## Chemical Science



#### **EDGE ARTICLE**

View Article Online
View Journal



Cite this: DOI: 10.1039/d5sc03537f

All publication charges for this article have been paid for by the Royal Society of Chemistry

# Molecular engineering of porphyrin dyes and copper complexes for enhanced dye regeneration toward high-performance dye-sensitized solar cells using copper(1/11) redox shuttles†

Porphyrin dyes have garnered significant attention as promising photosensitizers for dye-sensitized solar cells (DSSCs) due to their exceptional light-harvesting capabilities and remarkable power conversion efficiencies (PCEs) when paired with cobalt(II/III) complex-based redox shuttles. Meanwhile, copper(I/II) complexes have emerged as new generation redox shuttles, achieving impressive open-circuit voltages  $(V_{OC})$  exceeding 1.0 V. However, porphyrin-based DSSCs using copper(I/II) redox shuttles have struggled with low-to-moderate PCEs, primarily due to insufficient driving forces for the dye regeneration process. In this study, we introduce FL1, a novel porphyrin dye featuring a fluorene moiety with reduced electrondonating properties, designed to ensure a sufficient driving force for dve regeneration using copper(I/II) complexes. Under optimized conditions, DSSC incorporating FL1 with a copper(I/II) complex utilizing 4,4'-dimethoxy-6,6'-dimethyl-2,2'-bipyridine [Cu(2MeOby)<sub>2</sub>][TFSI]/[Cu(2MeOby)<sub>2</sub>][TFSI]<sub>2</sub> a notable PCE of 8.30% with a  $V_{\rm OC}$  of 0.890 V. Furthermore, our investigation into counterion effects revealed that DSSCs employing  $[Cu(2MeOby)_2][PF_6]/[Cu(2MeOby)_2][PF_6]_2$  as a redox shuttle delivered the highest PCE of 9.06% with a  $V_{\rm OC}$  of 0.900 V, attributed to its superior diffusion coefficient. Finally, cosensitized DSSCs featuring FL1 and XY1B achieved an outstanding PCE of 10.9%, while retaining a high  $V_{\rm OC}$  of 0.945 V, setting a new benchmark efficiency for porphyrin-based DSSCs utilizing copper(I/II) redox shuttles. This breakthrough highlights the immense potential of further refining porphyrin dyes and copper(I/II) redox shuttles through energy-level engineering to optimize the driving force for dye regeneration and propel advancements in DSSC technology.

Received 16th May 2025 Accepted 8th July 2025

DOI: 10.1039/d5sc03537f

rsc.li/chemical-science

#### Introduction

A substantial portion of global energy consumption still relies on fossil fuels, leading to substantial CO<sub>2</sub> emissions and pressing environmental concerns. This growing challenge has heightened the urgency for sustainable energy solutions, with solar energy emerging as a particularly compelling alternative, an abundant and virtually limitless resource derived from the sun. Among various solar technologies, organic solar cells

present a promising avenue due to their cost-effective production, design flexibility, and lightweight properties.<sup>1,2</sup> Notably, dye-sensitized solar cells (DSSCs) have captured considerable attention due to their impressive high power conversion efficiencies (PCEs), vibrant multicolor fabrication possibilities, and outstanding performance under low-light conditions.<sup>3-8</sup>

Given the fundamental working principle of DSSCs, the sensitizer is a key player in facilitating light-harvesting and electron transfer (ET) processes at the dye/TiO<sub>2</sub>/electrolyte interface. To push the boundaries of DSSC efficiency, the development of sensitizers with strong light-harvesting capabilities across the visible and near-infrared spectrum is highly desirable.<sup>9-13</sup> Moreover, the strategic incorporation of bulky substituents is crucial for suppressing charge recombination (CR) processes, thereby optimizing device performance. Porphyrin dyes have emerged as highly promising photosensitizers for DSSCs, thanks to their exceptional light-harvesting properties, adaptable electronic structures achieved through molecular engineering, and outstanding stability compared to conventional dyes.<sup>14-32</sup> Notably, single push-pull-type porphyrin

<sup>&</sup>lt;sup>a</sup>Department of Molecular Engineering, Graduate School of Engineering, Kyoto University, Kyoto, 615-8510, Japan. E-mail: t-higa@scl.kyoto-u.ac.jp; imahori@scl.kyoto-u.ac.jp

<sup>&</sup>lt;sup>b</sup>Institute for Integrated Cell-Material Sciences (WPI-iCeMS), Kyoto University, Kyoto, 606-8501, Japan

<sup>\*\*</sup>Tinstitute for Liberal Arts and Sciences (ILAS), Kyoto University, Kyoto, 606-8316, Japan † Electronic supplementary information (ESI) available: Experimental section, synthetic details, optical and electrochemical properties of porphyrin dyes and copper complexes, photovoltaic properties, HR-MS spectra, and NMR spectra. CCDC 2451025-2451027. For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.org/10.1039/d5sc03537f

dyes designed with effective steric hindrance from bulky substituents have demonstrated impressive PCEs, reaching up to 13% when paired with tris(bipyridyl)Co $^{\Pi/\Pi}$  ([Co(bpy)<sub>3</sub>] $^{2+/3+}$ ) as a redox shuttle. This progress underscores the immense potential of porphyrin-based sensitizers in advancing high-performance DSSCs.

Meanwhile, the redox shuttle serves as a crucial component in DSSCs, as the open-circuit voltage  $(V_{OC})$  is primarily determined by the potential difference between the quasi-Fermi level of  $TiO_2$  and the redox potential ( $E_{redox}$ ) of the redox shuttle.<sup>33-35</sup> Over the past few decades, iodide/triiodide (I-/I3-) and [Co(bpy)<sub>3</sub>]<sup>2+/3+</sup> have been the most widely utilized redox shuttles. However, DSSCs incorporating these traditional shuttles have faced a  $V_{\rm OC}$  limitation, with values restricted to below 1.0 V. To address this constraint, copper(I/II) complexes have emerged as new-generation redox shuttles, offering a more positive  $E_{\rm redox}$  and significantly enhancing performance. DSSCs employing suitable complementary dyes and [Cu(tmby)<sub>2</sub>]<sup>+/2+</sup> (tmby = 4,4'-6,6'-dimethyl-2,2'-bipyridine) have achieved PCEs of up to 15%, coupled with a higher  $V_{\rm OC}$  exceeding 1.0 V.<sup>8,36-44</sup> This progress emphasizes the transformative potential of copper-based redox shuttles in propelling DSSC efficiency to new heights.

In this context, DSSCs incorporating highly efficient porphyrin dyes alongside copper redox shuttles present a promising pathway for further enhancing PCE. However, despite their potential, the PCEs of porphyrin-based DSSCs using copper(I/II) complex-based redox shuttles remain below 7%, even in the case of LG4 (Fig. 1),45-47 falling significantly behind other organic dye-based DSSCs in performance. These low-to-moderate PCEs are primarily attributed to the challenge of achieving a sufficient driving force for effective dye regeneration ( $-\Delta G_{\text{reg}} \ge 0.2 \text{ eV}$ ). Conventional push-pull-type porphyrin dyes typically exhibit relatively low oxidation potentials  $(E_{ox})$  of 0.7–0.8 V vs. NHE, limiting their compatibility with copper(I/II) complexes. In our previous studies, we demonstrated that increasing the  $-\Delta G_{\text{reg}}$  value through energy-level engineering is an effective strategy for boosting PCEs in porphyrin-based DSSCs. 46 Thus, porphyrin dyes with elevated  $E_{ox}$  values (≥0.9 V vs. NHE) would be well-suited for DSSCs utilizing copper(I/II) complexes. Additionally, ligand modification of copper(I/II) complexes emerges as another promising approach for optimizing  $-\Delta G_{\text{reg}}$  values (Fig. 2).<sup>47,48</sup> Incorporating electrondonating substituents into the bipyridine framework shifts the  $E_{\rm redox}$  value negatively, a favorable adjustment for improving porphyrin-based DSSC performance. These advancements collectively pave the way for further refining porphyrin-copper systems, driving DSSCs toward higher efficiencies.

Herein, we present **FL1**, a novel porphyrin dye incorporating a 9,9-dioctylfluorene moiety, along with its photovoltaic

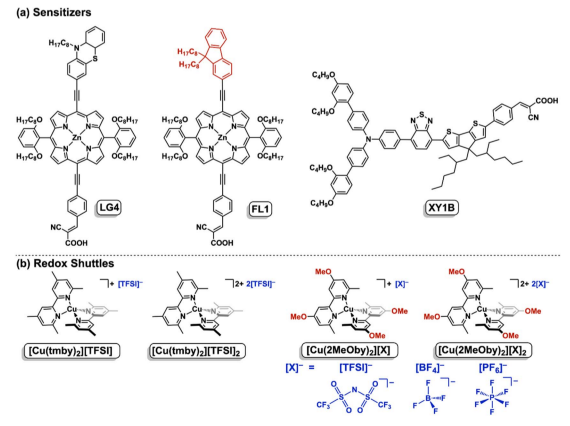


Fig. 1 Molecular structures of (a) porphyrin dyes LG4 and FL1 and co-sensitizer XY1B, and (b) copper complexes [Cu(tmby)<sub>2</sub>][TFSI], [Cu(tmby)<sub>2</sub>] [TFSI]<sub>2</sub>, [Cu(2MeOby)<sub>2</sub>][X], and [Cu(2MeOby)<sub>2</sub>][X]<sub>2</sub>. The counterions ([X]<sup>-</sup>) are trifluoromethanesulfonimide ([TFSI]<sup>-</sup>), hexafluorophosphate ([PF<sub>6</sub>]<sup>-</sup>), and tetrafluoroborate ([BF<sub>4</sub>]<sup>-</sup>).

