Computational studies have suggested that the integration of secondary amine as donor groups in the structure of unsymmetrical zinc phthalocyanine (ZnPc) should have positive effects on photovoltaic performance, once the molecule is integrated as light harvester in dye sensitized solar cells (DSSCs).

Aiming at obtaining experimental confirmation, we synthesized a peripherally substituted push-pull ZnPc bearing three electron donating diphenylamine substituents and a carboxylic acid anchoring group and integrated it as sensitizer in TiO$_2$-based DSSCs. Detailed functional characterization of solar energy converting devices resulted in ruling out the original hypothesis. The causes of this discrepancy have been highlighted, leading to a better understanding of the conditions for an effective design of push-pull diarylamino substituted ZnPcs for DSSCs.

Many research endeavors to spur the efficiency in solar energy conversion of dye sensitized solar cells (DSSCs) are currently directed to the development of optimized light harvesters, capable of exploiting the whole range of the solar spectrum, including the infrared portion lost in present silicon-based PV technology, in order to eventually obtain the so-called panchromatic sensitization. Investigations from diverse standpoints, spanning from the comprehension of the underlying fundamental phenomena to the assessment of the life cycle and economic effectiveness of these devices, are as well pursued, but special attention is paid to the design of new photosensitizers that could ensure optimal photovoltaic parameters and acceptable lifetime service to DSSCs. Ruthenium-based sensitizers had a prominent role in early developments of DSSCs and are still widely employed, but they show only moderate light harvesting capability, in particular in the red/near IR spectral regions of the solar radiation. In this respect, other families of dyes, such as phthalocyanine derivatives, offer better perspectives especially in view of the architectural integration of DSSCs in energy efficient buildings. Indeed phthalocyanines (Pcs) are thermally, photochemically and electrochemically stable compounds, with unparalleled light-harvesting capability in the far-red and near-IR spectral regions where a substantial solar fluence occurs. These peculiar properties can be fine-tuned by relatively simple synthetic modifications. The electronic absorption spectrum of Pcs shows two main bands at 300–400 nm (B band) and 620–700 nm (Q band with extinction coefficients in the order of $10^5$ M$^{-1}$ cm$^{-1}$). These bands can be shifted or broadened depending on the presence and nature of substituents.
in the peripheral and non-peripheral positions (see Fig. 1) of the macrocycle. Other relevant characteristics, as the HOMO-LUMO energy level and the distribution of the electronic density, can be varied following the same strategy. Finally, metallation of the macrocycle also allows to tune the behavior of Pc-based photosensitizers and Zn(II) complexes of Pcs (ZnPcs) with their long-lived singlet excited states proved to be remarkably useful in DSCCs.

The most successful ZnPcs sensitizers reported so far are unsymmetrically substituted molecules characterized by the presence of identical bulky, electron-donating groups on three of the four isoindole subunits (referred to as A) which constitute the Pc macrocycle, while the fourth one (referred to as B) bears one or more anchoring groups able to interact with the metal oxide semiconductor. This A3B molecular design aims to combine steric suppression of dye aggregation, favorable electronic push-pull effects and optimized binding mode of the dye to the surface of the semiconductor layer. It was originally suggested by Nazeeruddin, Torres and coworkers, who were able to achieve power conversion efficiencies up to 4.1% in the case of peripherally substituted A3B ZnPcs bearing tert-butyl donors and -COOH anchoring groups. The advent of A3B ZnPcs featuring bulky, electron rich aryl substituents rotated with respect to the macrocycle plane represented a second significant step forward towards efficient DSSCs based on Pcs. For instance, Mori, Kimura and coworkers have shown that power conversion efficiencies up to 6.4% under simulated one-sun illumination can be obtained using a peripherally substituted A3B ZnPc bearing electron-rich 2,6-bis(n-butoxy)phenoxy groups on A and one electron withdrawing -COOH group on B. Despite the steady progress made in the field, the full potential of ZnPcs as photosensitizers in DSSCs has yet to be realized and further improvements are conceivable. In particular, the application of predictive computational techniques for the rational molecular design of efficient unsymmetrical push-pull ZnPcs has been proposed. According to these theoretical studies, the use of -NH2, -NHR, -NR2, NHAr or -NAr2 donor groups instead of alkyl or phenoxy donor groups should improve the photovoltaic performance of unsymmetrical A3B ZnPcs thanks to higher LUMO levels, smaller energy gaps and red-shifted absorption bands. New absorption bands are also expected to emerge in the 400–600 nm region for A3B ZnPcs bearing diphenylamino substituents, which should make them good candidates as panchromatic sensitizers.

