

Promoting charge-separation in p-type dye-sensitized solar cells using bodipy†

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The viability of applying bodipy sensitisers to NiO-based p-type dye-sensitised solar cells (p-DSCs) has been successfully demonstrated. The triphenylamine donor–bodipy acceptor design promotes a long-lived charge-separated state which is difficult to achieve with NiO-based devices. The current was above 3 mA cm⁻² and the IPCE was 28%.

Tandem dye-sensitised solar cells could provide a route to high efficiency solar energy conversion at low cost.¹ Lindquist *et al.* proposed in 1999 that replacing the platinised counter electrode in Grätzel cells with a dye-sensitized photocathode (in series) should greatly improve the device efficiency by collecting high energy photons at the top electrode and lower energy photons at the bottom electrode.² This method increases the light collection and reduces thermal energy losses; the theoretical efficiency may be as high as 43%. To date, however, there has not been a tandem cell reported with power conversion efficiency higher than that of a state-of-the-art TiO₂-based DSC.

Since the rapid charge recombination reactions at the photocathode limit the photoelectric conversion efficiency of the p-DSCs, the main difficulty of enhancing the performance of tandem DSCs exists in raising the current generated at the photocathode to match that generated at the TiO₂ anode.³ The precise reason for fast recombination is still unclear but it has been demonstrated that the dye can be engineered to promote charge-separation, thereby improving the photocurrent.¹ A myriad of different dyes have been reported with TiO₂, but very few have been designed for p-type materials such as NiO.⁴ The most efficient sensitisers for NiO have both high extinction coefficients for maximum light harvesting on thin electrodes (*e.g.* dyes with favourable π - π^* electronic transitions⁵⁻⁷) and charge-transfer character to promote and maintain charge-separation (*e.g.* π -donor-acceptor

(“push-pull”) functionality,⁵ or with pendant NDI or C₆₀ electron acceptors^{8,9}).

Boradiazaindacene (bodipy) dyes are popular in a range of applications but have been used infrequently and with moderate success in n-DSCs.¹⁰⁻¹³ However, their large extinction coefficients, electrochemical stability and tuneable absorption properties (*i.e.* to optically match photoanodes in tandem cells) prompted us to test them in p-DSCs. Initially, we have chosen to use a triphenylamine-thiophene motif used in one of the most successful NiO photosensitisers (“P1”) coupled to a highly substituted bodipy (Fig. 1).⁶ The dye was prepared by reacting 4-carboxy-4',4"-di(5-formyl-2-thienyl)triphenylamine with 2,4-dimethyl-3-ethylpyrrole in the presence of trifluoroacetic acid (see ESI† for details). The resulting dipyrrene was oxidised with *p*-chloranil before reacting with boron trifluoride etherate to give **1**.

The optical and electrochemical properties are summarised in Table S1 (ESI†). The electronic spectrum of **1** contains absorption maxima at 365 nm ($\epsilon = 56\,000\text{ L mol}^{-1}\text{ cm}^{-1}$) and 540 nm ($\epsilon = 112\,000\text{ L mol}^{-1}\text{ cm}^{-1}$, almost double that of the push-pull sensitiser P1 at $\lambda_{\text{max}} = 481\text{ nm}$, Fig. S3 in the ESI†).¹⁴ The absorption and emission spectra are characteristic for a bodipy derivative, with narrow absorption and emission bands and a small Stokes shift when excited at 488 nm. λ_{max} was the same in CH₂Cl₂ or CH₃CN solution and when **1** was adsorbed onto NiO (Fig. S4 and S5, ESI†), suggesting that the

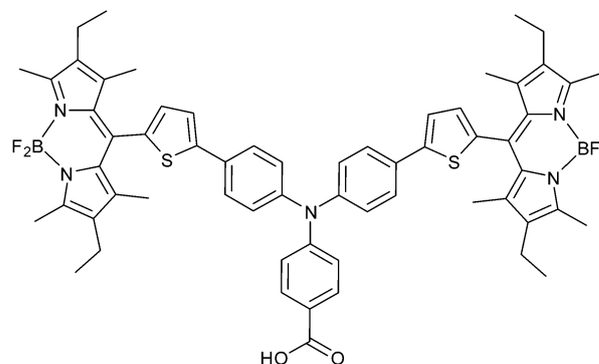


Fig. 1 Structure of bodipy dye **1**.