Edge Article Chemical Science

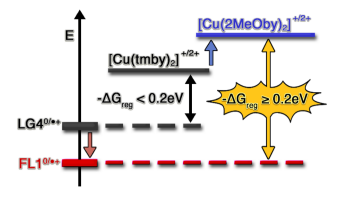


Fig. 2 Energy-level engineering of both dyes and redox shuttles for achieving an efficient driving force for dye regeneration  $(-\Delta G_{\text{reg}})$  toward enhanced cell performance.

performance in DSSCs employing copper(1/II) complex-based redox shuttles (Fig. 1). The reduced electron-donating nature of the 9,9-dioctylfluorene moiety, compared to the phenothiazine moiety in previously reported LG4,45 was expected to enhance the  $-\Delta G_{\text{reg}}$  value (Fig. 2), while the two bulky octyl chains at the 9-position of the fluorene skeleton were anticipated to provide a strong blocking effect against the redox shuttle. Furthermore, we aimed to further increase the  $-\Delta G_{\text{reg}}$ value by pairing FL1 with a copper(1/11) complex incorporating 4,4'-dimethoxy-6,6'-dimethyl-2,2'-bipyridine (2MeOby), [Cu(2MeOby)<sub>2</sub>]<sup>+/2+</sup>, featuring electron-donating methoxy groups (Fig. 1). Additionally, we also investigated the effect of counterions because the counterion effect of the ruthenium dyes (i.e. N719) has been reported.49 Actually, counterions play a critical role in the diffusion behavior of copper(I/II) complexes in organic solvents,50 significantly influencing the PCE of DSSCs, as high diffusion coefficients are essential for efficient charge transport within the electrolyte. 50,51 To optimize charge transport, we examined three counterions, trifluoromethanesulfonimide ([TFSI]<sup>-</sup>), tetrafluoroborate ([BF<sub>4</sub>]<sup>-</sup>), and hexafluorophosphate ([PF<sub>6</sub>]<sup>-</sup>). Through rational molecular engineering and a systematic exploration of the counterion effect, we successfully enhanced the PCE, with FL1-based DSSCs achieving an impressive PCE of 9.06%. Moreover, cosensitization with XY1B further pushed the PCE beyond 10% while retaining a high  $V_{\rm OC}$  of 0.945 V, marking the first instance of porphyrin-based DSSCs with copper(I/II) complex-based redox shuttles surpassing this milestone.

#### Results and discussion

#### Synthesis and characterization

The synthesis of the porphyrin dye **FL1** followed the procedure outlined in Scheme S1.† First, the Sonogashira coupling of 2-bromo-9,9'-dioctyl-9*H*-fluorene **1** (ref. 52) with triisopropylsilyl acetylene yielded ethynyl-substituted fluorene **2**. After deprotecting the silyl group with *n*-tetrabutylammonium fluoride (TBAF), subsequent Sonogashira coupling with *meso*-bromoporphyrin **3** (ref. 19) afforded bis-ethynyl substituted porphyrin **4**. Another round of triisopropylsilyl group deprotection

enabled a final Sonogashira coupling with 4-iodobenzaldehyde to synthesize porphyrin 5, which then underwent Knoevenagel condensation with cyanoacrylic acid to yield FL1.

Additionally, copper complexes incorporating [TFSI]- as a counterion were synthesized (Scheme S2†). The reaction of CuI with the ligand 2MeOby,53 followed by counterion exchange using LiTFSI, produced the copper(1) complex [Cu(2MeOby)2] [TFSI]. In contrast, the direct reaction of Cu(TFSI)2 with 2MeOby resulted in the copper(II) complex [Cu(2MeOby)2] [TFSI]2 without counterion exchange. Copper(I) complexes [Cu(2MeOby)2][X] (X  $= BF_4 \text{ or } PF_6$ ) were obtained by reacting  $[Cu(CH_3CN)_4][BF_4]$  or [Cu(CH<sub>3</sub>CN)<sub>4</sub>][PF<sub>6</sub>] with 2MeOby (Scheme S3†). Meanwhile, the copper(II) complexes  $[Cu(2MeOby)_2][X]_2$  (X = BF<sub>4</sub> or PF<sub>6</sub>) were prepared via a two-step process: first, CuSO<sub>4</sub>·5H<sub>2</sub>O was reacted with 2MeOby, followed by counterion exchange with the respective ammonium salts (NH<sub>4</sub>BF<sub>4</sub> or NH<sub>4</sub>PF<sub>6</sub>). Characterization of these compounds, including their copper complexes, was performed using <sup>1</sup>H and <sup>13</sup>C NMR, FT-IR spectroscopy, and high-resolution mass spectrometry (Fig. S1-S9†). Notably, the as-prepared copper(II) complexes contained varying proportions of their copper(1) counterparts due to unavoidable autoreduction under ambient conditions, as Cl ions were absent.54,55 Since Kitagawa et al. reported that the autoreduction of copper(II) complexes occurred in ethanol,<sup>54</sup> methanol in the reaction solvent may be a reductant for autoreduction. On the other hand, we obtained pure copper(II) complexes by the bulk electrolysis of the corresponding copper(1) complexes, following established literature protocols.56,57 The progress of electrochemical oxidation was monitored using the current, and the electrolysis was considered complete when the relative current approached zero, indicating that the copper(i) complexes had been entirely converted to their copper(ii) counterparts (Fig. S16 and S17†). Since the as-prepared copper(II) complexes contain no other impurities except for the corresponding copper(1) complexes, the actual compositions of the as-prepared copper(II) complexes were determined as  $[Cu(2MeOby)_2][X]_n$  (n = 1.6-1.9; X = TFSI, BF<sub>4</sub>, or PF<sub>6</sub>) by absorbance at 748 nm in acetonitrile (Fig. S19†), using the absorption coefficient of pure copper(II) complexes (vide infra). It is noteworthy that the actual compositions of the as-prepared copper(II) complexes in our laboratory were also determined as [Cu(tmby)<sub>2</sub>][TFSI]<sub>1.8</sub> in a similar manner.

Single crystals suitable for X-ray diffraction analysis were successfully grown *via* vapor diffusion of Et<sub>2</sub>O into acetonitrile solutions of [Cu(2MeOby)<sub>2</sub>][TFSI] or [Cu(2MeOby)<sub>2</sub>][BF<sub>4</sub>] and by vapor diffusion of *n*-butanol into acetonitrile solutions of [Cu(2MeOby)<sub>2</sub>][PF<sub>6</sub>]. The resulting crystal structures (Fig. S10†) revealed Cu–N bond distances of 2.01–2.07 Å, confirming that the counterions had negligible influence on the geometry of the copper(1) complexes. The crystal data of the copper(1) complexes are also summarized in Table S1.†

#### Optical and electrochemical properties of porphyrin dyes

The UV/vis/NIR absorption spectra of the porphyrin dyes in tetrahydrofuran (THF) are presented in Fig. 3, with their optical properties summarized in Table 1. The absorption spectrum of

**Chemical Science** working electrodes. The Eox values of FL1/TiO2 and LG4/TiO2 3.0

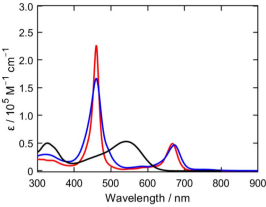


Fig. 3 UV/vis/NIR absorption spectra of FL1 (red), LG4 (blue), and XY1B18 (black) in THF.

FL1 exhibits light-harvesting capabilities comparable to those of LG4, with a slight 5 nm blue-shift in its lowest-energy Q-band. This shift is attributed to the fluorene moiety's weaker electrondonating ability compared to the phenothiazine moiety in LG4. The steady-state fluorescence spectrum of FL1 in THF also shows a 7 nm blue-shifted emission relative to LG4, aligning with the Q-band blue-shift observed in absorption spectra (Fig. S11†). The fluorescence lifetimes  $(\tau)$ , determined by timecorrelated single-photon counting (TCSPC), are 1.3 ns for FL1 and 1.2 ns for LG4 (Fig. S12†). The almost identical lifetimes suggest that the replacement of the phenothiazine moiety with the fluorene moiety does not cause unfavorable aggregation tendency. The optical HOMO-LUMO gaps  $(E_{0-0})$  of FL1 and LG4, calculated from the intersection of their normalized absorption and emission spectra, are 1.90 eV and 1.87 eV, respectively (Table 1). Although the fluorene moiety is a weaker electron donor than the phenothiazine moiety, it has a minimal impact on FL1's overall optical properties.

