REVIEW ARTICLE

View Article Online

Porphyrin-sensitized solar cells

Cite this: Chem. Soc. Rev., 2013, 42, 291

Lu-Lin Li and Eric Wei-Guang Diau*

Nature has chosen chlorophylls in plants as antennae to harvest light for the conversion of solar energy in complicated photosynthetic processes. Inspired by natural photosynthesis, scientists utilized artificial chlorophylls - the porphyrins - as efficient centres to harvest light for solar cells sensitized with a porphyrin (PSSC). After the first example appeared in 1993 of a porphyrin of type copper chlorophyll as a photosensitizer for PSSC that achieved a power conversion efficiency of 2.6%, no significant advance of PSSC was reported until 2005; beta-linked zinc porphyrins were then reported to show promising device performances with a benchmark efficiency of 7.1% reported in 2007. Meso-linked zinc porphyrin sensitizers in the first series with a push-pull framework appeared in 2009; the best cell performed comparably to that of a N3-based device, and a benchmark 11% was reported for a porphyrin sensitizer of this type in 2010. With a structural design involving long alkoxyl chains to envelop the porphyrin core to suppress the dye aggregation for a push-pull zinc porphyrin, the PSSC achieved a record 12.3% in 2011 with co-sensitization of an organic dye and a cobalt-based electrolyte. The best PSSC system exhibited a panchromatic feature for light harvesting covering the visible spectral region to 700 nm, giving opportunities to many other porphyrins, such as fused and dimeric porphyrins, with near-infrared absorption spectral features, together with the approach of molecular co-sensitization, to enhance the device performance of PSSC. According to this historical trend for the development of prospective porphyrin sensitizers used in PSSC, we review systematically the progress of porphyrins of varied kinds, and their derivatives, applied in PSSC with a focus on reports during 2007-2012 from the point of view of molecular design correlated with photovoltaic performance.

Received 12th July 2012 DOI: 10.1039/c2cs35257e

www.rsc.org/csr

Department of Applied Chemistry and Institute of Molecular Science, National Chiao Tung University, Hsinchu 30010, Taiwan, E-mail: diau@mail.nctu.edu.tw: Fax: +886-3-5723764: Tel: +886-3-5131524



Lu-Lin Li

Lu-Lin Li is an Assistant Research Fellow in National Chiao Tung University (Taiwan). She received her PhD in Chemistry from National Chung Hsing University, Taichung, Taiwan, in 2006. After two years as a post-doctoral fellow at Center of Nanoscience and Nanotechnology in NCHU, she joined NCTU as a postdoctoral fellow with Prof. Diau in 2008. Her main research interest is fabrication and characterization

of novel functional materials for dye-sensitized solar cells, with a focus on using electrochemical methods to synthesize nanostructured materials relevant for solar energy conversion, and to characterize their interfacial properties using impedance spectroscopy.



Eric Wei-Guang Diau

Eric Diau is a Professor in Department of Applied Chemistry of NCTU (Taiwan). He received his PhD in Chemistry from NTHU (Taiwan) in 1991 and was a postdoctoral fellow with Prof. Ahmed Zewail at Caltech (1997– 2001). He is interested in studying relaxation dynamics of organic and bio-organic molecules in solutions, as well as interfacial electron transfer and energy transfer dynamics in many dye/ semiconductor systems. Starting

from 2006, he devoted to the developments of novel functional materials for dye-sensitized solar cells (DSSC), with a focus on design, synthesis, processing and characterization of photosensitizers and nanostructured materials for DSSC applications. He has published more than 120 peer-reviewed papers.

1. Introduction

Review Article

Dye-sensitized solar cells (DSSC) have attracted much attention because they present a highly promising alternative to conventional photovoltaic devices based on silicon.¹⁻⁵ Fig. 1 shows a schematic diagram revealing the operation of a typical DSSC device, which consists of a dye-sensitized mesoporous working electrode (TiO2, anode) and a counter electrode (Pt-coated, cathode), and has an electrolyte (iodine-based or cobalt complexes, redox mediator) filling the space between the anode and the cathode. Upon light illumination, excited electrons in the LUMO level of a sensitizer are rapidly injected into the conduction band (CB) of TiO2, and then transferred to the platinized counter electrode through exterior electric circuits. The holes are reduced by the redox couple, either I^-/I_3^- or Co^{2^+}/Co^{3^+} , and the oxidized dye thereby regenerated by the I⁻ species (or a Co²⁺ complex) to produce the I_3 species (or a Co³⁺ complex). The diffusion of the oxidized species, the I₃ or Co³⁺ complex, to the surface of the cathode completes the circuit.

In nanocrystalline TiO2 solar cells sensitized with a dye, efficiencies of conversion of light to electric power (η) greater than 11% have been obtained with polypyridyl ruthenium complexes.6-11 The cost, rarity and environmental issues of ruthenium complexes limit, however, their wide application, and encourage exploration of cheaper and safer sensitizers. Scientists have made tremendous efforts in seeking new and efficient sensitizers suitable for practical use in DSSC. 8-44 Fig. 2 shows the evolution of photovoltaic performances of DSSC from 1991 to 2012 based on three types of potential photosensitizers being widely investigated.5 For the ruthenium-based sensitizers, the efficiency of the N3 dye reached 10.0% already since 1993;² the benchmark performance of the N3/N719 family was reported to be 11.2% since 2005,6 and the record efficiency of the C106 dye reached 11.7% reported in 2010.10 The efficiencies of metal-free organic dyes were reported to be 9-10% during recent five years with the best-performed organic dye

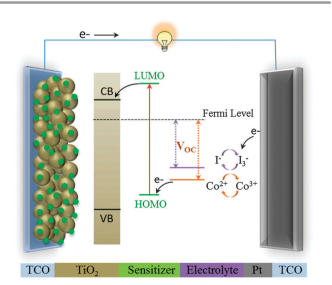


Fig. 1 Schematic representation of the composition and the operating principle of a DSSC.

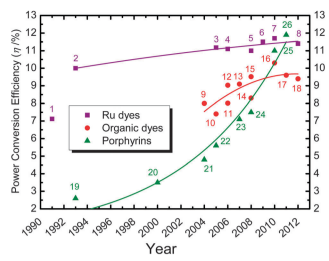


Fig. 2 Efficiency progressing records of DSSC from 1991 to 2012 on the basis of three representative sensitizers labelled 1-8 for Ru-based complexes (I), 9-18 for organic dyes (•) and 19-26 for porphyrin dyes (🛕). The labelled sensitizers may have an alternative given name or a specific code given by the authors: (1) trinuclear Ru dye, 1 (2) N3, 2 (3) N3, 6 (4) N719, 7 (5) C101, 25 (6) CYC-B11, 8 (7) C106,¹⁰ (8) black dye,¹¹ (9) indoline dye,²⁶ (10) NKX-2677,²⁷ (11) JK2,²⁸ (12) D149, ²⁹ (13) TA-St-CA, ³⁰ (14) MK-2, ³¹ (15) D205, ³² (16) C219, ³³ (17) Y123, ³⁴ (18) C218,³⁵ (19) Cu-2-a-oxymesoisochlorin,³⁶ (20) TCPP,³⁷ (21) Zn-1a,³⁸ (22) Zn-3,³⁹ (23) GD2, 40 (24) tda-2b-bd-Zn, 41 (25) YD-2, 42 and (26) YD2-oC8. 24

(C219) reaching $\eta = 10.3\%$. The trend in the performance progress of metal-free organic sensitizers seems to reach a bottleneck for their further development. However, recent developments on porphyrin-based solar cells exhibit a promising advance with the progress curve showing a feature of exponential rise. Moreover, porphyrin sensitizers have drawn great interest because of their excellent light-harvesting function mimicking photosynthesis. 16-24,36-44

In the photosynthetic cores of bacteria and plants, solar energy is collected at chromophores based on porphyrin;²³ the captured radiant energy is converted efficiently to chemical energy. Inspired by this efficient energy transfer in naturally occurring photosynthetic reaction centres, numerous porphyrins have been designed and synthesized for DSSC applications. 16-22 The intrinsic advantages of porphyrin-based dyes are their rigid molecular structures with large absorption coefficients in the visible region and their many reaction sites, i.e., four meso and eight β positions, available for functionalization: fine tuning of the optical, physical, electrochemical and photovoltaic properties of porphyrins thus becomes feasible. In particular, advances in optimization of the device performance for a zinc porphyrin sensitizer (YD2-oC8) co-sensitized with an organic dye (Y123) using a cobalt-based electrolyte to enhance photovoltage of the device (as demonstrated in Fig. 1) attained an unprecedented power conversion efficiency of $\eta = 12.3\%$, ²⁴ which is superior to devices based on Ru complexes, 6-10 stimulating investigation of the development of further porphyrin sensitizers to promote the device performance of porphyrin-sensitized solar cells (PSSC). In this report, we systematically review the recent development of PSSC and introduce a strategy to design potential porphyrin sensitizers for highly efficient DSSC applications.

Fig. 3 Typical structure of a porphyrin showing the four meso- and the eight β positions to be functionalized for porphyrin-sensitized solar cells.

2. Porphyrins with varied linkers

In DSSC, an anchoring group is required at the edge of the dye to link the dye with the semiconductor through a chemical bond. A carboxylic acid is so far considered to be the best binding group for porphyrins, 43 but other promising anchoring groups such as 8-hydroxylquinoline (HQ) have been reported. 44

The structure of a porphyrin exhibits two major points - four meso-positions and eight β -positions, as shown in Fig. 3, that can serve to functionalize one or multiple linkers containing carboxylic acids or HQ substitutes as anchoring groups that attach to the surface of TiO2. In what follows we introduce the historical development of PSSC beginning with β-linked porphyrin sensitizers because we learned that lesson first from natural porphyrins and chlorophylls.

