

Novel Blue Organic Dye for Dye-Sensitized Solar Cells Achieving High Efficiency in Cobalt-Based Electrolytes and by Co-Sensitization

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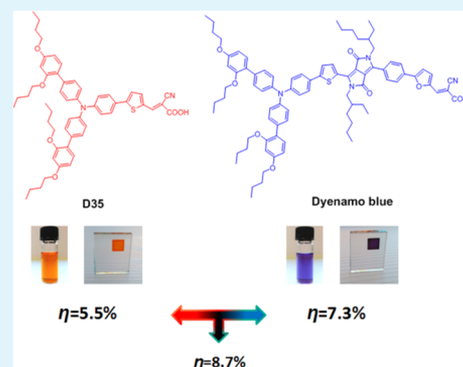
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Supporting Information

ABSTRACT: Blue and green dyes as well as NIR-absorbing dyes have attracted great interest because of their excellent ability of absorbing the incident photons in the red and near-infrared range region. A novel blue D- π -A dye (Dyename Blue), based on the diketopyrrolopyrrole (DPP)-core, has been designed and synthesized. Assembled with the cobalt bipyridine-based electrolytes, the device with Dyename Blue achieved a satisfying efficiency of 7.3% under one sun (AM1.5 G). The co-sensitization strategy was further applied on this blue organic dye together with a red D- π -A dye (D35). The successful co-sensitization outperformed a panchromatic light absorption and improved the photocurrent density; this in addition to the open-circuit potential result in an efficiency of 8.7%. The extended absorption of the sensitization and the slower recombination reaction between the blue dye and TiO₂ surface inhibited by the additional red sensitizer could be the two main reasons for the higher performance. In conclusion, from the results, the highly efficient cobalt-based DSSCs could be achieved with the co-sensitization between red and blue D- π -A organic dyes with a proper design, which showed us the possibility of applying this strategy for future high-performance solar cells.

KEYWORDS: blue color, organic dyes, co-sensitization, cobalt electrolyte, dye-sensitized solar cells



1. INTRODUCTION

Dye-sensitized solar cells (DSSCs) have attracted great interest as a low-cost alternative to silicon solar cells, exhibiting moderate power conversion efficiencies. One crucial component of the DSSC devices is the monolayer of dye molecules, which is located at the interface of a mesoporous wide-bandgap semiconductor electrode and an electrolyte.^{1–5} This dye monolayer takes the function of light harvesting in the devices, and determines which part of the photons can be absorbed in the solar spectrum. So far, a large number of red dyes with strong absorption below 600 nm have been practiced in DSSCs successfully. However, this leaves a major part of the incident photons in the red and near-infrared (NIR) range unused. Therefore, the blue and green dyes as well as the NIR-absorbing dyes attracted particular interest. Additionally, the semitransparent solar cell devices with different colors are an extremely attractive technology for various applications

including building integration. Although great efforts have been accomplished to design the blue or green dyes, few efficient sensitizers, compared to red dyes, have been developed. Squaraine-based sensitizers can reach a green or blue color, which absorbs the NIR region. However, the solar cell devices with this type of dyes always lead to low efficiencies. This is primarily due to the dye aggregation, poor electron injection, and low stability.^{6–12} Zinc phthalocyanines and perylene dyes are also favorable for their blue color and NIR absorption. However, both kinds of dyes did not show good performance until now.^{13–19} Recently, Holcombe et al. reported a series of diketopyrrolopyrrole (DPP)-based high-performance blue sensitizers for DSSCs, among which is the

Received: August 17, 2016

Accepted: October 28, 2016

Published: October 28, 2016

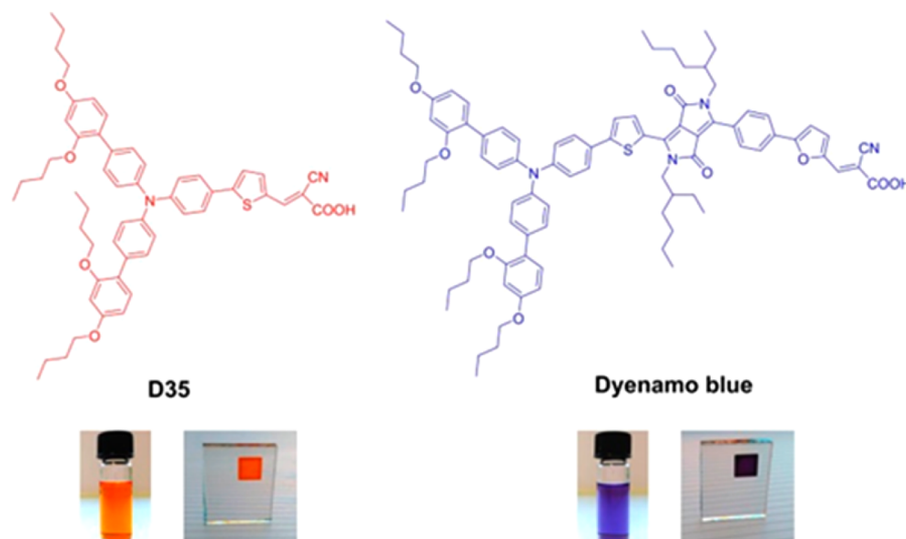


Figure 1. Chemical structures of the D35 and DB dyes and the dyes in solutions and on TiO₂ films.

device with DPP17 sensitizer that achieved over 10% PCE in combination with the [Co(bpy)₃]^{3+/2+} electrolyte at full sun (AM1.5 G).²⁰

