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Ultrafast dye regeneration in a core-shell NiO-dye-TiO₂ mesoporous film†

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In this study, a core-shell NiO-dye-TiO2 mesoporous film was fabricated for the first time, utilizing atomic layer deposition technique and a newly designed triphenylamine dye. The structure of the film was confirmed by SEM, TEM, and EDX. Excitation of the dye led to efficient and fast charge separation, by hole injection into NiO, followed by an unprecedentedly fast dye regeneration ($t_{1/2} \le 500$ fs) by electron transfer to TiO2. The resulting charge separated state showed a pronounced transient absorption spectrum caused by the Stark effect, and no significant decay was found within 1.9 ns. This indicates that charge recombination between NiO and TiO2 is much slower than that between the NiO and the reduced dye in the absence of the TiO_2 layer ($t_{1/2} \approx 100$ ps).

One strategy to obtain efficient dye-sensitized solar cells (DSSCs), is that dye regeneration has to outcompete charge recombination processes between injected electrons/holes and the oxidized/reduced dye. 1,2 In the standard DSSCs with liquid electrolyte, the dye is regenerated by a redox mediator in the electrolyte. The regeneration time typically ranges from $t_{1/2}$ = 100 ns to $t_{1/2} = 10 \ \mu s^{3-5}$ due to diffusion limited regeneration kinetics. By using a small electron donor tris(p-anisyl)amine (TPAA) in a cobalt complex electrolyte, the dye regeneration time can be shortened to 100 ps, and the performance of the DSSC device was therefore significantly improved.⁶ In some studies, the initial dye regeneration by I⁻/I₃⁻ redox couple was found to take place within ~ 50 ps. This was explained by the formation of a pre-association complex between the dye and the electrolyte. This process was found in both p-type⁵ and n-type⁷⁻¹⁰ DSSCs.

Another strategy is to design solid state DSSCs, which deliberately eliminates diffusion limitation and therefore accelerates dye regeneration. Here, the electrolyte is replaced by hole/electron conductors in DSSCs. By this, the regeneration of the dye could be accelerated to ~ 50 ps¹¹ due to the close and fixed proximity between the dye and the solid state electron/hole conductor. In p-type DSSCs, the recombination between the reduced dye and the hole in NiO often occurs on the time scale of hundreds of picoseconds. 12-15 This implies that a fast regeneration rate of the dye is required to outcompete the charge recombination process and obtain a satisfactory device performance.

NiO and TiO2 have been successfully used as large band gap semiconductors in p-type and n-type DSSCs, respectively.1,2 Both hole injection into NiO and electron injection into TiO₂ from sensitized dyes have been proven as ultrafast processes, 12,16,17 which inspired us to use a dye to connect NiO and TiO2. It could be a good method to obtain ultrafast dye regeneration. Previously, few attempts have been made to connect NiO and TiO2 via a dye. 18-20 However, all these relevant studies prepared the mesoporous NiO-TiO2 film first, and sensitized with dyes afterwards. 18-20 This procedure led to dye adsorption on both NiO and TiO2 surface (Scheme S1, ESI†). Therefore, the film configuration and sensitization order inhibit complete charge separation from every dye molecule into each semiconductor (h⁺ into NiO and e⁻ into TiO₂), as most dyes lack the alignment allowing both electron and hole injection by a single dye molecule.

We aim for a method to make a so-called "well-oriented" configuration featuring a NiO-dye-TiO2 structure, in which the electron donor part (D) of the charge transfer dye should be close to p-type semiconductor NiO and the electron acceptor part (A) should be nearby the layer of n-type semiconductor TiO2. With this kind of design, we anticipate to get fast and efficient hole and electron injection from the dye upon light illumination. Here, atomic layer deposition (ALD) can provide us with a method to make such an ideal "well-oriented" dye sensitized film, i.e. burying the dye molecules by ALD TiO2 on the dye sensitized mesoporous NiO film. This technique has

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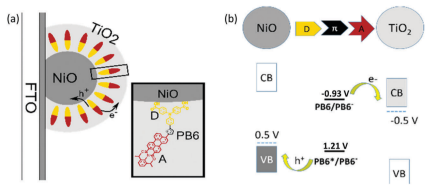