2.1 Linkers at β-positions

When Kay and Grätzel³⁶ reported in 1993 the first PSSC based on a copper chlorophyllin sensitizer with efficiency $\eta = 2.6\%$, a Ru-based DSSC had already attained $\eta = 10\%$.² In the succeeding decade there were almost no progress in developing β-linked porphyrins for PSSC until 2004: Officer, Grätzel and their co-workers38,43 reported zinc porphyrins in a series of which a promising sensitizer (Zn-1a, Fig. 4a) attained $\eta = 4.8\%$. They subsequently investigated 13 prospective zinc porphyrins and reported the best one (Zn-3, Fig. 4b) to attain $\eta = 5.6\%$ in the presence of co-adsorbent chenodeoxycholic acid (CDCA).³⁹ In 2007, the same groups reported porphyrin sensitizers in another series with the best one (GD2, Fig. 4c) reaching $\eta =$ 7.1% for PSSC, 40 opening a great opportunity for the development of various porphyrin sensitizers to enhance the efficiency of a DSSC.

The porphyrin sensitizers shown in Fig. 4 were designed to have a π -conjugated link at the β -position of the porphyrin ring; the best dye (GD2) featured a malonic acid as an anchoring group to enhance the electronic coupling of the dye with the surface of TiO2. Such a concept to design β-functionalized porphyrin sensitizers was then applied by Kim and co-workers,41 who demonstrated that zinc porphyrin 2b-bdta-Zn (Fig. 5a) with two equivalent π -conjugated malonic-acid linkers effectively enhanced the efficiency of electron injection and retarded charge recombination. Kim and co-workers⁴⁵ reported the β-functionalized zinc porphyrins with a diarylamino group for which the porphyrin coded as tda-2b-bd-Zn (Fig. 5b) exhibits the best performance with $\eta = 7.5\%$ that is comparable to the performance of a N3 dye $(\eta = 7.7\%)$ under the same conditions.

2.2 Linkers at meso-positions

The concept for the design of meso-ethynyl linked porphyrins was first given by Anderson⁴⁶ and Therien.⁴⁷ Although Cherian and Wamser³⁷ reported in 2000 the first meso-substituted PSSC with $\eta \sim 3.5\%$ based on a free-base porphyrin, no significant progress was made for porphyrin sensitizers of this type until 2007: Galoppini and co-workers⁴⁸ reported tetrachelated zinc porphyrins with four meta-substituted linkers at four mesopositions of the porphyrin to suppress dye aggregation; Lindsey, Meyer and their co-workers⁴⁹ reported *meso*-substituted porphyrin and chlorin derivatives to extend the absorption spectra to larger wavelengths; Imahori and co-workers⁵⁰ examined the meso-substituted porphyrins with five-membered heteroaromatic linkers. The strategy of Galoppini et al. 48,49 was to use a phenylethynyl (PE) unit as a bridging moiety to control the distance between the sensitizer and the semiconductor to retard charge recombination, which works well for pyrene⁵¹ and Ru complexes⁵² in model systems, but this approach failed when three or four PE units were combined to form a long bridge at one meso-position of a zinc porphyrin because of serious dye aggregation involved in porphyrins with a long PE linker. 53-55

To solve the aggregation problem of porphyrins, 3,5-di-tertbutylphenyl groups were introduced at the meso-positions of the porphyrin ring. Based on this molecular design, Yeh, Diau and co-workers⁵⁶ reported *meso*-substituted zinc porpyrin derivatives, for which YD0 (Fig. 6a) served as a reference compound. With the diarylamino group attached at the meso-position of

Fig. 4 Molecular structures of (a) Zn-1a, 38 (b) Zn-339 and (c) GD2.40

$$(a) \qquad \qquad (b) \qquad \qquad (b)$$

Fig. 5 Molecular structures of (a) 2b-bdta-Zn⁴¹ and (b) tda-2b-bd-Zn.⁴⁵

the porphyrin core, YD1 (Fig. 6b) attained $\eta = 6.0\%$, which was comparable to that of a N3 dye under the same experimental conditions ($\eta = 6.1\%$). The superlative cell performance of YD1 reflects its extraordinarily great short-circuit current density (I_{SC}) that arises from the large efficiency of conversion of incident photons to current (IPCE) broadly extending beyond 700 nm. The electron donor in YD1 plays a role not only spectrally to extend the absorption to a region of greater wavelength but also spatially pushing the excited electrons toward TiO₂ for an improved separation of charge. The appearance of porphyrins with a push-pull framework such as YD1 indicates the arrival of the era of PSSC.

Yeh and Diau^{57,58} designed another promising push-pull porphyrin, YD2 (Fig. 6c) based on the structure of YD1 in that two tert-butyl groups in the diarylamino substituent were replaced with two long alkyl chains to improve the thermal and photochemical stability of a device. This design mimics the strategy applied in an amphiphilic ruthenium polypyridyl sensitizer (Z907) that has shown excellent stability toward water-induced desorption under both thermal stress and light soaking.⁵⁹ The device performance of YD2 was further improved to η = 6.8%, slightly better than that of YD1 (η = 6.5%) but slightly less than that of N719 ($\eta = 7.3\%$);⁵⁷ the electron-donating feature of amino substituents in YD1 and YD2 seems to be responsible for the $V_{\rm OC}$ value being larger than that of the reference cell (YD0). 57,58 Using GD2 as an example, Mozer et al.⁶⁰ noted that the smaller V_{OC} of PSSC is due to the decreased electron lifetime related to a more rapid recombination of electrons with I_3^- ions. The observed V_{OC} of YD1 and YD2 larger than that of YD0 was thus expected to be due to a diminished recombination between I_3^- and conduction-band electrons, because I₃ might be attached to the positively

charged diarylamino moiety far from the TiO2 surface in the former case.57

In 2010, the device performance of YD2 was further improved by Grätzel and co-workers, 42 giving J_{SC}/mA cm⁻² = 18.6, $V_{\rm OC}/V = 0.77$, FF = 0.764, and $\eta = 10.9\%$, which lasted as a record for PSSC until 2011.

Multiple donor groups at the meso-positions

Because the diarylamino group plays a key role in promoting the device performance of YD2, the effects of the substituted number and position of electron-donating groups (EDG) became an interesting subject to test. Accordingly, Yeh and Diau⁶¹ reported push-pull zinc porphyrins with more than one EDG at the meso-positions of the porphyrin ring and found that YD2 was still the best sensitizer for PSSC. Almost at the same period Imahori and co-workers⁶² designed other pull-pull porphyrins with zero, one and two EDG, in which the two EDG are located at *meso*-positions of porphyrin to form a *cis* or a trans isomer. The same conclusion was made by Imahori that the mono-substituted porphyrin with a structure similar to that of YD2 gives the best device performance.

2.4 Thiophene substitutes in the linkers

The thiophene group has been widely utilized in rutheniumbased⁸⁻¹⁰ and metal-free organic solar cells¹²⁻¹⁵ to enhance the absorption coefficient of the dye and red-shift its absorption spectrum. This concept has been adopted for porphyrin-based DSSC. Thiophene units were used in both β-linked⁶³ and *meso*linked⁶⁴ porphyrin sensitizers to give efficiencies 4.0% and 5.1%, respectively. The IPCE results indicated that increasing the number of thiophene units does not extend the spectra; the spectral edge reaches ~ 650 nm in both cases. Hung and co-workers⁶⁵ found that the number of thiophene units substituted at the meso-position has a negative effect on device performance. Unlike photosensitizers of other types, thiophene substitutes play a lesser role in promoting the efficiencies of PSSC.

Porphyrins with extended spectral feature

The most viable way to enhance $J_{\rm SC}$ is to harvest a broader region of the solar spectrum. In general, porphyrins show a Soret band at 400-450 nm and Q bands at 500-650 nm. To extend the absorption of porphyrin dyes to the near infrared region, the energy gap between the highest occupied molecular

Fig. 6 Molecular structures of (a) YD0, (b) YD1 and (c) YD2.⁵⁷

orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) levels must decrease, for which purpose there are two approaches: one is to introduce a highly conjugated π -extended chromophore coupled with the porphyrin ring, and another is to make fused or dimeric porphyrins.