Co-sensitization of two or more dyes with complementary absorption regions is an effective approach to extend the absorption spectrum of the solar cell and enhance the photovoltaic properties of organic dyes based on DSSCs.^{21–33} Some successful co-sensitized DSSCs have been reported. In 2011, a porphyrin dye co-sensitized with an organic dye with cobalt-based electrolyte has been studied by Grätzel's group; an efficiency of 12.3% has been achieved.³¹ Recently, a record DSSC efficiency above 14% has been obtained with collaborative sensitization of silyl-anchor and carboxy-anchor dyes.³² Most successful co-sensitization systems with cobalt electrolyte employed one metal-complex sensitizer (such as ruthenium or porphyrin dyes) or other dyes as main light absorber, and another dye took little function to slightly fill up the absorption gap.^{21,22,27–30,33} The co-sensitization of two pure organic dyes with very different absorption spectra has not been well-studied yet. Here, a pure organic co-sensitization system with very different color dyes, one red and one blue D- π -A organic dyes, has been studied combined with cobalt-based electrolyte.

Herein, a new blue organic D- π -A dye with broad absorption spectra, Dyenamo Blue (DB), has been designed and synthesized for the co-sensitization. The structure of the blue dye is designed on the basis of the red dye D35 (shown in Figure 1), which was previously found to be suitable with the cobalt-based redox couples.³⁴ The introduction of the DPP structure into the blue dye gives a promising performance of 7.3% with cobalt electrolyte. The DB dye was found to have successful combination with D35 for co-sensitization. An improved efficiency of 8.7% was obtained for devices containing co-sensitized pure organic D- π -A sensitizers and cobalt mediator.

2. EXPERIMENTAL SECTION

2.1. Synthesis of Dyenamo Blue (DB) Dye. Methods and materials follow. All reagents from commercial sources were used without further purification. ¹H and ¹³C NMR spectra were recorded on Bruker 500 and 400 MHz instruments by using the residual signal acetone-*d*₆ δ = 2.05, 29.84, and 206.26 ppm as an internal reference for

¹H and ¹³C, respectively. Mass spectroscopy data were obtained using a Dionex UltiMate 3000 system with an electrospray ionization method. The solvent mixture used for direct injection was DCM:MeOH (1:1).

2.2. Solar Cell Assembly. Solar cells are assembled by the procedure which has been described in a previous report of our work.³⁵ The TiO₂ photoelectrodes, platinum decorated counter electrodes, and cobalt electrolyte were prepared as described and used in the previous report. The difference is mainly preparation of dye solutions. At a temperature of 90 °C, the sintered TiO₂ electrodes were immersed in a dye bath for 18 h containing D35 (0.2 mM), DB (0.025 mM), or a mixture of D35 and DB by varied molar ratio in *tert*-butanol:acetonitrile (1:1 v/v), respectively.

2.3. Solar Cells Characterization. All of the solar cell characterization methods performed in the present study are also described in our previous report.³⁵ Basically, current–voltage (*IV*) characteristics were measured under a solar simulator (Newport, model 91160), which is controlled by a source measurement unit (Keithley 2400). A certified silicon solar cell (Fraunhofer ISE) was calibrated to an intensity of 100 mW cm⁻² of a solar simulator (AM1.5 G spectral). Also, a black metal mask with area of 0.25 cm² was used for all of the solar cells' characterization.

Incident photon-to-current conversion efficiency (IPCE) spectra measurements were conducted by a setup combining a xenon light source (Spectral Products ASB-XE-175), a monochromator (Spectral Products CM110), and a Keithley multimeter (model 2700). The silicon solar cell mentioned above is also calibrated with the IPCE spectra.

Electron lifetime and charge extraction measurements as a function of varied voltage at different bias light intensities was investigated in a “toolbox setup” as described previously.^{36,37} Mainly, the setup comprises a white LED (Luxeon Star 1W) as the light source, a 16-bit resolution digital acquisition board (National Instruments) in combination with a current amplifier (Stanford Research Systems SRS70), and a custom-made system using electromagnetic switches.

2.4. Electrochemical Measurements. A three-electrode cell was used for the cyclic voltammetry experiments. The supporting electrolyte used for electrochemical measurements was 0.10 M LiN(CF₃SO₂)₂ in acetonitrile. The counter electrode is a stainless steel plate with an area of 3 cm². The reference electrode was Ag/AgCl; a salt bridge electrolyte was interposed between working and reference electrodes, containing 0.10 M LiN(CF₃SO₂)₂ in acetonitrile. The working electrode is a dye-sensitized SnO₂ film with the thickness of 4 μ m and area of 0.25 cm². The reference electrode was calibrated by recording the cyclic voltammogram of ferrocene in the same

electrolyte; the potential values are on the basis of the estimated value of the ferrocene redox potential in acetonitrile 0.624 V versus SHE.³⁸

2.5. Steady-State Absorption. TiO₂ films for steady-state absorption measurements were prepared by 30 nm of TiO₂ paste (Dyesol) with thickness of 2 μm on microscope glass slides and sintering at 500 °C. Absorption spectra were recorded on a HR-2000 Ocean Optics spectrophotometer with baseline correction.

For dye-loading calculation, transparent TiO₂ films with the same properties as those used in the DSSCs were sensitized with the four different dye mixtures for 18 h. Each film was then carefully washed and dried with air flow before being desorbed in 1 mL of 0.05 M TBAOH (in 1:1 ACN:tBuOH). The absorbance was then measured with a HR-2000 Ocean Optics spectrophotometer and the amount of each dye calculated from standard curves obtained for this specific condition (see SI).