Fig. 1 (a) The proposed well-oriented mesoporous NiO-dye-TiO₂ core-shell structure and the donor-acceptor dye (PB6) reported in this work. D: electron donor part, triphenylamine (TPA); A: electron acceptor, perylene monoimide (PMI). (b) The schematic drawing of charge transfer processes of PB6 between NiO and TiO2 including the electrochemical potentials vs. normal hydrogen electrode (NHE). CB: conduction band; VB: valence band;

been previously applied to prevent dye desorption, ^{21–23} increase the electron injection efficiency by replacing mesoporous TiO2 film with a thin TiO2 overlayer,24,25 and reduce the charge recombination rate by depositing Al₂O₃ as passivation layer.^{26,27} In the present study, good orientation between the semiconductors and the dyes was achieved by ALD of TiO2 on an organic dye sensitized NiO film (see Fig. 1a). Because the dye has anchoring groups on D to covalently bind to NiO, the D unit is close to NiO. The A moiety is buried in ALD TiO₂ layer; therefore, it is close to TiO2 by physical contact. This "well-oriented" NiOdye-TiO2 film was subsequently used to study its light induced charge transfer processes.

To avoid dye degradation during the ALD process and to make the reduced and oxidized states of the dye spectrally distinguishable in the following study, a new organic dye (PB6) was designed, featuring an electron donor-π linker-electron acceptor structure (see inset in Fig. 1(a)). A triphenylamine (TPA) moiety was chosen as D, and a thiophene group and a perylene monoimide (PMI) group was used as linker and A, respectively. Perylene-based dyes have been reported to have high chemical and thermal stability, as well as distinguishable and reversible photooxidation and photoreduction properties.²⁸⁻³⁰ The carboxylic acid groups in PB6 act as the anchoring units to sensitize the dye on the surface of NiO. The detailed synthesis and characterization of PB6 can be found in the ESI.†

The NiO mesoporous layer was fabricated by doctor blading³¹ on fluorine-doped tin oxide (FTO) glass with a pre-prepared compact NiO blocking layer (60 nm) to prevent unwanted direct contact between the ALD TiO₂ and the FTO glass. The NiO film was immersed into a dye solution for sensitization. ALD of TiO2 was performed on the PB6 sensitized NiO film to form the coreshell dye sensitized film. For more details, see Fig. 1 and Scheme S2 (ESI†).

An absorption peak situated at 530 nm was detected from a PB6 sensitized NiO film (NiO-PB6), which is similar to the PB6 absorption in solution (Fig. S4, ESI†). The absorption spectrum of the film after ALD of TiO2 (NiO-PB6-TiO2) showed the same features as the film before ALD except for a 45 nm red-shift of the spectrum without significant decrement of magnitude

(Fig. S4, ESI†). This shift is attributed to the change of dielectric strength of the surrounding of PB6 after coating by high-dielectric TiO2.21 It is encouraging that no significant degradation of PB6 after ALD was observed.

The configuration of a prepared core-shell dye sensitized film (NiO-PB6-TiO2) was imaged by SEM and TEM (Fig. 2). A more compact surface (Fig. 2b) was formed after ALD of TiO2 compared to the typical mesoporous characteristic of NiO (Fig. 2a) due to the increased size of the nanoparticles after ALD coating. TEM in Fig. 2c and d show that the crystalline NiO nanoparticles (black dots) were coated by a conformal amorphous TiO₂ layer (gray layer around NiO particles, ca. 7 nm). The results obtained prove the core-shell configuration of the proposed dye sensitized film. The SEM of the cross section of the NiO-PB6-TiO₂ film (Fig. 2e) displays the thickness of the mesoporous photoactive film of ca. 1.2 µm. EDX element analysis of the corresponding cross section reveals that the Ti signal was found inside the NiO layer, indicating that the TiO₂ layer penetrates the mesoporous NiO film (Fig. 2e).