3.1 Porphyrins functionalized with π -extended chromophores

The early results^{39,41} indicated that the π -conjugation in the β-substituted porphyrins has a limited effect to extend the absorption spectra to greater wavelength. The best strategy to extend the π -conjugation is thus to functionalize the target porphyrin at the *meso*-positions. A promising chromophore is the acene family, for which the π -conjugation can be effectively extended with an increasing number of aromatic rings. As a result, Lin and Diau⁶⁶ designed acenes, from benzene to pentacene, as π -extended chromophores to couple to the porphyrin core through the linkage of the acenyl-ethynyl group. Among those porphyrins, the anthracene-functionalized porphyrin (LAC-3, Fig. 7a) showed the best performance, which reached $\sim 80\%$ of a N3-based device under the same conditions; the tetracene-functionalized porphyrin (LAC-4) featured a broad IPCE spectrum extending to ~800 nm, but the IPCE values were too small (<50%) to give a satisfactory performance; the pentacene-substituted porphyrin (LAC-5) even showed an extended absorption spectrum beyond 900 nm, but almost no photocurrents were produced.⁶⁶ When Lin and Diau^{67,68} further designed cyclic aromatic substituents attached at the meso-position of the macrocycle opposite to the anchoring group, they found that the fluorene-functionalized porphyrin (LD22, Fig. 7b) featured an impressive device performance $\eta = 8.1\%$, ⁶⁷ and the pyrene-functionalized porphyrin (LD4, Fig. 7c) attained η = 10.1%, ⁶⁸ which was superior to that of a N719 dye ($\eta = 9.3$ %) under the same conditions. The superior photovoltaic performance of the LD4-based PSSC was attributed to its enhanced ability to harvest light with the IPCE action spectrum covering the

entire visible spectral region and extending beyond 800 nm, which outperforms N719. $V_{\rm OC}$ of device LD4 was much smaller than that of cell N719, because of a much smaller electron lifetime of porphyrin-based solar cells than that of N719 cells. 60 $V_{\rm OC}$ of LD4 was even smaller than that of YD2, which might be rationalized in that electron interception is more efficient for LD4 than for YD2 because of improved charge separation for the latter.⁵⁷

Based on the structure of YD2, Yeh and Diau⁶⁹ designed the acenyl-ethynyl unit as a link in porphyrins YD11-YD13. As shown in Fig. 8a-c, the bridge between ethyne and carboxyl groups is varied from phenylene for YD11 to naphthylene for YD12, and to anthracenylene for YD13. Without an added scattering layer, the photovoltaic performances of both YD11 and YD12 exhibited performance superior relative to that of N719 dye with I_{SC} of the two promising porphyrin-based devices being significantly greater than those of N719 devices, making the overall efficiencies of power conversion of YD11 ($\eta = 6.7\%$) and YD12 ($\eta = 6.8\%$) outperform that of the N719 device ($\eta =$ 6.1%).69 YD11 (an analogue of YD2) and YD12 are thus two green sensitizers remarkable for their outstanding cell performances relative to that of N719 without an added scattering layer for light-penetrable DSSC applications. When the TiO2 films (~10 μm) were covered with a scattering layer, the cell performance of N719 significantly improved to $\eta = 7.3\%$, whereas the performances of the porphyrin dyes increased only marginally ($\eta = 6.8\%$ and 7.0% for YD11 and YD12, respectively).⁶⁹ These results indicate that a substantial increase in $J_{\rm SC}$ for the N719 device is a key factor for the improvement of the cell performance with the addition of a scattering layer. 70 Based on those observations, the involvement of the partially allowed triplet MLCT states of ruthenium complexes was concluded to be responsible for the enhanced efficiency in the red shoulder of the IPCE spectrum of N719, whereas the effect of spin-orbit coupling in zinc porphyrins was insufficient for the $S_0 \rightarrow T_1$

Fig. 7 Molecular structures of (a) LAC-3, 66 (b) LD22 67 and (c) LD4. 68

Fig. 8 Molecular structures of (a) YD11, (b) YD12 and (c) YD13.69

transitions to occur the additional scattering layer provided no (5.0% under

transitions to occur; the additional scattering layer provided no improvement in the IPCE spectra of YD11–YD13 beyond the Q-band absorptions. $^{69}\,$

Functionalized chromophore anthracene plays an important role to extend the π -conjugation in LAC-3 for an enhanced overall device performance, ⁶⁶ but anthracene in YD13 with a link shorter than that in LAC-3 exhibited a notable effect to deteriorate significantly the device performance of YD13. ⁶⁹ Results obtained from femtosecond measurements of fluorescence decay indicate that the presence of the anthracene group in the bridge from YD13 to TiO_2 did not hamper the rate of interfacial electron transfer for the observed small injection yield of YD13; rather, it was the anthracene-induced rapid relaxation of intermolecular energy due to dye aggregation that gave the poor device performance of YD13, which was also evident in experiments in the absence and presence of co-adsorbent CDCA. ⁶⁹

Apart from the cyclic aromatic hydrocarbon systems introduced above, perylene is a promising functional chromophore to modify a porphyrin sensitizer for these two reasons: the strong absorptions of the perylene derivatives in the spectral region 500–650 nm⁷¹ are complementary to those of porphyrins, and a device made of a push–pull perylene anhydride sensitizer yields an impressive performance, $\eta = 6.8\%$. Efforts to integrate the perylene chromophores into porphyrin sensitizers showed, however, negative effects for the improvement of device performance of PSSC, ^{73–75} mainly because the co-planar structural nature between perylene and porphyrin moieties caused a serious problem of dye aggregation.

3.2 Fused porphyrins

Review Article

According to many above examples, porphyrins and their derivatives are promising photosensitizers for DSSC because of their intense absorption in the Soret and Q bands to harvest solar energy efficiently over a broad spectral region, but the existence of a gap between the Soret and Q bands in monoporphyrins limits their cell performance. One strategy to improve the light-harvesting ability of the sensitizer is to fuse a chromophore with a porphyrin to make a π -elongated macrocycle for PSSC. This idea became mature and practical when Imahori and co-workers^{76–78} reported their first examples, of which a device made of a naphthalene-based *meso*-β-edge fused zinc porphyrin (fused-Zn-1, Fig. 9a) attained $\eta = 4.1\%$

Fig. 9 Molecular structures of (a) fused-Zn-1^{76,77} and (b) ZnQMA.⁷⁹

(5.0% under co-sensitization), which was improved by half relative to the reference cell with an unfused porphyrin.^{76,77} Such an unsymmetrical π -elongation was achieved by the same group⁷⁹ to construct two quinoxaline-based β - β' -edge fused zinc porphyrins, of which a fused porphyrin with one anchoring group (ZnQMA, Fig. 9b) exhibited $\eta = 5.2\%$, attaining 80% performance of a N719 device under the same conditions. Compare the IPCE spectra of the two fused porphyrins: even though the fused-Zn-1 dye involves a broad light-harvesting feature extending the IPCE action spectrum to nearly 800 nm, a large gap in the middle of the spectrum limits the growth of photocurrents to an optimal condition. In contrast, the IPCE spectrum of the ZnQMA dye extends to only ~700 nm, but an effective electronic coupling between quinoxaline and porphyrin moieties diminishes the gap between the Soret and Q bands of the spectrum, leading to improved J_{SC} and a performance to that of fused-Zn-1.

Wang and Wu⁷⁵ designed and synthesized two perylene anhydride fused nickel porphyrin sensitizers (WW1 in Fig. 10a and WW2 in 10b) to extend the light absorption towards the near-infrared (~1000 nm) region. As those fused porphyrins suffer from dye aggregation, the IPCE values did not exceed 30%, which was unable to make the devices attain a notable performance ($\eta \sim 1.3\%$). Yeh and Diau⁸⁰ designed and synthesized two fused porphyrins (YDD2 in Fig. 10c and YDD3 in Fig. 10d), but they exhibited poor cell performance. For YDD2, the absorption spectrum extends even beyond 1200 nm, but essentially no photocurrent was observed because the energy level of LUMO was substantially lower than the conduction band edge of TiO2. For YDD3, although a small response was observed in the IPCE action spectrum corresponding to the contribution of broad bands I and II of the fused porphyrin, nearly no response was observed for the broad band III in region 700-900 nm. 80 WW2 and YDD3 are, nevertheless, two interesting panchromatic porphyrin sensitizers with the potential to extend the light-harvesting ability toward the near-infrared region for PSSC.

3.3 Dimeric porphyrins

Another strategy to improve light-harvesting ability of the sensitizer is to combine two porphyrin moieties through a chemical bond. The first attempt using dimeric porphyrins as potential sensitizers for PSSC was made in 2009 by Officer and co-workers; they reported the photovoltaic properties of porphyrin dimers comprising two monoporphyrin units connected in either a linear *anti* or a 90° *syn* fashion. The dimeric porphyrin dyes exhibited light-harvesting efficiencies slightly improved relative to the corresponding monomeric porphyrin dyes when the devices were fabricated on a thin TiO₂ film (thickness $\sim 3~\mu m$), but the effect of π -conjugation for the red shift of the IPCE spectra due to porphyrin dimerization was small as the link between the two porphyrins was made at the β -position.

The first effort to link the two porphyrins at the *meso*-position was made in 2009 by Kim and Osuka;⁸² their polyethanediol (PEG)-modified dimer with two β -substituted linkers similar to those of the tda-2b-bd-Zn monomer (Fig. 5b) showed

Fig. 10 Molecular structures of fused perylene-porphyrins (a) WW1 and (b) WW2;⁷⁵ fused porphyrin dimers (c)YDD2 and (d)YDD3.⁸⁰

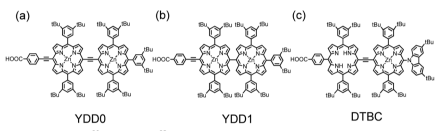


Fig. 11 Molecular structures of (a) YDD0, (b) YDD180 and (c) DTBC.83

the best device performance of the various dimers, giving an overall efficiency η = 4.2%. Similarly, the link between two porphyrins with only a single CC bond showed no effect of significant spectral shift to enhance the light-harvesting power for the porphyrin dimers.

Based on the structure of YD0 (Fig. 6a), Yeh and Diau⁸⁰ designed two porphyrin dimers, YDD0 (Fig. 11a) and YDD1 (Fig. 11b). Similar to the results of Kim and Osuka,82 the spectrum of YDD1 exhibited a spectral feature only slightly red-shifted relative to that of the monomeric porphyrin YD0, but, because of effective excitonic coupling between the two nearly perpendicular porpyrin units in YDD1, the gap shown in the IPCE spectrum of YD0 was completely filled in the spectrum of YDD1, making J_{SC} of the YDD1 device become much greater than that of the YD0 device. With the link of an ethynyl group between the two porphyrin units, YDD0 showed split Soret bands in the range of 400-500 nm, and red shifts and broadening of the Q bands extending to nearly 800 nm. The broad IPCE spectrum of YDD0 showed much smaller efficiency, < 40%, than those of YD0 and YDD1 (\sim 70%) because of the co-planar structural feature of YDD0 resulting in serious dye aggregation.

