#### [www.acsaem.org](www.acsaem.org?ref=pdf) Article

# **Orange/Red Benzo[1,2‑***b***:4,5‑***b***′]dithiophene 1,1,5,5-Tetraoxide-Based Emitters for Luminescent Solar Concentrators: Effect of Structures on Fluorescence Properties and Device Performances**

Matteo [Bartolini,](https://pubs.acs.org/action/doSearch?field1=Contrib&text1="Matteo+Bartolini"&field2=AllField&text2=&publication=&accessType=allContent&Earliest=&ref=pdf) Cosimo [Micheletti,](https://pubs.acs.org/action/doSearch?field1=Contrib&text1="Cosimo+Micheletti"&field2=AllField&text2=&publication=&accessType=allContent&Earliest=&ref=pdf) [Alberto](https://pubs.acs.org/action/doSearch?field1=Contrib&text1="Alberto+Picchi"&field2=AllField&text2=&publication=&accessType=allContent&Earliest=&ref=pdf) Picchi, Carmen [Coppola,](https://pubs.acs.org/action/doSearch?field1=Contrib&text1="Carmen+Coppola"&field2=AllField&text2=&publication=&accessType=allContent&Earliest=&ref=pdf) Adalgisa [Sinicropi,](https://pubs.acs.org/action/doSearch?field1=Contrib&text1="Adalgisa+Sinicropi"&field2=AllField&text2=&publication=&accessType=allContent&Earliest=&ref=pdf) [Mariangela](https://pubs.acs.org/action/doSearch?field1=Contrib&text1="Mariangela+Di+Donato"&field2=AllField&text2=&publication=&accessType=allContent&Earliest=&ref=pdf) Di Donato, Paolo [Foggi,](https://pubs.acs.org/action/doSearch?field1=Contrib&text1="Paolo+Foggi"&field2=AllField&text2=&publication=&accessType=allContent&Earliest=&ref=pdf) [Alessandro](https://pubs.acs.org/action/doSearch?field1=Contrib&text1="Alessandro+Mordini"&field2=AllField&text2=&publication=&accessType=allContent&Earliest=&ref=pdf) Mordini, Gianna [Reginato,](https://pubs.acs.org/action/doSearch?field1=Contrib&text1="Gianna+Reginato"&field2=AllField&text2=&publication=&accessType=allContent&Earliest=&ref=pdf) [Andrea](https://pubs.acs.org/action/doSearch?field1=Contrib&text1="Andrea+Pucci"&field2=AllField&text2=&publication=&accessType=allContent&Earliest=&ref=pdf) Pucci,[\\*](#page-15-0) [Lorenzo](https://pubs.acs.org/action/doSearch?field1=Contrib&text1="Lorenzo+Zani"&field2=AllField&text2=&publication=&accessType=allContent&Earliest=&ref=pdf) Zani,[\\*](#page-15-0) and Massimo [Calamante](https://pubs.acs.org/action/doSearch?field1=Contrib&text1="Massimo+Calamante"&field2=AllField&text2=&publication=&accessType=allContent&Earliest=&ref=pdf)



of optical devices able to harvest, downshift, and concentrate sunlight, thanks to the presence of emitting materials embedded in a polymer matrix. Use of LSCs in combination with silicon-based photovoltaic (PV) devices has been proposed as a viable strategy to enhance their ability to harvest diffuse light and facilitate their integration in the built environment. LSC performances can be improved by employing organic fluorophores with strong light absorption in the center of the solar spectrum and intense, redshifted emission. In this work, we present the design, synthesis, characterization, and application in LSCs of a series of orange/red organic emitters featuring a benzo[1,2-*b*:4,5-*b*′]dithiophene 1,1,5,5-tetraoxide central core as an acceptor (A) unit. The latter



was connected to different donor  $(D)$  and acceptor  $(A')$  moieties by means of Pd-catalyzed direct arylation reactions, yielding compounds with either symmetric (D−A−D) or non-symmetric (D−A−A′) structures. We found that upon light absorption, the compounds attained excited states with a strong intramolecular charge-transfer character, whose evolution was greatly influenced by the nature of the substituents. In general, symmetric structures showed better photophysical properties for the application in LSCs than their non-symmetric counterparts, and using a donor group of moderate strength such as triphenylamine was found preferable. The best LSC built with these compounds presented photonic (external quantum efficiency of  $8.4 \pm 0.1\%$ ) and PV (device efficiency of 0.94  $\pm$  0.06%) performances close to the state-of-the-art, coupled with a sufficient stability in accelerated aging tests.

KEYWORDS: *luminescent solar concentrators, organic emitters, benzo[1,2-b:4,5-b*′*]dithiophene 1,1,5,5-tetraoxide, DFT calculations, transient absorption spectroscopy, photovoltaics*

# **1. INTRODUCTION**

Luminescent solar concentrators (LSCs) are a class of optical devices invented in the late  $1970s$ ,  $^{1,2}$  able to harvest, downshift, and concentrate sunlight on a small-surface area. $3$  Being capable of decoupling the emission direction from the incident light direction, while simultaneously achieving high concentration gains, they have recently been employed for several different purposes, including lighting, driving of chemical reactions, and optical communication.<sup>4</sup> Besides that, they have also been extensively examined for the application in the field of photovoltaics (PVs), thanks to the possibility of coupling them with crystalline silicon solar cells, as well as other PV technologies.<sup>5,6</sup> Indeed, based on their ability to harvest diffuse or indirect light, $7$  accompanied by their colorful appearance and light weight, LSCs have been proposed as a potential solution to overcome the traditional limitations of Si-based PV technologies, such as their bulky and heavy structure or the need for direct illumination, and foster their diffusion in the built environment.<sup>[8](#page-16-0)</sup>

An LSC is usually made of a sheet of a common plastic material [e.g., poly(methyl methacrylate) PMMA] containing a fluorescent dopant, able to absorb sunlight and emit it at longer wavelengths. Typically used emitters comprise quantum dots, perovskites, rare-earth complexes, and organic compounds.[3](#page-16-0) Due to the different refractive indexes of air and the plastic material, the emitted radiation is mainly concentrated, via total internal reflection, at the edges of the panel, where Si

Received: February 9, 2023 Accepted: April 11, 2023 Published: April 20, 2023





© <sup>2023</sup> The Authors. Published by American Chemical Society **<sup>4862</sup>**

#### Benzodithiophene tetraoxide (BDT) core

<span id="page-1-0"></span>

Figure 1. Structures of organic emitter BDT-H2 (TPA-BDTO) and the other compounds designed in this work.

PV cells can be placed. Clearly, LSC performances are critically related to the nature and properties of their constituting materials, among which the fluorescent emitter plays a key role.

The most important properties an emitter should have for a fruitful application in LSCs are a strong light harvesting ability, a high fluorescence quantum yield  $(FQY, \Phi_f)$ , and a large Stokes shift to minimize re-absorption of emitted light, coupled with a good compatibility with the chosen polymer matrix.<sup>[3](#page-16-0)</sup> In recent years, the use of organic fluorescent compounds as emitters for LSCs has been intensely investigated since they allow the tuning of the abovementioned photophysical properties by careful structural modifications, while keeping relatively simple synthetic procedures and low cost.<sup>[9](#page-16-0)</sup> Most organic emitters present a molecular structure characterized by alternating electron-donating (D) and -accepting (A) moieties since such arrangement usually favors strong intramolecular charge transfer (ICT) processes upon photoexcitation, helping achieve the desired optical properties.<sup>[10](#page-16-0),[11](#page-16-0)</sup>

Notably, since LSCs are designed to work under natural sunlight, an optimal organic emitter should absorb light in the 500−600 nm range, where solar radiation presents the maximum irradiance, $12$  and display a Stokes shift large enough to induce fluorescence emission in the red/NIR region of the spectrum. The latter would be beneficial to improve the match with the absorption of Si-based solar cells, limiting energy losses due to thermalization and improving the overall power conversion efficiency (PCE) of the system. Nevertheless, achieving high FQY with this kind of emitters has proven challenging, due to the so-called "energy gap law", stating that nonradiative recombination becomes more and more likely as the energy gap between the first excited state and the ground state becomes smaller.<sup>13</sup> Therefore, careful structural design is mandatory to achieve the best possible compromise between all these different parameters, while at the same time ensuring sufficient solubility and good processability in the conditions of use.

In continuation of our recent activity dedicated to the investigation of new organic emitters for  $LSCs$ ,<sup>[14](#page-16-0)−[17](#page-16-0)</sup> we therefore sought to identify suitable heterocyclic structures that could potentially combine all the required optical and physicochemical properties. Upon careful inspection of the literature, we identified a series of fluorophores containing the strongly electron-withdrawing benzo[1,2-*b*:4,5-*b*′]dithiophene 1,1,5,5 tetraoxide core (BDT-tetraoxide) flanked by different electrondonating groups, originally reported by Tang and co-workers for applications in bio-imaging technologies.<sup>[18](#page-16-0)</sup> In particular, among the compounds described in the paper, we focused on the TPA-BDTO emitter (Figure 1, here also denoted as BDT-H2, see below) because besides showing aggregation-induced emission properties,<sup>[19](#page-16-0)</sup> it displayed the most red-shifted emission spectra both in solution and in the solid state, with maxima well beyond 600 nm. Indeed, thanks to their excellent emission features, compound TPA-BDTO and close derivatives thereof were widely applied by Tang's and other groups in the bio-diagnostic and theranostic fields,<sup>[20](#page-16-0)−[24](#page-17-0)</sup> as well as in high FQY PMMA dispersions for the fabrication of microlasers to be used in tensile strain sensors.<sup>[25](#page-17-0)</sup> On the other hand, compounds incorporating the BDT-tetraoxide core have never been reported as emitters for  $LSCs<sub>1</sub><sup>26</sup>$  $LSCs<sub>1</sub><sup>26</sup>$  $LSCs<sub>1</sub><sup>26</sup>$  and only the use of fluorophores based on the reduced BDT unit was described in the literature.<sup>[27](#page-17-0)</sup> For the above reasons, **TPA-BDTO** appeared as a promising candidate for use in LSCs, as well as an ideal starting point for developing other LSC fluorophores by introducing targeted modifications in its molecular structure.

Accordingly, in this paper, we report the design, synthesis, and spectroscopic characterization of a series of organic emitters featuring a benzo[1,2-*b*:4,5-*b*′]dithiophene 1,1,5,5 tetraoxide system as a central acceptor unit connected to

<span id="page-2-0"></span>different electron-donating and electron-withdrawing substituents. The spectroscopic properties of the compounds in organic solutions were investigated both in static conditions and by means of transient absorption techniques, to monitor the evolution of their excited states. Selected members of the series were then dispersed in PMMA films at different concentrations, and the optical and PV performances of the corresponding LSC were determined, together with a preliminary stability assessment by means of an accelerated aging protocol.

# **2. EXPERIMENTAL SECTION**

**2.1. General Synthetic and Experimental Remarks.** All reactions carried out under an inert nitrogen atmosphere were performed in flame- or oven-dried glassware using Schlenk techniques. Tetrahydrofuran (THF) was purified by distillation over metallic sodium in the presence of benzophenone as an indicator, while aniline was distilled over KOH at  $5 \times 10^{-2}$  atm of pressure. Toluene, dichloromethane, *N*,*N*-dimethylformamide (DMF), diethylether, and acetonitrile were dried on a resin exchange Solvent Purification System (Innovative Technology) and stored under nitrogen over 4 Å molecular sieves. Degassing of solvents was carried out using the "freeze−pump−thaw" method. Compound 2b was prepared according to a known procedure.<sup>[28](#page-17-0)</sup> Unless otherwise stated, all other chemicals employed were commercially available and were used as received. Petroleum ether was the 40−60 °C boiling fraction. Thin-layer chromatography (TLC) analyses were carried out using aluminum-supported silica gel plates containing a fluorescent indicator, and detection was carried out by exposure to UV light (*λ* = 254 and 365 nm) and/or treatment with permanganate or *p*anisaldehyde solutions followed by heating. Flash column chromatography was performed using Merck Kieselgel 60 (300−400 mesh) as the stationary phase. <sup>1</sup>H NMR spectra were recorded at 200, 300, or 400 MHz, and 13C NMR spectra were recorded at 100.6 MHz, with Bruker Avance or Varian Mercury series instruments. Chemical shifts are referenced to the residual solvent peak (CHCl<sub>3</sub>,  $\delta$  7.26 ppm for  $^1\rm H$ NMR and *δ* 77.0 ppm for 13C NMR; THF-*d*8, *δ* 1.72 and 3.58 ppm for  $^1\mathrm{H}$  NMR and  $\delta$  67.21 and 25.31 ppm for  $^{13}\mathrm{C}$  NMR). Coupling constants (*J*) are expressed in Hz, while the used abbreviations are s (singlet), d (doublet), dd (doublet of doublets), t (triplet), td (triplet of doublets), and m (multiplet). Multiplets are indicated as chemical shift intervals. ESI-MS spectra were obtained by the direct injection of the sample solution, with a Thermo Scientific LCQ-FLEET instrument and are reported in the form  $m/z$ . UV/vis spectra in different solvents were recorded with a Shimadzu UV-2600 spectrometer. Fluorescence spectra on solutions were measured at room temperature with a Jasco FP-8300 spectrofluorometer equipped with a 450 W Xenon arc lamp, while quantum yield analysis was conducted using an ILF-835 integrating sphere (*ø* = 100 mm) connected to the instrument.