Electrochemical reduction and oxidation potentials of PB6 were obtained from differential pulse voltammetry (DPV) of NiO-PB6 and PB6 sensitized TiO2 film (TiO2-PB6), respectively, in which reversible features involved in both processes were observed (Fig. S6, ESI†). The obtained energy alignment indicated that the hole ($\Delta G^0 = -710 \text{ mV}$) and electron injection ($\Delta G^0 = -490 \text{ mV}$) into the NiO valence band (VB) and the TiO₂ conduction band (CB), respectively, from the excited PB6 ($E_{0-0} = 2.14$ eV) were thermodynamically favorable, as indicated in Fig. 1b and Table S1 (ESI†).

In order to investigate the possibility of electron injection from PB6 into the TiO₂, photo-induced absorption (PIA) measurements were performed. The PB6 sensitized TiO2 sample (TiO2-PB6) and the TiO2 coated PB6 sample on non-reactive ZrO2 (ZrO2-PB6-TiO2) have the same pronounced peak formed at 920 nm, which by spectroelectrochemistry (Fig. 3 black curve) is assigned to the absorption of the oxidized PB6. It also proves the feasibility of electron injection from excited PB6 into amorphous TiO2 only by the physical contact. This is consistent with recent reports that amorphous TiO₂ can be utilized as electron extraction layer. 32-34 Interestingly, in the PIA of ZrO₂-PB6-TiO₂, there is a negative peak **PCCP**

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10 nm $^{-}$ II

Fig. 2 Micro-structure of NiO-PB6-TiO₂. (a and b) SEM surface images before and after ALD; (c and d) TEM figures, the insert figure is selected area electron diffraction (SAED). (e) SEM image of cross section: I, mesoporous NiO layer coated by TiO2; II, NiO blocking layer; III, FTO layer, and the corresponding EDX element mapping analysis.

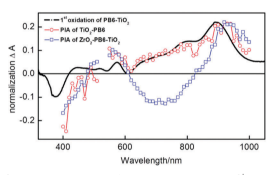


Fig. 3 Spectroelectrochemistry of the 1st oxidation cation PB6+ on PB6-TiO₂ shown as difference spectrum (black curve); photo induced absorption spectra (PIA) of TiO₂-PB6 (red curve) and ZrO₂-PB6-TiO₂ (blue curve) with 530 nm excitation light.

around 600-900 nm. However, the PIA of ZrO2-PB6 sample (Fig. S18, ESI†) did not have such a negative peak, so the fluorescence should not be the explanation for this peak. One hypothesis is that the PIA of ZrO₂-PB6-TiO₂ is an overlapped spectrum between absorption of oxidized PB6 and Stark effect of PB6. The Stark effect in ZrO₂-PB6-TiO₂ is probably formed from holes in oxidized dye or holes trapped on the ZrO2 surface³⁵ and electrons injected in TiO₂. Actually, we did see a weak Stark effect of ZrO₂-PB6-TiO₂ from transient absorption spectroscopy (TAS) (Fig. S13 and S16, ESI†). To further verify this, we introduced one layer of Al₂O₃ (1 nm) between ZrO₂ and TiO₂ after dye sensitization (Fig. S9, ESI†). If the peak around 600-900 nm is from Stark effect, increasing distance between the donor of dye or ZrO2 and TiO2 should weaken the electric field in the presence of identical amount charges, resulting in relatively weaker Stark effect. After normalizing the absorption peak of oxidized PB6 in PIAs, the peak at 600-900 nm in ZrO₂-PB6-TiO2 is weakened in the presence of an extra Al2O3 layer,

see Fig. 3 and Fig. S19 (ESI†). Therefore, we could assign the negative peak at 600-900 nm in ZrO2-PB6-TiO2 from Stark effect.

Time-correlated single photon counting (TCSPC) was employed to further experimentally investigate hole/electron injection into NiO VB and TiO₂ CB, respectively. In the presence of NiO or TiO₂, the decay of photoluminescence after excitation at 470 nm was accelerated as compared to ZrO2 only (see Fig. S8 and Table S2, ESI†). The resulting injection rates (\sim 98% hole injection in NiO-PB6 within \sim 35 ps and 70% electron injection in TiO₂-PB6 within 0.46 ns) suggest that the hole injection occurs first in the NiO-PB6-TiO₂ film (Fig. S8 and Table S2, ESI†). However, the relative fluorescence quantum yield calculated from the lifetimes obtained by TCSPC was higher than those obtained from the steady state fluorescence intensity measurements (see Fig. S9 and Table S3, ESI†). This indicates an additional quenching process on a shorter time scale than that resolved by TCSPC.