2.6. Nanosecond Transient Absorption Spectroscopy (TAS).

Samples are prepared according to previous procedures.³⁹ Basically, transparent TiO₂ films were sensitized to act as the photocathode, while nonplatinized FTOs were used as the counter electrode. Two types of sensitized films were used in the present study: one is sensitized by DB dye, and another is sensitized by co-sensitized dyes (DB:D35 = 3:4), all of which assembled with either inert (0.1 M LiClO₄, 0.2 M TBP) or redox electrolytes (the same electrolyte as described in the solar cell fabrication details).

TAS measurements were performed with a laser flash photolysis spectrometer (Edinburgh Instrument LP920), using a continuous wave xenon light as a probe light. Laser pulses were generated and tuned to 670 nm, using a frequency tripled Nd:YAG laser (Continuum Surelight II, 10 Hz repetition rate, 10 ns pulse width) in combination with an OPO (Continuum Surelight). The pulse intensity was attenuated on the order of 0.1 mJ cm⁻² per pulse using neutral density filters; therefore, only few electrons were injected into every TiO₂ nanoparticle upon the pulse excitation of dye molecules. Kinetic traces were monitored at 750 nm, where oxidized DB dye shows strong absorption indicated by PIA measurement (see Figure S7 in the Supporting Information). The signal was averaged from 100 to 500 measurements to reduce the noise.

2.7. Photoinduced Absorption Spectroscopy (PIA).

PIA measurements were performed using the same setup as described in previous literature.⁴⁰ Briefly, the absorption changes of the sample were probed by a 20 W tungsten-halogen lamp. A square-wave modulated blue LED (460 nm) with a frequency of 9.33 Hz was utilized to periodically excite the samples. The transmitted light through the samples was then focused on the monochromator, and detected by a Si photodiode during the time. The signal was further amplified with a current preamplifier and connected to a lock-in amplifier (Stanford Research Systems model 830 and 570, respectively). The excitation LED intensity was set to ~8 mW/cm². This measurement is conducted to confirm the absorption spectrum of oxidized D35, DB, and co-sensitized TiO₂ electrode (seen in Figure S7).

3. RESULTS AND DISCUSSION

3.1. Design and Synthesis of DB Dye.

Prior to the synthesis, quantum chemical calculations (Figure 2) were performed to predict the properties of the sensitizers and provide guidance in the choice of target sensitizers. All of the alkyl chains were truncated to balance the efficiency of calculation and its accuracy. Gas-phase geometries were optimized by density functional theory at the B3LYP/6-311G(d, p) level using Gaussian09 software. Analytic frequency calculations were executed to ensure that the lowest vibrational frequencies were real, and an energy minimum was achieved. Time-dependent density functional theory (TD-DFT) was then utilized at the same theoretical level to find vertical transitions. The most likely transitions were found to be HOMO to LUMO ($f = 0.71$), with HOMO mainly located on the triphenylamine moiety and LUMO largely located at the

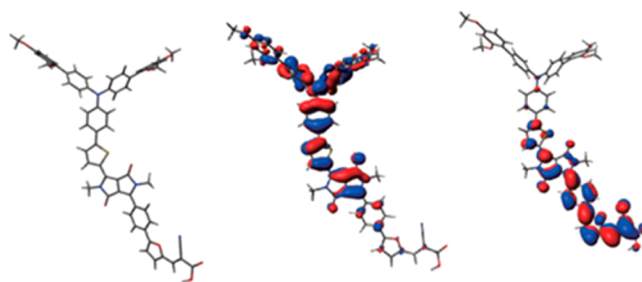


Figure 2. Optimized structure (left) of DB dye (with truncated alkyl chains), and the frontier molecular orbitals of the HOMO (middle) and LUMO (right) calculated with DFT at the B3LYP/6-311G(d,p) level in the gas phase.

carboxylic anchoring group, indicating a pronounced intramolecular charge separation after excitation.