**2.2. Experimental Synthetic Procedures.** *2.2.1. 4-Bromo-N,Nbis(4-(methylthio)phenyl)aniline (2c).* First step: In a Schlenk tube,  $Pd(OAc)_{2}$  (7 mg, 0.031 mmol, 0.03 equiv) and dppf (18 mg, 0.031 mmol, 0.03 equiv) were dissolved in dry toluene (10 mL), and the solution was kept stirring for 10 min under an inert atmosphere, until it turned orange. Then, aniline (100 mg, 1.04 mL, 1.1 mmol, 1.0 equiv), 4-bromothioanisole (478 mg, 2.32 mmol, 2.2 equiv) and, finally, *t*-BuONa (309 mg, 3.2 mmol, 3.0 equiv) were added. The resulting dark blue solution was heated up to 135 °C and stirred for 16 h, during which it turned yellow. The reaction mixture was cooled to room temperature, diluted with  $CH_2Cl_2$ , and washed with  $H_2O$  (20 mL) and brine (20 mL). The combined organic layers were dried over Na2SO4 and concentrated under vacuum. The residue was purified by flash column chromatography (SiO<sub>2</sub>, petroleum ether/CH<sub>2</sub>Cl<sub>2</sub> 6:1 to 4:1) to yield pure *N*,*N*-bis(4-(methylthio)phenyl)aniline (260 mg, 0.77 mmol, 70%) as a colorless oil. <sup>1</sup>H NMR (CDCl<sub>3</sub>, 200 MHz): *δ* 7.22−7.25 (m, 2 H), 7.16 (dd, *J* = 8.6, 2.0 Hz, 4 H), 6.97−7.10 (m, 7

H), 2.47 (s, 6 H) ppm. The analytical data are in agreement with those reported in the literature.<sup>[29](#page-17-0)</sup>

Second step: In a round-bottom flask, *N*,*N*-bis(4-(methylthio) phenyl)aniline (260 mg, 0.77 mmol, 1.0 equiv) was dissolved in DMF (5 mL), and the mixture was cooled to 0 °C. Then, a solution of *N*bromosuccinimide (NBS, 137 mg, 0.77 mmol, 1.0 equiv) in DMF (5 mL) was added dropwise, and the resulting mixture was allowed to warm up to room temperature while shielded from the light. After 3 h, the reaction was quenched with H<sub>2</sub>O and extracted with EtOAc (3  $\times$ 10 mL). The combined organic layers were dried over  $Na<sub>2</sub>SO<sub>4</sub>$  and concentrated under vacuum. The residue was purified by flash column chromatography (SiO<sub>2</sub>, petroleum ether/CH<sub>2</sub>Cl<sub>2</sub> 6:1 to 4:1) to yield pure product  $2c$  (221 mg, 0.53 mmol, 69%) as a white solid.  $\rm ^1H$  NMR (CDCl3, 200 MHz): *δ* 7.32 (d, *J* = 8.7 Hz, 2 H), 7.17 (d, *J* = 8.7 Hz, 4 H), 6.99 (d, *J* = 8.7 Hz, 4 H), 6.92 (d, *J* = 8.7 Hz, 2 H), 2.47 (s, 6 H) ppm. The analytical data are in agreement with those reported in the literature.<sup>[29](#page-17-0)</sup>

*2.2.2. General Procedure for the Symmetric Direct Arylation of Compound* 1. In a Schlenk tube,  $Pd_2(dba)$ <sub>3</sub> (0.05 equiv),  $P(o-a)$  $MeOPh$ )<sub>3</sub> (0.10 equiv), and pivalic acid (0.3 equiv) were dissolved in dry toluene (10 mL per 0.1 mmol of the substrate), and the mixture was kept under stirring for 10 min under an inert  $N_2$  atmosphere. Then, compound 1 (1.0 equiv), the appropriate aryl halide  $(2a-c,$ 2.2−4.0 equiv), and  $Cs<sub>2</sub>CO<sub>3</sub>$  (2.4 equiv) were added, and the resulting suspension was heated up to 110 °C and stirred for 16 h. The reaction mixture was cooled to room temperature and filtered over a short pad of Celite©. Then, the solution was diluted with  $CH<sub>2</sub>Cl<sub>2</sub>$  (equal volume of the reaction solvent) and washed with  $H<sub>2</sub>O$  $(2 \times$  volume of the reaction solvent) and brine  $(2 \times$  volume of the reaction solvent). The combined organic layers were dried over Na2SO4 and concentrated under vacuum. The residue was purified by flash column chromatography to yield the pure product.

*2.2.2.1. 2,6-Bis(4-(diphenylamino)phenyl)-4,8-bis((2-ethylhexyl) oxy)benzo[1,2-b:4,5-b*′*]dithiophene 1,1,5,5-Tetraoxide (BDT-H2).* Compound BDT-H2 was synthesized from 1 (207 mg, 0.41 mmol) and 4-bromophenyl(diphenyl)amine (2a, 289 mg, 0.89 mmol) in accordance with the general procedure. The crude product was purified by flash column chromatography  $(SiO<sub>2</sub>)$ , petroleum ether/  $CH_2Cl_2$  2:1 to 1:1), after which the desired compound was still contaminated by ca. 5% dibenzylideneacetone (dba). Recrystallization from a petroleum ether/ $CH_2Cl_2$  20:1 mixture yielded the pure product BDT-H2 (227 mg, 0.23 mmol, 56%) as <sup>a</sup> deep purple solid. <sup>1</sup> <sup>1</sup>H NMR (CDCl<sub>3</sub>, 400 MHz):  $\delta$  7.65 (d, *J* = 8.6 Hz, 4 H), 7.28–7.35 (m, 8 H), 7.24 (s, 2 H), 7.05−7.19 (m, 16 H), 4.39 (d, *J* = 5.1 Hz, 4 H), 1.79−1.89 (m, 2 H), 1.46−1.67 (m, 8 H), 1.32−1.41 (m, 8 H), 1.00 (t, *J* = 7.4 Hz, 6 H), 0.92 (t, *J* = 6.4 Hz, 6 H) ppm. 13C NMR (CDCl3, 100 MHz): *δ* 150.0, 146.6, 145.0, 142.2, 131.3, 129.5, 127.6, 127.4, 125.6, 124.4, 121.7, 119.3, 115.0, 79.0, 40.4, 30.3, 29.0, 23.7, 23.0, 14.1, 11.2 ppm. ESI-MS:  $m/z$  calcd for  $C_{62}H_{64}N_2O_6S_2$  [M<sup>+•</sup>], 996.4; found, 996.1. The analytical data are in agreement with those reported in the literature.<sup>18</sup>

*2.2.2.2. 2,6-Bis(4-(bis(4-methoxyphenyl)amino)phenyl)-4,8-bis- ((2-ethylhexyl)oxy)benzo[1,2-b:4,5-b*′*]dithiophene 1,1,5,5-Tetraoxide (BDT-O2).* Compound BDT-O2 was synthesized from 1 (30 mg, 0.06 mmol) and 2b (50 mg, 0.13 mmol) in accordance with the general procedure. The crude product was purified by flash column chromatography (SiO<sub>2</sub>, petroleum ether/CH<sub>2</sub>Cl<sub>2</sub> 2:1 to 1:1) to yield pure product BDT-O2 (84 mg, 0.075 mmol, 58%) as a deep purple solid. <sup>1</sup>H NMR (CDCl<sub>3</sub>, 400 MHz): *δ* 7.58 (d, *J* = 8.9 Hz, 4 H), 7.16 (s, 2 H), 7.10 (d, *J* = 8.5 Hz, 8 H), 6.91 (d, *J* = 8.5 Hz, 4 H), 6.86 (d, *J* = 8.9 Hz, 8 H), 4.36 (d, *J* = 5.5 Hz, 4 H), 3.82 (s, 12 H), 1.79−1.89 (m, 2 H), 1.53−1.65 (m, 8 H), 1.31−1.40 (m, 8 H), 0.98 (t, *J* = 7.4 Hz, 6 H), 0.90 (t,  $J = 7.4$  Hz, 6 H) ppm. <sup>13</sup>C NMR (CDCl<sub>3</sub>, 100 MHz): *δ* 156.8, 143.3, 139.4, 133.5, 130.5, 129.0, 128.4, 127.5, 126.6, 125.4, 118.8, 117.5, 114.9, 78.9, 55.5, 40.4, 30.2, 29.7, 29.0, 23.7, 23.0, 14.1, 11.2 ppm. ESI-MS:  $m/z$  calcd for  $C_{66}H_{72}N_2O_{10}S_2$  [M<sup>+•</sup>], 1116.5; found, 1116.1.

*2.2.2.3. 2,6-Bis(4-(bis(4-(methylthio)phenyl)amino)phenyl)-4,8 bis((2-ethylhexyl)oxy)benzo[1,2-b:4,5-b*′*]dithiophene 1,1,5,5-Tetraoxide (BDT-S2).* Compound BDT-S2 was synthesized from 1 (50 mg, 0.10 mmol) and 2c (92 mg, 0.22 mmol) in accordance with the general procedure. The crude product was purified by flash column chromatography (SiO<sub>2</sub>, petroleum ether/CH<sub>2</sub>Cl<sub>2</sub> 2:1 to 1:2) to yield pure product BDT-S2 (83 mg, 0.07 mmol, 70%) as a deep purple solid. <sup>1</sup> H NMR (THF-*d*8, 400 MHz): *δ* 7.73 (d, *J* = 8.9 Hz, 4 H), 7.42 (s, 2 H), 7.24 (d, *J* = 8.6 Hz, 8 H), 7.09 (d, *J* = 8.5 Hz, 8 H), 7.06 (d, *J* = 8.9 Hz, 4 H), 4.41 (d, *J* = 5.8 Hz, 4 H), 2.46 (s, 12 H), 1.85−1.92 (m, 2 H), 1.54−1.60 (m, 8 H), 1.35−1.41 (m, 8 H), 1.01 (t, *J* = 7.4 Hz, 6 H), 0.84-0.96 (m, 6 H) ppm. <sup>13</sup>C NMR (THF- $d_8$ , 100 MHz): *δ* 150.5, 145.8, 144.7, 143.4, 135.3, 132.9, 128.8, 128.4, 128.2, 126.6, 121.9, 120.6, 115.6, 79.6, 41.2, 32.7, 30.4, 24.3, 23.7, 15.9, 14.2, 11.3 ppm. ESI-MS:  $m/z$  calcd for  $C_{66}H_{73}N_2O_6S_6$  [M + H<sup>+•</sup>], 1181.4; found, 1181.3.

*2.2.2.4. 4,4*′*-(4,8-Bis((2-ethylhexyl)oxy)-1,1,5,5-tetraoxidobenzo- [1,2-b:4,5-b*′*]dithiophene-2,6-diyl)dibenzonitrile (BDT-AA).* Compound BDT-AA was synthesized from 1 (50 mg, 0.10 mmol) and 4-bromobenzonitrile (3, 72 mg, 0.40 mmol) in accordance with the general procedure. The crude product was purified by flash column chromatography (SiO<sub>2</sub>, petroleum ether/CH<sub>2</sub>Cl<sub>2</sub> 1:1 to sole CH<sub>2</sub>Cl<sub>2</sub>) to yield product BDT-AA (50 mg, 0.07 mmol, 70%) as an orange solid. <sup>1</sup>H NMR (CDCl<sub>3</sub>, 400 MHz): *δ* 7.92 (d, *J* = 8.4 Hz, 4 H), 7.79 (d, *J* = 8.4 Hz, 4 H), 7.57 (s, 2 H), 4.47 (d, *J* = 5.5 Hz, 4 H), 1.84− 1.92 (m, 2 H), 1.46−1.66 (m, 8 H), 1.33−1.43 (m, 8 H), 1.01 (t, *J* = 7.4 Hz, 6 H), 0.93 (t, *J* = 7.1 Hz, 6 H) ppm. <sup>13</sup>C NMR (CDCl<sub>3</sub>, 100 MHz): *δ* 145.4, 140.9, 133.0, 131.2, 127.5, 127.2, 121.1, 117.9, 114.2, 110.0, 79.2, 40.3, 30.3, 29.0, 23.7, 23.0, 14.1, 11.1 ppm. ESI-MS: *m*/*z* calcd for  $C_{80}H_{88}N_4NaO_{12}S_4$  [2M + Na<sup>+•</sup>], 1447.5; found, 1446.8.

*2.2.3. 2-(4-(Diphenylamino)phenyl)-4,8-bis((2-ethylhexyl)oxy) benzo[1,2-b:4,5-b*′*]dithiophene 1,1,5,5-Tetraoxide (4a).* In a Schlenk tube, Pd<sub>2</sub>(dba)<sub>3</sub> (4 mg, 0.004 mmol, 0.05 equiv), P(o- $MeOPh$ )<sub>3</sub> (6 mg, 0.016 mmol, 0.2 equiv), and pivalic acid (2 mg, 0.023 mmol, 0.3 equiv) were dissolved in dry toluene (5 mL), and the mixture was kept stirring for 10 min under an inert atmosphere. Then, 1 (40 mg, 0.078 mmol, 1.0 equiv), 2a (25 mg, 0.078 mmol, 1.0 equiv), and  $Cs_2CO_3$  (31 mg, 0.094 mmol, 1.2 equiv) were added, and the resulting suspension was heated up to 80 °C. After 16 h, the suspension was diluted with  $CH_2Cl_2$  (10 mL) and washed with  $H_2O$  $(2 \times 10 \text{ mL})$  and brine  $(10 \text{ mL})$ . The combined organic layers were dried over  $Na<sub>2</sub>SO<sub>4</sub>$  and concentrated under vacuum. The crude product was purified by flash column chromatography  $(SiO<sub>2</sub>,$ petroleum ether/CH<sub>2</sub>Cl<sub>2</sub> 2:1 to 1:1) to yield product 4a (20 mg, 0.027 mmol, 34%) as a deep purple solid.  ${}^{1}H$  NMR (CDCl<sub>3</sub>, 400 MHz): *δ* 7.65 (d, *J* = 8.6 Hz, 2 H), 7.38 (d, *J* = 7.0 Hz, 1 H), 7.27− 7.35 (m, 4 H), 7.23 (s, 1 H), 7.03−7.20 (m, 8 H), 6.65 (d, *J* = 7.0 Hz, 1 H), 4.39 (d, *J* = 5.6 Hz, 2 H), 4.36 (d, *J* = 5.6 Hz, 2 H), 1.76−1.85 (m, 2 H), 1.42−1.63 (m, 8 H), 1.31−1.38 (m, 8 H), 0.95−1.02 (m, 6 H), 0.87−0.95 (m, 6 H) ppm. 13C NMR (CDCl3, 100 MHz): *δ* 150.3, 146.6, 145.3, 145.0, 143.1, 131.4, 131.2, 130.0, 129.8, 129.7, 127.8, 127.5, 126.0, 125.8, 124.6, 121.6, 119.1, 114.5, 79.1, 78.9, 40.5, 30.4, 30.3, 29.2, 29.1, 23.9, 23.8, 23.1, 14.2, 11.3 ppm. ESI-MS: *m*/*z* calcd for  $C_{44}H_{51}NO_6S_2$  [M + H<sup>+•</sup>], 754.3; found, 754.3.

*2.2.4. 4-(6-(4-(Diphenylamino)phenyl)-4,8-bis((2-ethylhexyl) oxy)-1,1,5,5-tetraoxidobenzo[1,2-b:4,5-b*′*]dithiophen-2-yl) benzonitrile (BDT-H1)*. In a Schlenk tube,  $Pd_2(dba)$ <sub>3</sub> (1.4 mg, 0.0013 mmol, 0.05 equiv), P( $o$ -MeOPh)<sub>3</sub> (2 mg, 0.0054 mmol, 0.2 equiv), and pivalic acid (1 mg, 0.008 mmol, 0.3 equiv) were dissolved in dry toluene (5 mL), and the mixture was stirred for 10 min under an inert atmosphere. Then, compound 4a (20 mg, 0.027 mmol, 1.0 equiv), 4 bromobenzonitrile (3, 6 mg, 0.032 mmol, 1.2 equiv), and  $Cs_2CO_3$  (10 mg, 0.032 mmol, 1.2 equiv) were added, and the resulting suspension was heated up to 110 °C and stirred under an inert atmosphere for 16 h. The suspension was diluted with  $CH_2Cl_2$  (10 mL) and washed with  $H<sub>2</sub>O$  (2 × 10 mL) and brine (10 mL). The combined organic layers were dried over  $Na<sub>2</sub>SO<sub>4</sub>$  and concentrated under vacuum. The crude product was purified by flash column chromatography  $(SiO<sub>2</sub>)$ petroleum ether/CH<sub>2</sub>Cl<sub>2</sub> 2:1 to 1:1) to yield product BDT-H1 (6 mg, 0.007 mmol, 26%) as a deep purple solid. <sup>1</sup>H NMR (CDCl<sub>3</sub>, 400 MHz): *δ* 7.91 (d, *J* = 8.6 Hz, 2 H), 7.77 (d, *J* = 8.2 Hz, 2 H), 7.65 (d, *J* = 8.6 Hz, 2 H), 7.55 (s, 1 H), 7.29−7.35 (m, 4 H), 7.24 (s, 1 H), 7.05−7.18 (m, 8 H), 4.46 (d, *J* = 5.5 Hz, 2 H), 4.39 (d, *J* = 5.5 Hz, 2

H), 1.82−1.91 (m, 2 H), 1.54−1.65 (m, 8 H), 1.32−1.44 (m, 8 H), 1.01 (t, *J* = 7.4 Hz, 3 H), 1.00 (t, *J* = 7.4 Hz, 3 H), 0.88−0.98 (m, 6 H) ppm. <sup>13</sup>C NMR (CDCl<sub>3</sub>, 100 MHz): δ 150.4, 146.4, 145.6, 144.8, 143.4, 143.1, 139.9, 133.0, 131.2, 130.5, 129.6, 129.0, 128.4, 127.8, 127.0, 125.8, 125.1, 124.6, 121.6, 121.4, 118.7, 118.1, 114.2, 79.1, 79.0, 40.4, 30.3, 29.0, 23.7, 23.0, 14.1, 11.2 ppm. ESI-MS: *m*/*z* calcd for  $C_{51}H_{54}N_2O_6S_2$  [M<sup>+•</sup>], 854.3; found, 854.1.