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† Electronic supplementary information (ESI) available: Experimental details, electronic spectra, electrochemical characterisation, DFT calculations, *J-V* curves for **1** vs. P1, transient absorption spectra, spectroelectrochemistry of **1**/NiO. See DOI: 10.1039/c3cc46133e



electronic properties of the chromophore were unchanged on binding. However, when **1** was excited in the higher energy band (406 nm) the emission spectrum was solvent dependent: emitting from the higher energy triphenylamine (475 nm) when in CH₃CN; emitting from the lower energy bodipy (560 nm) when in CH₂Cl₂.

The electrochemical properties of **1** were determined by cyclic and square wave voltammetry and calibrated vs. FeCp₂^{+/0}/FeCp₂ (Fig. S13 and S14, ESI†). A reversible reduction process at -1.46 V and two closely spaced irreversible oxidation processes at 0.65 and 0.75 V were observed. The estimated driving forces‡ for both injection (ΔG_{inj}) and regeneration (ΔG_{reg}) are both negative, indicating that **1** is a suitable NiO sensitizer.^{15,16} DFT calculations were also performed on **1** to further explore the geometry and electronic properties. $E_{(D^*/D)^-}$ = 0.81 V, is suitably positive to allow for efficient charge-separation between geometry and electronic properties. The results supported experimental evidence that the bodipy was decoupled from the triphenylamine-thiophene donor and the calculated optimised geometry placed the two at an angle of $84 \pm 0.4^\circ$. Fig. S16 (ESI†) shows the electronic distribution in the frontier orbitals: the HOMO level of **1** is located on the triphenylamine; two degenerate HOMO - 1(-2) orbitals and two degenerate LUMO(+1) levels are located on the bodipy. TD-DFT suggested that the lowest energy electronic transitions were bodipy-localised $\pi-\pi^*$ in nature, with some contribution from a charge shift from the HOMO to the LUMO.

NiO p-DSCs were assembled with **1** as the sensitizer and a I₃⁻/I⁻ redox shuttle (see ESI† for details). The photocurrent density–photovoltage curve and spectral response of a typical cell are given in Fig. 2. The cell parameters were $J_{SC} = 3.15 \text{ mA cm}^{-2}$, $V_{OC} = 79 \text{ mV}$, FF = 0.31, $\eta = 0.08\%$. The maximum incident photon-to-current conversion efficiency (IPCE) was 28%. These values are superior to related triphenylamine-based dyes reported previously,^{16,17} but fall short of those achieved with the most successful dye, P1, ($J_{SC} = 5.48 \text{ mA cm}^{-2}$, IPCE = 64%).⁶ The photocurrents obtained with our bodipy-based dyes are extremely encouraging compared to bodipy-sensitized TiO₂ devices.^{10,18,19} TiO₂-DSCs are usually more efficient than p-DSCs, but to our knowledge the highest

IPCE for a bodipy-sensitized n-DSC is less than 30%.¹¹ This suggests that the bodipy chromophore may be better suited to the NiO system than conventional TiO₂-DSCs. Further improvements in efficiency are likely to come from broadening the absorption band by increasing the electronic coupling between the π -linker and the bodipy and extending the conjugation by functionalising the dye periphery.

To probe the charge-transfer dynamics of this new dye–NiO system, we performed ps- and ns-transient absorption spectroscopy (TA) both with **1** in CH₂Cl₂ and adsorbed on NiO. Excitation of **1** in solution (Fig. 3 black line) at 532 nm generated an excited state which absorbed between 400–500 nm and decayed over ca. 400 ps to form a new intermediate with narrower absorption bands at 425 and 650 nm which decayed much more slowly ($\tau_{425} \approx 300 (\pm 10) \text{ ns}$ in air, $\tau_{425} \approx 860 \text{ ns}$, $\tau_{650} \approx 730 \text{ ns}$ in Ar) to the ground state. We have assigned this long-lived transient to a triplet bodipy (³1*) excited state (see ESI† for further details).^{20,21}