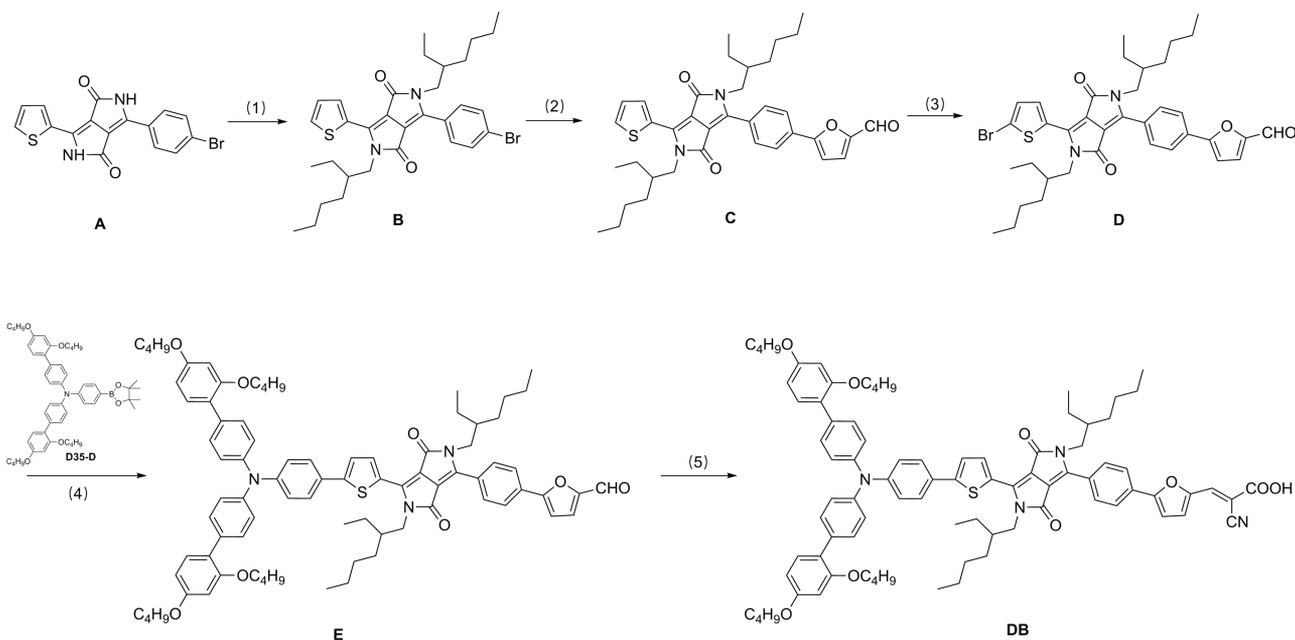
The Dyanamo Blue dye was synthesized in the general synthetic route as shown in Scheme 1. The linker part which employed the DPP-core was synthesized first according to the method reported by Jun-Ho Yum, Thomas W. Holcombe, etc.²⁰ Then, the donor part of D35 (D35-D) was combined with the linker part (D) by Suzuki coupling reaction. Further, the final dye was synthesized by condensation of aldehydes with cyanoacetic acid by the Knoevenagel reaction. The characterizations of the new compounds and also the details of synthesis are described in the Supporting Information section.

3.2. Photophysical and Electrochemical Properties.

The absorption spectra of the DB dye dissolved in *tert*-butanol:acetonitrile (1:1 v/v) solution with different concentrations are illustrated in Figure S1, while the spectrum of the sensitized TiO₂ film is shown in Figure 3a (blue line). The maximum absorption peaks of the DB dye are located at 571 nm in solution and 564 nm on TiO₂ film, respectively. The absorption of the dye ranges from the UV to about 700 nm, both in solution and on the film, with a pronounced dip around 450 nm. The data showed the wide absorption of the DB dye and the capability of absorbing the photons in the long wavelength which is almost up to the NIR region. The photophysical and electrochemical details of the novel DB dye can be found in Table 1.

The absorbance spectra obtained from the D35-sensitized (red line) and dye-mixture-sensitized (DB 0.075 mM, D35 0.10 mM, black line) TiO₂ films were presented in Figure 3a. With the co-sensitization strategy, the whole visible range was successfully covered. The improvement of the absorption spectra promoted the utilization of light, which further suggested a better DSSC performance.

The formal reduction and oxidation potentials of D35, DB, and co-sensitized (DB:D35 = 3:4) dyes adsorbed on nanocrystalline Sb-doped SnO₂ electrodes were measured and shown in Figure 3b and Table 1. The novel DB dye has a slightly less positive HOMO level than that of D35, which is 1.08 V versus SHE compared to 1.12 V versus SHE for D35. The lower HOMO of DB could be ascribed to the larger conjugated system of DB, whereas the slightly lower driving force (the potential difference between the dye and cobalt mediator) is still enough for fast regeneration kinetics between DB dye and cobalt mediator in DSSCs.⁴¹ Moreover, the co-sensitized D35 and DB dyes in a ratio of 3:4 show a HOMO level similar to that of the DB dye, which also satisfies the requirement of electron transfer between the oxidized dye and redox mediator.

Scheme 1. Synthetic Route of DB Dye^a

^a(1) 3-(Bromomethyl)heptane, K_2CO_3 in DMF, 90 °C, overnight, 73%; (2) (5-formylfuran-2-yl)boronic acid, toluene:MeOH, $Pd(dppt)Cl_2$, 70 °C, 1 h, 80%; (3) NBS, $CHCl_3$, RT, 4 h, 53%; (4) 2',4'-dibutoxy-N-(2',4'-dibutoxy-[1,1'-biphenyl]-4-yl)-N-(4-(4,4,5,5-tetramethyl-1,3,2-dioxaborolan-2-yl)phenyl)-[1,1'-biphenyl]-4-amine (D35-D), K_3PO_4 , toluene, Pd_2dba_3 , X-Phos (2-dicyclohexylphosphino-2',4',6'-triisopropylbiphenyl), XPhos Pd G3 ((2-dicyclohexylphosphino-2',4',6'-triisopropyl-1,1'-biphenyl)[2-(2'-amino-1,1'-biphenyl)]palladium(II) methanesulfonate), 80 °C, 6 h, 53%; (5) cyanoacetic acid, MeCN:CHCl₃, piperidine, 80 °C, 2 h, 44%.