*2.2.5. 4-(6-(4-(Diphenylamino)phenyl)-4,8-bis((2-ethylhexyl) oxy)-1,1,5,5-tetraoxidobenzo[1,2-b:4,5-b*′*]dithiophen-2-yl) benzonitrile (BDT-H1)*�*One-Pot Procedure.* In a Schlenk tube,  $Pd_2(dba)$ <sub>3</sub> (10 mg, 0.009 mmol, 0.05 equiv),  $P(o-MeOPh)$ <sub>3</sub> (6 mg, 0.018 mmol, 0.10 equiv), and pivalic acid (6 mg, 0.054 mmol, 0.3 equiv) were dissolved in dry toluene (5 mL), and the resulting mixture was stirred for 10 min under an inert atmosphere. Then, 1 (94 mg, 0.18 mmol, 1.0 equiv), 4-bromophenyl(diphenyl)amine (2a, 60 mg, 0.18 mmol, 1.0 equiv), and  $Cs_2CO_3$  (72 mg, 0.22 mmol, 1.2 equiv) were added, and the resulting suspension was heated up to 80 °C. After the aryl halide was consumed according to a TLC check (approx. 16 h of reaction, eluent petroleum ether/ $CH_2Cl_2$  2:1 to 1:1), a second aliquot of  $Cs_2CO_3$  (72 mg, 0.22 mmol, 1.2 equiv) was added to the mixture, followed by 4-bromobenzonitrile (3, 50 mg, 0.28 mmol, 1.5 equiv). The temperature was raised to 110 °C, and the reaction was stirred under an inert atmosphere for an additional 16 h. Then, the suspension was diluted with  $CH_2Cl_2$  (10 mL) and washed with H<sub>2</sub>O ( $2 \times 10$  mL) and brine ( $10$  mL). The combined organic layers were dried over  $Na<sub>2</sub>SO<sub>4</sub>$  and concentrated under vacuum. The residue was purified by flash column chromatography  $(SiO<sub>2</sub>)$ petroleum ether/CH<sub>2</sub>Cl<sub>2</sub> 2:1 to 1:1) to yield product  $\overrightarrow{BDT-H1}$  (43 mg, 0.05 mmol, 28%) as a deep purple solid. The spectroscopic characterization was identical to that of the product obtained via the two-step procedure (see above, paragraph 2.2.4).

*2.2.6. 2-(4-(Bis(4-methoxyphenyl)amino)phenyl)-4,8-bis((2 ethylhexyl)oxy)benzo[1,2-b:4,5-b*′*]dithiophene 1,1,5,5-Tetraoxide (4b)*. In a Schlenk tube,  $Pd_2(dba)_3$  (3 mg, 0.003 mmol, 0.05 equiv),  $P(o$ -MeOPh)<sub>3</sub> (4 mg, 0.012 mmol, 0.2 equiv), and pivalic acid (2 mg, 0.018 mmol, 0.3 equiv) were dissolved in dry toluene (5 mL), and the mixture was kept stirring for 10 min under an inert atmosphere. Then, 1 (30 mg, 0.06 mmol, 1.0 equiv), 2b (23 mg, 0.06 mmol, 1.0 equiv), and  $Cs_2CO_3$  (23 mg, 0.07 mmol, 1.2 equiv) were added, and the resulting suspension was heated up to 80 °C. After 16 h, the suspension was diluted with  $\mathrm{CH_2Cl_2}$  (10 mL) and washed with  $\mathrm{H_2O}$  $(2 \times 10 \text{ mL})$  and brine (10 mL). The combined organic layers were dried over  $Na<sub>2</sub>SO<sub>4</sub>$  and concentrated under vacuum. The crude product was purified by flash column chromatography  $(SiO<sub>2</sub>)$ , petroleum ether/CH<sub>2</sub>Cl<sub>2</sub> 2:1 to 1:1) to yield product 4b (19 mg, 0.023 mmol, 38%) as a deep purple solid.  ${}^{1}H$  NMR (CDCl<sub>3</sub>, 400 MHz): *δ* 7.59 (d, *J* = 8.4 Hz, 2 H), 7.37 (d, *J* = 6.9 Hz, 1 H), 7.15 (s, 1 H), 7.02−7.13 (m, 4 H), 6.80−7.00 (m, 6 H), 6.64 (d, *J* = 6.9 Hz, 1 H), 4.38 (d, *J* = 5.4 Hz, 2 H), 4.34 (d, *J* = 5.4 Hz, 2 H), 3.81 (s, 6 H), 1.76−1.85 (m, 2 H), 1.46−1.63 (m, 8 H), 1.29−1.41 (m, 8 H), 0.98 (t, *J* = 7.4 Hz, 3 H), 0.97 (t, *J* = 7.4 Hz, 3 H), 0.87−0.93 (m, 6 H) ppm. 13C NMR (CDCl3, 100 MHz): *δ* 157.0, 151.1, 145.3, 144.9, 143.5, 139.3, 131.4, 131.1, 130.7, 130.2, 129.8, 129.1, 128.5, 127.7, 125.6, 118.9, 115.0, 113.3, 79.1, 78.9, 55.6, 40.5, 31.1, 30.4, 29.2, 23.8, 23.2, 14.2, 11.3 ppm. ESI-MS:  $m/z$  calcd for  $C_{46}H_{56}NO_8S_2$  [M + H<sup>+</sup>• ], 814.3; found, 814.2.

*2.2.7. 4-(6-(4-(Bis(4-methoxyphenyl)amino)phenyl)-4,8-bis((2 ethylhexyl)oxy)-1,1,5,5-tetraoxidobenzo[1,2-b:4,5-b*′*]dithiophen-2 yl)benzonitrile* (*BDT-O1*). In a Schlenk tube,  $Pd_2(dba)_3$  (6 mg, 0.006 mmol, 0.05 equiv), P( $o$ -MeOPh)<sub>3</sub> (8 mg, 0.024 mmol, 0.2 equiv), and pivalic acid (4 mg, 0.035 mmol, 0.3 equiv) were dissolved in dry toluene (10 mL), and the mixture was stirred for 10 min under an inert atmosphere. Then, compound 4b (94 mg, 0.12 mmol, 1.0 equiv), 4-bromobenzonitrile (3, 33 mg, 0.18 mmol, 1.5 equiv), and  $Cs<sub>2</sub>CO<sub>3</sub>$  (16 mg, 0.048 mmol, 2.4 equiv) were added, and the resulting suspension was heated up to 110 °C and stirred under an inert atmosphere for 16 h. Then, the suspension was diluted with  $CH_2Cl_2$  (10 mL) and washed with H<sub>2</sub>O (2  $\times$  10 mL) and brine (10 mL). The combined organic layers were dried over  $Na<sub>2</sub>SO<sub>4</sub>$  and concentrated under vacuum. The crude product was purified by flash

column chromatography (SiO<sub>2</sub>, petroleum ether/CH<sub>2</sub>Cl<sub>2</sub> 2:1 to 1:1) to yield product BDT-O1 (80 mg, 0.088 mmol, 73%) as a deep purple solid. <sup>1</sup> H NMR (CDCl3, 400 MHz): *δ* 7.90 (d, *J* = 8.6 Hz, 2 H), 7.75−7.78 (m, 2 H), 7.72 (s, 1 H), 7.58−7.66 (m, 4 H), 7.54 (s, 1 H), 7.38−7.48 (m, 4), 6.86−6.92 (m, 4 H), 4.46 (d, *J* = 5.5 Hz, 2 H), 4.37 (d, *J* = 5.5 Hz, 2 H), 3.84 (bs, 6 H), 1.81−1.90 (m, 2 H), 1.48− 1.64 (m, 8 H), 1.33−1.42 (m, 8 H), 0.98−1.01 (m, 6 H), 0.90−0.94 (m, 6 H) ppm. <sup>13</sup>C NMR (CDCl<sub>3</sub>, 100 MHz): δ 145.6, 143.3, 139.8, 134.9, 134.8, 132.9, 131.3, 130.5, 129.0, 128.4, 127.8, 127.6, 127.0, 125.4, 124.7, 121.6, 118.7, 118.1, 115.0, 114.9, 114.8, 113.6, 113.0, 79.1, 79.0, 55.5, 40.4, 30.3, 29.7, 29.0, 23.7, 23.0, 14.1, 11.2 ppm. ESI-MS:  $m/z$  calcd for  $C_{53}H_{58}N_2O_8S_2$  [M<sup>+</sup>\*], 914.4; found, 914.1.

*2.2.8. 2-(4-(Bis(4-(methylthio)phenyl)amino)phenyl)-4,8-bis((2 ethylhexyl)oxy)benzo[1,2-b:4,5-b*′*]dithiophene 1,1,5,5-Tetraoxide (4c)*. In a Schlenk tube,  $Pd_2(dba)_3$  (5 mg, 0.005 mmol, 0.05 equiv), P( $o$ -MeOPh)<sub>3</sub> (7 mg, 0.02 mmol, 0.2 equiv), and pivalic acid (3 mg, 0.03 mmol, 0.3 equiv) were dissolved in dry toluene (5 mL), and the mixture was stirred for 10 min under an inert atmosphere. Then, 1 (50 mg, 0.1 mmol, 1.0 equiv), 2c (42 mg, 0.1 mmol, 1.0 equiv), and  $Cs<sub>2</sub>CO<sub>3</sub>$  (65 mg, 0.2 mmol, 2.0 equiv) were added, and the resulting suspension was heated up to 80 °C and stirred for 16 h. Then, the suspension was diluted with  $CH_2Cl_2$  (10 mL) and washed with  $H_2O$  $(2 \times 10 \text{ mL})$  and brine  $(2 \times 10 \text{ mL})$ . The combined organic layers were dried over  $Na<sub>2</sub>SO<sub>4</sub>$  and concentrated under vacuum. The residue was purified by flash column chromatography  $(SiO<sub>2</sub>,$  petroleum ether/CH<sub>2</sub>Cl<sub>2</sub> 2:1 to 1:2) to yield product 4c (31 mg, 0.036 mmol, 36%) as a deep purple solid. <sup>1</sup> H NMR (CDCl3, 400 MHz): *δ* 7.63 (d, *J* = 9.0 Hz, 2 H), 7.37 (d, *J* = 6.6 Hz, 1 H), 7.22 (s, 1 H), 7.20 (d, *J* = 8.6 Hz, 4 H), 7.03−7.08 (m, 6 H), 6.66 (d, *J* = 7.0 Hz, 1 H), 4.38 (d, *J* = 5.5 Hz, 2 H), 4.35 (d, *J* = 5.5 Hz, 2 H), 2.48 (s, 6 H), 1.76−1.85 (m, 2 H), 1.43−1.62 (m, 8 H), 1.31−1.39 (m, 8 H), 0.98 (t, *J* = 7.4 Hz, 3 H), 0.97 (t, *J* = 7.4 Hz, 3 H), 0.87−0.93 (m, 6 H) ppm. 13C NMR (CDCl3, 100 MHz): *δ* 149.8, 145.2, 144.9, 143.7, 142.9, 134.1, 131.2, 131.0, 129.8, 129.7, 128.2, 127.8, 127.4, 126.0, 125.9, 121.3, 119.0, 114.5, 79.0, 78.9, 40.3, 30.3, 30.2, 29.01, 28.96, 23.7, 23.6, 23.0, 16.4, 14.10, 14.07, 11.1 ppm. ESI-MS:  $m/z$  calcd for C<sub>46</sub>H<sub>55</sub>NO<sub>6</sub>S<sub>4</sub>  $[M^+$ ], 845.3; found, 845.0.

*2.2.9. 4-(6-(4-(Bis(4-(methylthio)phenyl)amino)phenyl)-4,8-bis- ((2-ethylhexyl)oxy)-1,1,5,5-tetraoxidobenzo[1,2-b:4,5-b*′*] dithiophen-2-yl)benzonitrile (BDT-S1)*. In a Schlenk tube, Pd<sub>2</sub>(dba)<sub>3</sub> (6 mg, 0.006 mmol, 0.05 equiv), P(o-MeOPh)<sub>3</sub> (8 mg, 0.024 mmol, 0.20 equiv), and pivalic acid (4 mg, 0.036 mmol, 0.3 equiv) were dissolved in dry toluene (10 mL), and the mixture was stirred for 10 min under an inert atmosphere. Then, 4c (100 mg, 0.12 mmol, 1.0 equiv), 4-bromobenzonitrile (3, 88 mg, 0.48 mmol, 0.4 equiv), and  $Cs<sub>2</sub>CO<sub>3</sub>$  (78 mg, 0.24 mmol, 2.0 equiv) were added, and the resulting suspension was heated up to 110 °C and stirred under an inert atmosphere for 16 h. Then, the suspension was diluted with  $CH_2Cl_2$ (10 mL) and washed with H<sub>2</sub>O (2  $\times$  10 mL) and brine (2  $\times$  10 mL). The combined organic layers were dried over  $Na<sub>3</sub>SO<sub>4</sub>$  and concentrated under vacuum. The residue was purified by flash column chromatography (SiO<sub>2</sub>, petroleum ether/CH<sub>2</sub>Cl<sub>2</sub> 2:1 to 1:3) to yield product BDT-S1 (80 mg, 0.085 mmol, 70%) as a deep purple solid. <sup>1</sup>H NMR (CDCl<sub>3</sub>, 400 MHz): *δ* 7.91 (d, *J* = 8.6 Hz, 2 H), 7.77 (d, *J* = 8.6 Hz, 2 H), 7.61−7.69 (m, 2 H), 7.55 (s, 1 H), 7.17−7.25 (m, 5 H), 6.95−7.13 (m, 6 H), 4.45 (d, *J* = 5.5 Hz, 2 H), 4.38 (d, *J* = 5.5 Hz, 2 H), 2.51 (bs, 6 H), 1.81−1.91 (m, 2 H), 1.50−1.65 (m, 8 H), 1.33−1.41 (m, 8 H), 1.00 (t, *J* = 7.4 Hz, 3 H), 0.99 (t, *J* = 7.4 Hz, 3 H), 0.88–0.95 (m, 6 H) ppm. <sup>13</sup>C NMR (CDCl<sub>3</sub>, 100 MHz): δ 145.6, 144.8, 143.4, 139.9, 132.9, 131.3, 131.2, 130.9, 130.5, 129.7, 128.9, 128.4, 128.0, 127.9, 127.7, 126.9, 126.2, 125.4, 125.2, 121.5, 118.1, 113.7, 108.5, 79.1, 78.9, 40.3, 30.2, 29.0, 23.7, 23.1, 16.3, 14.1, 11.2 ppm. ESI-MS:  $m/z$  calcd for  $C_{53}H_{58}N_2O_6S_4$  [M<sup>+•</sup>], 946.3; found, 946.3.