Results and discussion

We recently developed an efficient procedure for the preparation of 4-(diarylamino)phthalonitriles and demonstrated their use as precursors of symmetrical A4 ZnPcs. Zn(II)-templated cyclization performed on a mixture of the simplest of these building-blocks, 4-(diphenylamino)phthalonitrile 1, and pentyl 3,4-dicyanobenzoate 2 (molar ratio 3:1) followed by hydrolysis of the ester functionality afforded the A3B ZnPc BI54 in 12% overall yield (Scheme S1, Supporting Information). Having proved the synthetic feasibility of A3B ZnPcs with diarylamino substituents, we next examined the possible use of BI54 as a sensitizer in DSSCs.

The architecture of a DSSC requires the favourable energy alignment of all functional components the electrochemical circuit, namely the electron transport material (ETM), the sensitizer and the redox couple responsible for the regeneration of the photooxidized dye. The energy levels of the HOMO-LUMO orbitals of the new dye were thus assessed by electrochemical measurements combined with the analysis of the optical properties (See SI, Figures S1, S2 and S3 and Table S1). The heterogeneous electron transfer kinetics at glassy carbon interfaces resulted in quasi-reversible, diffusion-limited waves, with chemical reversibility attained for scan speed > 100 mV/s. Figure 2 sketches the energy alignment of the semiconducting metal oxide (MOx) used as ETM in this work (TiO2) with BI54. The energy levels of an efficient Ru-based dye (N719), widely employed in DSSCs, are also reported for comparison. BI54 presents the needed requirements to be applied as light harvester in DSSCs, with energy levels of about −5.4 eV for the HOMO orbital and −3.72 eV for the LUMO orbital, which...
would imply a thermodynamically favoured electronic injection from the dye to the TiO₂ conduction band (CB) \(\Delta G_{\text{INJ BI54-TiO}_2} \approx 0.28 \text{ eV}\).

Two kinds of ETM have been prepared and tested as photoanodes: bare commercial TiO₂ nanoparticles and composite multi-walled carbon nanotubes-TiO₂ nanoparticles (MWCNT-TiO₂), which have been reported to enhance the overall device functional performances\(^1\)\(^9\). Fig. 3a, d and g show the functional characterization of two typical DSSCs whose photoanodes are made by TiO₂ and MWCNT-TiO₂ sensitized for 20 hours, which is the commonly used soaking time for Ru-based dyes.

Low photocurrent density values (in both cases lower than 0.4 mA/cm\(^{-2}\)) were recorded and a depleted open-circuit voltage \(V_{\text{OC}}\) was as well observed (350 mV and 370 mV for the devices using bare TiO₂ and MWCNT-TiO₂ as ETM, respectively). Moreover, no significant difference between the photoanodes with and without MWCNTs was recorded. Analysis of open circuit voltage decay (OCVD) highlights slow charging kinetics: after 20 s irradiation the \(V_{\text{OC}}\) value was not reached (Fig. 3d), indicating slow charge accumulation, accompanied by electron recombination (both with BI54\(^+\) and I\(_3\)\(^-\)). Decay dynamics were essentially the same for both devices. As expected from OCVD observation, electron lifetime (Fig. 3g) is the same in all the voltage range analyzed for both devices.

Pcs aggregation due to \(\pi-\pi\) stacking has been previously reported as a serious issue affecting these molecular dyes\(^2\)\(^0\)\(^-\)\(^2\)\(^2\), with consequent lowering of associated device performances: co-adsorption of a molecular spacer on TiO₂ has then been proposed as an effective countermeasure. Deoxycholic acid (DCA) was thus added during the sensitization of bare TiO₂ photoanodes and the soaking time reduced from 20 to 13 h. Decreased sensitization time resulted in dramatically enhanced short-circuit current density \(J_{\text{SC}}\) values (Figure 3b and Table 1, Entry 3 vs 1), which were almost doubled, compared with those recorded for devices obtained after 20 hours soaking. However, the presence of DCA was found to be not significant in terms of enhanced solar energy conversion efficiency (PCE), which was calculated as high as 0.17% and 0.18% with and without DCA. A further shortening of the photoanode staining, down to 5 hours resulted in no improvement of cell efficiency in the absence of DCA (Fig. 3c, blue line). Actually, when no DCA is co-adsorbed on TiO₂ photoanode the \(V_{\text{OC}}\) value is not changed with respect to 13 hours soaking and only a slight increase in \(J_{\text{SC}}\) value, from 0.715 mA/cm\(^{-2}\) to 0.764 mA/cm\(^{-2}\) can be appreciated (Table 1 Entry 5 vs 3).