The electrochemical properties of the porphyrin dyes were evaluated using cyclic voltammetry (CV) and differential pulse voltammetry (DPV) in THF with 0.1 M tetrabutylammonium hexafluorophosphate (n-Bu<sub>4</sub>NPF<sub>6</sub>) as a supporting electrolyte. The first oxidation potential  $(E_{ox})$  of **FL1** was measured at +0.93 V vs. NHE (Fig. S13†), which is more positive than that of LG4 (+0.86 V vs. NHE), reinforcing the fluorene moiety's weaker electron-donating ability. To further assess the ET processes at the dye/TiO<sub>2</sub>/electrolyte interface, CV measurements were performed for dye-adsorbed TiO2 films (FL1/TiO2 and LG4/TiO2) as

were determined to be +1.07 V and +0.98 V vs. NHE, respectively (Fig. S14† and Table 1). Considering the redox potential  $(E_{redox})$ of the copper(I/II) complexes [Cu(tmby)<sub>2</sub>][TFSI]/[Cu(tmby)<sub>2</sub>] [TFSI]<sub>2</sub> (+0.88 V vs. NHE, Fig. S20a†), the driving forces for dye regeneration  $(-\Delta G_{reg})$  of the porphyrin radical cation  $(ZnP^{*+})$ were calculated to be 0.19 eV for FL1 and 0.10 eV for LG4. Since previous studies suggest that  $-\Delta G_{reg}$  should be comparable to or larger than 0.20 eV for efficient dye regeneration via copper(1/ II) redox shuttles,<sup>36</sup> the larger  $-\Delta G_{\text{reg}}$  value for **FL1** than for **LG4** is expected to enhance dye regeneration efficiency, contributing to improved PCE. Additionally, the driving forces for electron injection  $(-\Delta G_{\rm inj})$  from the porphyrin excited singlet state ( $^{1}$ ZnP\*) to the conduction band (CB) of TiO<sub>2</sub> (-0.5 V vs. NHE) were estimated using their excited-state oxidation potentials calculated using their oxidation potentials and optical HOMO-LUMO gaps. Both FL1 and LG4 possess sufficient  $-\Delta G_{\rm ini}$  values  $(\geq 0.30 \text{ eV})$  to facilitate efficient electron injection into the CB of TiO<sub>2</sub>.

To obtain insight into the ground state geometries and electronic structures of the porphyrin dyes, we performed density functional theory (DFT) calculations on model porphyrins at the B3LYP/6-31G(d) level (Fig. 4). Both FL1 and LG4 adopt a highly planar structure, facilitating an extended  $\pi$ system for effective charge delocalization. The two alkyl chains at the 9-position of the fluorene moiety in FL1 serve a dual purpose: providing a blocking effect and suppressing dye aggregation due to the enhanced steric hindrance (Fig. S15†). The highest occupied molecular orbitals (HOMOs) display significant orbital distribution on the porphyrin core and donor moieties, while the orbital distributions of the lowest unoccupied molecular orbitals (LUMOs) are mainly localized on the electron-withdrawing cyanoacrylic acid anchoring groups and porphyrin core. The HOMO level of FL1 (-4.81 eV) is slightly more negative than that of LG4 (-4.74 eV), agreeing with the weaker electron-donating ability of the fluorene moiety than the phenothiazine moiety. The calculated HOMO-LUMO energy gaps were determined to be 2.01 eV for LG4 and 2.06 eV for FL1, aligning well with the optical HOMO-LUMO gaps and electrochemical properties of these porphyrin dyes.

#### Optical and electrochemical properties of copper complexes

The UV/vis/NIR absorption spectra of the copper complexes  $[Cu(2MeOby)_2][X]$  and  $[Cu(2MeOby)_2][X]_2$  (X = TFSI, BF<sub>4</sub>, or PF<sub>6</sub>) in acetonitrile are illustrated in Fig. S18,† with their optical

Table 1 Optical and electrochemical properties of porphyrin dyes

Dye	$\lambda_{\rm abs}{}^a/{\rm nm} \left( \epsilon/10^3 \ { m M}^{-1} \ { m cm}^{-1} \right)$	$\lambda_{\mathrm{em}}{}^{b}/\mathrm{nm}$	$E_{\rm ox}^{c}/{\rm V}$	$E_{0-0}/\text{eV}$	$E_{\rm ox}^{*\ d}/{ m V}$	$-\Delta G_{ m inj}^{\ e}/{ m eV}$	$-\Delta G_{\mathrm{reg}}^{}f}/\mathrm{eV}$	$-\Delta G_{\mathrm{reg}}^{}g}/\mathrm{eV}$
FL1	460 (226), 668 (72)	681	1.07	1.90	-0.83 $-0.89$	0.33	0.19	0.25
LG4	460 (206), 673 (69)	688	0.98	1.87		0.39	0.10	0.16

<sup>&</sup>lt;sup>a</sup> Wavelengths for Soret and Q-band maxima in THF. <sup>b</sup> Wavelengths for emission maxima in THF by excitation at Soret band maxima. <sup>c</sup> Determined by cyclic voltammetry (CV) measurements of the porphyrin-adsorbed TiO<sub>2</sub> films (vs. the normal hydrogen electrode (NHE)). d Determined by adding the  $E_{0-0}$  value to the  $E_{0x}$  (vs. NHE). Driving forces for electron injection from the porphyrin singlet excited state ( ${}^{1}\text{ZnP}^{*}$ ) to the conduction band (CB) of TiO<sub>2</sub> ( $-0.5 \text{ V} \nu s$ . NHE). Driving forces for the regeneration of oxidized porphyrin dyes ( $\text{ZnP}^{*+}$ ) by the [Cu(tmby)<sub>2</sub>]<sup>+/2+</sup> redox couple (+0.88 V  $\nu s$ . NHE). Triving forces for the regeneration of oxidized porphyrin dyes (ZnP++) by the [Cu(2MeOby)<sub>2</sub>]+<sup>1/2+</sup> redox couple (+0.82 V vs. NHE).

**Edge Article** 

LUMO -2.73 eV -2.75 eV

Fig. 4 Selected Kohn–Sham orbitals for (a) LG4 and (b) FL1 obtained using DFT calculations at the B3LYP/6-31G(d) level. To simplify the calculations, octyl groups on the donor side were replaced with ethyl groups and octyloxy groups on the *meso*-aryl groups were replaced with methoxy groups.

Table 2 Optical properties and diffusion coefficients of  $[Cu(2MeOby)_2][X]$  and  $[Cu(2MeOby)_2][X]_2$  (X = TFSI, BF<sub>4</sub>, or PF<sub>6</sub>) in acetonitrile

	$\lambda_{\rm abs}^{}/{\rm nm} \left( \epsilon/{\rm M}^{-1} \ {\rm cm}^{-1} \right)$	$D^b/10^{-5} {\rm cm}^2 {\rm s}^{-1}$		
[Cu(2MeOby) <sub>2</sub> ][TFSI]	334 (4700), 445 (4500)	1.23		
[Cu(2MeOby) <sub>2</sub> ][TFSI] <sub>2</sub>	385 (1800), 748 (110)	_		
[Cu(2MeOby) <sub>2</sub> ][BF <sub>4</sub> ]	334 (4500), 445 (4500)	1.30		
$[Cu(2MeOby)_2][BF_4]_2$	385 (1700), 748 (130)	_		
[Cu(2MeOby) <sub>2</sub> ][PF <sub>6</sub> ]	334 (4700), 445 (4600)	2.41		
$[Cu(2MeOby)_2][PF_6]_2$	385 (1700), 748 (110)	_		

 $<sup>^</sup>a$  Wavelength of the absorption peaks in acetonitrile.  $^b$  Diffusion coefficients determined via chronoamperometry.

properties detailed in Table 2. Although we obtained the pure copper( $\pi$ ) complexes via electrochemical oxidation of the corresponding copper( $\pi$ ) complexes ( $vide\ supra$ ), we could not isolate the pure copper( $\pi$ ) complexes after electrolysis. Thus, we directly used the resulting solution for absorption and electrochemical measurements of the copper( $\pi$ ) complexes. The pronounced absorption bands spanning 300–500 nm are characteristic of metal-to-ligand charge transfer (MLCT) transitions,

a defining feature of copper complexes. In contrast, the faint absorption observed at 748 nm arises from d-d transitions, a hallmark of copper(II) complexes.<sup>54</sup> Interestingly, [Cu(2MeOby)<sub>2</sub>][X]/[Cu(2MeOby)<sub>2</sub>][X]<sub>2</sub> (X = TFSI, BF<sub>4</sub>, or PF<sub>6</sub>) exhibits identical absorption spectra, indicating that counter anions exert minimal influence on their optical properties.

To further explore the electrochemical behavior, we conducted CV measurements for both copper(1) and copper(11) complexes in acetonitrile (Fig. S20†). The copper(1) complex [Cu(2MeOby)2][TFSI] displayed a reversible oxidation peak at +0.82 V vs. NHE. Notably, the copper(II) complex [Cu(2MeOby)<sub>2</sub>] [TFSI]<sub>2</sub> exhibited a reversible reduction peak at precisely the same potential, +0.82 V, identical to the oxidation potential of its copper(1) counterpart. Consequently, the redox potential of [Cu(2MeOby)<sub>2</sub>][TFSI]/[Cu(2MeOby)<sub>2</sub>][TFSI]<sub>2</sub> was determined to be +0.82 V vs. NHE, which is lower than that of [Cu(tmby)<sub>2</sub>] [TFSI]/[Cu(tmby)2][TFSI]2 attributable to the electron-rich nature of the 2MeOby ligand with methoxy substitutions. Furthermore, the redox potentials of [Cu(2MeOby)2][BF4]/  $[Cu(2MeOby)_2][BF_4]_2$  and  $[Cu(2MeOby)_2][PF_6]/[Cu(2MeOby)_2]$ [PF<sub>6</sub>]<sub>2</sub> were similarly determined to be +0.82 V vs. NHE, reinforcing the notion that counterions have negligible impact on the redox behavior of the copper(I/II) complexes. Considering the  $E_{\rm ox}$  values of FL1/TiO<sub>2</sub> (+1.07 V vs. NHE) and LG4/TiO<sub>2</sub> (+0.98 V vs. NHE), the  $-\Delta G_{\text{reg}}$  values for FL1 and LG4 with  $[Cu(2MeOby)_2]^{+/2+}$  were estimated to be 0.25 and 0.16 eV, respectively. The increase in  $-\Delta G_{\mathrm{reg}}$  values is expected to accelerate the dye regeneration process, which, in turn, should enhance the PCE employing [Cu(2MeOby)<sub>2</sub>]<sup>+/2+</sup> as a redox shuttle.