**2.3. Femtosecond Transient Absorption Spectroscopy Experiments.** The apparatus used for the transient absorption spectroscopy (TAS) measurements has been described in detail before. Briefly, 40 fs pulses centered at 800 nm were produced by an integrated Ti/sapphire oscillator (Micra-Coherent) coupled with a regenerative amplifier system (Legend-Coherent). The excitation

wavelength was set at 500 or 400 nm, and excitation power was set at about 200 nJ for all measurements. Visible pulses at 500 nm were generated by frequency mixing the idler and signal output produced by pumping a commercial optical parametric amplifier (TOPAS, Light Conversion) with a portion of the fundamental 800 nm radiation in a *β*-barium borate (BBO) crystal. Excitation pulses at 400 nm were obtained by the second harmonic generation of the fundamental laser radiation using a 2 mm BBO crystal. The pump beam polarization has been set to magic angle with respect to the probe beam by rotating a *λ*/2 plate to exclude rotational contributions.<sup>[30](#page-17-0)−[32](#page-17-0)</sup> The white light probe pulse was generated by focusing a small portion of the fundamental laser radiation on a 3 mm thick  $CaF<sub>2</sub>$  window. A portion of the generated white light was sent to the sample through a different path and used as a reference signal. After passing through the sample, the white light probe and reference pulses were both directed to a flat field monochromator coupled to a home-made detector. Transient signals were acquired in a time interval spanning up to 1500 ps. The sample was contained in a 2 mm quartz cuvette, mounted on a movable holder in order to minimize photodegradation. Measurements were performed at room temperature. Concentrations were adjusted to an absorbance of 0.9−1.0 OD (for the respective optical path) at the absorption maximum which amounted to about 0.3−0.5 OD at the excitation wavelength. Before and after the measurements, the integrity of the sample was checked on a PerkinElmer LAMBDA 950 spectrophotometer. Data analysis has been performed applying a global analysis procedure<sup>[33](#page-17-0)</sup> using software GLOTARAN<sup>[34](#page-17-0)</sup> and employing a linear unidirectional kinetic scheme. Global analysis allows the simultaneous fit of all the measured wavelengths with a combination of exponential decay functions and retrieves the kinetic constants describing the dynamic evolution of the system and the corresponding spectral component, called evolution-associated difference spectra (EADS).

**2.4. LSC Preparation and Film Characterization.** Fluorophore/PMMA thin films were prepared by drop casting according to the following procedure. Chloroform solutions (1.5 mL) containing 60 mg of the polymer and the proper amount of the fluorophore to obtain concentrations in the range 0.4−2.8 wt % were poured on a 50  $\times$  50  $\times$  3 mm<sup>3</sup> optically pure glass substrate (Edmund Optics Ltd. BOROFLOAT window 50  $\times$  50 TS). The glass slides were cleaned with chloroform and immersed in 6 M HCl for at least 12 h, and were then rinsed with water, acetone, and isopropanol. The resulting LSCs were stored for 24 h to allow the solvent to evaporate completely. The film thickness was measured by a Starrett micrometer to be  $25 \pm 5$ *μ*m. After LSC characterization, the polymer films were carefully detached from the glass surface by immersing the LSC in water, stored in a desiccator, and then analyzed by means of absorption and emission spectroscopies.

UV/vis spectra were recorded on polymer films at room temperature with a Cary 5000 UV−Vis−NIR spectrophotometer (Agilent). The fluorescence spectra were measured on polymer films at room temperature with a Fluorolog-3 spectrofluorometer (Horiba Jobin-Yvon, Horiba Italy) equipped with a 450 W Xenon arc lamp and double-grating excitation and single-grating emission monochromators. Quantum yield measurements were carried out using an external integration sphere (Quanta-*ϕ* F-3029, Horiba), equipped with solid or liquid sample holder, and connected to the spectrofluorometer by optical fibers and a fiber-optics adaptor (FL-3000, Horiba). Photographs of the film emissions under illumination were obtained by placing them in a Dark Reader 46B transilluminator equipped with a 450 nm LED source (Clare Chemical Research).

**2.5. Optical Efficiency.** Internal  $(\eta_{int})$  and external  $(\eta_{ext})$  photon and device ( $η<sub>dev</sub>$ ) efficiencies were determined according to recently agreed protocols, and details are reported in the [Supporting](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) [Information.](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf)<sup>[35,36](#page-17-0)</sup>  $\eta_{int}$  and  $\eta_{ext}$  were measured by using a commercially available system (Arkeo, Cicci research *s.r.l.*, Grosseto, Italy) containing a complementary metal-oxide semiconductor (CMOS) based spectrometer with a symmetric Czerny−Turner optical bench connected to an integrating sphere as reported in a recent publication.<sup>[16](#page-16-0)</sup> As an illumination source, an ORIEL LCS-100 solar simulator 94011A S/N: 322 was utilized under controlled

<span id="page-5-0"></span>

Figure 2. Bond lengths (Å) and dihedral angles (degrees) of  $S_0$ - and  $S_1$  (in brackets)-optimized geometries of compounds BDT-H2, BDT-H1, and BDT-AA.



Figure 3. DFT (B3LYP/6-31G\*) ground-state FMOs of BDT-H2, H1, and AA in toluene, with indication of their computed energies (see [Table](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) [S1](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf)).

illumination (1 sun, AM 1.5G). For the determination of  $\eta_{\text{dev}}$ , two PV cells IXYS KXOB25-12  $\times$  1F (22  $\times$  7 mm,  $V_{oc}$  = 0.69 V,  $I_{sc}$  = 46.7 mA, FF > 70%, and PCE = 25%) were connected in series, and the current/voltage characteristics were determined with a precision source/measure unit (B2900 Series, Keysight Technologies). Silicon was used to grease the LSC edge. The other three edges of the LSC were covered with a reflective aluminum tape. A black matte layer was placed beneath the LSC with an air gap of about 2.5 mm during the measurements.

#### **3. RESULTS AND DISCUSSION**

**3.1. Design of the Emitters.** Before delving into the description of the emitters' structures, we point out that, from here on, fluorophore TPA-BDTO will be denoted as BDT-H2 so that a homogeneous nomenclature can be used across the entire series of compounds, facilitating the immediate identification of their structural differences.

The first set of structures was designed based on a symmetric architecture. Considering the marked electronwithdrawing character of the central BDT-tetraoxide core, attaching two electron-donating triarylamine (TAA) moieties on both of its sides gives rise to a D−A−D arrangement. We reasoned that the electron-donating ability of the TAA group could be modulated by the introduction of different substituents on the 4′-position of the terminal phenyl rings, namely, a methoxy- or a thiomethyl-group, giving compounds BDT-O2 and BDT-S2, respectively ([Figure](#page-1-0) 1, left). As mentioned earlier, D−A−D compounds of this kind were expected to give rise to an intramolecular charge transfer (ICT) upon photoexcitation, helping achieve a polar excited

state with emission in the red/NIR region and a large Stokes shift.

We also envisioned a second array of compounds based on a non-symmetric structure, in which the same donor groups were introduced only on one side of the molecules, while on the other an additional electron-withdrawing cyanobenzene unit was placed, yielding a D−A−A′ architecture (BDT-H1,O1,S1, [Figure](#page-1-0) 1, right). Such arrangement was investigated because it was reported to enhance the bathochromic shift of the emission, while retaining moderate-to-good FQY, at least in less polar solvents. $37,38$  Furthermore, although nonsymmetric emitters have been previously employed in  $\text{LSCs},$ <sup>[39](#page-17-0)−[42](#page-17-0)</sup> to the best of our knowledge a direct comparison between the performances of devices built with symmetric and non-symmetric analogues, measured under the same conditions, is still missing, probably due to the more complicated synthetic procedures usually required to prepare the latter.

Finally, to check the effect of the foreseen ICT on the compound properties, a further symmetric emitter (named, BDT-AA, [Figure](#page-1-0) 1, bottom) was also designed as a reference, i.e., by introducing two cyanobenzene units at the sides of the central BDT-tetraoxide system. Indeed, its peculiar A′−A−A′ arrangement should not be capable of yielding ICT transitions, thus providing a potentially interesting comparison to the other compounds featuring alternating donor−acceptor moieties.

**3.2. Computational Investigation.** To assess if the photophysical properties of the designed compounds could be suitable for their employment in LSCs, we carried out a computational investigation based on density functional theory

<span id="page-6-0"></span> $(DFT)^{43,44}$  $(DFT)^{43,44}$  $(DFT)^{43,44}$  and time-dependent DFT  $(TD-DFT)^{45,46}$  $(TD-DFT)^{45,46}$  $(TD-DFT)^{45,46}$  methods, using Gaussian 16, Revision C.01 suite of programs.<sup>47</sup> The  $S_0$ -optimized geometries have been obtained at the B3LYP/6- $31G^*$  level of theory<sup>[48,49](#page-17-0)</sup> in vacuo, while the S<sub>1</sub>-optimized geometries have been computed at the TD-CAM-B3LYP/6-  $31G^*$  level of theory<sup>[50](#page-17-0)</sup> including the effects of the chosen solvent, toluene ([Figure](#page-5-0) 2, where only those of compounds BDT-H2, BDT-H1, and BDT-AA are shown for brevity; for the complete series, see [Figure](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S1, Supporting Information).

In the ground state, dihedral angles of 21−26° were found between the central oxidized BDT core and the flanking phenyl rings of all compounds, leading to a partial loss of co-planarity of their structures. This can explain the relatively small differences observed in the experimental absorption spectra of the compounds in different solvents (see below) as the torsion limits the dipole moment of the molecules in the ground state.<sup>38</sup> As already observed for other classes of emitters,<sup>[15](#page-16-0)</sup> in the first excited state the torsional angles were reduced to values comprised between 6 and 17° and thus were expected to favor the electronic delocalization across the molecules, which should be visible in the corresponding emission spectra.

The ground-state energies of frontier molecular orbitals (FMOs) and their electron density distributions were assessed by means of DFT calculations at the B3LYP/6-31G\* level of theory including the solvent effects ([Table](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S1 and [Figure](#page-5-0) 3, where, once again, only compounds BDT-H2, BDT-H1, and BDT-AA are shown; for the complete series, see [Figure](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S2). In terms of energy, the destabilizing effect of the stronger donor groups on the occupied orbitals (and in particular the HOMOs) is evident in both the symmetric and non-symmetric emitter series, while the effect on the LUMOs was less pronounced. Remarkably, the energy levels of non-symmetric compounds were all clearly stabilized due to the presence of the additional benzonitrile acceptor; such an effect was even more pronounced for compound BDT-AA featuring two lateral EWG groups.

Concerning the electron density distribution, in the case of the D−A−D compounds, the HOMOs were mostly localized on the lateral donor groups, with a minor contribution of the central core, while the LUMOs were centered on the acceptor BDT unit. A similar tendency was also observed for the nonsymmetric compounds, although in such cases, the HOMOs were mainly located on their single donor unit, while the LUMOs were widely delocalized over both the BDT central core and the additional acceptor. As expected, the frontier orbitals of BDT-AA did not present the same features since they all stretched along the entire conjugated scaffold of the molecule.

TD-MPW1K<sup>[51](#page-17-0)</sup>/6-311+G(2d,p) absorption maxima (*λ*cal abs,max), vertical excitation (*E*exc) energies, oscillator strengths (*f*), and composition (%) in terms of molecular orbitals for the lowest singlet–singlet excitations  $(S_0 \rightarrow S_1)$  were then calculated and are reported in Table 1. Solvent effects have been included by using the polarizable continuum model  $(PCM).$ <sup>52</sup>

For D−A−D and D−A−A<sup>'</sup> compounds,  $\lambda_{\rm cal}^{\rm abs,max}$  values were comprised between 496 and 525 nm, while a shorter wavelength was calculated for BDT-AA. Importantly, for all compounds the lowest energy excitations were largely associated with the HOMO−LUMO transitions (>80%). Given the overall FMOs electron density described above, this supported the hypothesis of an ICT photoexcitation process in the BDT series of compounds, except for BDT-AA,

Table 1. TD-MPW1K/6-311+G(2d,p) Absorption Maxima (*λ*cal abs,max in nm), Excitation Energies (*E*exc in eV), Oscillator Strengths (*f*), and Orbital Contributions (%) to the Lowest Energy Transition of BDT Compounds in Toluene Solution

compound	$\lambda_{\text{cal}}^{\text{abs,max}}$ $\lceil nm \rceil$	$E_{\text{exc}}$ [eV]	$\mathfrak{f}$	contribution $\lceil \% \rceil$
BDT-H <sub>2</sub>	501	2.48	1.64	85 $(H \rightarrow L)$
BDT-02	524	2.37	1.67	$86$ (H $\rightarrow$ L)
BDT-S2	521	2.38	1.70	83 $(H \rightarrow L)$
BDT-H1	496	2.50	1.27	82 $(H \rightarrow L)$
<b>BDT-01</b>	525	2.36	1.22	84 $(H \rightarrow L)$
BDT-S1	521	2.38	1.18	82 $(H \rightarrow L)$
<b>BDT-AA</b>	420	2.95	1.21	94 $(H \rightarrow L)$

whose frontier orbitals were evenly distributed along the molecule due to its different structure.

To evaluate the emission behavior of the compounds, the first excited-state FMO energies and spatial distributions were then assessed in toluene at the TD-CAM-B3LYP/6-31G\* level ([Table](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S2 and [Figure](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S3), revealing the same qualitative features already observed for the ground state, possibly with a slightly more pronounced delocalization, as a consequence of the more planar structures. Then, emission maxima ( $λ<sub>cal</sub><sup>emi,max</sup>$ ), vertical emission (*E*emi) energies, oscillator strengths (*f*), and composition (%) in terms of molecular orbitals for the lowest singlet−singlet emissions (S<sub>1</sub> → S<sub>0</sub>) were calculated, following a previously developed computational protocol, $53$  at the TD-MPW1K/6-311+G(2d,p) level, using the linear-response implementation (LR-PCM) (Table 2).