These findings indicated once more the critical role played by phthalocyanine aggregation on photoanode surface, even for reduced uptake times. Indeed, by decreasing the soaking time to 5 hours and co-adsorbing DCA (Fig. 3c, red line), a remarkable enhancement of device working parameters \((J_{\text{SC}} = 1.376 \text{ mA/cm}^2, V_{\text{OC}} = 540 \text{ mV, PCE} = 0.43\%)\) is recorded, corresponding to an increase of 80%, 13% and 140%, respectively, with respect to the analogous device with no DCA (Table 1, Entry 6 vs 5).

The increase of \(V_{\text{OC}}\) is ostensibly related to the negative shift of Fermi level, which may be induced upon the insertion of co-absorbent in the dye\(^2\)\(^3\), while the enhancement in the photocurrent density \(J_{\text{SC}}\) is ascribable to the increased electron injection and charge collection efficiency\(^9\).

The improvements achieved by the combined reduction of sensitization time and use of DCA as co-adsorbent also involve the electron lifetime as depicted in Fig. 3g–i. For devices fabricated in the absence of DCA, no difference is observed using sensitization times of 20 and 13 hours (Fig. 3g, black line, and Fig. 3h, blue line, respectively). For 13 hours sensitization (Fig. 3h) the presence of DCA results in a small, but appreciable difference. Finally, the electron lifetime is highly enhanced by the presence of DCA for 5 hours sensitization (Fig. 3i).

In order to gain a deeper insight on the relationship between device performances and sensitization time with/without DCA, a kinetic study of the uptake of BI54 from DCM solutions has been carried out by following the variation of optical density of the photoanode at 725 nm (Fig. 4a). The commercially available Ru-based dye
N719, for which a detailed uptake kinetic analysis (monitored at 500 nm) has been previously reported, was used as a reference.

Dye uptake on MOx surface is usually a two-step process that can be properly analyzed as a fast exponential growth followed by a second slower step, possibly featuring a pseudo-0-order kinetic, moving towards a plateau. The first part of the uptake curve can be suitably fitted by the relationship:

Table 1. Functional parameters of solar energy converting devices based on TiO2 uploaded with dye BI54 for different sensitization times. aBare TiO2; b MWCNT-TiO2; c DCA added.

| Entry | Sensitization time (h) | Jsc (mA/cm²) | Voc (mV) | FF (%) | PCE (%) |
|-------|------------------------|--------------|----------|--------|---------|
| 1a    | 20                     | 0.361        | 350      | 47     | 0.060   |
| 2b    | 20                     | 0.392        | 370      | 47     | 0.068   |
| 3c    | 13                     | 0.715        | 480      | 49     | 0.17    |
| 4c    | 13                     | 0.777        | 480      | 47     | 0.18    |
| 5a    | 5                      | 0.764        | 480      | 51     | 0.18    |
| 6c    | 5                      | 1.376        | 540      | 58     | 0.43    |
| 7a    | 4                      | 3.02         | 520      | 58     | 0.92    |
| 8c    | 4                      | 3.39         | 510      | 66     | 1.13    |

Figure 3. Analysis of DSSCs using BI54 dye as light harvester. (a), (b) and (c): J-V curves of DSSCs sensitized for 20 h, 13 h and 5 h, respectively. (d), (e) and (f): normalized open circuit photovoltage decay (sensitization time: 20 h, 13 h, 5 h). (g), (h) and (i) calculated electron lifetime (sensitization time: 20 h, 13 h, 5 h, respectively). Black and green lines in (a), (d) and (g) refer to photoanodes made by TiO2 nanoparticles and MWCNT-TiO2, respectively. Blue and red lines in (b), (c), (e), (f), (h), (i) refer to devices sensitized with BI54 dye alone and BI54 dye with DCA, respectively.
Where \([T_0]\) refers to the sites originally available on TiO\(_2\) for dye uptake and TD to the sites occupied (dye onto TiO\(_2\)). The second part of the uptake curve is properly fitted by linear regression, and follows the relationship:

\[
[TD] = [T_0] \left(1 - e^{-k_1t}\right)
\]  

(1)

where \(k_1\) and \(k_2\) are the kinetic constants and \([TD]_I\) refers to the dye concentration at the time in which the curve changes from fast to low uptake.