Similar to the structural design of YDD0, Segawa and co-workers83 added a carbazole EDG at the meso-position of the porphyrin edge to form a push-pull porphyrin dimer, DTBC (Fig. 11c). DTBC has an absorption spectrum similar to that of YDD0, but its IPCE spectrum showed much greater efficiency than that of YDD0. In the absence of a TBP additive, the DTBC device exhibited IPCE values up to 80%, yielding a remarkable $J_{\rm SC}/{\rm mA~cm}^{-2}$ = 18.2 but also causing a poor $V_{\rm OC}/{\rm V} \sim 0.4$; the poor $V_{\rm OC}$ was significantly improved in the presence of TBP to attain an optimized device performance $\eta = 5.2\%$, 83 comparable to the performance of YDD0, $\eta = 4.1\%$.

4. Strategies to suppress dye aggregation for porphyrins

The reported near-infrared dyes such as fused porphyrins⁷⁵⁻⁸⁰ and dimeric porphyrins⁸⁰⁻⁸³ introduced herein are particularly interesting, but the planar structural feature of those porphyrins might facilitate the formation of dye aggregates that significantly decrease the efficiency of electron injection. In the following sections we describe two plausible strategies to suppress effectively the dye aggregation for PSSC.

4.1 Enveloping porphyrins with long alkoxyl chains

The great performance of porphyrins such as YD2, 42,57,58 LD4⁶⁸ and YD12⁶⁹ was due to their superior light-harvesting ability through introduction of an EDG or a π -extended chromophore at the meso-position of the porphyrin ring. V_{OC} of those highly efficient porphyrin dyes was reported, however, to be significantly

Fig. 12 Molecular structures of (a) LH3⁸⁴ and (b) ZnPDCA.⁸⁵

Fig. 13 Molecular structures of (a) LD12,89 (b) LD1489 and (c) LD16.88

less than that of the commonly used ruthenium dye N719. The significantly diminished electron lifetime was reported to account for the smaller $V_{\rm OC}$ of porphyrins, and I_3 in the electrolyte might become attached to the positively charged Zn-center of the porphyrin core for efficient electron interception from the TiO2 surface.60 Tian and co-workers concluded that $V_{\rm OC}$ can be improved on decreasing the charge recombination and increasing the efficiency of electron injection with an appropriate design of an organic dye. 15 To tackle this problem for porphyrins, a new concept was introduced to design a zincporphyrin sensitizer with long alkoxyl chains to protect the porphyrin core for retarded charge recombination and also to decrease effectively the dye aggregation for an efficient electron injection.

The examples based on such a molecular design were first given in 2010 by Hupp and co-workers;84,85 they reported porphyrin sensitizers with two phenyl groups attached at the 5,15-meso-positions bearing two dodecoxyl (-OC₁₂H₂₅) chains at the ortho-position of each phenyl group. Among those orthosubstituted porphyrins, LH3 (Fig. 12a)84 and ZnPDCA (Fig. 12b)⁸⁵ displayed the best performance of each series, with the device made of ZnPDCA attaining efficiency η = 5.5%, comparable to that of N719 under the same conditions. 85 Before Hupp's results were published, Imahori and co-workers had announced their results based on ortho-substituted porphyrins, 86,87 but the alkyl chains (only methyl and ethyl groups were considered) substituted at the ortho-positions were too short to protect effectively the porphyrins from dye aggregation.⁸⁸

The concept for the design of ortho-substituted porphyrins was first demonstrated by Lin, Diau, and their co-workers. 88,89 Fig. 13a-c show the molecular structures of porphyrins LD12, LD14 and LD16, respectively, based on this design. The photovoltaic results indicate that J_{SC} becomes significantly improved on incorporating the π -conjugated EDG for both LD14 and LD16 with respect to LD12, and $V_{\rm OC}$ of those ortho-substituted porphyrins is significantly greater than that of a para- or metasubstituted counterpart; the best devices (LD14 and LD16) attained power conversion efficiencies beyond 10%. 88,89 Results from molecular-dynamics (MD) simulation⁸⁵ indicate that the porphyrin core of LD14 is fully enveloped by the four dodecoxyl chains to provide more effective insulation of the dye, resulting in diminished molecular aggregation and increased solubility in non-coordinating organic solvents. The transient photoelectric results indicate that the upward shift of the TiO2 potential and the retarded charge recombination are two important factors for the enhanced $V_{\rm OC}$ of the *ortho*-substituted porphyrins.⁸⁹ The long alkoxyl chains thus play a key role to prevent the approach of I₃ in the electrolyte to the surface of TiO₂ so as to retard the electron interception at the electrolyte/TiO2 interface. A similar idea for the molecular design of ortho-substituted porphyrins YD20-YD22 with noticeable device performances was reported elsewhere.90

Work developed along this track led to an advance for a device made of an ortho-substituted push-pull zinc porphyrin, YD2-oC8 (Fig. 14a).24 Its design was based on YD2, for which the four tert-butyl side chains in the meso-phenyls of YD2 were replaced by four ortho-substituted alkoxyl chains so as to

(a) (b)
$$C_6H_{17}OC_6H_{13}$$
 $C_6H_{13}OC_6H_{13}$ $C_6H_{13}OC_6H_{13}$ $C_6H_{13}OC_6H_{13}$ $C_6H_{13}OC_6H_{13}$ $C_6H_{13}OC_6H_{13}$ $C_6H_{13}OC_6H_{13}OC_6H_{13}$ $C_6H_{13}OC_$

Fig. 14 Molecular structures of (a) YD2-oC8²⁴ and (b) Y123.^{34,91}

envelope effectively the porphyrin ring to decrease the degree of dye aggregation. The molecular design of YD2-oC8 is similar to that of LD1489 and LD1688 of which outstanding device performances were reported based on the iodide/triiodide redox electrolyte.

In 2011, Grätzel and co-workers²⁴ reported an optimized performance of the YD2-oC8 device under AM1.5 one-sun irradiation: $J_{SC}/\text{mA cm}^{-2} = 17.7$, $V_{OC}/V = 0.935$, FF = 0.74, and η = 12.3%, obtained when YD2-oC8 was co-sensitized with an organic dye (Y123, Fig. 14b) 34,91 on a TiO₂ film of (12 + 5) μ m in a cobalt-based redox electrolyte.24 The efficiency of power conversion attained η = 13.1% under 0.5-sun irradiation. Light-induced photoelectric measurements of the YD2-oC8 devices support that the enhanced V_{OC} of the ortho-substituted devices is due to the upward shift of the TiO2 conduction band and the enhanced electron lifetimes. The long alkoxyl chains in those ortho-substituted porphyrins thus play an important role to diminish effectively the degree of dye aggregation for an improved device performance. These results 24,84,85,88-90 might provide a conceptual basis for the molecular design of porphyrin sensitizers to attain even greater efficiencies of power conversion in the near future.

4.2 Co-sensitizations

Co-sensitization is an effective approach to enhance the device performance through a combination of two or more dyes with complementary spectral features sensitized on semiconductor films together, extending the light-harvesting ability so as to increase the photocurrents of the devices. Many co-sensitization systems, such as ruthenium complex plus organic dye, 11,92 metalfree organic dye, 93,94 phthalocyanine plus organic dye, 95-97 and dye

Fig. 15 Molecular structures of (a) HC-A1¹⁰⁴ and (b) HC-A3.¹⁰⁶

co-sensitization in separate layers, 98-100 have been reported to show enhanced photovoltaic performance relative to their individual single-dye systems. Based on the organic dye systems, Robertson¹⁰¹ stated that co-sensitization with strongly absorbing dyes would give sufficient space on the surface of TiO2 to allow absorption of other dyes with a complementary absorption spectrum. Porphyrin sensitizers are hence perfect candidates to improve the device performance through co-sensitization.

The first porphyrin/organic dye co-sensitization system was reported in 2010 by Grätzel and co-workers42 who demonstrated an instance of YD2 co-sensitized with an organic dye (D205)102 on a thin TiO2 film (2.4 µm), which showed an enhanced device performance η = 6.9% relative to those of their individual dyes, $\eta = 5.6-5.7\%$. Although two mixed β-linked porphyrins served to enhance the IPCE values by 300%, 103 the overall performance of the co-sensitized system was poor. For the best DSSC system with co-sensitization of YD2-oC8 with Y123 (Fig. 14), ²⁴ the enhancement in J_{SC} was limited – 17.3 mA cm⁻² with YD2-oC8 alone and 17.7 mA cm⁻² with YD2-oC8 + Y123 - because the gap between the Soret and the Q bands in the IPCE spectrum of the porphyrin-alone device was small. Moreover, in not only the YD2-related systems²⁴ but also in many other systems aforementioned, 42,93-97 the values of $V_{\rm OC}$ were between those of the individual single-dye sensitized devices, which limits the device performance of the co-sensitization system to improve further.