#### Diffusion behavior of copper(1) complexes

The diffusion behavior of copper(1) complexes is closely linked to mass transport in DSSCs. Since charge transport in the electrolyte involves the movement of electroinactive counterions to maintain electroneutrality, the overall mass transport of copper(I/II) complexes is predominantly governed by their counterions. 51,58 The diffusion coefficients (D) of the copper(I) complexes were determined via chronoamperometry (Table 3 and Fig. S21†). For [Cu(tmby)2][TFSI] and [Cu(2MeOby)2][TFSI], the measured diffusion coefficients were  $1.28 \times 10^{-5}$  and 1.23 $\times$  10<sup>-5</sup> cm<sup>2</sup> s<sup>-1</sup>, respectively, aligning well with previous rotating disk electrode measurement for [Cu(tmby)2] TFSI] (1.12  $\times$  10<sup>-5</sup> cm<sup>2</sup> s<sup>-1</sup>).<sup>39</sup> This consistency suggests that the ligand structure has little influence on diffusion behavior. However, a notable deviation was observed for [Cu(2MeOby)2][PF6], which exhibited a significantly higher diffusion coefficient of 2.41 ×  $10^{-5}$  cm<sup>2</sup> s<sup>-1</sup>, compared to [Cu(2MeOby)<sub>2</sub>][TFSI] and  $[Cu(2MeOby)_2][BF_4]$  (1.30 × 10<sup>-5</sup> cm<sup>2</sup> s<sup>-1</sup>). This enhanced diffusion is likely attributed to weak electrostatic interaction between the positively charged copper center and the negatively charged [PF<sub>6</sub>] counterion, facilitating superior mobility. Such favorable diffusion properties are essential for efficient charge transport within the electrolyte, even at low redox shuttle ultimately supporting improved concentrations, performance.

Table 3 Photovoltaic performances of the DSSCs using copper(I/II) complexes as redox shuttles under AM 1.5 illumination<sup>6</sup>

Dye	Electrolyte	$J_{\rm IPCE}^{b}/{\rm mA~cm^{-2}}$	$J_{\rm SC}/{\rm mA~cm^{-2}}$	$V_{ m OC}/{ m V}$	ff	PCE/%
LG4	0.2 M [Cu(tmby) <sub>2</sub> ][TFSI]	9.50 (0.7%)	$9.57~(9.35\pm0.4)$	$0.842 \ (0.844 \pm 0.011)$	$0.716 \ (0.716 \pm 0.003)$	$5.77~(5.65\pm0.2)$
	0.05 M [Cu(tmby) <sub>2</sub> ][TFSI] <sub>2</sub>					
LG4	0.2 M [Cu(2MeOby) <sub>2</sub> ][TFSI]	10.0 (0.7%)	$10.07~(9.96\pm0.2)$	$0.821 \ (0.819 \pm 0.002)$	$0.762 \ (0.763 \pm 0.004)$	$6.30~(6.22\pm0.1)$
	0.05 M [Cu(2MeOby) <sub>2</sub> ][TFSI] <sub>2</sub>					
FL1	0.2 M [Cu(tmby) <sub>2</sub> ][TFSI]	10.8 (3.5%)	$11.2~(10.7\pm0.6)$	$0.902 \; (0.895 \pm 0.013)$	$0.767 \ (0.757 \pm 0.019)$	$7.75~(7.28\pm0.5)$
	0.05 M [Cu(tmby) <sub>2</sub> ][TFSI] <sub>2</sub>					
FL1	0.2 M [Cu(2MeOby) <sub>2</sub> ][TFSI]	12.3 (2.4%)	$12.6~(12.1\pm0.4)$	$0.887~(0.893\pm0.01)$	$0.730 \ (0.719 \pm 0.009)$	$8.15~(7.78\pm0.3)$
	0.05 M [Cu(2MeOby) <sub>2</sub> ][TFSI] <sub>2</sub>					
FL1	0.1 M [Cu(tmby) <sub>2</sub> ][TFSI]	10.2 (1.9%)	$10.4~(10.1\pm0.4)$	$0.935 \ (0.919 \pm 0.024)$	$0.720 \ (0.735 \pm 0.03)$	$7.00~(6.84\pm0.2)$
	$0.025 \text{ M } [\text{Cu(tmby)}_2][\text{TFSI}]_2$					
FL1	0.1 M [Cu(2MeOby) <sub>2</sub> ][TFSI]	11.6 (6.5%)	$12.4~(12.3~\pm~0.5)$	$0.890 \ (0.895 \pm 0.006)$	$0.752 \ (0.748 \pm 0.008)$	$8.30~(8.20\pm0.2)$
	0.025 M [Cu(2MeOby) <sub>2</sub> ][TFSI] <sub>2</sub>					
FL1	$0.1 \text{ M } [\text{Cu}(2\text{MeOby})_2][\text{BF}_4]$	$12.0 \ (4.7\%)$	$12.6~(12.4\pm0.6)$	$0.913 \ (0.916 \pm 0.003)$	$0.777~(0.762\pm0.02)$	$8.94~(8.63\pm0.3)$
	0.025 M [Cu(2MeOby) <sub>2</sub> ][BF <sub>4</sub> ] <sub>2</sub>					
FL1	0.1 M [Cu(2MeOby) <sub>2</sub> ][PF <sub>6</sub> ]	$13.0\ (1.5\%)$	$13.2~(12.7\pm0.4)$	$0.900~(0.910\pm0.01)$	$0.763 \ (0.762 \pm 0.006)$	$9.06~(8.80\pm0.2)$
	0.025 M [Cu(2MeOby) <sub>2</sub> ][PF <sub>6</sub> ] <sub>2</sub>					
XY1B	0.1 M [Cu(2MeOby) <sub>2</sub> ][PF <sub>6</sub> ]	$14.1\ (2.1\%)$	$14.4~(14.6\pm0.2)$	$0.961 \ (0.963 \pm 0.002)$	$0.752 \ (0.73 \pm 0.02)$	$10.4~(10.3\pm0.1)$
	$0.025 \text{ M } [\text{Cu}(2\text{MeOby})_2][\text{PF}_6]_2$					
FL1 + XY1B	0.1 M [Cu(2MeOby) <sub>2</sub> ][PF <sub>6</sub> ]	14.4 (5.3%)	$15.2\ (15.3\ \pm\ 0.1)$	$0.945~(0.940\pm0.01)$	$0.756 \ (0.752 \pm 0.005)$	$10.9~(10.8\pm0.1)$
	$0.025 \text{ M } [\text{Cu}(2\text{MeOby})_2][\text{PF}_6]_2$					

 $<sup>^</sup>a$  Photovoltaic parameters derived from the highest PCEs. The values in parentheses denote average values from three or four independent experiments. Error bars represent the standard error of the mean.  $^b$  The  $J_{\rm IPCE}$  values were obtained by integrating the IPCE plots, and the relative discrepancies from the corresponding  $J_{\rm SC}$  values obtained from the J-V characteristics are provided in the parentheses.

## Photovoltaic properties of DSSCs with FL1 and LG4 using [Cu(tmby)<sub>2</sub>][TFSI]/[Cu(tmby)<sub>2</sub>][TFSI]<sub>2</sub> and [Cu(2MeOby)<sub>2</sub>][TFSI]/[Cu(2MeOby)<sub>2</sub>][TFSI]<sub>2</sub> as redox shuttles

To optimize the fabrication conditions for DSSCs with FL1 under standard AM 1.5 conditions, we first utilized [Cu(tmby)<sub>2</sub>] [TFSI]/[Cu(tmby)<sub>2</sub>][TFSI]<sub>2</sub> as a redox shuttle. Because the asprepared copper(II) complex contains no other impurities except for the corresponding copper(1) species (vide supra), we can prepare the electrolyte solution with an accurate concentration by adjusting the amounts of copper(I/II) complexes based on the actual composition of the copper(II) complex [Cu(tmby)<sub>2</sub>] [TFSI]<sub>1.8</sub>. The electrolyte solution consisted of 0.2 M [Cu(tmby)<sub>2</sub>] [TFSI], 0.05 M [Cu(tmby)<sub>2</sub>][TFSI]<sub>2</sub>, 0.1 M LiTFSI, and 0.5 M 4-tertbutylpyridine (TBP) in acetonitrile. We screened immersion solvents while maintaining a fixed immersion time of 3 hours, ultimately selecting a mixture of toluene and ethanol (EtOH) in a 1:5 v/v ratio. We then evaluated the effect of immersion time, observing that PCEs increased with extended immersion periods, reaching a peak value of 6.94% for FL1 at 3 hours (Fig. S22†). To further enhance performance, we investigated chenodeoxycholic acid (CDCA) as a co-adsorbent to suppress dye aggregation on TiO<sub>2</sub> (Fig. S23†). The DSSC employing FL1 achieved a maximum PCE of 7.75% upon the addition of 1 equivalent of CDCA. For a comparative analysis, we fabricated DSSCs with LG4-sensitized TiO2 films under previously optimized conditions.46 The DSSC with LG4 using [Cu(tmby)2][TFSI]/ [Cu(tmby)<sub>2</sub>][TFSI]<sub>2</sub> exhibited a lower PCE of 5.77%, significantly below that with FL1. Notably, the optimized CDCA concentration was 1 equivalent for FL1 but 4 equivalents for LG4, likely due to FL1's reduced aggregation tendency arising from steric hindrance by its two alkyl chains at the 9-position of the

fluorene moiety. Additionally, we fabricated DSSCs utilizing  $[Cu(2MeOby)_2][TFSI]/[Cu(2MeOby)_2][TFSI]_2$  as a redox shuttle, benefiting from its larger  $-\Delta G_{\rm reg}$  values due to the less positive  $E_{\rm redox}$  value (Table 1). The DSSCs employing  $[Cu(2MeOby)_2][TFSI]/[Cu(2MeOby)_2][TFSI]_2$  demonstrated superior PCEs (8.15% for FL1 and 6.30% for LG4) compared to those using  $[Cu(tmby)_2][TFSI]/[Cu(tmby)_2][TFSI]_2$ . While DSSCs with  $[Cu(2MeOby)_2][TFSI]/[Cu(2MeOby)_2][TFSI]_2$  exhibited slightly lower  $V_{\rm OC}$  values due to the less positive  $E_{\rm redox}$  value, the enhanced  $J_{\rm SC}$  values effectively compensated for this decrease, ultimately resulting in improved PCEs.