**BI54** does not reach a plateau in the uptake curve, not even after 25 hours impregnation. The analysis of apparent kinetic constants (Table 2) reveals other significant details. In the absence of DCA, the initial rate of **BI54** anchoring on TiO\(_2\) sites (exponential part of the uptake curve) is comparable to that shown by **N719**, but the second, linear step is dramatically faster (almost four times). A different trend is observed in the case of addition of DCA. The uptake curve of **BI54** reaches a plateau after a few hours of soaking, thus confirming the role exerted by DCA, which competes with the dye for TiO\(_2\) anchoring sites. This can be as well inferred by the much slower increase of the exponential growth, well visible in Fig. 4a, and by the comparison of the apparent kinetic constants (Table 2). The presence of DCA, indeed, slows down the uptake kinetics by occupying available sites on TiO\(_2\) nanoparticles surface, thus limiting the amount of **BI54** molecules anchoring on the photoanode: the rate of anchoring in the first step is 1.5 times slower in presence of DCA, while the rate in the second step is decreased of almost six times.

The kinetics of the anchoring process of **BI54** is affected by the presence of DCA even at very reduced dyeing times (Fig. 4b): this provides a further evidence that DCA competes very efficiently with **BI54** for TiO\(_2\) available anchoring sites\(^{24}\), a critical factor for reducing photogenerated charge recombination in solar energy converting devices\(^{23}\).

**Table 2.** Kinetics data for dye uptake onto TiO\(_2\). \(^{4}\)Determined by fitting of eqn (1); \(^{b}\)Determined by fitting of eqn (2); \(^{c}\)Values retrieved from ref\(^{25}\).

| Dye          | \(k_1^a (s^{-1}) \times 10^3\) | \(k_2^b (M^{-1}s^{-1}) \times 10^3\) |
|--------------|-------------------------------|--------------------------------------|
| **BI54**     | 13.06                         | 1.03                                 |
| **BI54 (+DCA)** | 8.144                         | 0.1805                               |
| **N719**     | 16.15                         | 0.2808                               |

**Figure 4.** Dye uptake analysis: (a) Overview of dyeing process and (b) detail for uptake of **BI54** in the first hour. Blue circles = **BI54** without DCA; red squares = **BI54** with DCA; empty triangles = **N719**. Markers are experimental data, lines are fitting curves as discussed in the text.
by enhancing the photocurrent density (from 3.02 to 3.39 mA cm$^{-2}$), while both V$_{OC}$ and fill factor (FF) present similar values (Fig. 5a and Table 1, entries 7 and 8). The increased photocurrent density results in an improvement of the photoconversion efficiency of about 23% (from 0.92% to 1.13%). Consistent with J-V curves, incident photon to electron conversion efficiency (IPCE) spectra, reported in Fig. 5b, show an enhancement in all the visible range when DCA is present. In particular IPCE values are increased in the NIR region (730 nm) from 4.6% to 5.2% (14%) and at lower wavelengths (480 nm) from 2.8% to 4.2% (50%). IPCE data recorded in cells having a comparable optical density indicate that the factor $\phi_{inj} \times \eta_{collection}$, where $\phi_{inj}$ is the injection quantum yield and $\eta_{collection}$ is the electron collection efficiency across the photoanode, is improved. The increased quantum yield may result from reduced aggregation, which would otherwise lead to excited state quenching or exciton annihilation. It should be herein reminded that only the first monolayer of adsorbed dye is effective in generating useful photocharges, while subsequent adsorption, via dispersive forces, of dye molecules to form molecular aggregates decoupled from the titania surface is detrimental, resulting in parasitic light absorption and quenching. OCVD analysis, reported in Fig. 5c, shows a drastically reduced drop of voltage for the cell in which DCA is used. By fitting the OCVD curves with an exponential function, the time decay constants may be estimated at 0.45 s and 2.18 s for the cells without and with DCA, respectively, indicating an increase of almost five times of the decay rate when BI54 is left free to aggregate on TiO$_2$.

The smoother V$_{OC}$ decay is reflected in an increased electron lifetime (Fig. 5d): after 0.10 s the voltage is quite reduced (0.11 V) if DCA is not employed, while it is twice this value (0.23) when DCA is present.

Open circuit photovoltage observed in all the investigated DSSCs show reduced values as compared with the expectation for TiO$_2$ based cells in the presence of the iodide redox couple. Such a decrease may origin from either a decrease in the band offset or an increased recombination rate constant. A shift in the band edge (which would alter the relative energy positions of TiO$_2$ CB and standard redox potential of the electrolyte redox couple) can be estimated through electrochemical impedance spectroscopy (EIS) by a shift in the chemical capacitance of the cell as a function of voltage.