Table 2. TD-MPW1K/6-311+G(2d,p) Emission Maxima (*λ*cal emi,max in nm), Emission Energies (*E*emi in eV), Oscillator Strengths (*f*), and Orbital Contributions (%) to the Lowest Energy Transition of BDT Compounds in Toluene Solution

compound	1 emi, max $\lceil nm \rceil$	$E_{\text{emi}}$ [eV]	$\mathfrak{f}$	contribution $[\%]$
BDT-H <sub>2</sub>	644	1.92	1.66	92 $(L \rightarrow H)$
BDT-02	667	1.86	1.72	91 $(L \rightarrow H)$
BDT-S2	663	1.87	1.78	$86(L \rightarrow H)$
BDT-H1	626	1.98	1.46	90 $(L \rightarrow H)$
BDT-01	653	1.90	1.45	88 $(L \rightarrow H)$
BDT-S1	649	1.91	1.47	$86(L \rightarrow H)$
<b>BDT-AA</b>	571	2.17	1.23	96 $(L \rightarrow H)$

Donor−acceptor compounds provided computed maximum emission values well above 600 nm, in agreement with expectations, with a very pronounced effect of the more electron-donating side groups. Once again, except for BDT-AA, the lowest energy emissions were mostly associated with ICT transitions involving the two frontier orbitals (LUMO  $\rightarrow$ HOMO), in good agreement with the shifts in the maximum values and the solvatochromic effect observed experimentally (see below).

Based on the results of the computational investigation described above, we concluded that both the D−A−D and D− A−A′ series of BDT-containing compounds presented appropriate photophysical properties to work as emitters in LSCs, with emission in the red-orange part of the spectrum. Even BDT-AA, despite its different electronic properties, was computed to have a robust emission at around 570 nm, which made it potentially able to give devices with different color and light conversion ability. For this reason, all compounds

# <span id="page-7-0"></span>Scheme 1. Synthesis of the New Symmetric and Non-symmetric Emitters



described above were synthesized, and their photophysical properties were experimentally determined.

**3.3. Synthesis.** The synthesis of the new compounds started with the preparation of the oxidized central BDT core (1), which was carried out starting from commercially available benzo[1,2-*b*:4,5-*b*′]dithiophene-4,8-dione according to some previously reported procedures [\(Scheme](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S1, see Supporting Information for experimental details).

Since compound 1 is endowed with two oxidized thiophene rings with free 2-positions (Scheme 1), its further functionalization could be carried out by means of direct arylation procedures (differently from the original synthesis of TPA-BDTO, which was achieved by Suzuki-Miyaura crosscouplings).[18](#page-16-0) Indeed, direct arylation protocols were already

shown to be effective on this class of heterocycles<sup>[54](#page-18-0),[55](#page-18-0)</sup> and have been employed several times by our research group to prepare highly conjugated photoactive compounds,<sup>[56](#page-18-0)–[58](#page-18-0)</sup> allowing to shorten the synthetic sequences and reduce the amount of waste produced compared to traditional cross-coupling procedures.[59](#page-18-0)

Considering the symmetric compounds first (Scheme 1, top), central core 1 was thus simply reacted with an excess of the required bromides 2a−c or 3 in the presence of  $[\text{Pd}_{2}(\text{dba})_{3}]$  as the catalyst precursor,  $P(\text{2-MeOPh})_{3}$  as the ligand, PivOH as the acid additive, and  $Cs_2CO_3$  as the base in refluxing toluene for 16 h. The desired compounds of the BDT series were obtained in 56−70% yield, with the MeSsubstituted BDT-S2 and BDT-AA providing the best results.

In the case of the non-symmetric derivatives, the synthesis required two consecutive direct arylation steps with different bromides [\(Scheme](#page-7-0) 1, bottom). First, central core 1 was desymmetrized by insertion of the three different donor groups seen above. Reaction conditions were analogous to those employed in the preparation of the symmetric emitters, except for the use of a strictly stoichiometric amount of bromides and application of a lower temperature, to minimize double arylation. Then, the isolated intermediates 4a−c were reacted with an excess of 4-bromobenzonitrile (3) under similar conditions to provide the final products.

Interestingly, we could demonstrate that compound BDT-H1 could be directly prepared in a moderate yield of 28% from central core 1 by means of a one-pot process without isolation of intermediate 4a, as already shown by us for compounds with a different heterocyclic core.<sup>56</sup> During optimization of this protocol, it was found that the yield of BDT-H1 was limited by the need to stop the first arylation step before completion, as demonstrated by the concomitant formation of a certain amount of BDT-AA, clearly stemming from the double arylation of unreacted starting material 1 with 4-bromobenzonitrile (3).

Although some of the reactions described above proceeded with only moderate yields, all symmetric and non-symmetric compounds were obtained in quantities large enough for the subsequent characterization and device fabrication stages.

**3.4. Steady-State Spectroscopic Characterization in Solution.** The characterization of the new emitters started by recording the corresponding UV/vis absorption spectra in toluene solution (Table 3). This solvent was chosen because

# Table 3. Spectroscopic Properties of BDT-Series Compounds in Toluene Solution



<sup>*a*</sup>In good agreement with the literature data.<sup>[18](#page-16-0)</sup> <sup>*b*</sup>Absolute QY</sub> determined using an integrating sphere, see the [Experimental](#page-2-0) Section for details. <sup>*c*</sup> Stokes shifts. <sup>*d*</sup> For these compounds, characterized by low fluorescence QY and longer emission wavelengths, the *λ*<sub>max</sub> and Stokes shift values are likely overestimated as a result of the application of the correction function necessary to compensate the sensitivity loss of the photomultiplier in the red wavelength region.

its refractive index (1.496) is similar to that of PMMA (1.491), which is the most commonly used polymer in LSCs, thus providing a relevant reference for the prospected application. Symmetric D−A−D compounds [\(Figure](#page-9-0) 4a) exhibited relatively broad absorption features, peaked above 500 nm, which, in agreement with the results of the computational investigation (see above), were attributed to the ICT transition from the donor moiety to the acceptor core. In particular, while BDT-H2 presented a maximum absorption wavelength (*λ*max abs ) at 530 nm, substituted analogues displayed 11−25 nm red-shifted maxima and absorption onsets in the order of donor group strength  $(BDT-O2 > BDT-S2)$ , with very similar molar attenuation coefficients around  $5 \times 10^4$  M<sup>-1</sup> cm<sup>-1</sup>.

Symmetric compound BDT-AA presented a different absorption profile, with a maximum at 449 nm and a lower light harvesting ability, potentially related to its different photoexcitation process, as suggested above. As far as nonsymmetric analogues are concerned [\(Figure](#page-9-0) 4b), the absorption properties were almost identical to those of the symmetric compounds, the only noticeable difference being a generalized reduction of molar attenuation coefficients, not exceeding  $3 \times 10^4$  M<sup>-1</sup> cm<sup>-1</sup>. In general, the experimental findings were in reasonable agreement with the computed data, with differences in terms of  $E_{\text{exc}}$  smaller than 0.2 eV, and the same  $\lambda_{\text{max}}^{\text{abs}}$  trend was observed within the two series of compounds  $(H < S < 0)$ .

Regarding emission spectra, parent symmetric compound BDT-H2 presented a strong emission with maximum wavelength  $(\lambda_{\max}^{\text{emi}})$  well above 600 nm and high FQY  $(\Phi_{\text{f}})$  of 77%, in good agreement with the literature data [\(Figure](#page-9-0) 4c).<sup>[18](#page-16-0)</sup> The presence of a shoulder at longer wavelength suggested the presence of different vibronic transitions. Compounds BDT-O2 and BDT-S2 gave rise to red-shifted emissions compared to **BDT-H2**, with  $\lambda_{\text{max}}^{\text{emi}}$  differences of 56 and 39 nm, respectively, therefore larger than those found in the corresponding absorption spectra; moreover, the  $\Phi_f$  was much smaller, being comprised in the 31−38% range. These observations are once again consistent with the attainment of a CT excited state upon light absorption. Indeed, in the presence of a very strong acceptor group such as BDT, the increased electron-donor capacity of the lateral groups of BDT-S2 and, especially, BDT-O2 compared to BDT-H2 could further emphasize the degree of ICT of the  $S_0 \rightarrow S_1$  transition, with the consequent need of an extended structural reorganization of the excited state. The resulting energy loss due to vibrational relaxation could lead to the observed red shift and decrease of the quantum yield. Finally, compound BDT-AA presented a blue-shifted emission spectrum compared to the other symmetric species, accompanied by a good FQY of 66%.

The emission spectra of the non-symmetric derivatives presented some differences compared to those of the symmetric ones. First, compound BDT-H1 displayed a largely red-shifted transition in comparison to BDT-H2, accompanied by a slightly decreased  $\Phi_{\scriptscriptstyle{\theta}}$  probably as a result of the more pronounced CT character of the excited state imparted by the D−A−A′ architecture [\(Figure](#page-9-0) 4d).<sup>37</sup> Due to the very similar light absorption properties of the two compounds, this resulted in a larger Stokes shift for compound BDT-H1, which is potentially beneficial to reduce the amount of self-absorption losses in LSC devices. Unfortunately, by changing the donor group from simple triphenylamine to the methoxy- and thiomethyl-substituted analogues, the same tendency toward a reduction of FQY observed above for the symmetric compounds was evidenced. In this case, however, it was much more pronounced, resulting in an almost complete emission quenching for compounds BDT-O1 and BDT-S1 in toluene solution (Table 3). It must be pointed out that, due to their weak emission, the  $\lambda_{\rm max}^{\rm emi}$  values determined for these compounds are likely overestimated as a result of spectral broadening induced by the application of the correction function necessary to compensate for the sensitivity loss of the photomultiplier at long wavelengths. This is indirectly demonstrated by the large discrepancy observed between the experimental data and the computed emission maxima of BDT-O1,S1 (see [Table](#page-6-0) 2), while a much better agreement,

<span id="page-9-0"></span>

Figure 4. UV/vis absorption  $(a,b)$  and normalized fluorescence emission  $(c,d)$  spectra in toluene solution of the compounds prepared in this work.



Figure 5. (a,c) Transient absorption spectra registered for compound BDT-H1 in toluene and CH<sub>2</sub>Cl<sub>2</sub>, respectively [the peak at 500 nm in panel (c) is due to scattered pump light]; (b,d) EADS obtained from global analysis of the transient spectra reported in panel (a,c), respectively; and (e) kinetic traces at 532 nm registered for molecule BDT-H1 in toluene (hollow circles) and CH<sub>2</sub>Cl<sub>2</sub> (red squares). The continuous lines represent the fittings obtained by global analysis.

with differences in transition energies <0.2 eV, was found in all other cases.

To further probe the photophysical properties of the new compounds, the absorption and emission spectra of BDT-H2, BDT-H1, and BDT-AA, as typical representatives of each series of emitters, were measured in solvents of different polarities [\(Figure](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S4, [Table](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S3). In agreement with previous reports,<sup>38</sup> the absorption spectra were generally consistent in all solvents, and a trend toward a slight red shift of the absorption maxima with increasing solvent polarity could be recognized only for compounds BDT-H2 and BDT-H1, albeit with some exceptions (e.g., diethyl ether), possibly due to solubility reasons. The situation for the emission spectra was clearly different: as previously noted, BDT-H2 presented a strong red shift of the emission in more polar solvents, coupled with progressively larger Stokes shifts and decreasing FQY values, all indicative of an excited state with a significant CT character. In the case of BDT-H1, this behavior was even emphasized, producing an emission maximum close to 700 nm and a Stokes shift >150 nm even in diethyl ether. Regrettably, in more polar solvents such as THF or  $CH_2Cl_2$ , the  $\Phi_f$  values swiftly fell below 5%, causing the same issues seen above with the emission spectra of BDT-O1/S1 in toluene. Finally, in the case of BDT-AA, no clear shift of the spectra with solvent polarity was observed, confirming the localized nature of the electronic transitions undergone by this compound.

Based on the results of the photophysical characterization in solution, all compounds presenting moderate to good FQY in toluene, namely, all symmetric derivatives (BDT-H2/O2/S2/ AA) plus BDT-H1, were employed for the preparation of emissive PMMA films.

**3.5. TAS Studies.** Before starting the studies on the lightconcentrating devices, we carried out some TAS experiments with sub-picosecond time resolution on selected compounds, in order to investigate the relaxation dynamics of their excited states in solution and, possibly, to rationalize the different emission quantum yields observed for some of them as a function of solvent polarity (see above). The measurements were carried out both in toluene and  $CH_2Cl_2$  solution exciting the samples at 500 nm, with the exception of BDT-AA, whose excitation took place at 400 nm. To extract the kinetic constants describing the excited-state evolution of the samples, global analysis was used to interpret the data, employing a unidirectional linear decay scheme with three kinetic components (four in the case of BDT-AA). The analysis allows us to obtain both the excited-state lifetimes and the EADS, associated to each kinetic constant (see the [Experimental](#page-2-0) Section for details).

We begin our discussion with compound BDT-H1 since it did show the largest differences in  $\lambda_{\max}^{\text{emi}}$  and  $\Phi_{\text{f}}$  values in going from non-polar to polar solvents [\(Table](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S3). Its transient spectra in toluene and  $CH_2Cl_2$  are reported in [Figure](#page-9-0) 5, together with the EADS retrieved from global analysis.

In toluene, the transient spectrum of molecule BDT-H1 presents two negative bands peaked at 540 and 630 nm ([Figure](#page-9-0) 5a), which, by comparison with its steady-state absorption and emission spectra [\(Figure](#page-9-0) 4), can be assigned to ground-state bleaching (GSB) and stimulated emission (SE), respectively. Two excited-state absorption (ESA) bands can also be noted, one in the blue region of the investigated spectral interval (<470 nm) and the other peaked at about 750 nm. The comparison of the spectra measured for BDT-H1 with those of compounds BDT-H2 and BDT-AA (see below) allows us to assign the red-shifted ESA band to the presence of the nitrile-substituted benzene ring: this band is indeed observed in the case of BDT-AA but missing for BDT-H2.

The initial spectral component reported in [Figure](#page-9-0) 5b, with a lifetime of 0.85 ps (black line), represents the transient spectrum of the system immediately after excitation. It presents a very broad negative signal, ascribed to the convolution of GSB and SE of the unrelaxed  $S_1$  excited state. The system rapidly evolves toward the second spectral component (red line). Here, we notice a red shift of the SE band, whose minimum moves toward 625 nm. This spectral evolution is interpreted in terms of a rapid stabilization of the excited state, whose electronic distribution evolves from that of the initially excited Frank−Condon state toward the minimum of the potential energy surface of the  $S_1$  state. From there, a further evolution is observed to occur in 9.3 ps, characterized by an additional red shift of the SE band (blue line). This further evolution can be interpreted in terms of a stabilization of the excited state associated to the solvent reorganization and to vibrational cooling. The lifetime of this final spectral component is longer than the investigated time range (>1.5 ns), which is consistent with the good FQY of BDT-H1 in toluene.