The photocurrent-voltage characteristics of DSSCs under optimized conditions are presented in Fig. 5, with the corresponding photovoltaic data listed in Table 3. The superior PCEs observed for FL1 compared to LG4 primarily stem from its higher  $J_{SC}$  and  $V_{OC}$  values. The enhanced  $J_{SC}$  values are attributed to improved incident photon-to-current efficiencies (IPCEs), as illustrated in Fig. 6b. The IPCE values follow the trend: LG4 with  $[Cu(tmby)_2][TFSI]/[Cu(tmby)_2][TFSI]_2 < LG4$ with  $[Cu(2MeOby)_2][TFSI]/[Cu(2MeOby)_2][TFSI]_2 < FL1$  with  $[Cu(tmby)_2][TFSI]_2/[Cu(tmby)_2][TFSI]_2$  $[Cu(2MeOby)_2][TFSI]/[Cu(2MeOby)_2][TFSI]_2$ . Indeed, the  $J_{IPCE}$ values derived from photocurrent action spectra are compatible with the measured  $J_{SC}$  values. To further elucidate the differences in IPCE, we analyzed light-harvesting efficiency (LHE), electron injection efficiency ( $\phi_{\rm inj}$ ), and charge collection efficiency  $(\eta_{\rm col})$ , based on the equation: IPCE = LHE  $\times \phi_{\rm inj} \times \eta_{\rm col}$ . The absorption spectra of dye-adsorbed TiO<sub>2</sub> films, recorded without a light-scattering layer, revealed nearly identical absorption profiles for FL1 and LG4 (Fig. 6a). Taking into account the excellent absorption coefficients of the porphyrin dyes and the presence of a light-scattering layer in the actual Edge Article Chemical Science

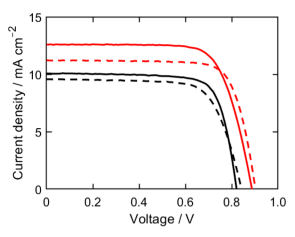


Fig. 5 Photocurrent–voltage characteristics of the DSSCs with FL1 (red) and LG4 (black) using [Cu(tmby)<sub>2</sub>][TFSI]/[Cu(tmby)<sub>2</sub>][TFSI]<sub>2</sub> (dashed lines) and [Cu(2MeOby)<sub>2</sub>][TFSI]/[Cu(2MeOby)<sub>2</sub>][TFSI]<sub>2</sub> (solid lines) redox shuttles for achieving the highest PCEs under the optimized conditions. The composition of the electrolyte solution: 0.2 M [CuL<sub>2</sub>][TFSI], 0.05 M [CuL<sub>2</sub>][TFSI]<sub>2</sub>, 0.1 M LiTFSI, and 0.5 M TBP in acetonitrile (L = tmby or 2MeOby).

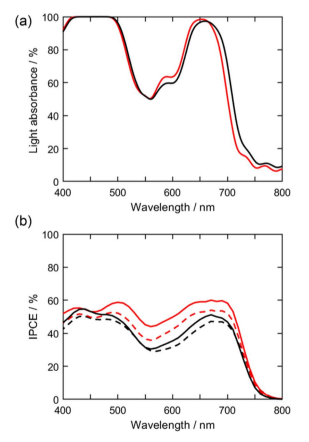


Fig. 6 (a) Absorption spectra of the porphyrin-adsorbed  $TiO_2$  film (2  $\mu$ m) of FL1 (red) and LG4 (black), and (b) photocurrent action spectra of the DSSCs with FL1 (red) and LG4 (black) using [Cu(tmby)<sub>2</sub>][TFSI]/[Cu(tmby)<sub>2</sub>][TFSI]<sub>2</sub> (dashed lines) and [Cu(2MeOby)<sub>2</sub>][TFSI]/[Cu(2MeOby)<sub>2</sub>][TFSI]<sub>2</sub> (solid lines) redox shuttles. The light-scattering  $TiO_2$  layers were not used to obtain an accurate absorption profile for the light-harvesting efficiency.

device, the LHE values approach 100% for both dyes. Although self-quenching of the excited state due to dye aggregation on  ${\rm TiO_2}$  could potentially reduce  $\phi_{\rm inj}$ , co-adsorption of CDCA effectively mitigates dye aggregation effects. Consequently, the higher IPCE values observed for **FL1** compared to **LG4** are primarily attributed to its superior  $\eta_{\rm col}$  values.

Given that the  $\eta_{col}$  value correlates with the dye regeneration efficiency ( $\phi_{reg}$ ), we conducted microsecond time-resolved transient absorption (TA) measurements on the porphyrinsensitized TiO2 films. The decay profiles of the porphyrin radical cation (ZnP<sup>•+</sup>) at 800 nm were analyzed, as the electrochemical oxidation of the porphyrin dyes distinctly revealed the characteristic absorption at 700–850 nm (Fig. S24†). The  $\varphi_{reg}$ values were derived from the lifetimes of ZnP\*+ in the absence and presence of copper(1/II) redox shuttles (Fig. S25 and Table S2†). The estimated  $\varphi_{\text{reg}}$  values follow the trend: **LG4** with  $[Cu(tmby)_2][TFSI]/[Cu(tmby)_2][TFSI]_2$  (58%) < LG4 with  $[Cu(_2 MeOby)_2$  [TFSI]/[Cu(2MeOby)<sub>2</sub>][TFSI]<sub>2</sub> (66%) <  $[Cu(tmby)_2][TFSI]/[Cu(tmby)_2][TFSI]_2 (71\%) <$ FL1  $[Cu(2MeOby)_2][TFSI]/[Cu(2MeOby)_2][TFSI]_2$  (84%). This order aligns well with the trend observed for the  $-\Delta G_{\text{reg}}$  values. The highest  $\varphi_{reg}$  value of 84% for FL1 with [Cu(2MeOby)<sub>2</sub>][TFSI]/ [Cu(2MeOby)2][TFSI]2 underscores the critical role of achieving a  $-\Delta G_{reg}$  value of 0.2 eV for an efficient dye regeneration process, even in DSSCs employing copper(1/II) redox shuttles.

To further assess CR between electrons in the CB of TiO<sub>2</sub> and the redox shuttle, we examined current-voltage characteristics under dark conditions (Fig. S26†). The more positive voltage onsets observed for FL1 compared to LG4 indicate an effective suppression of the CR process, likely due to an enhanced blocking effect. Dye loading amounts ( $\Gamma$ ) for FL1 and LG4, under the optimized sensitization conditions, were determined to be  $9.5 \times 10^{-11}$  and  $1.1 \times 10^{-10}$  mol cm<sup>-2</sup>, respectively. These values were obtained by measuring the absorbance of porphyrin dves dissolved from the dve-sensitized TiO2 films into a THF/  $H_2O$  (v/v = 1/1) solution with 0.1 M NaOH. The smaller  $\Gamma$  value for FL1 compared to LG4 suggests that the increased steric hindrance of FL1 plays a critical role in enhancing the blocking effect and suppressing dye aggregation on TiO2, primarily due to the presence of alkyl chains on the fluorene moiety. Additionally, the ET process at the dye/TiO2/electrolyte interface was investigated using electrical impedance spectroscopy (EIS) under AM 1.5 illumination under open-circuit conditions (Fig. S27†). The charge transfer resistance (R<sub>D</sub>) at the TiO<sub>2</sub>/dye/ electrolyte interface followed this trend: (32.2  $\Omega$  for LG4 with  $[Cu(2MeOby)_2][TFSI]/[Cu(2MeOby)_2][TFSI]_2 < 36.8 \Omega \text{ for } LG4$ with  $[Cu(tmby)_2][TFSI]/[Cu(tmby)_2][TFSI]_2 < 42.5 \Omega$  for FL1 with  $[Cu(2MeOby)_2][TFSI]/[Cu(2MeOby)_2][TFSI]_2 < 45.0 \Omega$  for FL1 with [Cu(tmby)<sub>2</sub>][TFSI]/[Cu(tmby)<sub>2</sub>][TFSI]<sub>2</sub>). This order is consistent with that of the observed  $V_{\rm OC}$  values, reinforcing the correlation between increased  $R_p$  values for FL1 and higher  $V_{OC}$ values in the DSSC utilizing FL1 compared to LG4.