Figure 6a shows the chemical capacitance of DSSCs sensitized with BI54 (with DCA) and N719 (used as reference). In the low potential region (100–300 mV), the trend of the internal capacitances of the cells is quite constant and is originated by the interfaces of the platinum counter electrode and of the FTO back contact of the photoanode$^{26}$. In the high potential region the chemical capacitance dominates the capacitive effects inside the cells, showing a linear trait on the logarithmic scale. The chemical capacitance is directly related to the density of electronic states (DOS) in the metal oxide$^{25,26}$. The trend of the chemical capacitance could then give information about the displacement of the CB of the metal oxide. Since the thickness and the porosity of the oxide are the same for both cells, the effect on the CB is ascribable to the interaction between the dye and the metal oxide$^{27,28}$. 

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**Figure 5.** Functional characterization of DSSCs using BI54 dye as light harvester with (red markers) and without DCA (blue markers) (sensitization time: 4 h). (a) J−V characteristics; (b) IPCE; (c) photovoltage decay and (d) electron lifetime. Markers are experimental points, lines are guide for the eye.
The net effect in the case of N719 is the downward shift of the $E_C$ (by about 150 mV), resulting in an enhancement of the charge injection from the Ru(II) dye to the metal oxide. This contributes to explain the lower photogenerated current observed for BI54, as compared with N719. Moreover, the slopes of the chemical capacitance vs potential curves are different for the two sensitzers, and in particular it appears less steep for BI54. As shown by Barea et al. this means that the interaction between N719 and the TiO$_2$ modify the trap states.

It is also worth noting that recombination resistances $R_{REC}$ (Fig. 6b and c) of devices sensitized with N719 and BI54 show slightly higher value for N719 (once the as retrieved data, Fig. 6b, are recalculated for the same electron density, Fig. 6c). This finding is not related with the low performance of the devices, as clearly reported by Barea et al. These data suggest that low performances recorded for devices sensitized with BI54 are ascribable to impaired charge injection from BI54 LUMO to TiO$_2$ CB and are not due to enhanced recombination processes. Indeed, the driving force for charge injection from the LUMO of BI54 to the CB of the TiO$_2$ is reduced compared to the case of N719 (Fig. 2). By shifting the photocurrent density of BI54 sensitized cell to simulate the same $J_{SC}$ value (namely, similar charge injection condition) recorded for the N719 sensitized device, it is possible to observe (Fig. 7) that the two solar energy converting devices perform in the very similar way, confirming that charge injection is the limiting factor for BI54.

Conclusions

Despite the optimistic evaluations pointed out by computational predictions, the push-pull A3B ZnPc BI54 showed rather low solar to electric power conversion efficiencies once integrated as light harvester in DSSCs. As predicted by models, BI54 fulfilled the mandatory requirements for energy alignment with respect to TiO$_2$ used as ETM and the iodide/iodine redox couple, but the successful implementation of this ZnPc requires a careful control of adsorption conditions and possibly the search for other semiconductors offering a better driving force for charge injection. A major issue with this dye design is its tendency to aggregation, which proved drastically detrimental for cell efficiency. This drawback can be partially alleviated by modulating the sensitization time and using an anti-aggregation agent such as DCA that acts as a molecular spacer on the ETM surface. Also, theoretical
calculations did not take into account the possible shift in the CB of TiO₂, which was highlighted by our experiments and results in a decrease in charge injection efficiency.

Based on these warnings, the design of new push-pull ZnPcs bearing diarylamino substituents is now underway in our laboratories.

Methods

Synthesis of B154. Commercially available reagents were used as received. 4-(diphenylamino)phthalonitrile 1 and pentyl 3,4-dicyanobenzoate 2 were prepared as described in our preceding works. ⇢ Solvents were purified by standard methods and dried if necessary. Reactions were monitored by thin layer chromatography (TLC) that was conducted on plates precoated with silica gel Si 60-F254 (Merck, Germany). Column chromatography was conducted using silica gel Si 60, 0.063–0.200 mm (Merck, Darmstadt, Germany). UV-Vis measurements in solution were performed on a Nicolet Evolution 500 spectrophotometer (Thermo Electron Corporation). FTIR were performed on a Cary 530 instrument (Agilent Technologies). ESI mass spectra were obtained with a ICR-FTMS APEX II (Bruker Daltonics) mass spectrometer. MALDI mass spectra were obtained with a TOF-TOF mass spectrometer Microflex LT (Bruker). Elemental analyses were carried out by the Departmental Service of Microanalysis (University of Milano).