The measurements were then repeated with BDT-H1 dissolved in  $CH_2Cl_2$ , in which a significant decrease of FQY was observed (see above). The transient spectra registered in that solvent ([Figure](#page-9-0) 5c) appear quite different from those measured in toluene. In this case, the SE band is not observed since an intense ESA band is present in the region between 550 and 750 nm. On the contrary, an intense bleaching band peaked at ca. 540 nm is observed also in this case, as well as a weak ESA band around 400 nm. Concerning the excited-state evolution, the initial EADS ([Figure](#page-9-0) 5d, black line) somehow recalls that observed in toluene since it presents an intense bleaching band, a very weak SE band peaked at about 630 nm, and a further ESA band in the red spectral region. This EADS rapidly evolves in about 0.8 ps toward the second spectral component (red line). Here, we observe the appearance of an intense ESA peaked at about 650 nm, which completely compensates the emission band. Such a notable spectral evolution indicates that the system moves toward an excited state different from that reached immediately after light absorption. By combining this and all the previous observations, it can be inferred that the excited state reached in about 0.8 ps has a very strong CT character, with the electron density being mostly relocated on the acceptor groups present in the molecule. The following evolution (from red to blue line), occurring in 5.9 ps, can be interpreted in terms of relaxation and stabilization of such a CT state, causing a significant energy loss. In agreement with the observed evolution, the excited-state lifetime is clearly reduced in  $CH<sub>2</sub>Cl<sub>2</sub>$  compared to toluene: in this case, the final spectral component (blue line) relaxes in about 870 ps. All these observations perfectly agree with the strong reduction of the FQY measured in  $CH_2Cl_2$  as compared to toluene. The difference in the excited-state lifetimes of BDT-H1 in the two solvents can also be noticed by comparing the kinetic traces measured on the maximum of the bleaching band, as reported in [Figure](#page-9-0) 5e.

We now turn our attention to the measurements conducted on compound BDT-H2. Indeed, while the appearance of the transient spectra of BDT-H2 in toluene is similar to those of BDT-H1, some significant differences emerge in  $CH_2Cl_2$ 



Figure 6. (a,c) Transient absorption spectra registered for compound BDT-H2 in toluene and CH<sub>2</sub>Cl<sub>2</sub>, respectively [the peak at 500 nm in panel (c) is due to scattered pump light]; (b,d) EADS obtained from global analysis of the transient spectra reported in panel (a,c), respectively; and (e) kinetic traces registered on the bleaching bands of compound BDT-H2 in toluene (hollow circles) and  $CH_2Cl_2$  (red squares). The continuous lines represent the fittings obtained by global analysis.

(Figure 6a,c) as it can also be noted by comparing the EADS obtained in the two solvents (Figure 6b,d). In toluene (Figure  $(6a,b)$ , besides the GSB (ca. 530 nm) and ESA band (<470) nm), a SE band is clearly visible also for **BDT-H2** at around 625 nm, and the evolution between the different spectral components takes place on a timescale similar to what previously observed for BDT-H1 in the same solvent [\(Figure](#page-9-0) [5](#page-9-0)b), including the survival of the last spectral component beyond the investigated time range. In  $CH_2Cl_2$  (Figure 6c,d), contrarious to the case of BDT-H1 [\(Figure](#page-9-0) 5c,d), the SE band is still visible, but it does indeed recover much faster compared to toluene because of the rise of an ESA band peaked in the same spectral region, on the picosecond timescale. Nevertheless, the excited-state lifetime in the two solvents is more similar in this case as compared to BDT-H1, as can be noticed by comparing the kinetic traces registered on the maximum of the corresponding bleaching signals (Figure 6e).

Clearly, also in the case of BDT-H2, this behavior can be explained by invoking, after the initial light absorption, the attainment of a low-lying CT excited state which gets more stabilized in the more polar solvent: however, for BDT-H2, the decay of this excited state seems to be much slower than for BDT-H1, probably due to its symmetric structure devoid of additional electron-withdrawing groups, that reduces the strength of its donor−acceptor character. This is in good agreement with its superior QY values in polar solvents.

As expected, in the case of BDT-AA, the same analysis gave rise to a different result since both the transient spectra, the EADS and the excited-state lifetimes, were found to be very similar in the two investigated solvents [\(Figure](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S5, where the GSB at ca. 450 nm and the SE at ca. 500−550 nm are clearly visible in both panels). This was in agreement with the less pronounced CT character of the excited state of BDT-AA and the relatively invariable QY values observed in different media. Finally, the same analysis was also conducted on compounds BDT-O2/S2, giving results qualitatively analogous to those obtained with BDT-H1/H2 [\(Figures](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S6 and S7). Nevertheless, it must be pointed out that for these compounds in  $CH<sub>2</sub>Cl<sub>2</sub>$  solution, the CT excited state was found to decay with much faster kinetics compared to the previous cases, and no sign of a SE band could be spotted after the initial state evolution, in agreement with the very pronounced donoracceptor character of these compounds conferred by their strong electron-donating groups. The decay time constants obtained for all molecules in the two solvents are summarized in [Table](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S4.

**3.6. Emitters Characterization in the PMMA Matrix.** Fluorophore-containing thin films were prepared employing PMMA, a widely used matrix in LSC systems. $60,61$  PMMA is 100% amorphous, transparent, cheap, and commercially available, making this polymer a perfect candidate for large-scale LSC applications.<sup>[62](#page-18-0)</sup> PMMA films with a thickness of 25  $\pm$ 5 *μ*m were obtained by drop-casting a solution of the polymer in  $CHCl<sub>3</sub>$ , varying the amount of dye between approximately 0.4 and 2.8% as a percentage of the total weight of the film (see the [Experimental](#page-2-0) Section for details).

In agreement with the optical properties in solution, all films appeared grayish to purple when illuminated with natural light, while their emission ranged from yellow to dark orange when put in a transilluminator equipped with a 450 nm light source ([Figure](#page-12-0) 7). In general, the molecules showed good dispersion in the films, which appeared smooth and did not present macroscopic phase separation. Nevertheless, some significant

<span id="page-12-0"></span>

Figure 7. Appearance of PMMA films (portion of  $2 \times 2 \text{ cm}^2$ ) doped with BDT fluorophores at 0.4−0.6 wt % concentration upon excitation in a transilluminator with a 450 nm light source.

differences emerged when analyzing their microscopic structures, as it will be highlighted below.

The films were characterized by means of UV/vis absorption and fluorescence emission spectroscopy. Their relevant properties at the lowest concentrations are reported in Table 4. In general, absorption and emission maxima were consistent

Table 4. Spectroscopic Properties of BDT Compounds in the PMMA Matrix at the Lowest Concentrations

compound	conc [wt $%$ ]	$\lambda_{\max}^{\text{abs}}$ [nm]	$\lambda_{\rm max}^{\rm emi}$ [nm]	SS [nm] $\{eV\}^a$
BDT-H <sub>2</sub>	0.6	533	656	$123 \{0.44\}$
BDT-O <sub>2</sub>	0.4	560	651	$91 \{0.31\}$
BDT-S2	0.4	542	633	$91 \{0.33\}$
BDT-H1	0.5	523	618	$95 \{0.36\}$
<b>BDT-AA</b>	0.4	447	550	103 {0.52}
<sup>a</sup> Stokes shifts.				

with the values obtained in toluene, resulting in significant Stokes shifts close or superior to 100 nm in all cases. On the other hand, differences in the behavior of the single emitters were observed depending on their concentration in the films, and thus they will be discussed individually. As a representative example, the absorption and emission spectra of BDT-H2 containing films are reported in Figure 8, while data for the other compounds are presented in the Supporting Information ([Figure](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S8).

As can be seen from Figure 8, the absorption intensity of BDT-H2-containing films regularly increased with the concentration, without any change of the spectral shape, denoting a good dispersion in the polymer. Regarding fluorescence, the intensity increased up to 1.7 wt %, then a partial quenching was observed for higher concentrations, accompanied by a slight red shift of the main peak; furthermore, the presence of a shoulder peak at higher energy could be spotted. This behavior is consistent both with the insurgence of re-absorption phenomena due to the increased

fluorophore concentration (inner filter effect) $^{63}$  $^{63}$  $^{63}$  and the possible formation of microscopic aggregates<sup>[64](#page-18-0)</sup> emitting at a shorter wavelength. The latter phenomenon was indeed confirmed by inspection of the corresponding epifluorescence microscopy images [\(Figure](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S9a), revealing the presence of a few fluorophore aggregates with sizes in the 30−40 *μ*m range. Despite that, no significant decrease in the efficiency of the corresponding LSC devices was later observed (see below).

Interestingly, symmetric D−A−D compounds BDT-O2 and BDT-S2 and non-symmetric BDT-H1 presented a qualitatively similar behavior to that of BDT-H2, albeit with some significant differences. First of all, no formation of microscopic aggregates was observed in the PMMA films of BDT-O2,S2 ([Figure](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S9b−d), possibly due to the substituents present on the 4-positions of the benzene rings of their donor groups, potentially altering their geometry in the solid state and hindering intermolecular interactions. Moreover, emissions of all these compounds appeared much weaker than that of BDT-H2, especially for BDT-O2, whose fluorescence intensity was even found to decrease progressively with fluorophore doping ([Figure](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S8).

On the other hand, the properties of BDT-AA-containing films were quite different from those of the other samples: already at the minimum concentration of 0.4 wt %, the absorption spectrum displayed two shoulder peaks at the sides of the main transition at approx. 447 nm, which became very evident above 1.2 wt % ([Figure](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S8). Based on the results of the epifluorescence microscopy experiments, this was attributed to the extensive formation of microscopic aggregates, resulting in an almost complete phase segregation between the fluorophore and the polymer matrix ([Figure](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S9e). Accordingly, a dramatic fall in emission intensity was observed at concentrations higher than 0.4 wt %, clearly due to significant light scattering by the microcrystalline aggregates.

To provide a quantitative assessment of the matching between the fluorophores absorption spectra and the emission of the solar simulator lamp used for LSC characterization (see below), the absorption efficiency parameter ( $\eta_{\text{abs}}$ ) in the 300− 800 nm range was calculated for all films at each concentration, according to the definition given by Debije et al. ([Figure](#page-13-0) 9a, see Supporting [Information](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) for details). $35$ 

All donor−acceptor compounds gave excellent *η*abs values close to or higher than 0.5 at their maximum concentration (a value of 1 indicating a perfect match), following an exponential trend vs maximum absorbance, as previously reported.<sup>65</sup> The best result was given by BDT-O2, due to its red-shifted and broader absorption band compared to the other compounds.



Figure 8. UV/vis absorption (a) and fluorescence emission (b) spectra of PMMA films containing compound BDT-H2 at different concentrations.

<span id="page-13-0"></span>

Figure 9. (a) Absorption efficiency in the 300−800 nm range and (b) FQYs of BDT compounds in the PMMA matrix. BDT-H2, red squares; BDT-O2, blue circles; BDT-S2, green upside triangles; BDT-H1, orange rhombs; and BDT-AA, purple downside triangles. In panel (a), fitting according to an exponential function is shown as a dotted line, see the Supporting [Information](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) for details.



Figure 10. External (a, full symbols) and internal (b, hollow symbols) photon efficiencies of LSCs built with BDT compounds. BDT-H2, red squares; BDT-O2, blue circles; BDT-S2, green upside triangles; BDT-H1, orange rhombs; and BDT-AA, purple downside triangles.

**BDT-H2**, on the other hand, gave slightly lower  $\eta_{\text{abs}}$  than the other emitters at the same maximum absorbance level, but the very high absorbance values of its films ensured an optimal light harvesting ability of the corresponding devices. As for **BDT-AA**, apparently, it gave high  $\eta_{\text{abs}}$  values at low maximum absorbance, but the analysis in this case was surely affected by the abovementioned aggregation phenomena, which caused excessive tailing of the spectrum at longer wavelengths due to scattering [\(Figure](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S8), leading to an overestimation of  $\eta_{\text{abs}}$ .

Finally, FQYs were measured for all fluorophore-doped films using an integrating sphere (see the [Experimental](#page-2-0) Section for details) and are shown in Figure 9b. In agreement with the measurements in toluene solution, the highest  $\Phi_f$  values were shown by compound BDT-H2, with a maximum of 55% at 0.6 wt %, which moderately decreased up to approx. 42% at the highest concentration, likely due to inner filter effects combined with the possible formation of less emissive supramolecular (amorphous) chromophoric aggregates.<sup>[66](#page-18-0)</sup> Although these values were inferior to that recorded in toluene (77%), they were still compatible with the fabrication of efficient LSC devices. Slightly lower quantum yields were obtained for emitter BDT-H1 and even BDT-AA, despite the abovementioned formation of micro-aggregates, denoting the excellent light emission properties of the latter compound. Unfortunately, as suggested by their emission spectra, the other symmetric D−A−D compounds presented much lower  $\Phi_f$ values, once again in agreement with results in solution (see above). In particular, the FQYs of BDT-O2 were always equal or inferior to 10% at all concentrations, suggesting that the corresponding LSC could present unsatisfying optical efficiencies.

**3.7. Performances of LSC Devices.** The performances of the fluorophore-containing polymer films as LSCs were first characterized by recording their external and internal photon efficiencies under simulated solar light (Figure 10, see Supporting [Information](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) for details).

The external photon efficiency  $(\eta_{\text{ext}})$  corresponds to the ratio between the output photon flux measured at the edges of the LSC with respect to the incident photon flux, and it allows us to evaluate both the light-harvesting ability of the LSC and the effectiveness of the waveguide propagation within the devices. As expected from the spectroscopic characterization, the best *η*ext figures were obtained with compound BDT-H2, which reached the maximum value of approx. 8.4% for the two films with the highest fluorophore concentrations. This was probably due to an optimal balance between the excellent light-absorption ability of BDT-H2 and the losses due to dissipative phenomena and light scattering, which are both enhanced at increasing fluorophore concentrations. This result suggests that the large Stokes shifts and small spectral overlaps of the BDT-series compounds can indeed be successfully exploited to maximize the device efficiency. The performance of **BDT-H2** in terms of  $η<sub>ext</sub>$  is remarkable, appearing superior to those of other organic donor−acceptor fluorophores recently reported both by us and other research groups.<sup>[16](#page-16-0),[42](#page-17-0),[65,67](#page-18-0)</sup>

The other compounds provided inferior results compared to **BDT-H2, but the general tendency toward higher**  $\eta_{\text{ext}}$  **values at** higher fluorophore concentration was confirmed, except for BDT-O2, for which a slight decrease was observed instead. Indeed, its overall performance was quite poor, clearly due to its very low FQY compared to the other compounds of the series. Interestingly, **BDT-H1** provided fair  $η<sub>ext</sub>$  values with a maximum of around 4.4%, but was still quite far from BDT-H2, despite its slightly higher  $η<sub>abs</sub>$  (see above). We can attribute this behavior to its much weaker light harvesting ability compared to BDT-H2, as evidenced by the lower absorbance values of the corresponding films (as compared in [Figure](#page-12-0) 8 and [Figure](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S8), together with its lower FQY and smaller Stokes shift in the polymer matrix [\(Table](#page-12-0) 4).