3.3. DSSCs. The photovoltaic performances of DSSCs based on the individual dyes as well as the best co-sensitization composition (DB:D35 = 3:4 (0.075 mM:0.1 mM in dye bath)) are presented in Table 2 and Figure 4a. A relatively thin, transparent TiO_2 film (4 μm) with scattering TiO_2 layer (4 μm) was used in this work to decrease mass transport problems that happened in cobalt complex mediator-based DSSCs. According to the data, the DSSC devices with DB dye generate 40% more photocurrent than that of the D35-sensitized devices for their broader absorption. However, the open-circuit potential is lower for the devices with DB dye, which suggested the higher recombination between the electrons in TiO_2 and Co(III) species in the electrolyte, and/or to oxidized dyes, indicating the less efficient blocking effect of DB dye. The overall power conversion efficiency (PCE) of the device with DB is higher (7.3%) than that of D35 (5.5%). Further, the improved performance of solar cells, 8.7%, was reached by co-sensitization with the molar ratio of 3:4 (DB:D35).

Different molar ratios and concentrations of DB:D35 in the dye bath have been performed, before the best composition of the co-sensitized system for photovoltaic performance was achieved. The solar cell parameters from different combinations are shown in Table S1. Initially, the concentration of D35 varied from 0.025 to 0.2 mM with a constant concentration of DB as 0.025 mM. As a result of the increasing concentration of D35 in the dye bath, the V_{OC} of the devices increased markedly (rows 1–3 in Table S1). The reason for this might be that, with D35 as co-sensitizer, the blocking effect was improved; therefore, the recombination between the electrons in TiO_2 and the electrolyte was reduced. Then, in contrast, the concentration of D35 was fixed (0.1 mM), and the concentration of DB was increased from 0.025 to 0.1 mM (rows 2 and 4–6 in Table S1). This time, there are only minor

changes in the V_{OC} , while the photocurrents are very different; the optimum photocurrent was found with 0.075 mM DB. This could be explained by the broader absorption spectrum of DB. With higher concentration of DB dye in the dye bath, an improved IPCE can be observed in the further red part of the IPCE spectrum (the discussion could be found in below, see Figure S6).

Figure 4b illustrates the incident photon-to-current efficiency (IPCE) spectra of the DSSCs, which are sensitized by the individual dyes D35, DB, and the co-sensitized dyes at the best condition (DB:D35 = 3:4). The IPCEs of the DSSCs sensitized by D35 and DB are located in the ranges 400–650 and 500–750 nm, respectively. The co-sensitized IPCE curve, showing the complementary behavior of the D35 and DB dyes, gives an enhanced and broader IPCE response compared to the single dye, resulting in an improved J_{SC} and the photovoltaic efficiency. The IPCE values obtained in Figure 4b are consistent with the photovoltaic performance presented in Figure 4a, showing the advantage of co-sensitized strategy.

Photovoltage response time measurements were performed in dependence of the light intensity in order to determine electron lifetime. As shown in Figure 5a, at a given potential, the electron lifetime of DB-sensitized solar cells is much shorter (by about 2 orders of magnitude) than that of D35. This corresponds to a higher recombination rate of electrons in the TiO_2 with Co(III) species in the electrolyte in DB-based devices.⁴² No significant shift of the conduction band edge was found (by charge extraction measurement seen in Figure 5b). The electron-blocking properties of DB are thus poor in comparison to those of D35. This is likely related to the relatively lower dye coverage of DB in comparison to D35 (details can be found in the latter part of this article, see Table

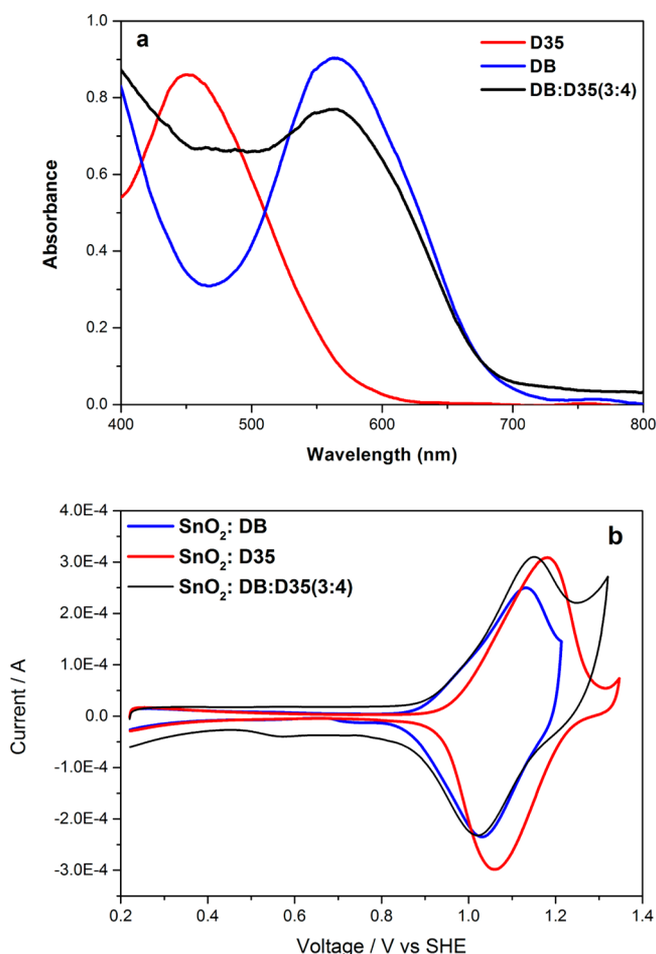


Figure 3. (a) UV-vis spectra of D35, DB, and co-sensitized (DB:D35 = 3:4) dyes on TiO₂ mesoporous film (2 μm). (b) Cyclic voltammetry of D35, DB, and co-sensitized (DB:D35 = 3:4) dyes adsorbed on nanocrystalline Sb-doped SnO₂ electrodes.