ZnPc B154. In a flame dried Schlenk tube equipped with a stir bar a mixture of phthalonitriles 1 (266 mg, 0.90 mmol), 2 (73 mg, 0.30 mmol) and Zn(OAc)₂·2H₂O (55 mg, 0.16 mmol) was suspended in dimethylaminoethanol (12 mL). The mixture was stirred for 24 h at 150 °C under a nitrogen atmosphere. The solvent was then evaporated under vacuum and the residue was treated with MeOH (50 mL). The precipitated black solid was collected by filtration on a Büchner funnel, thoroughly washed with MeOH. The raw material was purified twice by column chromatography (silica gel, THF) to give the dimethylaminomethyl ester of B154 (MW = 1194) as determined by MALDI-TOF mass analysis (32 mg, 17% with respect to the limiting Zn(II) reagent). The ester was dissolved in THF (4 mL) and treated with 1 M aqueous KOH (1 mL). The mixture was stirred at 70 °C for 8 h, then overnight at room temperature. The organic solvent was removed under reduced pressure and H₂O (1 mL) was added to the residue. The pH of the aqueous phase was adjusted to 3 by dropwise addition of aqueous 10% HCl and the mixture was extracted twice with DCM (5 mL). The combined organic layers were washed with H₂O, brine and dried over MgSO₄. Evaporation of the solvent under reduced pressure afforded the title compound as a mixture of positional isomers (22 mg, 71%). UV-Vis (THF): λ max/nm (log ε) = 359 (5.01), 496 (4.37), 640 (4.71), 711 (5.35). FTIR (KBr): ν max/cm⁻¹ = 3429, 2922, 1679, 1613, 1590, 1490, 1446, 1385, 1331, 1260, 1088, 1046, 745, 695. M (MALDI-TOF): m/z calc for [C₆₉H₄₃N₁₁O₂Zn]+ 1121.3. Found: 1121.3. Anal. calcd for C₆₉H₄₃N₁₁O₂Zn: C 73.76, H 3.86, N 13.71. Found: C 73.70, H 3.89, N 13.67.

Dye-sensitized solar cells. TiO₂ paste (20 nm anatase nanoparticles 18NR-T) was purchased from Dyesol and multi-walled carbon nanotubes (average length 5–20 μm) were purchased from Sigma-Aldrich. The redox electrolyte was composed of 0.1 M LiI, 0.05 M I₂, 0.6 M 1,2-dimethyl-3-n-propylimidazolium iodide, and 0.5 M 4-tert-butylpyridine dissolved in acetonitrile. All chemicals were purchased from Sigma-Aldrich. Photoanode thickness for all the photoanodes was about 15 μm and was evaluated without masking by using a Keithley 2400 SourceMeter under simulated sunlight using an ABET2000 solar simulator at AM 1.5 G (100 mW cm⁻²) calibrated using a reference silicon cell. The electrochemical impedance spectroscopy (EIS) was carried out in dark conditions using a SOLARTRON 1260 A Impedance/Gain-Phase Analyzer, with an AC signal 20 mV in amplitude, in the frequency range between 100 mHz and 300 kHz. External bias applied was from 0 to 100 mV above the V OC.

Electron lifetime of the photogenerated charges present in the CB of the dyed semiconductor is calculated from open circuit voltage decay using the following equation:

\[ \text{Electron lifetime} = \frac{1}{\text{Open circuit voltage decay}} \]

Device fabrication. B154 dye paste was deposited onto FTO glass (sheet resistance 10 Ω/sq) by tape casting technique. The paste was dried 15 min at ambient conditions and then fired on hot plate for 6 min (130 °C). Two layers of transparent TiO₂ and one of scattering TiO₂ were deposited. All the samples were finally annealed at 500 °C for 30 min under ambient atmosphere. Photoanode thickness for all the photoanodes was about 15 μm and was evaluated by stylus profilometry.

B154 dye was dissolved in tetrahydrofuran and the sensitization of photoanodes was performed for 20, 13, 5 and 4 hours in the dark. Concentration of B154 was kept constant in all the experiments at 5.0 × 10⁻⁴ M; DCA was used at the same concentration.

DSSCs were fabricated using dye sensitized oxide photoanodes and platinized FTO glass as a counter electrode (5 nm Pt thin film deposited by sputtering) separated by 25 μm-thick plastic spacers (Surlin from Soloranox). Electrolyte composition was as follows: 0.1 M LiI, 0.05 M I₂, 0.6 M 1,2-dimethyl-3-n-propylimidazolium iodide, and 0.5 M 4-tert-butylpyridine dissolved in acetonitrile. All chemicals were purchased from Sigma–Aldrich and multi-walled carbon nanotubes (average length 5–20 μm) were purchased from Sigma-Aldrich. The redox electrolyte was composed of 0.1 M LiI, 0.05 M I₂, 0.6 M 1,2-dimethyl-3-n-propylimidazolium iodide, and 0.5 M 4-tert-butylpyridine dissolved in acetonitrile. All chemicals were purchased from Sigma–Aldrich and multi-walled carbon nanotubes (average length 5–20 μm) were purchased from Sigma-Aldrich.
where $k_b T$ is the thermal energy, $e$ is the positive elementary charge.