Then, to evaluate the photon transport process within the waveguide, the internal photon efficiency ( $η<sub>int</sub>$ ) was determined. This parameter can be obtained from the edge-emitted power spectra, calculating the ratio between the average output power emitted from the four edges and the fraction of photons effectively absorbed by the LSC (see the [Supporting](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) [Information](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) for details). Being related to the number of absorbed photons (rather than the total incident power, as  $\eta_{\text{ext}}$ ),  $\eta_{\text{int}}$  is a key parameter that allows evaluating in detail all lightguide losses taking place in the device. In almost all cases,  $\eta_{\text{int}}$  was maximized at the lowest emitter concentration [\(Figure](#page-13-0) [10b](#page-13-0)) and moderately decreased as the concentration increased. This is a typical trend often observed for LSCs and is usually consistent with that observed for the  $\Phi_f$  values, suggesting that a progressively larger fluorescence dissipation occurred within the waveguide. Interestingly, in this case, the only exception was **BDT-AA**, for which relatively constant  $η<sub>int</sub>$  values were observed across the entire concentration range. It is possible that, due to the abovementioned formation of microaggregates, intense light scattering occurs even at low concentrations, resulting in smaller  $η<sub>int</sub>$  values compared to BDT-H1 even with slightly better FQYs [\(Figure](#page-13-0) 9), but without worsening too much with concentration, thus explaining the relatively constant  $η<sub>int</sub>$  trend.

To assess the LSC performances in the direct light-toelectricity conversion, we determined the device efficiency (*η*dev) parameter, connecting two Si-PV cells in series to an edge of the thin-film LSC by using silicone grease and measuring the electric power generated by such device under simulated solar irradiation (Figure 11, see the [Experimental](#page-2-0) [Section](#page-2-0) for details).

The obtained parameter can be defined as the electrical power effectively extracted from the PV cells  $(P_{\rm el}^{\rm out})$  relative to the luminous power hitting the top surface of the LSC  $(P_{\rm opt}^{\rm in})$ and is conceptually analogous to the PCE of a PV cell (eq 1)



Figure 11. Device efficiencies of LSC built with BDT compounds. BDT-H2, red squares; BDT-O2, blue circles; BDT-S2, green upside triangles; BDT-H1, orange rhombs; and BDT-AA, purple downside triangles.

el out opt in

 $v = \frac{1}{p}$ 

$$
= \frac{P_{\text{el}}^{\text{out}}}{P_{\text{opt}}^{\text{in}}} = \frac{I_{\text{sc}} \cdot V_{\text{oc}} \cdot \text{ff}}{P_{\text{opt}}^{\text{in}} \cdot A_{\text{LSC}}}
$$
(1)

(1)

where ff,  $I_{\rm sc}$  and  $V_{\rm oc}$  are the fill factor, short-circuit current, and open-circuit voltage of the edge-mounted PV cells, respectively,  $A_{\rm LSC}$  is the front-illuminated area of the LSC device, and  $P_{\rm opt}^{\rm in}$  is the incident solar power density expressed in mW cm<sup>-2</sup>.

Results were in general agreement with the trends observed for the  $\eta_{\text{ext}}$  parameter, with the best performance once again provided by BDT-H2, whose device reached a maximum value of 0.94% at 2.2 wt %. The efficiency reached by BDT-H1 was also noteworthy, with a maximum value of 0.8% at the relatively high doping levels of 1.9−2.4 wt %. On the other hand, much lower figures could be gathered from the other compounds due to the above-described issues of low  $\Phi_f$  values and extensive light scattering, which effectively decreased the amount of photons hitting the solar cells and, consequently, their electricity production.

The best results obtained for all compounds together with the corresponding doping concentrations are reported in Table 5, alongside those obtained with state-of-the-art commercial

Table 5. Best Efficiency Parameters of the LSC Built with BDT Compounds and LR305 in the PMMA Matrix

compound	conc [wt $%$ ]	$\eta_{\rm ext}$ [%]	$\eta_{\text{int}}$ [%]	$\eta_{\text{dev}}$ [%]
BDT-H <sub>2</sub>	0.6		$40 \pm 4$	
	2.2	$8.4 \pm 0.1$		$0.94 \pm 0.06$
BDT-O2	0.4	$1.4 \pm 0.2$	$10 \pm 1$	$0.23 \pm 0.02$
BDT-S2	0.4		$20 \pm 4$	
	1.6	$3.1 \pm 0.6$		
	2.0			$0.44 \pm 0.03$
BDT-H1	0.5		$23 \pm 3$	
	2.4	$4.4 \pm 0.4$		$0.80 \pm 0.02$
<b>BDT-AA</b>	0.4		$17 \pm 2$	
	1.6	$1.8 \pm 0.2$		$0.32 \pm 0.03$
<b>LR305</b>	0.4		$50 \pm 7$	
	1.6			$1.0 \pm 0.1$
	2.0	$9.1 \pm 0.4$		

emitter Lumogen F Red 305 (LR305), which, compared to the compounds discussed in this study, presents not only a higher  $\Phi_f$  value but also a more extensive re-absorption.<sup>[3](#page-16-0),7</sup>

While the parameters extracted from the LSC built with LR305 were still the best among those measured, it is remarkable that BDT-H2 provided similar performances, especially in terms of  $\eta_{\text{dev}}$ . Even in the case of **BDT-H1**, with much lower photon efficiency values, the best  $\eta_{\text{dev}}$  value was only approx. 20% lower than that of LR305. This highlights how the design of the BDT-series emitters and the low re-absorption losses associated with their large Stokes shifts can generate an emission profile highly matched with the light absorption of silicon solar cells.

Finally, we also assessed the photostability of a BDT-H2 containing PMMA film with an accelerated test, consisting of continuous irradiation under UV light (350−420 nm) at a constant temperature of 70 °C for 650 min (potentially corresponding to a much longer period of time at room temperature, see the Supporting [Information](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) for details). The selected film was that containing 2.2 wt % of fluorophore as it provided the best results among all those tested in terms of *η*ext and  $η_{\text{dev}}$  (Table 5). The emission spectrum of the incident light is reported in [Figure](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S15 together with that of the AM 1.5G <span id="page-15-0"></span>solar spectrum. Its power density was integrated to be 38.43 W  $m^{-2}$  (95% peak area), corresponding to irradiation conditions of 1.15 sun, considering that in the same region the AM 1.5G spectrum presents an irradiance of 33.45 W  $\text{m}^{-2}$ . Under these conditions, the film demonstrated approx. 20% loss in maximum absorption intensity during the entire experiment ([Figure](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S16a), which was mirrored by a progressive reduction of the area of its emission profile [\(Figure](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf) S16b). Nevertheless, the general shape of the absorption spectrum was not significantly altered, and only a small blue shift of approx. 9 nm was observed, indicating that most of the fluorophore was not degraded in the accelerated aging test.

# **5. CONCLUSIONS**

In this paper, we have described the design, synthesis, and spectroscopic characterization of a series of organic fluorescent emitters bearing an electron-withdrawing benzo[1,2-*b*:4,5 *b*′]dithiophene 1,1,5,5-tetraoxide unit as their central core. The compounds had either a symmetric (D−A−D or A′−A− A′) or non-symmetric (D−A−A′) structure, characterized by the presence of different electron-donating or electronaccepting groups at the sides of the central heterocyclic system. A DFT and TD-DFT computational investigation revealed that compounds containing both donor and acceptor groups were expected to attain excited states characterized by a large degree of ICT upon photoexcitation, potentially yielding emissions in the red part of the spectrum accompanied by large Stokes shifts.

Accordingly, the optical properties of the emitters in solution were found to be critically dependent on their general structure and on the nature of the substituents: in particular, the presence of strong electron-donating groups connected to the central BDT-tetraoxide unit caused a significant red shift of the emission, which was also observed when going from symmetric to the corresponding non-symmetric structures. Such bathochromic shifts were regrettably accompanied by a more or less pronounced decrease of FQY, which in some cases prevented the fluorophores use in LSCs (BDT-O1,S1). TAS studies confirmed that, after the initial photoexcitation, the compounds underwent a fast transition to lower-lying ICT excited states, whose lifetime was largely dictated both by their structures and the surrounding environment. Thus, nonsymmetric compounds (BDT-H1), as well as compounds with stronger electron-donating groups (BDT-O2,S2), showed a much faster excited-state decay compared to BDT-H2, especially in more polar solvents, in good agreement with the corresponding  $\Phi_f$  trends.

All emitters presenting moderate-to-good FQY in toluene solution were dispersed in a PMMA matrix at different concentrations for LSC studies. In general, LSC performances were enhanced in the case of a strong emission accompanied by an efficient minimization of self-absorption phenomena, as it is typical in this kind of devices. Thus, despite its slightly lower light absorption efficiency, BDT-H2 yielded the most efficient concentrators, thanks to its higher  $\Phi_f$  and smaller overlap compared to the other compounds of the series. Remarkably, both the photonic ( $\eta_{\text{ext}}$  of 8.4  $\pm$  0.1%) and PV  $(\eta_{\text{dev}}$  of 0.94  $\pm$  0.06%) efficiency figures of LSC built with BDT-H2 were higher than those recently reported for several other organic emitters and were very close to those of state-ofthe-art devices fabricated with standard fluorophore LR305.

This study thus highlights that, in the case of organic emitters for LSCs, finding the right balance between the relative strengths of the connected donor and acceptor groups, as well as the most appropriate molecular architecture, is key to achieve the best compromise between all the relevant photophysical properties (light harvesting ability, maximum absorption wavelength, FQY, and Stokes shift), leading to solar collectors with optimized performances.

# ■ **ASSOCIATED CONTENT**

# $\bullet$  Supporting Information

The Supporting Information is available free of charge at [https://pubs.acs.org/doi/10.1021/acsaem.3c00362](https://pubs.acs.org/doi/10.1021/acsaem.3c00362?goto=supporting-info).

Computational analysis; synthetic procedure for the preparation of compound 1; copies of the  ${}^{1}H$  and  ${}^{13}C$ NMR spectra of all new compounds; additional spectroscopic characterization of the compounds in solution and in the PMMA matrix; TAS analysis; experimental details on the optical and PV efficiency measurements; and experimental details on the accelerated lifetime tests ([PDF\)](https://pubs.acs.org/doi/suppl/10.1021/acsaem.3c00362/suppl_file/ae3c00362_si_001.pdf)

# ■ **AUTHOR INFORMATION**

#### **Corresponding Authors**

- Andrea Pucci − *Institute of Chemistry of Organometallic Compounds (CNR-ICCOM), 50019 Sesto Fiorentino, Italy; Department of Chemistry and Industrial Chemistry, University of Pisa, 56124 Pisa, Italy;* Orcid.org/0000-[0003-1278-5004](https://orcid.org/0000-0003-1278-5004); Email: [andrea.pucci@unipi.it](mailto:andrea.pucci@unipi.it)
- Lorenzo Zani − *Institute of Chemistry of Organometallic Compounds (CNR-ICCOM), 50019 Sesto Fiorentino, Italy;* [orcid.org/0000-0003-0621-2648;](https://orcid.org/0000-0003-0621-2648) Email: [lorenzo.zani@](mailto:lorenzo.zani@iccom.cnr.it) [iccom.cnr.it](mailto:lorenzo.zani@iccom.cnr.it)

#### **Authors**

- Matteo Bartolini − *Institute of Chemistry of Organometallic Compounds (CNR-ICCOM), 50019 Sesto Fiorentino, Italy* Cosimo Micheletti − *Department of Chemistry and Industrial*
- *Chemistry, University of Pisa, 56124 Pisa, Italy* Alberto Picchi − *Department of Chemistry and Industrial Chemistry, University of Pisa, 56124 Pisa, Italy*
- Carmen Coppola − *Department of Biotechnology, Chemistry and Pharmacy, R2 ES Lab, University of Siena, 53100 Siena, Italy; CSGI, Consorzio per lo Sviluppo dei Sistemi a Grande Interfase, 50019 Sesto Fiorentino, Italy;* [orcid.org/0000-](https://orcid.org/0000-0002-8050-7615) [0002-8050-7615](https://orcid.org/0000-0002-8050-7615)
- Adalgisa Sinicropi − *Institute of Chemistry of Organometallic Compounds (CNR-ICCOM), 50019 Sesto Fiorentino, Italy; Department of Biotechnology, Chemistry and Pharmacy, R2 ES Lab, University of Siena, 53100 Siena, Italy; CSGI, Consorzio per lo Sviluppo dei Sistemi a Grande Interfase, 50019 Sesto Fiorentino, Italy;* [orcid.org/0000-0001-](https://orcid.org/0000-0001-5605-6482) [5605-6482](https://orcid.org/0000-0001-5605-6482)
- Mariangela Di Donato − *Institute of Chemistry of Organometallic Compounds (CNR-ICCOM), 50019 Sesto Fiorentino, Italy; LENS, European Laboratory for Non-Linear Spectroscopy, 50019 Sesto Fiorentino, Italy;* [orcid.org/0000-0002-6596-7031](https://orcid.org/0000-0002-6596-7031)
- Paolo Foggi − *LENS, European Laboratory for Non-Linear Spectroscopy, 50019 Sesto Fiorentino, Italy; Department of Chemistry, Biology and Biotechnology, University of Perugia, 06123 Perugia, Italy; National Institute of Optics (CNR-INO), 50019 Sesto Fiorentino, Italy*

<span id="page-16-0"></span>Alessandro Mordini − *Institute of Chemistry of Organometallic Compounds (CNR-ICCOM), 50019 Sesto Fiorentino, Italy; Department of Chemistry "U. Schiff", University of Florence, 50019 Sesto Fiorentino, Italy*

Gianna Reginato − *Institute of Chemistry of Organometallic Compounds (CNR-ICCOM), 50019 Sesto Fiorentino, Italy;* [orcid.org/0000-0002-7712-3426](https://orcid.org/0000-0002-7712-3426)

Massimo Calamante − *Institute of Chemistry of Organometallic Compounds (CNR-ICCOM), 50019 Sesto Fiorentino, Italy; Department of Chemistry "U. Schiff", University of Florence, 50019 Sesto Fiorentino, Italy*

Complete contact information is available at: [https://pubs.acs.org/10.1021/acsaem.3c00362](https://pubs.acs.org/doi/10.1021/acsaem.3c00362?ref=pdf)

#### **Author Contributions**

M.B., C.M., and A.P. contributed equally. The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

#### **Notes**

The authors declare no competing financial interest.

# ■ **ACKNOWLEDGMENTS**

We thank "COLOURS" project, funded by Regione Toscana (POR FESR 2014 − 2020, grant no. 3553.04032020.158000411), "SUNNYSIDE" project, funded by Fondazione Cassa di Risparmio di Firenze (grant no. 2020.1408), and project "nuovi Concetti, mAteriali e tecnologie per l'iNtegrazione del fotoVoltAico negli edifici in uno scenario di generazione diffuSa" ("CANVAS"), funded by the Italian Ministry of the Environment and the Energy Security, through the Research Fund for the Italian Electrical System (type-A call, published on G.U.R.I. n. 192 on 18-08- 2022), for financial support. This research was also funded by the European Union's Horizon 2020 Research and Innovation program under grant agreement no. 871124 Laserlab-Europe. S. Desbrousses is acknowledged for his assistance in the preparation of compound BDT-H2.