3). This would leave uncovered TiO₂ surface sites, which can promote electron recombination.

By the co-sensitization strategy, significantly longer electron lifetimes were found in comparison with that of the individual DB sensitization. This was caused by the better TiO₂ blocking with additional D35, the dye which has been demonstrated to have excellent blocking properties with cobalt electrolyte in earlier work.³⁴ The new co-sensitization between DB and D35 leads to an improved blocking effect of the dye monolayer, which decreases the electron recombination rate between TiO₂/dye and the electrolyte. However, there is still much room for improvement, and better combinations of dyes are expected to be found in future work.

Table 1. Photophysical and Electrochemical Properties of the DB Dye

absorption			emission ^b			
λ_{\max}^a (nm)	ϵ at λ_{\max} (M ⁻¹ cm ⁻¹)	λ_{\max} on TiO ₂ (nm)	λ_{\max} (nm)	E_{0-0}^c (V)	E_{ox}^d (V vs NHE)	$E_{\text{ox}} - E_{0-0}$ (V vs NHE)
571	33640	564	725	1.95	1.08	-0.87

^aAbsorption spectra were measured in the solution of *tert*-butanol:acetonitrile = 1:1 with different concentrations at 25 °C. ^bEmission spectra were measured in *tert*-butanol:acetonitrile = 1:1 solutions (1 × 10⁻⁶ M) at 25 °C. ^cThe zeroth-zeroth transition (E_{0-0}) value was estimated from the intersection of the absorption and emission spectra (Figure S2). ^dCyclic voltammetry of the oxidation behavior of the dyes was measured on SnO₂ film containing 0.1 M LiN(CF₃SO₂)₂ in acetonitrile as supporting electrolyte (working electrode, dye-sensitized SnO₂ film; reference electrode, Ag/Ag⁺ calibrated with ferrocene/ferrocenium (Fc/Fc⁺) as an internal reference; counter electrode, a stainless steel plate).

Table 2. Photovoltaic Performance of DSSCs Based on the D35, DB, and Co-Sensitized Dyes (DB:D35 = 3:4) under 1000 W m⁻² AM1.5 G Illumination

dye	V_{OC} (mV)	J_{SC} (mA/cm ²)	FF (%)	η (%)
D35	853	9.14	70.3	5.5
DB	757	13.21	72.7	7.3
DB/D35 = 3:4	797	15.60	70.1	8.7

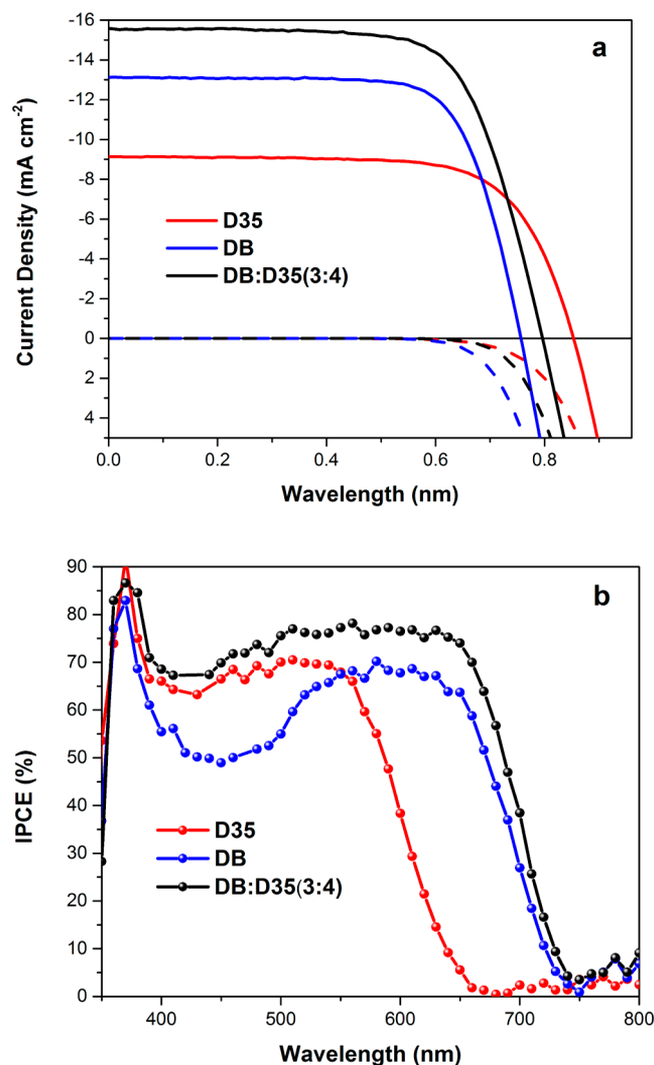


Figure 4. (a) Current density versus applied potential curves under 1000 W m⁻² AM1.5 G illumination (dot lines) and in darkness (dashed lines). (b) Spectra of incident photon-to-current conversion efficiency (IPCE) for DSSCs based on D35, DB, and co-sensitized dyes (DB:D35 = 3:4).