In order to capture this additional quenching process and the reduced dye formation upon light illumination, femtosecond TAS (fs-TAS) was performed. After excitation of the NiO-PB6-TiO₂ sample at 560 nm, a clear negative signal was observed in the range from 480 nm to 620 nm due to the ground state bleach (see Fig. 4a). A shoulder in the early spectra around 650 nm is present in the fs-TAS of NiO-PB6-TiO2, but not in the ZrO₂-PB6-TiO₂ (Fig. S13 and S16, ESI†). This shoulder is in agreement with the reduced PB6 as confirmed by spectroelectrochemistry of NiO-PB6 (Fig. S10, ESI†). On the NiO films, the reduced species PB6 is formed within the time resolution of the instrument (200 fs) due to ultrafast hole injection from excited PB6 into NiO. Intermolecular charge transfer between PB6 dyes on the surface can be excluded as the reduced dye spectrum only is detectable on the NiO samples but not on the non-injecting ZrO₂ samples. Previous studies reported hole

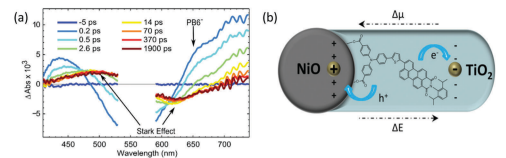


Fig. 4 (a) Transient absorption spectra of NiO-PB6-TiO₂ film from -5 ps to 1.9 ns excited by pulse laser of 560 nm, 300 nJ per pulse; (b) the schematic drawing of electric field formed after charge separation, $\Delta\mu$ change of dipole moment upon excitation and ΔE difference electrical field after charge separation.

injection on the same time scale in different NiO-dye systems. ^{11,12} In the NiO-PB6 sample, most of the transient absorption signal of the reduced PB6 decays within 100 ps without any significant spectral change. This is in agreement with efficient charge recombination between the hole in NiO and the reduced dye as commonly seen in p-type DSSCs. ¹²⁻¹⁴

In the NiO-PB6-TiO $_2$ film, the TA spectrum of the PB6⁻ decays quickly from 200 fs onwards. In contrast to NiO-PB6, a pronounced spectrum with a peak around 500 nm and a bleach around 620 nm was formed at the same time (Fig. 4a). This spectrum matches the first derivative of the steady state absorption spectrum of the NiO-PB6-TiO $_2$ film (Fig. S15, ESI†). Therefore, it can be attributed to the Stark effect of PB6 located within an electric field generated after electron injection from the reduced dye into TiO $_2$ (dye regeneration), *i.e.* NiO⁺-PB6-TiO $_2$ ⁻, as depicted in Fig. 4b.

From the kinetic traces at 530 nm (ground state bleach) and 650 nm (predominantly PB6⁻) in the NiO-PB6-TiO₂ film (Fig. 5), the formation of the Stark effect signal can be traced. By this, dye regeneration of PB6⁻ by the TiO₂ can be concluded to take place on an ultrafast time scale, $t_{1/2} \leq 500$ fs. Global analysis was performed by fitting the kinetic traces at several wavelengths to a sum of three exponentials and an offset (see Fig. 5 and Fig. S14, ESI†). The regeneration was found to be biphasic with time constants of 360 fs and 3.7 ps.

Hole and electron injection observed by fs-TAS are indeed occurring on a much faster time scale than the instrument

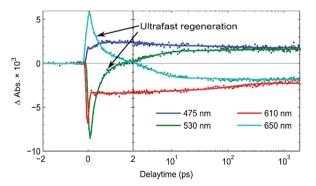


Fig. 5 Kinetics of the fs TAS signal of NiO-PB6-TiO $_2$ at significant wavelengths after excitation at 560 nm.

resolution of TCSPC (30 ps). It explains why steady state fluorescence intensity is much more quenched than expected from the TCSPC lifetime measurements. From the fs-TAS data we estimate hole and electron injection efficiencies up to 99.9% and 93%, respectively, assuming that all ultrafast quenching is due to charge separation (see Table S3, ESI†).