To demonstrate the potential of our new porphyrin dye **FL1** for DSSCs using copper(I/II) complex-based redox shuttles, we also measured photovoltaic performances of the DSSCs with **FL1** and **LG4** using the standard  $I^-/I_3^-$  redox shuttle (Fig. S28, S29 and Table S3†). Although the DSSCs using the  $I^-/I_3^-$  redox

**Chemical Science Edge Article** 

shuttle exhibited higher  $J_{SC}$  values than those using  $[Cu(L)_2]$  $[TFSI]/[Cu(L)_2][TFSI]_2$  (L = tmby or 2MeOby) due to their larger driving forces for dye regeneration, the less positive redox potential of I<sup>-</sup>/I<sub>3</sub><sup>-</sup> (+0.40 V vs. NHE) resulted in significantly lower  $V_{\rm OC}$  values. Overall, the PCEs of the DSSCs using the I $^-/I_3^$ redox shuttle (5.84% for LG4 and 6.74% for FL1) are lower than those using the [Cu(2MeOby)<sub>2</sub>][TFSI]/[Cu(2MeOby)<sub>2</sub>][TFSI]<sub>2</sub> redox shuttle. These results clearly corroborate that the combination of the porphyrin dye and the copper(I/II) complexbased redox shuttle with a sufficient  $-\Delta G_{\text{reg}}$  value is an effective means to achieve further enhancement in PCE.

#### Effect of electrolyte composition on photovoltaic performances

The electrolyte solution plays a crucial role in DSSCs, prompting an investigation into the effect of its composition, specifically the concentration and counterions, on the photovoltaic properties of DSSCs with FL1. We first examined the performance using a low-concentration electrolyte containing 0.1 M [CuL<sub>2</sub>] [TFSI] and 0.025 M [CuL<sub>2</sub>][TFSI]<sub>2</sub> (L = tmby or 2MeOby) for [Cu(tmby)<sub>2</sub>][TFSI]/[Cu(tmby)<sub>2</sub>][TFSI]<sub>2</sub> and [Cu(2MeOby)<sub>2</sub>][TFSI]/ [Cu(2MeOby)<sub>2</sub>][TFSI]<sub>2</sub>. The DSSC employing [Cu(2MeOby)<sub>2</sub>] [TFSI]/[Cu(2MeOby)<sub>2</sub>][TFSI]<sub>2</sub> exhibited a slight improvement in PCE (8.30%), whereas the DSSC with [Cu(tmby)<sub>2</sub>][TFSI]/ [Cu(tmby)<sub>2</sub>][TFSI]<sub>2</sub> showed a significant decrease in PCE (7.00%) (Fig. S30† and Table 3). Reducing the concentration of copper(I/II) complexes led to a slight increase in the  $V_{OC}$  values likely due to suppressed CR between electrons in TiO2 and the redox shuttle. Interestingly, the  $J_{SC}$  value of the DSSC using [Cu(2MeOby)<sub>2</sub>][TFSI]/[Cu(2MeOby)<sub>2</sub>][TFSI]<sub>2</sub> remained comparable across both low and high concentrations, suggesting its robustness. In contrast, the  $J_{SC}$  value for  $[Cu(tmby)_2][TFSI]/$ [Cu(tmby)<sub>2</sub>][TFSI]<sub>2</sub> decreased substantially, directly contributing to its lower PCE. Considering the similar diffusion coefficients of [Cu(tmby)2][TFSI] and [Cu(2MeOby)2][TFSI], the diminished  $J_{SC}$  value for  $[Cu(tmby)_2][TFSI]/[Cu(tmby)_2][TFSI]_2$ at low concentrations can likely be ascribed to an inefficient dye regeneration process. Therefore, achieving higher  $-\Delta G_{\text{reg}}$ values proves advantageous for DSSCs operating with lowconcentration electrolytes.

Since counterions significantly affect diffusion properties (vide supra), they are also expected to impact the photovoltaic performance of DSSCs utilizing copper(I/II) complexes. To investigate this effect, we examined different counterions at optimal concentrations of 0.1 M [Cu(2MeOby)<sub>2</sub>][X] and 0.025 M  $[Cu(2MeOby)_2][X]_2$  (X = TFSI, BF<sub>4</sub>, or PF<sub>6</sub>) (Fig. 7a and Table 3). DSSCs incorporating  $[Cu(2MeOby)_2][X]/[Cu(2MeOby)_2][X]_2$  (X =  $BF_4$  or  $PF_6$ ) attained higher PCEs (8.94% for  $[Cu(2MeOby)_2]$  $[BF_4]/[Cu(2MeOby)_2][BF_4]_2$  and 9.06% for  $[Cu(2MeOby)_2][PF_6]/[BF_4]_2$  $[Cu(2MeOby)_2][PF_6]_2$  compared to  $[Cu(2MeOby)_2][TFSI]/$ [Cu(2MeOby)<sub>2</sub>][TFSI]<sub>2</sub>. Remarkably, the PCE of the DSSC employing FL1 with  $[Cu(2MeOby)_2][PF_6]/[Cu(2MeOby)_2][PF_6]_2$  is the highest reported among porphyrin-based DSSCs with copper(I/II) redox shuttles. The superior PCE observed for [Cu(2MeOby)2][PF6]/[Cu(2MeOby)2][PF6]2 can be attributed to its significantly improved  $J_{SC}$  value, in line with its higher IPCE values relative to those of [Cu(2MeOby)2][BF4]/[Cu(2MeOby)2]  $[BF_4]_2$  and  $[Cu(2MeOby)_2][TFSI]/[Cu(2MeOby)_2][TFSI]_2$  (Fig. 7b). To further elucidate this enhancement, we estimated  $\varphi_{reg}$  values through time-resolved TA measurements conducted in the presence of copper(1/II) complexes at low concentrations (0.1 M  $Cu^{I}$  and 0.025 M  $Cu^{II}$ ) (Fig. S31 and Table S4†). The  $\varphi_{reg}$  values followed the trend:  $[Cu(2MeOby)_2][TFSI]/[Cu(2MeOby)_2][TFSI]_2$  $(83\%) < [Cu(2MeOby)_2][BF_4]/[Cu(2MeOby)_2][BF_4]_2 (84\%) <$  $[Cu(2MeOby)_2][PF_6]/[Cu(2MeOby)_2][PF_6]_2$  (87%). Since counterions do not directly impact the  $E_{redox}$  value, the highest diffusion coefficient of [Cu(2MeOby)2][PF6] is likely responsible for the superior  $\varphi_{reg}$ , IPCE, and  $J_{SC}$  values, ultimately leading to the highest PCE. Even though the maximum IPCE value of the DSSC using [Cu(2MeOby)<sub>2</sub>][PF<sub>6</sub>]/[Cu(2MeOby)<sub>2</sub>][PF<sub>6</sub>]<sub>2</sub> exceeded 60% due to the highest  $\varphi_{reg}$  value, there remains room for further improvement in the  $J_{SC}$  value. Considering that the enhanced blocking effect can suppress the unfavorable CR process, 17,18 the introduction of bulky substituents is expected to increase the  $\eta_{\rm col}$  value, leading to a higher  $J_{\rm SC}$  value.

We also measured current-voltage characteristics under dark conditions (Fig. S32†) to evaluate the impact of counterions on VOC values. The onsets of dark current for  $[Cu(2MeOby)_2][BF_4]/[Cu(2MeOby)_2][BF_4]_2$  and  $[Cu(2MeOby)_2]$ [PF<sub>6</sub>]/[Cu(2MeOby)<sub>2</sub>][PF<sub>6</sub>]<sub>2</sub> were more positive than those for [Cu(2MeOby)2][TFSI]/[Cu(2MeOby)2][TFSI]2. Thus, the slightly improved V<sub>OC</sub> values observed for [Cu(2MeOby)<sub>2</sub>][BF<sub>4</sub>]/  $[Cu(2MeOby)_2][BF_4]_2$  and  $[Cu(2MeOby)_2][PF_6]/[Cu(2MeOby)_2]$  $[PF_6]_2$  can be ascribed to more effective suppression of CR between electrons in the CB of TiO2 and the redox shuttle. Although the precise mechanism remains unclear at this stage, these findings suggest that counterions significantly influence interfacial behavior at the TiO2/dye/electrolyte interfaces, with anion potentially accelerating undesirable CR processes.

#### Photovoltaic properties of co-sensitized DSSCs with XY1B

The DSSC with FL1 using [Cu(2MeOby)<sub>2</sub>][PF<sub>6</sub>]/[Cu(2MeOby)<sub>2</sub>] [PF<sub>6</sub>]<sub>2</sub> achieved the highest PCE among porphyrin-based DSSCs with copper(I/II) redox shuttles. However, further improvement in  $J_{\rm SC}$  values and overall PCEs remains possible due to relatively low IPCE values at around 550 nm. Co-sensitization with additional dyes exhibiting complementary absorption properties presents an effective strategy for achieving panchromatic absorption by addressing absorption deficits. 59-61 To this end, we selected XY1B as a co-sensitizer, considering its excellent absorption profile ( $\varepsilon$  = 53 000 M<sup>-1</sup> cm<sup>-1</sup> at 541 nm)<sup>18</sup> and high PCE (10.4%) in DSSCs under our optimized conditions (Fig. 2, S33† and Table 3). The cosensitized TiO<sub>2</sub> films were prepared via a second immersion step, wherein FL1-sensitized TiO2 films were immersed in an XY1B solution (0.1 M, 50 equiv. CDCA) in THF and ethanol solution (v/v = 1/4). Following optimization of the second immersion conditions, the co-sensitized DSSC with FL1 + XY1B exhibited an enhanced PCE, driven by improved  $J_{SC}$  and  $V_{OC}$  values (Fig. 7a and Table 3). The photocurrent action spectra clearly demonstrate a substantial increase in IPCE values within the 500-600 nm range, reaching 69% after co-sensitization with XY1B (Fig. 7b).