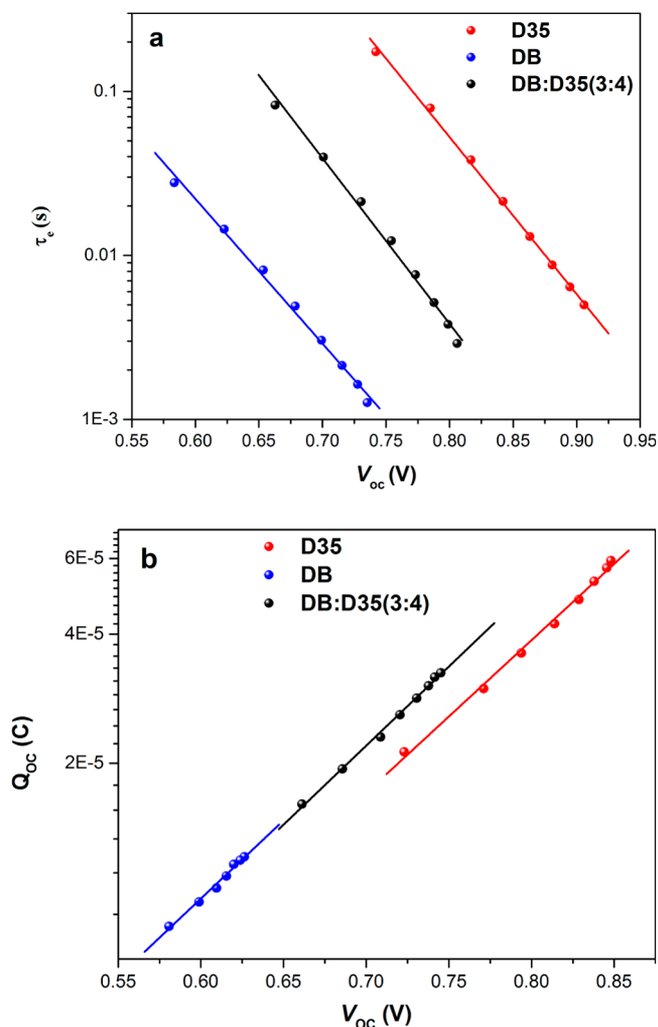


Figure 5. (a) Electron lifetime and (b) accumulation charge as a function of the V_{OC} under open-circuit conditions for DSSCs sensitized with D35, DB, and co-sensitized dyes employing cobalt electrolyte.

Table 3. Dye-Loading Measurements of Single and Co-Sensitized TiO_2 Films^a

dye concentration in solution (DB:D35)	nD35 (nmol)	nDB (nmol)	total dye (nmol)	dye ratio on films (DB:D35)
0 mM:0.2 mM (0:1)	20.2 ± 0.9	0	20.2 ± 0.9	
0.025 mM:0.2 mM (1:8)	16.6 ± 0.7	2.8 ± 0.1	19.4 ± 0.8	1:6
0.025 mM:0.1 mM (1:4)	15.5 ± 0.6	4.5 ± 0.2	20.0 ± 0.8	2:7
0.05 mM:0.1 mM (1:2)	13.4 ± 0.5	8.5 ± 0.3	21.9 ± 0.8	2:3
0.075 mM:0.1 mM (3:4)	11.9 ± 0.6	11.1 ± 0.4	23.0 ± 1.0	1:1
0.025 mM:0 (1:0)	0	8.2 ± 0.3	8.2 ± 0.3	

^aElectrode area 0.25 cm²; 4 μm TiO_2 film.

3.4. Dye-Loading Properties. By different co-sensitization conditions, the absorption spectrum of the film can be changed, as well as the dye stacking patterns. Some characterizations have been conducted to investigate how the co-sensitization

system works on the basis of the varied molar ratio of DB and D35. Figure S5 (top) shows the UV–vis absorption spectra of TiO_2 films (2 μm) with different co-sensitization conditions. It is obvious to see that the maxima of the absorption peaks of the individual dyes are related to the dye bath ratios; especially, the DB dye peak is becoming larger with an increased concentration of DB, accompanied with the declining absorption peak of D35. In order to gain a deeper insight about the actual dye load on the sensitized films, the sensitized dye was desorbed in 0.05 M TBAOH (1:1 *tert*-butanol:acetonitrile) followed by absorbance measurements of the solution (Figure S5 (bottom)). The results of the dye desorption measurements are summarized in Table 3, where the amount of each dye loaded on the film and the film ratio (DB:D35) are presented for each co-sensitization composition. Here it becomes even clearer that the DB dye has a larger affinity to the TiO_2 electrode than D35, since the relative amount of D35 on the film always becomes smaller than in the dye bath. At the molar ratio of 3:4 (DB:D35) in the dye bath, in which the best performance was achieved, the two dyes has almost equal amounts of molecules adsorbed on the TiO_2 film. The IPCE spectra of DSSCs which are co-sensitized give consistent hints of the relative amount of D35 and DB on the surface (Figure S6).

3.5. Kinetics. The transient absorption kinetics of DB and co-sensitized DB:D35 on TiO_2 electrodes were also investigated. The experiments were either conducted with inert electrolyte or in contact with the redox-couple-based electrolyte, on a nano- to millisecond time scale (see Figure 6 and Table 4). An excitation wavelength of 670 nm was used to exclusively excite the DB molecules. Under the conditions used, the transfer of electrons from the mesoporous TiO_2 to oxidized DB molecules occurred with a half time ($t_{1/2}$) of 5.75 μs. In comparison, this process occurred significantly slower with half time of 8.27 μs for co-sensitized TiO_2 electrodes. Here, slower electron-oxidized dye recombination kinetics is preferred. Under operational conditions, the concentration of electrons in the mesoporous TiO_2 is high. Therefore, the electron-oxidized dye recombination will result in significant energy losses, which is negative for the solar cell performance. The device with the co-sensitization system showed sluggish electron-oxidized dye recombination kinetics here. This is favorable for solar cell performance; moreover, this also confirms the exceptional blocking effect by additional D35 with DB dye. The better blocking effect may correlate to the binding mode of both DB and D35 dyes on the TiO_2 surface, which probably would be because one dye affects the orientation of another dye on the TiO_2 surface. The different orientation is likely to decrease the electron transfer reaction rate between electrons in TiO_2 and the location of the “hole” in the oxidized dye, in the triphenyl amine group. In combination with the cobalt-based electrolyte, similar regeneration kinetics were found for the DB and co-sensitized DSSCs. This indicates that the additional D35 did not slow down the electron transfer between oxidized sensitizer and redox species inside the electrolyte.