In 2011, Kim and co-workers¹⁰⁴ reported the organic dyes (HC-A1, Fig. 15a) as hole conductors to enhance significantly the device performance of a coumarin system. For porphyrins, Kim and co-workers¹⁰⁵ showed that the device performance of a push-pull porphyrin with the cyano-acrylic acid anchoring group co-adsorbed with HC-A1 attained $\eta = 7.2\%$, which was superior to that made of a single porphyrin only ($\eta = 4.4\%$) or porphyrin co-adsorbed with CDCA ($\eta = 6.6\%$). The effects of HC-A1 and HC-A3 (Fig. 15b)¹⁰⁶ on the photovoltaic performance were remarkable in improving not only J_{SC} but also V_{OC} to some extent. The absorptions of the hole conductors reported by Kim et al. are in the ultraviolet region, 104-106 so unable to fill the gap in the IPCE spectrum between the Soret and the Q bands of a porphyrin to improve the photocurrent generation. A superior approach is to use an HC organic dye with an absorption spectrum complementary to that of a porphyrin.

Fig. 16 shows three potential organic dyes that have served as co-sensitizers for PSSC applications. The boron dipyrromethene dye BET (Fig. 16a) belongs to the BODIPY family 107

Fig. 16 Molecular structures of (a) BET, 109 (b) CD4 and (c) CD5. 110

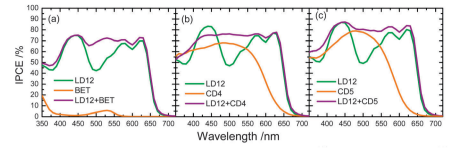


Fig. 17 IPCE spectra of LD12 co-sensitized with (a) BET, 109 (b) CD4 and (c) CD5. 111

with an absorption maximum at ~ 530 nm to compensate for the absorption loss of a porphyrin sensitizer; Lee and Hupp⁸⁴ used a BODIPY moiety as a light-harvesting molecule connected to the meso-position of a zinc porphyrin to make LH3 (Fig. 12a) for PSSC applications; Odobel and co-workers 108 used the BODIPY dye as an antenna attached to the metal centre of a zinc porphyrin derivative to form a supermolecular assembly for DSSC, and possibly also for organic photovoltaic (OPV) applications. When BET was used as a co-sensitizer with an ortho-substituted porphyrin LD12 (Fig. 13a), the gap in the IPCE spectrum of LD12 was completely filled as shown in Fig. 17a, and both J_{SC} = 14.7 mA cm⁻² and η = 7.5% were improved by \sim 12% relative to the device made of LD12 alone. The BETalone device exhibited a poor IPCE response that generated a small photocurrent, $J_{SC} = 0.5 \text{ mA cm}^{-2}$, but co-sensitization remarkably enhanced J_{SC} , which was much greater than the sum of the photocurrents generated from two individual devices. An energy-transfer mechanism was proposed to rationalize the observed exceptional phenomenon, 109 but $V_{\rm OC}$ of the LD12 + BET device was slightly less than that of the LD12 device, probably due to the lack of effective HC character for the BODIPY chromophore.

To design organic dyes with HC character and featuring absorption complementary to that of a porphyrin, Chen and co-workers¹¹⁰ synthesized spirally configured donor-acceptor organic dyes for DSSC applications. Among those dyes, CD4 (Fig. 16b) and CD5 (Fig. 16c) are candidates for co-sensitization with porphyrins such as LD12 (Fig. 13a). The co-sensitization of LD12 with CD4 or CD5 on a TiO₂ film of thickness $\sim 20 \,\mu m$ was achieved via a stepwise approach:111 the TiO2 electrode was immersed in the LD12 solution for 3 h, and then immersed in the CD4 or CD5 solution for 2 h. The co-sensitized film was afterwards assembled into a DSSC device of sandwich type with a Pt-coated counter electrode and filled with a traditional iodide/tri-iodide electrolyte. Fig. 17b and c show the IPCE spectra of LD12 co-sensitized with CD4 and CD5, respectively. Similar to the LD12 + BET system shown in Fig. 17a, the gap of the IPCE response about 500 nm for the LD12 device was filled in the action spectra of the LD12 + CD4 or LD12 + CD5 device with the contribution of CD4 or CD5 that shows its maximum photoresponse in that spectral region. Taking CD5 as an example, 111 $J_{\rm SC}$ of the co-sensitized device increased from 14.97 mA cm $^{-2}$ to 16.74 mA cm $^{-2}$ to contribute \sim 12% enhancement of the overall performance. Moreover, $V_{\rm OC}$ of the co-sensitized device increased from 0.711 V (LD12) and 0.689 V (CD5) to 0.736 V (LD12 + CD5), which takes advantage of the HC character for organic dyes of this series. The observed $V_{\rm OC}$ upon co-sensitization was enhanced through the retarded charge recombination overwhelming the downward shift of the TiO₂ potential according to the results obtained from measurements of charge extraction and intensity-modulated photovoltage spectra.111

We have shown that co-sensitization of complementary porphyrin and organic dyes produces a panchromatic spectral feature to promote the performance of PSSC. For the system LD12 + CD5, 111 the IPCE spectrum extended to \sim 650 nm; for the system YD2-oC8 + Y123,24 the efficiency response extended further to \sim 700 nm. The near-IR dyes previously introduced ^{75,79,80} are thus candidates to elevate the photocurrents and promote the overall cell efficiencies. Yeh and Diau¹¹² designed a dimeric porphyrin (YDD6, Fig. 18), in which eight tert-butyl side chains in YDD0 were replaced with eight ortho-substituted isoamyloxy chains, 113 and the meso-phenyl opposite the anchoring group was replaced

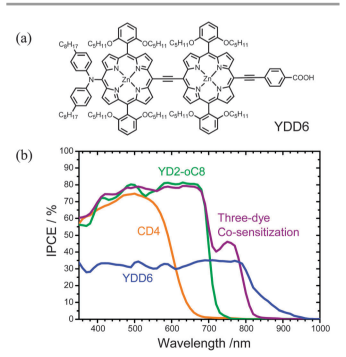


Fig. 18 (a) Molecular structure of YDD6 and (b) IPCE spectra of CD4, YD2-oC8, YDD6 and three-dye co-sensitized systems. 112

Table 1 Summary of current device performance for porphyrin sensitizers or co-sensitization systems

Sensitizers	Device efficiency (%)	Reference efficiency (%)	Ref.
Zn-1a	4.8	_	38
Zn-3	5.6	_	39
GD2	7.1	_	40
2b-bdta-Zn	3.0	N3, 5.9	41
tda-2b-bd-Zn	7.5	N3, 7.7	45
Fused-Zn-1	4.1	Zn-1, 2.8	76, 77
ZnQMA	5.2	N719, 6.5	79 [°]
WW2	1.4		75
KS-3	4.0	_	63
PZn-hT	5.1	N719, 8.0	64
LAC-3	5.4	N719, 6.7	66
LAC-4	2.8	N719, 6.7	66
LD4	10.1	N719, 9.3	68
LD12	7.4		89
LD14	10.2	_	89
LD16	10.2	_	88
LD22	8.1	N719, 9.2	67
YD0	4.5	N719, 7.3	57
YD1	6.5	N719, 7.3	57
YD2	6.8	N719, 7.3	57
YD2	11	_ ′	42
YD11	6.8	N719, 7.3	69
YD12	7.0	N719, 7.3	69
YD13	1.9	N719, 7.3	69
YD14	6.8	YD2, 7.1	61
YD15	4.2	YD2, 7.1	61
YD20	8.1	_ ′	90
YDD0	4.1	YD0, 5.1	80
YDD1	5.2	YD0, 5.1	80
DTBC	5.2	_ ′	83
LH3	1.6	_	84
ZnPDCA	5.5	N719, ~ 6	85
YD2-oC8	11.9	YD2, 8.4 (cobalt electrolyte)	24
YD2-oC8+Y123	12.3	YD2, 8.4 (cobalt electrolyte)	24
2Flu-ZnP-CN-COOH + HC-A1	7.2	N719, 8.6	105
LD12 + BET	7.5	LD12, 6.7	109
LD12 + CD5	9.0	LD12, 7.5	111
YDD6 + YD2-oC8 + CD4	10.4	YD2-oC8, 8.8	112

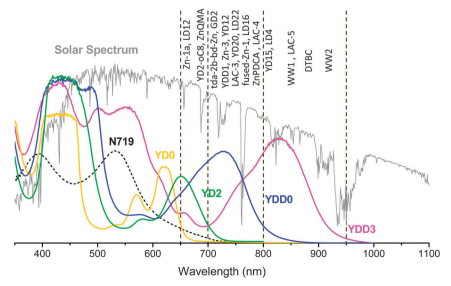


Fig. 19 Thin-film absorption spectra of typical porphyrin sensitizers with spectral edges indicated by dashed lines at 650, 700, 800 and 950 nm for YD0, YD2, YDD0 and YDD3, respectively. The approximate positions of the spectral edges of other porphyrins reviewed in this article are indicated.

with a diarylamino group, similar to the molecular design of YD2-oC8 (Fig. 14a) but with the YDD0 (Fig. 11a) diporphyrin core as a light-harvesting centre. The ortho-substituted alkoxyl chains in YDD6 failed to prevent dye aggregation for this porphyrin; molecular engineering of co-sensitization was implemented in a solution containing three spectrally complementary dyes -YDD6, YD2-oC8 and CD4 - at a specific molar ratio to optimize the device performance of the system. As shown in Fig. 18, the IPCE action spectrum of the co-sensitized device attained 75-80% in region 400-700 nm and 40-45% in region 700-800 nm, giving a short-circuit current density J_{SC} = 19.28 mA cm⁻², which is significantly improved over that of YD2-oC8, 16.60 mA cm⁻², and that of the YD2-oC8 + CD4 co-sensitized system, 16.91 mA cm⁻², because of the additional light-harvesting power in the near-infrared region without degradation of the performance in the visible region. Using an iodine-based electrolyte, the best co-sensitized system achieved a remarkable power conversion efficiency of 10.4%, which is much superior to those obtained from their individual single-dye devices and the two-dye co-sensitized systems. 112 These results indicate that the suppression of dye aggregation for YDD6 in the three-dye co-sensitized system is the key to improvement of the device performance. With progress using this approach of co-sensitization, a future challenge is to seek other potential near-IR co-sensitizers with not only a light-harvesting ability extending beyond 800 nm but also decreased dye aggregation to enhance IPCE above 80%.