**Edge Article Chemical Science** 

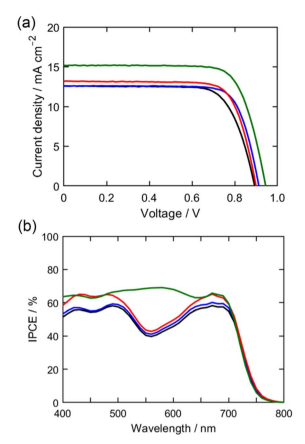


Fig. 7 (a) Photocurrent-voltage characteristics and (b) photocurrent action spectra of the DSSCs with FL1 using [Cu(2MeOby)2][TFSI]/  $[Cu(2MeOby)_2][TFSI]_2$  (black),  $[Cu(2MeOby)_2][BF_4]/[Cu(2MeOby)_2]$  $[BF_4]_2$  (blue), and  $[Cu(2MeOby)_2][PF_6]/[Cu(2MeOby)_2][PF_6]_2$  (red) and the DSSC with FL1 + XY1B using [Cu(2MeOby)<sub>2</sub>][PF<sub>6</sub>]/[Cu(2MeOby)<sub>2</sub>] [PF<sub>6</sub>]<sub>2</sub> (green) under the optimized conditions for achieving the highest PCEs. The composition of the electrolyte solution: 0.1 M  $[Cu(2MeOby)_2][X]$ , 0.025 M  $[Cu(2MeOby)_2][X]_2$ , 0.1 M LiX, and 0.5 M TBP in acetonitrile (X = TFSI,  $BF_4$ , or  $PF_6$ ).

This enhancement is corroborated by increased absorption observed in co-sensitized TiO2 films at approximately 550 nm (Fig. S34†). Dye loading amounts of FL1 and XY1B on the cosensitized TiO $_2$  were determined to be 4.7  $\times$   $10^{-11}$  and 6.2  $\times$  $10^{-11}$  mol cm<sup>-2</sup>, respectively, confirming that the  $J_{SC}$  enhancement stems from the complementary absorption of XY1B. Furthermore, current-voltage characteristics of the co-sensitized DSSC under dark conditions displayed a positive shift in the onset voltage, agreeing with the observed increase in  $V_{\rm OC}$  (Fig. S32†). Ultimately, the co-sensitized DSSC with FL1 + XY1B attained an outstanding PCE of 10.9%, surpassing the highest PCE recorded for porphyrin-based DSSCs employing copper(1/11) redox shuttles, marking the first instance of exceeding 10% PCE. Since the  $V_{\rm OC}$  value (0.945 V) is slightly higher than the highest V<sub>OC</sub> value in the co-sensitized porphyrin-based DSSCs with [Co(bpy)<sub>3</sub>]<sup>2+/3+</sup> (0.935 V), <sup>19</sup> further optimization of device fabrication conditions will achieve a  $V_{\rm OC}$  value higher than 1.0 V. In contrast, the co-sensitized DSSC with LG4 + XY1B delivered a PCE of 8.40%, which fell below that with XY1B as a single dye (Fig. S35 and Table S5†). This reduction may be attributed to LG4's aggregation tendency, which likely quenches XY1B's excited state, thereby impairing performance. Notably, the two alkyl chains on the fluorene moiety of FL1 play a crucial role in suppressing dye aggregation, proving advantageous not only for aggregation control but also for co-sensitization.

#### Stability of the DSSCs

The long-term stability of the DSSCs was assessed under whitelight illumination (100 mW cm<sup>-2</sup>) at 25 °C over a 500-hour period. The DSSC with FL1 + XY1B retained 84% of its initial efficiency, outperforming the DSSC with LG4 + XY1B, which maintained 73% (Fig. S36†). The superior stability of FL1 + XY1B may have resulted from the weaker electron-donating ability of the fluorene moiety in FL1 relative to the phenothiazine moiety in LG4, a characteristic beneficial for practical use.

#### Conclusions

We designed and synthesized a novel porphyrin dye, FL1, incorporating a fluorene skeleton with a weaker electrondonating nature relative to the phenothiazine skeleton. The  $E_{\rm ox}$  value of FL1/TiO<sub>2</sub> was determined to be +1.07 V vs. NHE, ensuring a  $-\Delta G_{\text{reg}}$  value comparable to or larger than 0.2 eV. Thanks to this increased  $-\Delta G_{reg}$  value, the DSSC with FL1 achieved a higher PCE of 7.75% than that with LG4 (5.77%) when using [Cu(tmby)<sub>2</sub>][TFSI]/[Cu(tmby)<sub>2</sub>][TFSI]<sub>2</sub> as the redox shuttle. Furthermore, the DSSC with FL1 employing [Cu(2MeOby)<sub>2</sub>][TFSI]/[Cu(2MeOby)<sub>2</sub>][TFSI]<sub>2</sub> exhibited an even higher PCE of 8.15%, attributed to an increased  $-\Delta G_{\text{reg}}$  value (0.25 eV). This finding underscores the importance of maintaining sufficient  $-\Delta G_{\text{reg}}$  values for achieving high-performance porphyrin-based DSSCs using copper(1/11) redox shuttles. To further explore performance optimization, we investigated the influence of counterions in [Cu(2MeOby)2][X]/[Cu(2MeOby)2]  $[X]_2$  (X = TFSI, BF<sub>4</sub>, or PF<sub>6</sub>) on diffusion behavior and identified [Cu(2MeOby)<sub>2</sub>][PF<sub>6</sub>] as having the highest diffusion coefficient. Consequently, the DSSC with FL1 using [Cu(2MeOby)2][PF6]/  $[Cu(2MeOby)_2][PF_6]_2$  attained the highest PCE of 9.06%, creating a new benchmark for porphyrin-based DSSCs employing copper(I/II) complex-based redox shuttles. Notably, co-sensitized DSSC with FL1 and the complementary dye XY1B led to an impressive PCE of 10.9%, while maintaining a high  $V_{\rm OC}$  of 0.945 V, setting a groundbreaking record as the first porphyrin-based DSSC employing copper(1/11) complex-based redox shuttles to exceed 10% PCE. Overall, our findings confirm that rational molecular engineering of porphyrin dyes and copper(1/11) complex-based redox shuttles with sufficient  $-\Delta G_{\text{reg}}$  values can significantly enhance photovoltaic performance. Additionally, selecting appropriate counterions with high diffusion coefficients is crucial for maximizing the charge transporting capabilities of copper(I/II) complexes. We anticipate that further refinement of porphyrin-based DSSCs, complementary dyes, and copper(I/II) redox shuttles through energy-level engineering will pave the way for next-generation solar cell technology, driving it toward top-tier performance.

12 Y. Ren, D. Sun, Y. Cao, H. N. Tsao, Y. Yuan, Data availability

The data supporting this article have been included as part of the ESI.†

#### **Author contributions**

T. Higashino and H. Imahori conceived and designed this work. Y. Zhang and K. Namikawa conducted the synthesis and characterization of the products. Y. Zhang, K. Namikawa, and W. R. Osterloh performed spectroscopic and electrochemical measurements. Y. Zhang fabricated the DSSCs and evaluated their photovoltaic performances. Y. Zhang, T. Higashino, and H. Imahori co-wrote the manuscript.

#### Conflicts of interest

There are no conflicts to declare.

### Acknowledgements

This work was supported by the JSPS (KAKENHI Grant Numbers JP20H05841 (T. H.), JP22K05066 (T. H.), JP25K01874 (T. H.), JP20H05832 (H. I.), and JP23H00309 (H. I.)). Y. Z. thanks the China Scholarship Council (CSC) (Chinese Government Scholarship 202006250042) for funding. In this work, DFT calculations were performed using the supercomputer of ACCMS under the collaborative research program for young/women scientists, Kyoto University.

#### Notes and references

- 1 E. K. Solak and E. Irmak, RSC Adv., 2023, 13, 12244-12269.
- 2 T. N. Murakami and N. Koumura, Adv. Energy Mater., 2019, 9, 1802967.
- 3 B. O'Regan and M. Grätzel, Nature, 1991, 353, 737-740.
- 4 A. Hagfeldt, G. Boschloo, L. Sun, L. Kloo and H. Pettersson, Chem. Rev., 2010, 110, 6595-6663.
- 5 A. Fakharuddin, R. Jose, T. M. Brown, F. Fabregat-Santiago and J. Bisquert, Energy Environ. Sci., 2014, 7, 3952-3981.
- 6 A. B. Muñoz-García, I. Benesperi, G. Boschloo, J. J. Concepcion, J. H. Delcamp, E. A. Gibson, G. J. Meyer, M. Pavone, H. Pettersson, A. Hagfeldt and M. Freitag, Chem. Soc. Rev., 2021, 50, 12450-12550.
- 7 H. Zhou, M. Aftabuzzaman, Masud, S. H. Kang and H. K. Kim, ACS Energy Lett., 2025, 10, 881-895.
- 8 Y. Ren, D. Zhang, J. Suo, Y. Cao, F. T. Eickemeyer, N. Vlachopoulos, S. M. Zakeeruddin, A. Hagfeldt and M. Grätzel, Nature, 2023, 613, 60-65.
- 9 Y. Wu, W.-H. Zhu, S. M. Zakeeruddin and M. Grätzel, ACS Appl. Mater. Interfaces, 2015, 7, 9307-9318.
- 10 P. Brogdon, H. Cheema and J. H. Delcamp, ChemSusChem, 2018, 11, 86-103.
- 11 J.-M. Ji, H. Zhou and H. K. Kim, J. Mater. Chem. A, 2018, 6, 14518-14545.