4. CONCLUSIONS

A novel blue D-π-A organic sensitizer (DB) was designed and synthesized for DSSCs with cobalt-based redox. With this blue dye, an energy conversion efficiency of 7.3% was yielded, under one sun AM1.5 G. The co-sensitization strategy was applied with an additional D35 dye. The device with co-sensitized dyes

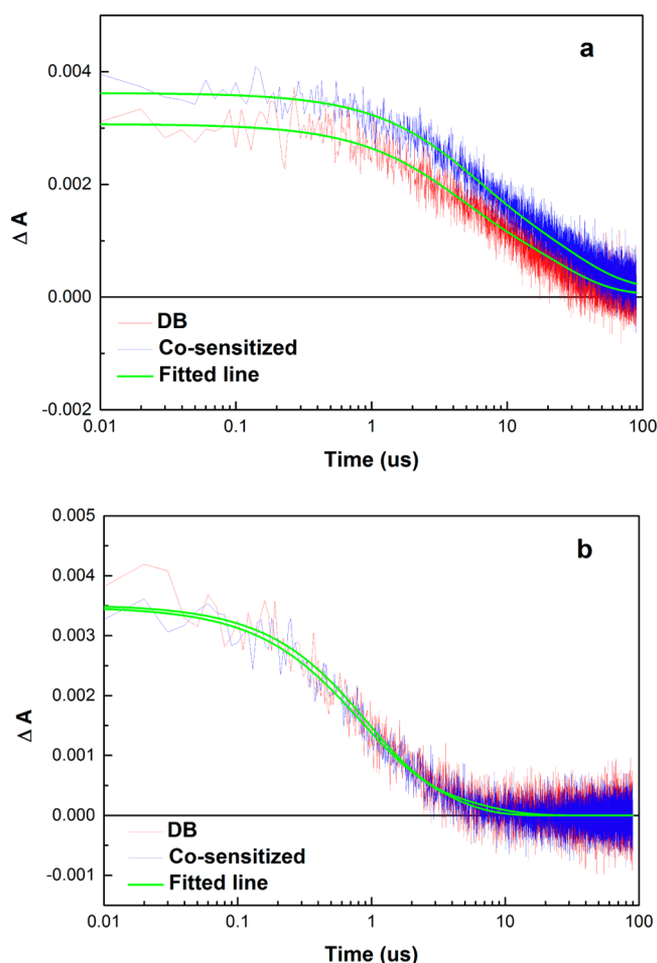


Figure 6. Transient absorption kinetics of TiO₂ sensitized with DB (blue line) and co-sensitized (DB:D35 = 3:4) (red line) under the condition of inert electrolyte (a) and cobalt bipyridine electrolyte (b).

Table 4. Measured Halftimes and Calculated Regeneration Efficiencies for DB-Sensitized and Co-Sensitized (DB:D35 = 3:4) DSSCs Employing Cobalt Bipyridine Redox Couple

sample	$t_{1/2 \text{ redox}}$ (μs)	$t_{1/2 \text{ inert}}$ (μs)	η_{reg} (%)
DB	0.7 ± 0.1	5.8 ± 0.5	89 ± 1
DB:D35 = 3:4	0.7 ± 0.1	8.3 ± 0.8	91 ± 1

achieved a PCE of 8.7% with a greater current density from the complementary absorption. This performance is superior to either of the individual devices sensitized by DB (7.3%) or D35 (5.5%) under the same fabrication condition. The additional D35 was found to decrease the recombination at the interface between TiO₂/dye and electrolyte, as well as the recombination between the electrons in TiO₂ and oxidized dyes. This work shows the possibility of using two pure organic dyes with very different absorbing spectra together with cobalt complex-based redox. This co-sensitization greatly uses the light in solar spectra, and at the same time increases the blocking effect on the TiO₂ surface. The results indicate that, by proper design of dyes, the performance of DSSCs can be further improved by the co-sensitization strategy.

■ ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsami.6b09671.

Detailed synthesis, characterization of synthesized mediators and sensitizer (DB dye), UV-vis absorption, solar cell performances and IPCE spectra, and PIA and TAS figures (PDF)

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Notes

The authors declare no competing financial interest.

■ ACKNOWLEDGMENTS

We gratefully acknowledge the Swedish Energy Agency, the Swedish Research Council (VR), the STandUP for Energy program, the Knut and Alice Wallenberg Foundation, Stiftelsen Olle Engkvist Byggmätare, and the Swiss National Science Foundation Contract 200021_157135/1 "Fundamental studies of mesoscopic devices for solar energy conversion" for financial support. Dr. Martin Karlsson and Dr. Palas Baran Pati are acknowledged for their help with characterization of the novel blue dye.

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