During the formation of the Stark effect signal, the transient absorption at 475 nm and 610 nm only shows minor changes in the fs-TAS measurement window. No clear decay has been observed in the fs-TAS and the kinetic traces. It indicates that relatively long-lived electrons and holes can be localized in TiO₂ and NiO semiconductors, respectively, for more than 1.9 ns without noticeable recombination. This is in contrast to the NiO-PB6 sample (see Fig. S12, ESI†), where most of the signal of the reduced PB6⁻ decays much faster; a bi-exponential fit gives lifetimes of 6 ps and 90 ps. The long lived signal for NiO-PB6-TiO₂ is surprising in comparison to the recent electron transfer study in amorphous TiO₂ system with a covalently linked dye, where up to 60% of the oxidized dye recombined in 220 ps.³⁴

Conclusions

A dye sensitized NiO–dye–TiO $_2$ core–shell architecture film was successfully fabricated for the first time. TEM measurements confirmed that the NiO nanoparticles were covered by a conformal TiO $_2$ layer, allowing both close contact between the dye and semiconductors and "well-oriented" dye alignment for optimal electron/hole injection. The dye was observed to be rather stable during ALD process. This film architecture renders ultrafast and efficient hole injection (>98%, \leq 200 fs) and dye regeneration (70–93%, \leq 500 fs) proven by the appearance of a spectrally pronounced charge separated state (Stark effect). The recombination in the core–shell NiO–dye–TiO $_2$ structure was found to be much slower than that of the conventional dye sensitized NiO film. The efficient charge separation and dye regeneration make this architecture interesting for further applications in solid state DSSCs as well as in dye sensitized solar fuel devices.

Conflicts of interest

The authors declare no competing financial interests.

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References

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- 1 B. O'regan and M. Grätzel, Nature, 1991, 353, 737.
- 2 J. He, H. Lindström, A. Hagfeldt and S.-E. Lindquist, J. Phys. Chem. B, 1999, 103, 8940.
- 3 A. Y. Anderson, P. R. Barnes, J. R. Durrant and B. C. O'Regan, J. Phys. Chem. C, 2011, 115, 2439.
- 4 J. N. Clifford, E. Palomares, M. K. Nazeeruddin, M. Grätzel and J. R. Durrant, J. Phys. Chem. C, 2007, 111, 6561.
- 5 A. Morandeira, G. Boschloo, A. Hagfeldt and L. Hammarström, J. Phys. Chem. C, 2008, 112, 9530.
- 6 Y. Hao, W. Yang, L. Zhang, R. Jiang, E. Mijangos, Y. Saygili, L. Hammarström, A. Hagfeldt and G. Boschloo, Nat. Commun., 2016, 7, 13934.
- 7 S. J. Simon, F. G. Parlane, W. B. Swords, C. W. Kellett, C. Du, B. Lam, R. K. Dean, K. Hu, G. J. Meyer and C. P. Berlinguette, J. Am. Chem. Soc., 2016, 138, 10406.
- 8 L. Casarin, W. B. Swords, S. Caramori, C. A. Bignozzi and G. J. Meyer, Inorg. Chem., 2017, 56, 7324.
- 9 W. B. Swords, G. Li and G. J. Meyer, *Inorg. Chem.*, 2015, 54, 4512.
- 10 L. J. Antila, P. Myllyperkio, S. Mustalahti, H. Lehtivuori and J. Korppi-Tommola, J. Phys. Chem. C, 2014, 118, 7772.
- 11 L. Zhang, G. Boschloo, L. Hammarström and H. Tian, Phys. Chem. Chem. Phys., 2016, 18, 5080.
- 12 A. Morandeira, G. Boschloo, A. Hagfeldt and L. Hammarström, J. Phys. Chem. B, 2005, 109, 19403.
- 13 P. Qin, J. Wiberg, E. A. Gibson, M. Linder, L. Li, T. Brinck, A. Hagfeldt, B. Albinsson and L. Sun, J. Phys. Chem. C, 2010, **114**, 4738.
- 14 E. Sheibani, L. Zhang, P. Liu, B. Xu, E. Mijangos, G. Boschloo, A. Hagfeldt, L. Hammarström, L. Kloo and H. Tian, RSC Adv., 2016, 6, 18165.
- 15 Y. Yu, K. A. Click, S. M. Polen, M. He, C. M. Hadad and Y. Wu, J. Phys. Chem. C, 2017, 121, 20720.