- S. M. Zakeeruddin, P. Wang and M. Grätzel, J. Am. Chem. Soc., 2018, 140, 2405-2408.
- 13 J.-M. Ji, H. J. Lee, H. Zhou, Y. K. Eom, C. H. Kim and H. K. Kim, ACS Appl. Mater. Interfaces, 2022, 14, 52745-52757.
- 14 H. Imahori, T. Umeyama and S. Ito, Acc. Chem. Res., 2009, 42, 1809-1818.
- 15 T. Higashino and H. Imahori, Dalton Trans., 2015, 44, 448-
- 16 Y. Zhang, T. Higashino and H. Imahori, J. Mater. Chem. A, 2023, 11, 12659-12680.
- 17 Y. Kurumisawa, T. Higashino, S. Nimura, Y. Tsuji, H. Iiyama and H. Imahori, J. Am. Chem. Soc., 2019, 141, 9910-9919.
- 18 Y. Zhang, T. Higashino, I. Nishimura and H. Imahori, ACS Appl. Mater. Interfaces, 2024, 16, 67761-67770.
- 19 M. Urbani, M. Grätzel, M. K. Nazeeruddin and T. Torres, Chem. Rev., 2014, 114, 12330-12396.
- 20 A. Yella, H.-W. Lee, H. N. Tsao, C. Yi, A. K. Chandiran, K. Nazeeruddin, E. W.-G. Diau, C.-Y. Yeh, S. M. Zakeeruddin and M. Grätzel, Science, 2011, 334, 629-634.
- 21 S. Mathew, A. Yella, P. Gao, R. Humphry-Baker, B. F. E. Curchod, N. Ashari-Astani, I. Tavernelli, U. Rothlisberger, M. K. Nazeeruddin and M. Grätzel, Nat. Chem., 2014, 6, 242-247.
- 22 A. Yella, C.-L. Mai, S. M. Zakeeruddin, S.-N. Chang, C.-H. Hsieh, C.-Y. Yeh and M. Grätzel, Angew. Chem., Int. Ed., 2014, 53, 2973-2977.
- 23 S. H. Kang, M. J. Jeong, Y. K. Eom, I. T. Choi, S. M. Kwon, Y. Yoo, J. Kim, J. Kwon, J. H. Park and H. K. Kim, Adv. Energy Mater., 2017, 7, 1602117.
- 24 J. Ji, H. Zhou, Y. K. Eom, C. H. Kim and H. K. Kim, Adv. Energy Mater., 2020, 10, 2000124.
- 25 H. Zhou, J.-M. Ji, H. S. Lee, Masud, M. Aftabuzzaman, D.-N. Lee, C. H. Kim and H. K. Kim, ACS Appl. Mater. Interfaces, 2023, 15, 39426-39434.
- 26 H. Song, Q. Liu and Y. Xie, Chem. Commun., 2018, 54, 1811-1824.
- 27 S. Huang, Q. Li, S. Li, C. Li, H. Tan and Y. Xie, Chem. Commun., 2024, 60, 4521-4536.
- 28 K. Zeng, Y. Lu, W. Tang, S. Zhao, Q. Liu, W. Zhu, H. Tian and Y. Xie, Chem. Sci., 2019, 10, 2186-2192.
- 29 K. Zeng, Y. Chen, W.-H. Zhu, H. Tian and Y. Xie, J. Am. Chem. Soc., 2020, 142, 5154-5161.
- 30 Z. Li, Q. Lu, Y. Zhang, Q. Li, W. Wu, S. Li, H. Wang, J. Jiang, C. Li and Y. Xie, J. Mater. Chem. A, 2025, 13, 4176-4185.
- 31 C.-C. Chen, J.-S. Chen, V. S. Nguyen, T.-C. Wei and C.-Y. Yeh, Angew. Chem., Int. Ed., 2021, 60, 4886-4893.
- 32 C.-C. Chen, Y.-H. Chen, V. S. Nguyen, S.-Y. Chen, M.-C. Tsai, J.-S. Chen, S.-Y. Lin, T.-C. Wei and C.-Y. Yeh, Adv. Energy Mater., 2023, 13, 2300353.
- 33 M. Wang, C. Grätzel, S. M. Zakeeruddin and M. Grätzel, Energy Environ. Sci., 2012, 5, 9394-9405.
- 34 J. Cong, X. Yang, L. Kloo and L. Sun, Energy Environ. Sci., 2012, 5, 9180-9194.

35 B. Pashaei, H. Shahroosvand and P. Abbasi, *RSC Adv.*, 2015, 5, 94814–94848.

**Edge Article** 

- 36 T. Higashino and H. Imahori, *ACS Energy Lett.*, 2022, 7, 1926–1938.
- 37 C. E. Housecroft and E. C. Constable, *Chem. Sci.*, 2022, **13**, 1225–1262.
- 38 Masud and H. K. Kim, ACS Omega, 2023, 8, 6139-6163.
- 39 Y. Saygili, M. Söderberg, N. Pellet, F. Giordano, Y. Cao, A. B. Muñoz-García, S. M. Zakeeruddin, N. Vlachopoulos, M. Pavone, G. Boschloo, L. Kavan, J.-E. Moser, M. Grätzel, A. Hagfeldt and M. Freitag, J. Am. Chem. Soc., 2016, 138, 15087–15096.
- 40 Y. Cao, Y. Liu, S. M. Zakeeruddin, A. Hagfeldt and M. Grätzel, *Joule*, 2018, **2**, 1108–1117.
- 41 Y. Ren, N. Flores-Díaz, D. Zhang, Y. Cao, J. Decoppet, G. C. Fish, J. Moser, S. M. Zakeeruddin, P. Wang, A. Hagfeldt and M. Grätzel, Adv. Funct. Mater., 2020, 30, 2004804.
- 42 D. Zhang, M. Stojanovic, Y. Ren, Y. Cao, F. T. Eickemeyer, E. Socie, N. Vlachopoulos, J.-E. Moser, S. M. Zakeeruddin, A. Hagfeldt and M. Grätzel, *Nat. Commun.*, 2021, 12, 1777.
- 43 H. Rui, J. Shen, Z. Yu, L. Li, H. Han and L. Sun, *Angew. Chem., Int. Ed.*, 2021, **60**, 16156–16163.
- 44 S. M. Meethal, S. C. Pradhan, J. Velore, S. Varughese, R. S. Pillai, F. Sauvage, A. Hagfeldt and S. Soman, *J. Mater. Chem. A*, 2024, 12, 1081–1093.
- 45 A. Colombo, G. Di Carlo, C. Dragonetti, M. Magni, A. Orbelli Biroli, M. Pizzotti, D. Roberto, F. Tessore, E. Benazzi, C. A. Bignozzi, L. Casarin and S. Caramori, *Inorg. Chem.*, 2017, 56, 14189–14197.
- 46 T. Higashino, H. Iiyama, I. Nishimura and H. Imahori, *Chem. Lett.*, 2020, **49**, 936–939.
- 47 Y.-H. Chen, C.-C. Chen, V. S. Nguyen, X.-T. Jiang, Y.-D. Chen, M.-Y. Li, S.-Y. Chen, T.-C. Wei and C.-Y. Yeh, *Cell Rep. Phys. Sci.*, 2024, 5, 102159.

- 48 T. Higashino, H. Iiyama, S. Nimura, Y. Kurumisawa and H. Imahori, *Inorg. Chem.*, 2020, **59**, 452–459.
- 49 G. C. Vougioukalakis, A. I. Philippopoulos, T. Stergiopoulos and P. Falaras, *Coord. Chem. Rev.*, 2011, 255, 2602–2621.
- 50 L. Li, L. Zhao, X. Jiang, Z. Yu, J. liu, H. Rui, J. Shen, W. Sharmoukh, N. K. Allam and L. Sun, *J. Mater. Chem. A*, 2022, 10, 4131–4136.
- 51 S. C. Pradhan, A. Hagfeldt and S. Soman, *J. Mater. Chem. A*, 2018, **6**, 22204–22214.
- 52 J. Ding, M. Day, G. Robertson and J. Roovers, *Macromolecules*, 2002, 35, 3474–3483.
- 53 M. Karpacheva, F. J. Malzner, C. Wobill, A. Büttner, E. C. Constable and C. E. Housecroft, *Dye. Pigment.*, 2018, 156, 410–416.
- 54 S. Kitagawa, M. Munakata and A. Higashie, *Inorg. Chim. Acta*, 1984, **84**, 79–84.
- 55 M. Giordano, G. Volpi, M. Bonomo, P. Mariani, C. Garino and G. Viscardi, *New J. Chem.*, 2021, **45**, 15303–15311.
- 56 P. Ferdowsi, Y. Saygili, S. M. Zakeeruddin, J. Mokhtari, M. Grätzel, A. Hagfeldt and L. Kavan, *Electrochim. Acta*, 2018, 265, 194–201.
- 57 L. Kavan, Y. Saygili, M. Freitag, S. M. Zakeeruddin, A. Hagfeldt and M. Grätzel, *Electrochim. Acta*, 2017, 227, 194–202.
- 58 G. Kalaignan and Y. Kang, J. Photochem. Photobiol. C Photochem. Rev., 2006, 7, 17-22.
- 59 N. V. Krishna, J. V. S. Krishna, M. Mrinalini, S. Prasanthkumar and L. Giribabu, *ChemSusChem*, 2017, 10, 4668–4689.
- 60 J. M. Cole, G. Pepe, O. K. Al Bahri and C. B. Cooper, *Chem. Rev.*, 2019, 119, 7279–7327.
- 61 Z. Li, Q. Li, C. Li and Y. Xie, *Mater. Chem. Front.*, 2024, **8**, 652–680.