References

1. Hagfeldt, A., Boschloo, G., Sun, L., Kloo, L. & Pettersson, H. Dye-sensitized solar cells. *Chem. Rev.* **110**, 6595–6663 (2010).
2. Lee, C.-T. et al. Recent progress in organic sensitizers for dye-sensitized solar cells. *RSC Adv.* **5**, 23810–23825 (2015).
3. Robertson, N. Catching the rainbow: light harvesting in dye-sensitized solar cells. *Angew. Chem. Int. Ed.* **47**, 1012–1014 (2008).
4. de la Torre, G., Claessen, G. C. & Torres, T. Phthalocyanines: old dyes, new materials. *Putting color in nanotechnology. Chem. Commun.* **20**, 2000–2015 (2007).
5. Ragoudis, M.-E. & Torres, T. Modern synthetic tools toward the preparation of sophisticated phthalocyanine-based photoactive systems. *Chem. Asian J.* **9**, 2676–2707 (2014).
6. Martínez-Díaz, M. V., de la Torre, G. & Torres, T. Lighting porphyrins and phthalocyanines for molecular photovoltaics. *Chem. Commun.* **46**, 7090–7108 (2010).
7. Singh, V. K., Kanaparthi, R. K. & Giribabu, L. Emerging molecular design strategies of unsymmetrical phthalocyanines for dye-sensitized solar cell applications. *RSC Adv.* **4**, 6970–6984 (2014).
8. Martín-Gomis, L., Fernández-Lázaro, F. & Sastre-Santos, Á. Advances in phthalocyanine-sensitized solar cells (PcSSCs). *J. Mater. Chem. A* **2**, 15672–15682 (2014).
9. Reddy, P. Y. et al. Efficient sensitization of nanocrystalline TiO$_2$ films by a near-IR absorbing unsymmetrical zinc phthalocyanine. *Angew. Chem. Int. Ed.* **46**, 373–376 (2007).
10. Cid, J. J. et al. Structure-function relationships in unsymmetrical zinc phthalocyanines for dye-sensitized solar cells. *Chem. Eur. J.* **15**, 5310–5317 (2009).
11. García-Iglesias, M. et al. Effect of anchoring groups in zinc phthalocyanine on the dye-sensitized solar cell performance and stability. *Chem. Sci.* **2**, 1145–1150 (2011).
12. Eu, S., Katoh, T., Uemeya, T., Matano, Y. & Imahori, H. Synthesis of sterically hindered phthalocyanines and their applications to dye-sensitized solar cells. *Dalton Trans.* **40**, 5476–5483 (2008).
13. Mori, S. et al. Enhancement of incident photon-to-current conversion efficiency for phthalocyanine-sensitized solar cells by 3D molecular structualization. *J. Am. Chem. Soc.* **132**, 4054–4055 (2010).
14. Heuchi, T. et al. Molecular engineering of zinc phthalocyanine sensitizers for efficient dye-sensitized solar cells. *Chem. Commun.* **50**, 1941–1943 (2014).
15. Yang, L. et al. Theoretical design and screening of panchromatic phthalocyanine sensitizers derived from TTA for dye-sensitized solar cells. *J. Mol. Graphics Modell.* **34**, 1–9 (2012).
16. Yang, L. et al. Substituent effects on zinc phthalocyanine derivatives: A theoretical calculation and screening of sensitizer candidates for dye-sensitized solar cells. *J. Mol. Graphics Modell.* **38**, 82–89 (2012).
17. Linares-Flores, C., Mendizabal, E., Arratia-Pérez, R., Inostroza, N. & Orellana, C. Substituents role in zinc phthalocyanine derivatives used as dye-sensitized solar cells. *Chem. Rev.* **115**, 23810–23825 (2015).
18. Cho, K. T. et al. Perovskite solar cells employing molecularly engineered Zn(II) phthalocyanines as hole-transporting materials. *Nanoscale* **30**, 853–857 (2016).
19. Dembele, K. T. et al. Hybrid carbon nanotubes–TiO$_2$ photoanodes for high efficiency dye-sensitized solar cells. *J. Phys. Chem. C* **128**, 14510–14517 (2013).
20. He, J. et al. Modified phthalocyanines for efficient near-IR sensitization of nanostructured TiO$_2$ electrode. *J. Am. Chem. Soc.* **124**, 4922–4932 (2002).
21. Morandera, A. et al. Ru(II)-phthalocyanine sensitized solar cells: the influence of co-adsorbents upon interfacial electron transfer kinetics. *J. Mater. Chem.* **19**, 5016–5026 (2009).