5. Summary and future perspectives

Inspired by the efficient energy transfer in naturally occurring photosynthetic reaction centers, numerous porphyrin and dimeric porphyrin sensitizers are introduced in this review of highly efficient dye-sensitized solar cells. At the current stage, the device fabricated using the push-pull zinc porphyrin, YD2oC8, co-sensitized with Y123 on a TiO₂ film and using a cobalt electrolyte has attained an unprecedented efficiency of 12.3% under standard AM 1.5 one-sun irradiation; such efficiency of power conversion has never been achieved with a ruthenium-free sensitizer. Similar to the design of YD2-oC8, ortho-substituted push-pull porphyrins such as LD14, LD16 and YD22 with extended light harvesting towards \sim 750 nm are expected to yield performance at least comparable to that of YD2-oC8. Functional chromophores such as anthracene in LAC-3, tetracene in LAC-4, pentacene in LAC-5, fluorene in LD22, pyrene in LD4, BODIPY in LH3, and perylene anhydride-linked porphyrin derivatives might also play a role to improve the light-harvesting ability when they can be integrated into the framework of an ortho-substituted porphyrin to build a new PSSC. Fused porphyrins such as fused-Zn-1, WW1, WW2, YDD3, and their derivatives, porphyrin dimers such as DTBC, YDD0, YDD6 and their derivatives, are prospective photosensitizers to extend the light-harvesting power toward the near-IR spectral region. In most cases those fused and dimeric porphyrins suffered from poor device performance due to their nearly co-planar structural feature favoring the formation of dye aggregates to deteriorate significantly the device performance - a solution to solve the aggregation problem for porphyrins is to

implement the approach of co-sensitization. Here we highlight the significance of molecular co-sensitization of multiple dyes with complementary absorption spectra on a semiconductor film, which would be an effective approach to enhance the light-harvesting ability and to retard the charge recombination for significant promotion of the overall performance of a PSSC. Table 1 summarizes the current status of device performance for porphyrin sensitizers or co-sensitization systems introduced in this review.

Hardin, Snaith and McGehee¹¹⁴ suggested that extending the dye absorption to 830 nm might increase the efficiency of the YD2-oC8 system to 13.6%; an ultimate device performance in a cobalt-electrolyte system is estimated to be $\sim 19\%$ if the total loss in potential can be decreased to 500 mV and the absorption spectrum of the dye can be extended to 920 nm. We show the absorption spectra of typical porphyrin sensitizers adsorbed on TiO2 thin films in Fig. 19, for which the spectral edges of YD0, YD2, YDD0 and YDD3 are approximately 650, 700, 800 and 950 nm, respectively; the edges of other porphyrins discussed here are also indicated. Both fused porphyrins WW2 and YDD3 have spectral edges of about 950 nm; they are perfect candidates to promote the efficiency of DSSC if suitable structural modifications and appropriate co-sensitization procedure are applied. We believe that this review provides guidance for strategies to design the best donor-acceptor porphyrin system and to apply an optimal co-sensitization for porphyrin and holeconducting organic dyes towards a new record for the future developments and applications of DSSC.

Acknowledgements

National Science Council of Taiwan (NSC101-2627-M-009-007) and Ministry of Education of Taiwan under the ATU program supported this work. We are indebted to many collaborators who provided invaluable porphyrin and organic dyes for PSSC applications reported in this review article: Prof. Chen-Yu Yeh of NCHU (Taiwan) for the YD-series porphyrins, Prof. Ching-Yao Lin of NCNU (Taiwan) for the LD-series porphyrins, Prof. Chien-Tien Chen of NTHU (Taiwan) for the CD-series dyes, Prof. Chen-Hsiung Hung of Academia Sinica (Taiwan) for thienyl porphyrins, Prof. Hwan Kyu Kim of Korea University (Korea) for HC-A1 and HC-A3 dyes, and Prof. Hongshen He of SDSU (USA) for the BET dye. We thank Prof. Michael Grätzel of EPFL (Switzerland) for optimization of the device performance of PSSC and for guiding us in the direction to design new porphyrin sensitizers. We are grateful to Prof. Juan Bisquert of UJI (Spain) for many enlightening discussions on impedance spectra. We thank the researchers for their works cited in this review.

References

- 1 B. Oregan and M. Grätzel, Nature, 1991, 353, 737-740.
- 2 M. K. Nazeeruddin, A. Kay, I. Rodicio, R. Humphrybaker, E. Muller, P. Liska, N. Vlachopoulos and M. Grätzel, J. Am. Chem. Soc., 1993, 115, 6382-6390.
- 3 M. K. Nazeeruddin, P. Pechy, T. Renouard, S. M. Zakeeruddin, R. Humphry-Baker, P. Comte, P. Liska, L. Cevey, E. Costa, V. Shklover, L. Spiccia, G. B. Deacon, C. A. Bignozzi and M. Grätzel, J. Am. Chem. Soc., 2001, 123, 1613-1624.

- 4 M. Grätzel, Acc. Chem. Res., 2009, 42, 1788-1798.
- 5 A. Hagfeldt, G. Boschloo, L. C. Sun, L. Kloo and H. Pettersson, Chem. Rev., 2010, 110, 6595-6663.
- 6 M. K. Nazeeruddin, F. De Angelis, S. Fantacci, A. Selloni, G. Viscardi, P. Liska, S. Ito, T. Bessho and M. Grätzel, J. Am. Chem. Soc., 2005, 127, 16835-16847.
- 7 Q. Wang, S. Ito, M. Grätzel, F. Fabregat-Santiago, I. Mora-Sero, J. Bisquert, T. Bessho and H. Imai, J. Phys. Chem. B, 2006, 110, 25210-25221.
- 8 C. Y. Chen, M. K. Wang, J. Y. Li, N. Pootrakulchote, L. Alibabaei, C. H. Ngoc-le, J. D. Decoppet, J. H. Tsai, C. Grätzel, C. G. Wu, S. M. Zakeeruddin and M. Grätzel, ACS Nano, 2009, 3, 3103-3109.
- 9 Y. M. Cao, Y. Bai, Q. J. Yu, Y. M. Cheng, S. Liu, D. Shi, F. F. Gao and P. Wang, J. Phys. Chem. C, 2009, 113, 6290-6297.
- 10 Q. J. Yu, Y. H. Wang, Z. H. Yi, N. N. Zu, J. Zhang, M. Zhang and P. Wang, ACS Nano, 2010, 4, 6032-6038.
- 11 L. Y. Han, A. Islam, H. Chen, C. Malapaka, B. Chiranjeevi, S. F. Zhang, X. D. Yang and M. Yanagida, Energy Environ. Sci., 2012, 5, 6057-6060.
- 12 N. Robertson, Angew. Chem., Int. Ed., 2006, 45, 2338-2345.
- 13 Y. Ooyama and Y. Harima, Eur. J. Org. Chem., 2009, 2903-2934.
- 14 A. Mishra, M. K. R. Fischer and P. Bäuerle, Angew. Chem., Int. Ed., 2009, 48, 2474-2499.
- 15 Z. J. Ning, Y. Fu and H. Tian, Energy Environ. Sci., 2010, 3, 1170-1181.
- 16 H. Imahori, T. Umeyama and S. Ito, Acc. Chem. Res., 2009, 42, 1809-1818.
- 17 I. Radivojevic, A. Varotto, C. Farley and C. M. Drain, Energy Environ. Sci., 2010, 3, 1897-1909.
- 18 M. V. Martínez-Díaz, G. de la Torrea and T. Torres, Chem. Commun., 2010, 46, 7090-7108.
- 19 M. G. Walter, A. B. Rudine and C. C. Wamser, J. Porphyrins Phthalocyanines, 2010, 14, 759-792.
- 20 M. J. Griffith, K. Sunahara, P. Wagner, K. Wagner, G. G. Wallace, D. L. Officer, A. Furube, R. Katoh, S. Mori and A. J. Mozer, Chem. Commun., 2012, 48, 4145-4162.
- 21 H. Imahori, T. Umeyama, K. Kurotobi and Y. Takano, Chem. Commun., 2012, 48, 4032-4045.
- 22 M. K. Panda, K. Ladomenou and A. G. Coutsolelos, Coord. Chem. Rev., 2012, 256, 2601-2627.
- 23 T. Hasobe, H. Imahori, P. V. Kamat, T. K. Ahn, S. K. Kim, D. Kim, A. Fujimoto, T. Hirakawa and S. Fukuzumi, J. Am. Chem. Soc., 2005, 127, 1216-1228.
- 24 A. Yella, H. W. Lee, H. N. Tsao, C. Yi, A. K. Chandiran, M. K. Nazeeruddin, E. W. Diau, C. Y. Yeh, S. M. Zakeeruddin and M. Grätzel, Science, 2011, 334, 629-634.
- 25 F. Gao, Y. Wang, D. Shi, J. Zhang, M. K. Wang, X. Y. Jing, R. Humphry-Baker, P. Wang, S. M. Zakeeruddin and M. Grätzel, J. Am. Chem. Soc., 2008, 130, 10720-10728.
- 26 T. Horiuchi, H. Miura, K. Sumioka and S. Uchida, J. Am. Chem. Soc., 2004, **126**, 12218–12219.
- 27 K. Hara, Z.-S. Wang, T. Sato, A. Furube, R. Katoh, H. Sugihara, Y. Dan-oh, C. Kasada, A. Shinpo and S. Suga, J. Phys. Chem. B, 2005, 109, 15476-15482.
- 28 S. Kim, J. K. Lee, S. O. Kang, J. Ko, J. H. Yum, S. Fantacci, F. De Angelis, D. Di Censo, M. K. Nazeeruddin and M. Grätzel, J. Am. Chem. Soc., 2006, 128, 16701-16707.
- 29 S. Ito, S. M. Zakeeruddin, R. Humphry-Baker, P. Liska, R. Charvet, P. Comte, M. K. Nazeeruddin, P. Pechy, M. Takata, H. Miura, S. Uchida and M. Grätzel, Adv. Mater., 2006, 18, 1202-1205.
- 30 S. Hwang, J. H. Lee, C. Park, H. Lee, C. Kim, C. Park, M.-H. Lee, W. Lee, J. Park, K. Kim, N.-G. Park and C. Kim, Chem. Commun., 2007, 4887-4889.
- 31 Z. S. Wang, N. Koumura, Y. Cui, M. Takahashi, H. Sekiguchi, A. Mori, T. Kubo, A. Furube and K. Hara, Chem. Mater., 2008, 20,
- 32 S. Ito, H. Miura, S. Uchida, M. Takata, K. Sumioka, P. Liska, P. Comte, P. Pechy and M. Grätzel, Chem. Commun., 2008, 5194-5196.
- 33 W. Zeng, Y. Cao, Y. Bai, Y. Wang, Y. Shi, M. Zhang, F. Wang, C. Pan and P. Wang, Chem. Mater., 2010, 22, 1915-1925.
- 34 H. N. Tsao, C. Yi, T. Moehl, J.-H. Yum, S. M. Zakeeruddin, M. K. Nazeeruddin and M. Grätzel, ChemSusChem, 2011, 4, 591-594.
- 35 M. Xu, M. Zhang, M. Pastore, R. Li, F. De Angelis and P. Wang, Chem. Sci., 2012, 3, 976-983.