The transient absorption results were different for **1** adsorbed on NiO (Fig. 3 red dashed line). On the ps timescale, the broad features between 400–470 nm corresponding to ¹1* that were observed in CH₂Cl₂ were less intense. Instead a peak at 575 nm formed rapidly ($\tau_{rise} \approx 1 \text{ ps}$) which was characteristic of 1⁻ by comparison with our spectroelectro-chemistry experiments (Fig. S26, ESI†). A broad featureless absorption persisted throughout the ps- and ns-transient spectra. This is attributed to oxidised NiO and indicates that the charge-separated state (NiO⁺/1⁻) had formed. The dynamics for each band were characteristically heterogeneous and we have fitted the rise and decay in each experiment to first order kinetics giving a short (τ_1 ps) and long (τ_2 ns) time constant (Table S4, ESI†) to enable us to compare the general behaviour of the intermediates. The rapid time constant for the formation of NiO⁺/1⁻ can be compared to the lifetime of ¹1* recorded in solution to give an estimate of the injection efficiency.‡ Using this approach gives a very high efficiency of 99% but it should be noted that any slower injection processes were masked by the subsequent decay of this peak. To try and provide a more realistic value for this process we have compared

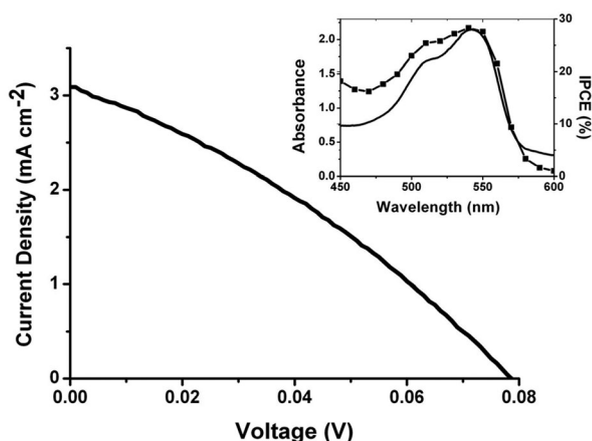


Fig. 2 J - V curve of **1**-sensitized NiO solar cell. Inset: comparison of IPCE (line + symbol) and absorption spectrum (line) of **1**-sensitized NiO solar cell to confirm the dye is contributing to the photocurrent. An absorbance of 2 means the light harvesting efficiency is ca. 99%.

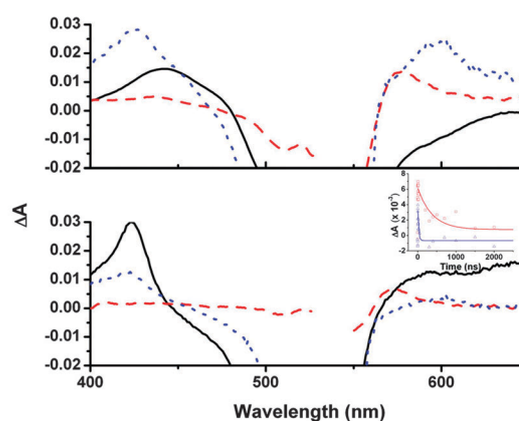


Fig. 3 Transient absorption spectra of **1** top: 10 ps after excitation (1.2 μJ), bottom: 2 ns after excitation (1.6 μJ); black solid line = **1** in CH₂Cl₂, red dashed line = **1**/NiO, blue dotted line = **1**/NiO and I₃⁻/I⁻. Inset: kinetic traces of **1**/NiO (red) at 575 nm and **1**/NiO and I₃⁻/I⁻ at 600 nm (blue). Symbols represent data, lines represent fit of single exponential decay.



the relative amplitudes after excitation of the signals at 425 nm (which contains contributions from both $^1\mathbf{1}^*$ and $\mathbf{1}^-$) and 575 nm ($\text{NiO}^+/\mathbf{1}^-$ only) at early (2 ps) and late (2 ns) times. This analysis suggests that the charge-separation efficiency is closer to 50% and this result is more in agreement with the maximum IPCE for the solar cells. The peak attributed to $\text{NiO}^+/\mathbf{1}^-$ at 575 nm ($\tau_2 \approx 180$ ns) lived three orders of magnitude longer than the charge-separated state of the widely studied P1 dye.¹⁶ This substantial increase in the charge-separated state lifetime prompted us to investigate the effect of substituting the I_3^-/I^- for the tris(4,4'-di-*tert*-butyl-2,2'-dipyridyl)cobalt(III/II) redox shuttle in our DSCs in an attempt to increase the photovoltage.⁷ Unfortunately the photocurrent was negligible in this case, suggesting that the lifetime was still not sufficient for re-oxidation of $\mathbf{1}^-$ by the cobalt electrolyte.[§]