22. Yum, J. H. et al. Effect of coadsorbent on the photovoltaic performance of zinc phthalocyanine-sensitized solar cells. *Langmuir* **24**, 5636–5640 (2008).
23. Lim, J., Kwon, Y. S. & Park, T. Effect of coadsorbent properties on the photovoltaic performance of dye-sensitized solar cells. *Chem. Commun.* **47**, 4147–4149 (2011).
24. Sharma, G. D., Kurchania, R., Ball, J. R., Roy, M. S. & Mikroyannidis, J. A. Effect of deoxygenic acid on the performance of liquid electrolyte dye-sensitized solar cells using a perylene monoimide derivative. *International Journal of Photoenergy* **2012**, 983081 (2012).
25. Concina, I. et al. On-line monitoring and active control of dye uptake in dye-sensitised solar cells. *Chem. Commun.* **47**, 11656–11658 (2011).
26. Bisquert, J. Chemical capacitance of nanostructured semiconductors: its origin and significance for nanocomposite solar cells. *Phys. Chem. Chem. Phys.* **5**, 5360–5364 (2003).
27. Howie, W. H., Claeyssens, F., Miura, H. & Peter, L. M. Characterization of solid-state dye-sensitized solar cells utilizing high absorption coefficient metal-free organic dyes. *J. Am. Chem. Soc.* **130**, 1367–1375 (2008).
28. O’Regan, B. C., Scully, S., Mayer, A. C., Palomares, E. & Durrant, J. The effect of Al$_2$O$_3$ barrier layers in TiO$_2$/dye/CuSCN photovoltaic cells explored by recombination and DOS characterization using transient photovoltage measurements. *J. Phys. Chem. B* **109**, 4616–4623 (2005).
29. Nazeeruddin, M. K., Humphry-Baker, R., Liska, P. & Grätzel, M. Investigation of sensitizer adsorption and the influence of protons on current and voltage of a dye-sensitized nanocrystalline TiO$_2$ solar cell. *J. Phys. Chem. B* **107**, 8981–8987 (2003).
30. Liu, Y., Hagfeldt, A., Xiao, X.-R. & Lindquist, S.-E. Investigation of influence of redox species on the interfacial energetics of a dye-sensitized nanophotocatalysis TiO$_2$ solar cell. *Sol. Energy Mater. Sol. Cells* **55**, 267–281 (1998).
31. Fabregat-Santiago, F., Bisquert, J., Garcia-Belmonte, G., Boschloo, G. & Hagfeldt, A. Influence of electrolyte in transport and recombination in dye-sensitized solar cells studied by impedance spectroscopy. *Sol. Energy Mater. Sol. Cells* **87**, 117–131 (2005).
32. Bisquert, J. et al. A review of recent results on electrochemical determination of the density of electronic states of nanocrystalline metal-oxide semiconductors and organic hole conductors. *Inorg. Chem. Acta* **361**, 684–698 (2008).
33. Rühle, S. et al. Molecular Adjustment of the Electronic Properties of Nanoporous Electrodes in Dye-Sensitized Solar Cells. *J. Phys. Chem. B* **109**, 18907–18913 (2005).
34. Barea, E. M. et al. Energetic factors governing injection, regeneration and recombination in dye solar cells with phthalocyanine sensitizers. *Energy Environ. Sci.* **3**, 985–94 (2010).
35. Pozzi, G. et al. Fluorocarbon molecules for dye-sensitized solar cells: synthesis and photoelectrochemistry of unsymmetrical zinc phthalocyanine sensitizers with bulky fluorophile donor groups. *J. Phys. Chem. C* **115**, 3777–3788 (2011).
36. Zaban, A., Greenshtein, M. & Bisquert, J. Determination of the electron lifetime in nanocrystalline dye solar cells by photovoltage decay measurements. *Chem. Phys. Chem.* **4**, 859–864 (2003).
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Author Contributions
S.O., M.C. and G.P. synthesized the new dye; R.B. and S.C. performed the optical and electrochemical characterization of the dye in solution; R.M., G.S.S. and I.C. prepared and characterized the solar cells; I.C., R.M. and G.P. wrote the first draft of the manuscript; I.C. and G.P. supervised the research project. All the authors contributed in the data analysis and in the discussion of the manuscript.

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