (see Figure 8). In Figure 8 the surface in blue indicates the positively charged regions of the cationic radical dye, which should be the more favorable points for interacting with iodide.
Iso away from the oxygen and sulfur atoms. The calculations indicate that iodide does not interact strongly with any cationic radical dye; the most favorable interaction energy between both species (computed as the electronic energy difference between the dye-I\(^-\) complex and the separated species) falls in a range of -0.20 to -0.30 eV for all dyes (See Table S1 in ESI). In all the optimized dye-I\(^-\) complex structures iodide moves around to find a more positively charged place to interact with, normally a terminal C-H bond (see Figure S5 in ESI), with quite long distances in the range of 3.0-3.5 Å. These calculations seem to rule out the formation of stable complexes between the oxidized dye and iodide and thus the regeneration issues should be attributed to other reasons.

However, the more positively charged groups in TPAF-Tiso and TPAT8-Tiso being located closer to the anchoring function may favor the approach of the iodide closer to the TiO\(_2\) surface thus leading to higher recombination rate. This hypothesis is fully consistent with the higher recombination kinetics deduced from TPV measurements for these dyes, resulting in the lower Voc values.

As may be observed, the positively charged surface corresponds to the more hydrophobic regions, usually far away from the oxygen and sulfur atoms. We decided to compute two (or more, up to six for TPAT8-Tiso and TPAT8-Iso) different conformations for the interaction of each oxidized dye and the iodide, where the latter is placed close to the positively charged regions of the dye, namely terminal positively charged methyl, phenyl and hydrogen groups. For completion reasons, we also included all the S and O-S pockets as plausible anchoring points.

The calculations for all the dyes show a similar trend with iodide more favorably positioned over the terminal carboxylic acid groups. Figure 9 shows the open-circuit voltage as a function of the dipole moment of the oxidized dyes. This phenomenon was mostly observed using additive molecules, however the large variation of the dipole moments within our family of molecules could explain the large variations of the Voc for the solar cells that we fabricated. To verify this hypothesis the variation of the Voc of the solar cells was plotted as a function of the dipole of the molecules, the graph is presented in Figure 9. We found a linear dipole dependence of the Voc with the dipole of the dyes, similarly to the observation by Rühle and coworkers with co-adsorbent...
molecules, the dye molecules showing the larger dipoles leading to the higher open circuit voltages. This interesting observation could help chemists for designing molecules leading to higher $V_{oc}$ in solar cells.

**Perspective of the work.**

Isoindigo-based dyes possess a high dipole in their oxidized state and consequently they give higher $Voc$ in solar cells. We also demonstrated from the TAS experiments that these dyes can be regenerated efficiency by the electrolyte and TPV experiments indicate that they lead to lower charge recombination rates. To highlight that our findings could drive the molecular design of more efficient dyes, we synthesized a new photosensitizer (coded 6OTPA-Iso) which is an analog of TPA8-Iso. This molecule was synthesized in only four steps from the isoindigo unit using a modified TPA unit bearing hexyloxy chains. The chemical structure of this dye is presented in Figure 10 and the synthetic route, and optoelectronic properties of this molecule are presented in the ESI file.

In dichrometane solution, 6OTPA-Iso shows a maximum wavelength of absorption in the visible at 579nm and an absorption edge located at 712nm. The experimental determination of the energy levels (by CV and DPV) indicates a HOMO at -5.1eV and a LUMO at -3.8eV. Solar cells were fabricated using this photosensitizers and TiO2 electrodes ([12µm-thick transparent layer and 4µm-thick scattering layer]. After optimization of the TBP ratio in the electrolyte and the CDCA amount in the dyeing bath, a device with a maximum power conversion efficiency of 7% was obtained. The device reach a Voc of 0.56V and a FF of 0.64, but more interestingly a high Jsc of 19.4 mA/cm² was obtained.

Figure to add

**Conclusions**

In conclusion, we investigated the potential of isoindigo, thieno-isoindigo or benzo-thieno-isoindigo chromophores to create organic dyes; absorbing in the UV-visible and near-infrared spectral range; for DSSCs applications. Interestingly some of the dyes showed absorption domains extended up to 900nm. Devices based on TPA8-Iso embedding an isoindigo chromophore attained a PCE of circa 5.84%. Despite of a panchromatic absorption and high absorption coefficients some of the dyes based on thieno-isoindigo cores led to devices with poor performances with quite low Jsc. Thanks to charge extraction, transient photovoltage, laser transient absorption spectroscopy experiments and computational investigations we unravel the interfacial processes that occurs with this family of dyes and we identify the detrimental processes that are responsible for the low performances. In particular we demonstrate that thieno-isoindigo-based dyes exhibit higher charge recombination kinetics in solar cells principally because this chromophore unit modify the electrostatic potential over the molecule allowing iodide to approach the TiO2 surface. We also found that the dipole of the oxidized molecules plays a crucial role on the $V_{oc}$ value, the dyes possessing the larger dipoles leading to the solar cells with higher open circuit voltages. Based on our observations, we have synthesized a new photosensitizer based on Isoindigo unit with a TPA. Consequently, this work unravel fundamental processes in DSSCs based on NIR dyes and could help define new strategies of molecular engineering to develop more efficient organic dyes for DSSCs.

**Conflicts of interest**

There are no conflicts to declare.

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**Notes and references**

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