τ_1 for the $\text{NiO}^+/\mathbf{1}^-$ signal differed little in the presence of I_3^-/I^- (Fig. 3 blue dotted line) and absence of I_3^-/I^- (Fig. 3 red dashed line, shifted to $\lambda_{\text{max}} = 600$ nm), hence there was no measurable effect of the electrolyte on a short timescale. However, on a ns timescale the signal for $\text{NiO}^+/\mathbf{1}^-$ decayed more rapidly ($\tau_2 \approx 23$ ns) in the presence of the electrolyte, indicating that interception of the electron by the redox shuttle efficiently competes with recombination. This re-oxidation is very fast compared to the μs timescale reported for re-reduction of dyes on TiO_2 .²² The mechanism for the dye regeneration pathway in n-DSCs is still under debate and it is currently believed that two equivalents of I^- per dye molecule are required.^{22–24} For p-DSCs, re-oxidation of the dye requires only one molecule of I_3^- per dye and this may be why we observe this process proceeding more quickly. Reported timescales for the regeneration reaction range from ps to μs .^{9,25} Reasons for the differing kinetics could be the different driving force for the reaction (reported $\Delta G_{\text{reg}} < 200$ meV to 1 eV; $\Delta G_{\text{reg}} = 640$ meV for $\mathbf{1}$) and the overlapping signals for reduced dye and I_2^- produced from the reduction of I_3^- ($\lambda = 420\text{--}475$ nm).

In our TA experiments on $\mathbf{1}/\text{NiO}$ in the presence of electrolyte we also observed an additional peak centred at 425 nm which decayed with $\tau = 500$ ns. At first we were concerned that the presence of the heavy atoms in the electrolyte had driven the $\text{S}_1 \rightarrow \text{T}_1$ conversion as observed by Morandeira *et al.* for Coumarin 343/ NiO ²⁵ but the band at 650 nm that accompanied the higher energy peak for $^3\mathbf{1}^*$ in CH_2Cl_2 was absent in this sample. Instead, we have assigned this transient as I_2^- . The extinction coefficient of I_2^- should be three times greater than that of the reduced bodipy. The different amplitudes at 420 nm vs. 600 nm in the TA spectrum 2 ns after excitation indicates that both reduced dye and reduced bodipy are present in similar quantities. Therefore we have compelling evidence that electron transfer from the photoreduced dye to I_3^- generating I_2^- contributes to the photocurrent in p-DSCs.

In conclusion, the three orders of magnitude increase in charge-separated state lifetime is a significant breakthrough in our efforts to improve the efficiency of dye-sensitized photocathodes. We anticipate that tuning the electronic coupling by modifying the substituents on the bodipy will increase the charge-separated state yield and lifetime further. This will enable higher photocurrents to be obtained and alternative electrolytes

to be used which increase the photovoltage. These results have wider implications to the field of “solar fuels” since photocatalysis requires long-lived charge-separated intermediates for the desired chemical reactions to take place.

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Notes and references

‡ $\Delta G_{\text{inj}} = e[E_{\text{VB}}(\text{NiO}) - E_{\text{D}^*/\text{D}^-}]$; $\Delta G_{\text{reg}} = e[E(\text{I}_3^-/\text{I}_2^{\bullet-}) - E_{\text{D}^*/\text{D}^-}]$; $E_{(\text{D}^*/\text{D})^-} = E_{(\text{D}/\text{D})^-} + E_{0,0}$; $E_{\text{VB}}(\text{NiO}) \approx -0.12$ V vs. $\text{Fe}(\text{Cp})_2^{+/0,15}$; $E^0(\text{I}_3^-/\text{I}_2^{\bullet-}) = -0.82$ V vs. $\text{Fe}(\text{Cp})_2^{+/0}$ in acetonitrile;¹⁵ $\eta_{\text{inj}} = 1 - (\tau_{1/\text{NiO}}/\tau_1)$.

§ This is consistent with the work of Le Pleux *et al.* who reported that the oxidation of the PMI-NDI sensitiser with $\text{Co}(\text{III/II})$ occurred on the μs timescale ($\tau = 3.5$ μs) and was competitive with charge recombination.

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