- 36 A. Kay and M. Grätzel, J. Phys. Chem., 1993, 97, 6272-6277.
- 37 S. Cherian and C. C. Wamser, J. Phys. Chem. B, 2000, 104, 3624-3629.
- 38 M. K. Nazeeruddin, R. Humphry-Baker, D. L. Officer, W. M. Campbell, A. K. Burrell and M. Grätzel, Langmuir, 2004, 20, 6514-6517.
- 39 Q. Wang, W. M. Carnpbell, E. E. Bonfantani, K. W. Jolley, D. L. Officer, P. J. Walsh, K. Gordon, R. Humphry-Baker, M. K. Nazeeruddin and M. Grätzel, J. Phys. Chem. B, 2005, 109, 15397-15409.
- 40 W. M. Campbell, K. W. Jolley, P. Wagner, K. Wagner, P. J. Walsh, K. C. Gordon, L. Schmidt-Mende, M. K. Nazeeruddin, Q. Wang, M. Grätzel and D. L. Officer, J. Phys. Chem. C, 2007, 111, 11760-11762.
- 41 J. K. Park, H. R. Lee, J. P. Chen, H. Shinokubo, A. Osuka and D. Kim, J. Phys. Chem. C, 2008, 112, 16691-16699.
- 42 T. Bessho, S. M. Zakeeruddin, C. Y. Yeh, E. W. G. Diau and M. Grätzel, Angew. Chem., Int. Ed., 2010, 49, 6646-6649.
- 43 W. M. Campbell, A. K. Burrell, D. L. Officer and K. W. Jolley, Coord. Chem. Rev., 2004, 248, 1363-1379.
- 44 H. He, A. Gurung and L. Si, Chem. Commun., 2012, 48, 5910-5912.
- 45 M. Ishida, S. W. Park, D. Hwang, Y. B. Koo, J. L. Sessler, D. Y. Kim and D. Kim, J. Phys. Chem. C, 2011, 115, 19343-19354.
- 46 T. E. O. Screen, K. B. Lawton, G. S. Wilson, N. Dolney, R. Ispasoiu, T. Goodson Iii, S. J. Martin, D. D. C. Bradley and H. L. Anderson, J. Mater. Chem., 2001, 11, 312-320.
- 47 V. Lin, S. DiMagno and M. Therien, Science, 1994, 264, 1105-1111.
- 48 J. Rochford, D. Chu, A. Hagfeldt and E. Galoppini, J. Am. Chem. Soc., 2007, 129, 4655-4665.
- 49 J. R. Stromberg, A. Marton, H. L. Kee, C. Kirmaier, J. R. Diers, C. Muthiah, M. Taniguchi, J. S. Lindsey, D. F. Bocian, G. J. Meyer and D. Holten, J. Phys. Chem. C, 2007, 111, 15464-15478.
- 50 S. Eu, S. Hayashi, T. Umeyama, A. Oguro, M. Kawasaki, N. Kadota, Y. Matano and H. Imahori, J. Phys. Chem. C, 2007, 111, 3528-3537.
- 51 O. Taratula, J. Rochford, P. Piotrowiak, E. Galoppini, R. A. Carlisle and G. J. Meyer, J. Phys. Chem. B, 2006, 110, 15734-15741.
- 52 C. C. Clark, G. J. Meyer, Q. Wei and E. Galoppini, J. Phys. Chem. B, 2006, 110, 11044-11046.
- 53 C.-Y. Lin, C.-F. Lo, L. Luo, H.-P. Lu, C.-S. Hung and E. W.-G. Diau, J. Phys. Chem. C, 2009, 113, 755-764.
- 54 L. Luo, C. J. Lin, C. Y. Tsai, H. P. Wu, L. L. Li, C. F. Lo, C. Y. Lin and E. W. G. Diau, Phys. Chem. Chem. Phys., 2010, 12, 1064-1071.
- 55 L. Y. Luo, C. J. Lin, C. S. Hung, C. F. Lo, C. Y. Lin and E. W. G. Diau, Phys. Chem. Chem. Phys., 2010, 12, 12973-12977.
- 56 C. W. Lee, H. P. Lu, C. M. Lan, Y. L. Huang, Y. R. Liang, W. N. Yen, Y. C. Liu, Y. S. Lin, E. W. G. Diau and C. Y. Yeh, Chem.-Eur. J., 2009, 15, 1403-1412.
- 57 H. P. Lu, C. Y. Tsai, W. N. Yen, C. P. Hsieh, C. W. Lee, C. Y. Yeh and E. W. G. Diau, J. Phys. Chem. C, 2009, 113, 20990-20997.
- 58 C.-P. Hsieh, H.-P. Lu, C.-L. Chiu, C.-W. Lee, S.-H. Chuang, C.-L. Mai, W.-N. Yen, S.-J. Hsu, E. W.-G. Diau and C.-Y. Yeh, J. Mater. Chem., 2010, 20, 1127-1134.
- 59 P. Wang, S. M. Zakeeruddin, J. E. Moser, M. K. Nazeeruddin, T. Sekiguchi and M. Grätzel, Nat. Mater., 2003, 2, 402-407.
- 60 A. J. Mozer, P. Wagner, D. L. Officer, G. G. Wallace, W. M. Campbell, M. Miyashita, K. Sunahara and S. Mori, Chem. Commun., 2008, 4741-4743.
- 61 S. L. Wu, H. P. Lu, H. T. Yu, S. H. Chuang, C. L. Chiu, C. W. Lee, E. W. G. Diau and C. Y. Yeh, Energy Environ. Sci., 2010, 3, 949-955.
- 62 H. Imahori, Y. Matsubara, H. Iijima, T. Umeyama, Y. Matano, S. Ito, M. Niemi, N. V. Tkachenko and H. Lemmetyinen, J. Phys. Chem. C, 2010, 114, 10656-10665.
- 63 V. A. Nuay, D.-H. Kim, S.-H. Lee and J.-J. Ko, Bull. Korean Chem. Soc., 2009, 30, 2871-2872.
- 64 Y. Liu, N. Xiang, X. Feng, P. Shen, W. Zhou, C. Weng, B. Zhao and S. Tan, Chem. Commun., 2009, 2499-2501.
- 65 R. Ambre, K.-B. Chen, C.-F. Yao, L. Luo, E. W.-G. Diau and C.-H. Hung, J. Phys. Chem. C, 2012, 116, 11907-11916.
- 66 C.-Y. Lin, Y.-C. Wang, S.-J. Hsu, C.-F. Lo and E. W.-G. Diau, J. Phys. Chem. C, 2010, 114, 687-693.
- 67 C. H. Wu, T. Y. Pan, S. H. Hong, C. L. Wang, H. H. Kuo, Y. Y. Chu, E. W. Diau and C. Y. Lin, Chem. Commun., 2012, 48, 4329-4331.
- 68 C.-L. Wang, Y.-C. Chang, C.-M. Lan, C.-F. Lo, E. Wei-Guang Diau and C.-Y. Lin, Energy Environ. Sci., 2011, 4, 1788-1795.
- 69 H. P. Lu, C. L. Mai, C. Y. Tsia, S. J. Hsu, C. P. Hsieh, C. L. Chiu, C. Y. Yeh and E. W. G. Diau, Phys. Chem. Chem. Phys., 2009, 11, 10270-10274.