- 16 Y. Tachibana, J. E. Moser, M. Grätzel, D. R. Klug and J. R. Durrant, J. Phys. Chem., 1996, 100, 20056.
- 17 N. J. Cherepy, G. P. Smestad, M. Grätzel and J. Z. Zhang, J. Phys. Chem. B, 1997, 101, 9342.
- 18 K. H. Wong, K. Ananthanarayanan, S. R. Gajjela and P. Balaya, Mater. Chem. Phys., 2011, 125, 553.
- 19 Y.-M. Lee and C.-H. Lai, Solid-State Electron., 2009, 53, 1116.
- 20 J. Bandara and H. Weerasinghe, Sol. Energy Mater. Sol. Cells, 2005, 85, 385.
- 21 H.-J. Son, C. Prasittichai, J. E. Mondloch, L. Luo, J. Wu, D. W. Kim, O. K. Farha and J. T. Hupp, J. Am. Chem. Soc., 2013, 135, 11529.
- 22 R. J. Kamire, M. B. Majewski, W. L. Hoffeditz, B. T. Phelan, O. K. Farha, J. T. Hupp and M. R. Wasielewski, Chem. Sci.,
- 23 K. Hanson, M. D. Losego, B. Kalanyan, D. L. Ashford, G. N. Parsons and T. J. Meyer, Chem. Mater., 2012, 25, 3.
- 24 L. Alibabaei, M. K. Brennaman, M. R. Norris, B. Kalanyan, W. Song, M. D. Losego, J. J. Concepcion, R. A. Binstead, G. N. Parsons and T. J. Meyer, Proc. Natl. Acad. Sci. U. S. A., 2013, 110, 20008.
- 25 L. Alibabaei, B. D. Sherman, M. R. Norris, M. K. Brennaman and T. J. Meyer, Proc. Natl. Acad. Sci. U. S. A., 2015, 112, 5899.
- 26 G. Natu, Z. Huang, Z. Ji and Y. Wu, Langmuir, 2011, 28, 950.
- 27 C. Prasittichai and J. T. Hupp, J. Phys. Chem. Lett., 2010, 1, 1611.
- 28 D. Gosztola, M. P. Niemczyk, W. Svec, A. S. Lukas and M. R. Wasielewski, J. Phys. Chem. A, 2000, 104, 6545.
- 29 A. Osuka, N. Mataga and T. Okada, Pure Appl. Chem., 1997,
- 30 P. Qin, H. Zhu, T. Edvinsson, G. Boschloo, A. Hagfeldt and L. Sun, J. Am. Chem. Soc., 2008, 130, 8570.
- 31 L. D'Amario, L. J. Antila, B. Pettersson Rimgard, G. Boschloo and L. Hammarström, J. Phys. Chem. Lett., 2015, 6, 779.
- 32 I. S. Kim, R. T. Haasch, D. H. Cao, O. K. Farha, J. T. Hupp, M. G. Kanatzidis and A. B. Martinson, ACS Appl. Mater. Interfaces, 2016, 8, 24310.
- 33 I. S. Kim, D. H. Cao, D. B. Buchholz, J. D. Emery, O. K. Farha, J. T. Hupp, M. G. Kanatzidis and A. B. Martinson, Nano Lett., 2016, 16, 7786.
- 34 M. K. Gish, A. M. Lapides, M. K. Brennaman, J. L. Templeton, T. J. Meyer and J. M. Papanikolas, J. Phys. Chem. Lett., 2016, 7, 5297.
- 35 J. Chavez, R. Devine and L. Koltunski, J. Appl. Phys., 2001, 90, 4284.