- 70 S. Ito, T. N. Murakami, P. Comte, P. Liska, C. Grätzel, M. K. Nazeeruddin and M. Grätzel, Thin Solid Films, 2008, 516, 4613-4619.
- 71 T. Edvinsson, C. Li, N. Pschirer, J. Schöneboom, F. Eickemeyer, R. Sens, G. Boschloo, A. Herrmann, K. Müllen and A. Hagfeldt, I. Phys. Chem. C, 2007, 111, 15137-15140.
- 72 C. Li, J.-H. Yum, S.-J. Moon, A. Herrmann, F. Eickemeyer, N. G. Pschirer, P. Erk, J. Schöneboom, K. Müllen, M. Grätzel and M. K. Nazeeruddin, ChemSusChem, 2008, 1, 615-618.
- 73 H. Y. Chen, H. P. Lu, C. W. Lee, S. H. Chuang, E. W. G. Diau and C. Y. Yeh, J. Chin. Chem. Soc., 2010, 57, 1141-1146.
- 74 C.-W. Lee, H.-P. Lu, N. M. Reddy, H.-W. Lee, E. W.-G. Diau and C.-Y. Yeh, Dyes Pigm., 2011, 91, 317-323.
- 75 C. Jiao, N. Zu, K.-W. Huang, P. Wang and J. Wu, Org. Lett., 2011, 13, 3652-3655.
- 76 M. Tanaka, S. Hayashi, S. Eu, T. Umeyama, Y. Matano and H. Imahori, Chem. Commun., 2007, 2069-2071.
- 77 S. Hayashi, M. Tanaka, H. Hayashi, S. Eu, T. Umeyama, Y. Matano, Y. Araki and H. Imahori, J. Phys. Chem. C, 2008, 112, 15576-15585.
- 78 S. Hayashi, Y. Matsubara, S. Eu, H. Hayashi, T. Umeyama, Y. Matano and H. Imahori, Chem. Lett., 2008, 37, 846-847.
- 79 S. Eu, S. Hayashi, T. Umeyama, Y. Matano, Y. Araki and H. Imahori, J. Phys. Chem. C, 2008, 112, 4396-4405.
- 80 C. L. Mai, W. K. Huang, H. P. Lu, C. W. Lee, C. L. Chiu, Y. R. Liang, E. W. Diau and C. Y. Yeh, Chem. Commun., 2010, 46, 809-811.
- 81 A. J. Mozer, M. J. Griffith, G. Tsekouras, P. Wagner, G. G. Wallace, S. Mori, K. Sunahara, M. Miyashita, J. C. Earles, K. C. Gordon, L. C. Du, R. Katoh, A. Furube and D. L. Officer, J. Am. Chem. Soc., 2009, 131, 15621-15623.
- 82 J. K. Park, J. P. Chen, H. R. Lee, S. W. Park, H. Shinokubo, A. Osuka and D. Kim, J. Phys. Chem. C, 2009, 113, 21956-21963.
- Y. Liu, H. Lin, J. T. Dy, K. Tamaki, J. Nakazaki, D. Nakayama, S. Uchida, T. Kubo and H. Segawa, Chem. Commun., 2011, 47, 4010-4012.
- 84 C. Y. Lee and J. T. Hupp, Langmuir, 2010, 26, 3760-3765.
- 85 C. Y. Lee, C. X. She, N. C. Jeong and J. T. Hupp, Chem. Commun., 2010, 46, 6090-6092.
- 86 H. Imahori, S. Hayashi, H. Hayashi, A. Oguro, S. Eu, T. Umeyama and Y. Matano, J. Phys. Chem. C, 2009, 113, 18406-18413.
- S. Mathew, H. Iijima, Y. Toude, T. Umeyama, Y. Matano, S. Ito, N. V. Tkachenko, H. Lemmetyinen and H. Imahori, J. Phys. Chem. C, 2011, **115**, 14415–14424.
- 88 C.-L. Wang, C.-M. Lan, S.-H. Hong, Y.-F. Wang, T.-Y. Pan, C.-W. Chang, H.-H. Kuo, M.-Y. Kuo, E. W.-G. Diau and C.-Y. Lin, Energy Environ. Sci., 2012, 5, 6933-6940.
- 89 Y. C. Chang, C. L. Wang, T. Y. Pan, S. H. Hong, C. M. Lan, H. H. Kuo, C. F. Lo, H. Y. Hsu, C. Y. Lin and E. W. Diau, Chem. Commun., 2011, 47, 8910-8912.
- 90 T. Ripolles-Sanchis, B. C. Guo, H. P. Wu, T. Y. Pan, H. W. Lee, S. R. Raga, F. Fabregat-Santiago, J. Bisquert, C. Y. Yeh and E. W. Diau, Chem. Commun., 2012, 48, 4368-4370.
- 91 A. Dualeh, F. De Angelis, S. Fantacci, T. Moehl, C. Yi, F. Kessler, E. Baranoff, M. K. Nazeeruddin and M. Grätzel, J. Phys. Chem. C, 2011, 116, 1572-1578.

- 92 R. Y. Ogura, S. Nakane, M. Morooka, M. Orihashi, Y. Suzuki and K. Noda, Appl. Phys. Lett., 2009, 94, 073308.
- 93 Y. Chen, Z. Zeng, C. Li, W. Wang, X. Wang and B. Zhang, New J. Chem., 2005, 29, 773-776.
- 94 D. Kuang, P. Walter, F. Nuesch, S. Kim, J. Ko, P. Comte, S. M. Zakeeruddin, M. K. Nazeeruddin and M. Grätzel, Langmuir, 2007, 23, 10906-10909
- 95 J. J. Cid, J. H. Yum, S. R. Jang, M. K. Nazeeruddin, E. M. Ferrero, E. Palomares, J. Ko, M. Grätzel and T. Torres, Angew. Chem., Int. Ed., 2007, 46, 8358-8362
- 96 J. N. Clifford, A. Forneli, H. J. Chen, T. Torres, S. T. Tan and E. Palomares, J. Mater. Chem., 2011, 21, 1693-1696.
- 97 M. Kimura, H. Nomoto, N. Masaki and S. Mori, Angew. Chem., Int. Ed., 2012, 51, 4371-4374.
- 98 H. Choi, S. Kim, S. O. Kang, J. J. Ko, M. S. Kang, J. N. Clifford, A. Forneli, E. Palomares, M. K. Nazeeruddin and M. Grätzel, Angew. Chem., Int. Ed., 2008, 47, 8259-8263.
- 99 F. Inakazu, Y. Noma, Y. Ogomi and S. Hayase, Appl. Phys. Lett., 2008, 93, 093304.
- 100 K. Lee, S. W. Park, M. J. Ko, K. Kim and N. G. Park, Nat. Mater., 2009, 8, 665-671.
- 101 N. Robertson, Angew. Chem., Int. Ed., 2008, 47, 1012-1014.
- 102 D. Kuang, S. Uchida, R. Humphry-Baker, S. M. Zakeeruddin and M. Grätzel, Angew. Chem., Int. Ed., 2008, 47, 1923-1927.
- 103 M. J. Griffith, A. J. Mozer, G. Tsekouras, Y. Dong, P. Wagner, K. Wagner, G. G. Wallace, S. Mori and D. L. Officer, Appl. Phys. Lett., 2011, 98, 163502.
- 104 B. J. Song, H. M. Song, I. T. Choi, S. K. Kim, K. D. Seo, M. S. Kang, M. J. Lee, D. W. Cho, M. J. Ju and H. K. Kim, Chem.-Eur. J., 2011, 17, 11115-11121.
- 105 M. S. Kang, S. H. Kang, S. G. Kim, I. T. Choi, J. H. Ryu, M. J. Ju, D. Cho, J. Y. Lee and H. K. Kim, Chem. Commun., 2012, 48, 9349-9351.
- 106 H. M. Song, K. D. Seo, M. S. Kang, I. T. Choi, S. K. Kim, Y. K. Eom, J. H. Ryu, M. J. Ju and H. K. Kim, J. Mater. Chem., 2012, 22, 3786-3794.
- 107 A. Loudet and K. Burgess, Chem. Rev., 2007, 107, 4891-4932.
- 108 J. Warnan, Y. Pellegrin, E. Blart and F. Odobel, Chem. Commun., 2012, 48, 675-677.
- 109 M. Shrestha, L. P. Si, C. W. Chang, H. S. He, A. Sykes, C. Y. Lin and E. W. G. Diau, J. Phys. Chem. C, 2012, 116, 10451-10460.
- 110 W. S. Chao, K. H. Liao, C. T. Chen, W. K. Huang, C. M. Lan and E. W. Diau, Chem. Commun., 2012, 48, 4884-4886.
- 111 C.-M. Lan, H.-P. Wu, T.-Y. Pan, C.-W. Chang, W.-S. Chao, C.-T. Chen, C.-L. Wang, C.-Y. Lin and E. W.-G. Diau, Energy Environ. Sci., 2012, 5, 6460-6464.
- 112 H.-P. Wu, Z.-W. Ou, T.-Y. Pan, C.-M. Lan, W.-K. Huang, H.-W. Lee, N. M. Reddy, C.-T. Chen, W.-S. Chao, C.-Y. Yeh and E. W.-G. Diau, Energy Environ. Sci., 2012, DOI: 10.1039/c2ee22870j.
- 113 M. P. Nikiforov, U. Zerweck, P. Milde, C. Loppacher, T. H. Park, H. T. Uyeda, M. J. Therien, L. Eng and D. Bonnell, Nano Lett., 2008, 8, 110-113.
- 114 B. E. Hardin, H. J. Snaith and M. D. McGehee, Nat. Photonics, 2012, 6, 162-169.