#### ■ **REFERENCES**

(1) Weber, W. H.; Lambe, J. [Luminescent](https://doi.org/10.1364/ao.15.002299) Greenhouse Collector for Solar [Radiation.](https://doi.org/10.1364/ao.15.002299) *Appl. Opt.* 1976, *15*, 2299−2300.

(2) Batchelder, J. S.; Zewai, A. H.; Cole, T. [Luminescent](https://doi.org/10.1364/ao.18.003090) Solar [Concentrators](https://doi.org/10.1364/ao.18.003090) 1: Theory of Operation and Techniques for Performance [Evaluation.](https://doi.org/10.1364/ao.18.003090) *Appl. Opt.* 1979, *18*, 3090−3110.

(3) Roncali, J. [Luminescent](https://doi.org/10.1002/aenm.202001907) Solar Collectors: Quo Vadis? *Adv. Energy Mater.* 2020, *10*, 2001907.

(4) Papakonstantinou, I.; Portnoi, M.; Debije, M. G. The [Hidden](https://doi.org/10.1002/aenm.202002883) Potential of Luminescent Solar [Concentrators.](https://doi.org/10.1002/aenm.202002883) *Adv. Energy Mater.* 2021, *11*, 2002883.

(5) Ferreira, R. A. S.; Correia, S. F. H.; Monguzzi, A.; Liu, X.; Meinardi, F. Spectral Converters for [Photovoltaics](https://doi.org/10.1016/j.mattod.2019.10.002) − What's Ahead. *Mater. Today* 2020, *33*, 105−121.

(6) Lee, S. H.; Baek, D.; Cho, W.; Lee, N.; Kim, K.; Kim, J.; Kim, H.; Kim, H. H.; Kim, H. J.; Lee, S.; Lee, S.-M. Tailoring [Luminescent](https://doi.org/10.1002/adfm.202210357) Solar Concentrators for [High-Performance](https://doi.org/10.1002/adfm.202210357) Flexible Double-Junction III-V [Photovoltaics.](https://doi.org/10.1002/adfm.202210357) *Adv. Funct. Mater.* 2023, *33*, 2210357.

(7) Li, Y.; Sun, Y.; Zhang, Y. Luminescent Solar [Concentrators](https://doi.org/10.1016/j.solener.2019.07.035) Performing under Different Light [Conditions.](https://doi.org/10.1016/j.solener.2019.07.035) *Sol. Energy* 2019, *188*, 1248−1255.

(8) Meinardi, F.; Bruni, F.; Brovelli, S. [Luminescent](https://doi.org/10.1038/natrevmats.2017.72) Solar Concentrators for [Building-Integrated](https://doi.org/10.1038/natrevmats.2017.72) Photovoltaics. *Nat. Rev. Mater.* 2017, *2*, 17072.

(9) Beverina, L.; Sanguineti, A. Organic Fluorophores for Luminescent Solar Concentrators. In *Solar Cell Nanotechnology*;

Tiwari, A., Boukherroub, R., Sharon, M., Eds.; Scrivener Publishing LLC: Beverly, MA, 2014; pp 317−356.

(10) Clarke, T. M.; Gordon, K. C.; Kwok, W. M.; Phillips, D. L.; Officer, D. L. Tuning from *π*,*π*\* to [Charge-Transfer](https://doi.org/10.1021/jp0600312?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as) Excited States in [Styryl-Substituted](https://doi.org/10.1021/jp0600312?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as) Terthiophenes: An Ultrafast and Steady-State [Emission](https://doi.org/10.1021/jp0600312?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as) Study. *J. Phys. Chem. A* 2006, *110*, 7696−7702.

(11) Kundu, R.; Kulshreshtha, C. Design, Synthesis and [Electronic](https://doi.org/10.1039/C5RA13416A) [Properties](https://doi.org/10.1039/C5RA13416A) of Push−Pull−Push Type Dye. *RSC Adv.* 2015, *5*, 77460− 77468.

(12) National Renewable Energy Laboratory (NREL). Reference Air Mass 1.5 Spectra. [https://www.nrel.gov/grid/solar-resource/spectra](https://www.nrel.gov/grid/solar-resource/spectra-am1.5.html)[am1.5.html](https://www.nrel.gov/grid/solar-resource/spectra-am1.5.html) (accessed Jan 10, 2023).

(13) Kim, H. U.; Kim, T.; Kim, C.; Kim, M.; Park, T. [Recent](https://doi.org/10.1002/adfm.202208082) Advances in Structural Design of Efficient [Near-Infrared](https://doi.org/10.1002/adfm.202208082) Light-Emitting Organic Small [Molecules.](https://doi.org/10.1002/adfm.202208082) *Adv. Funct. Mater.* 2023, *33*, 2208082.

(14) Papucci, C.; Geervliet, T. A.; Franchi, D.; Bettucci, O.; Mordini, A.; Reginato, G.; Picchioni, F.; Pucci, A.; Calamante, M.; Zani, L. [Green/Yellow-Emitting](https://doi.org/10.1002/ejoc.201800242) Conjugated Heterocyclic Fluorophores for Luminescent Solar [Concentrators.](https://doi.org/10.1002/ejoc.201800242) *Eur. J. Org. Chem.* 2018, *2018*, 2657−2666.

(15) Papucci, C.; Dessì, A.; Coppola, C.; Sinicropi, A.; Santi, G.; Di Donato, M.; Taddei, M.; Foggi, P.; Zani, L.; Reginato, G.; Pucci, A.; Calamante, M.; Mordini, A. Benzo[1,2-*d*:4,5-*d*′[\]Bisthiazole](https://doi.org/10.1016/j.dyepig.2021.109207) Fluorophores for Luminescent Solar [Concentrators:](https://doi.org/10.1016/j.dyepig.2021.109207) Synthesis, Optical [Properties](https://doi.org/10.1016/j.dyepig.2021.109207) and Effect of the Polymer Matrix on the Device [Performances.](https://doi.org/10.1016/j.dyepig.2021.109207) *Dyes Pigm.* 2021, *188*, 109207.

(16) Papucci, C.; Charaf, R.; Coppola, C.; Sinicropi, A.; Di Donato, M.; Taddei, M.; Foggi, P.; Battisti, A.; De Jong, B.; Zani, L.; Mordini, A.; Pucci, A.; Calamante, M.; Reginato, G. [Luminescent](https://doi.org/10.1039/d1tc02923a) Solar [Concentrators](https://doi.org/10.1039/d1tc02923a) with Outstanding Optical Properties by Employment of D-A-D Quinoxaline [Fluorophores.](https://doi.org/10.1039/d1tc02923a) *J. Mater. Chem. C* 2021, *9*, 15608−15621.

(17) Yzeiri, X.; Calamante, M.; Dessì, A.; Franchi, D.; Pucci, A.; Ventura, F.; Reginato, G.; Zani, L.; Mordini, A. [Synthesis](https://doi.org/10.3390/molecules26185428) and Spectroscopic Characterization of [Thienopyrazine-Based](https://doi.org/10.3390/molecules26185428) Fluorophores for Application in Luminescent Solar [Concentrators](https://doi.org/10.3390/molecules26185428) (LSCs). *Molecules* 2021, *26*, 5428.

(18) Zhen, S.; Wang, S.; Li, S.; Luo, W.; Gao, M.; Ng, L. G.; Goh, C. C.; Qin, A.; Zhao, Z.; Liu, B.; Tang, B. Z. Efficient [Red/Near-Infrared](https://doi.org/10.1002/adfm.201706945) Fluorophores Based on Benzo[1,2-*b*:4,5-*b*′[\]Dithiophene](https://doi.org/10.1002/adfm.201706945) 1,1,5,5- Tetraoxide for Targeted [Photodynamic](https://doi.org/10.1002/adfm.201706945) Therapy and In Vivo Two-Photon [Fluorescence](https://doi.org/10.1002/adfm.201706945) Bioimaging. *Adv. Funct. Mater.* 2018, *28*, 1706945.

(19) (a) Mei, J.; Hong, Y.; Lam, J. W. Y.; Qin, A.; Tang, Y.; Tang, B. Z. [Aggregation-Induced](https://doi.org/10.1002/adma.201401356) Emission: The Whole Is More Brilliant than the [Parts.](https://doi.org/10.1002/adma.201401356) *Adv. Mater.* 2014, *26*, 5429−5479. (b) Cai, X.; Liu, B. [Aggregation-Induced](https://doi.org/10.1002/anie.202000845) Emission: Recent Advances in Materials and Biomedical [Applications.](https://doi.org/10.1002/anie.202000845) *Angew. Chem., Int. Ed.* 2020, *59*, 9868−9886. (c) Zhao, Z.; Zhang, H.; Lam, J. W. Y.; Tang, B. Z. [Aggregation-](https://doi.org/10.1002/anie.201916729)Induced Emission: New Vistas at the [Aggregate](https://doi.org/10.1002/anie.201916729) Level. *Angew. Chem., Int. Ed.* 2020, *59*, 9888−9907. (d) Peng, Q.; Shuai, Z. [Molecular](https://doi.org/10.1002/agt2.91) mechanism of [aggregation-induced](https://doi.org/10.1002/agt2.91) emission. *Aggregate* 2021, *2*, No. e91.

(20) Yi, X.; Dai, J.; Han, Y.; Xu, M.; Zhang, X.; Zhen, S.; Zhao, Z.; Lou, X.; Xia, F. A High [Therapeutic](https://doi.org/10.1038/s42003-018-0204-6) Efficacy of Polymeric Prodrug [Nano-Assembly](https://doi.org/10.1038/s42003-018-0204-6) for a Combination of Photodynamic Therapy and [Chemotherapy.](https://doi.org/10.1038/s42003-018-0204-6) *Commun. Biol.* 2018, *1*, 202.

(21) Zhen, S.; Yi, X.; Zhao, Z.; Lou, X.; Xia, F.; Tang, B. Z. [Drug](https://doi.org/10.1016/j.biomaterials.2019.119330) Delivery Micelles with Efficient Near-Infrared [Photosensitizer](https://doi.org/10.1016/j.biomaterials.2019.119330) for Combined Image-Guided Photodynamic Therapy and [Chemotherapy](https://doi.org/10.1016/j.biomaterials.2019.119330) of [Drug-Resistant](https://doi.org/10.1016/j.biomaterials.2019.119330) Cancer. *Biomaterials* 2019, *218*, 119330.

(22) Dai, J.; Xu, M.; Wang, Q.; Yang, J.; Zhang, J.; Cui, P.; Wang, W.; Lou, X.; Xia, F.; Wang, S. [Cooperation](https://doi.org/10.7150/thno.41708) Therapy between Anti-Growth by [Photodynamic-AIEgens](https://doi.org/10.7150/thno.41708) and Anti-Metastasis by Small Molecule [Inhibitors](https://doi.org/10.7150/thno.41708) in Ovarian Cancer. *Theranostics* 2020, *10*, 2385− 2398.

(23) Dai, J.; Li, Y.; Long, Z.; Jiang, R.; Zhuang, Z.; Wang, Z.; Zhao, Z.; Lou, X.; Xia, F.; Tang, B. Z. Efficient [Near-Infrared](https://doi.org/10.1021/acsnano.9b07972?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as) Photo-

<span id="page-17-0"></span>sensitizer with [Aggregation-Induced](https://doi.org/10.1021/acsnano.9b07972?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as) Emission for Imaging-Guided [Photodynamic](https://doi.org/10.1021/acsnano.9b07972?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as) Therapy in Multiple Xenograft Tumor Models. *ACS Nano* 2020, *14*, 854−866.

(24) Middha, E.; Chen, C.; Manghnani, P. N.; Wang, S.; Zhen, S.; Zhao, Z.; Liu, B. Synthesis of Uniform Polymer [Encapsulated](https://doi.org/10.1002/smtd.202100808) Organic Nanocrystals through Ouzo [Nanocrystallization.](https://doi.org/10.1002/smtd.202100808) *Small Methods* 2022, *6*, 2100808.

(25) Guo, X.; Zhen, S.; Ouyang, T.; Zhou, S.; Pan, Q.; Yang, D.; Chen, J.; Dong, G.; Zhao, Z.; Qiu, J.; Tang, B. Z. An [Organic](https://doi.org/10.1039/d1tc00323b) Microlaser Based on an [Aggregation-Induced](https://doi.org/10.1039/d1tc00323b) Emission Fluorophore for Tensile Strain [Sensing.](https://doi.org/10.1039/d1tc00323b) *J. Mater. Chem. C* 2021, *9*, 4888−4894.

(26) For the application of oxidized benzothieno-benzothiophene (BTBT) emitters, see: Mattiello, S.; Sanzone, A.; Bruni, F.; Gandini, M.; Pinchetti, V.; Monguzzi, A.; Facchinetti, I.; Ruffo, R.; Meinardi, F.; Mattioli, G.; Sassi, M.; Brovelli, S.; Beverina, L. [Chemically](https://doi.org/10.1016/j.joule.2020.08.006) Sustainable Large Stokes Shift Derivatives for [High-Performance](https://doi.org/10.1016/j.joule.2020.08.006) Large-Area Transparent Luminescent Solar [Concentrators.](https://doi.org/10.1016/j.joule.2020.08.006) *Joule* 2020, *4*, 1988−2003.

(27) Albano, G.; Colli, T.; Biver, T.; Aronica, L. A.; Pucci, A. [Photophysical](https://doi.org/10.1016/j.dyepig.2020.108368) Properties of New *p*-Phenylene- and Benzodithiophene-Based Fluorophores for Luminescent Solar [Concentrators](https://doi.org/10.1016/j.dyepig.2020.108368) [\(LSCs\).](https://doi.org/10.1016/j.dyepig.2020.108368) *Dyes Pigm.* 2020, *178*, 108368.

(28) Castriotta, L. A.; Infantino, R.; Vesce, L.; Stefanelli, M.; Dessì, A.; Coppola, C.; Calamante, M.; Reginato, G.; Mordini, A.; Sinicropi, A.; Di Carlo, A.; Zani, L. Stable [Methylammonium-Free](https://doi.org/10.1002/eem2.12455) *p-i-n* Perovskite Solar Cells and [Mini-Modules](https://doi.org/10.1002/eem2.12455) with Phenothiazine Dimers as Hole [Transporting](https://doi.org/10.1002/eem2.12455) Materials. *Energy Environ. Mater.* 2023, No. e12455.

(29) Planells, M.; Abate, A.; Hollman, D. J.; Stranks, S. D.; Bharti, V.; Gaur, J.; Mohanty, D.; Chand, S.; Snaith, H. J.; Robertson, N. Diacetylene Bridged [Triphenylamines](https://doi.org/10.1039/c3ta11417a) as Hole Transport Materials for Solid State Dye [Sensitized](https://doi.org/10.1039/c3ta11417a) Solar Cells. *J. Mater. Chem. A* 2013, *1*, 6949−6960.

(30) Di Donato, M.; Iagatti, A.; Lapini, A.; Foggi, P.; Cicchi, S.; Lascialfari, L.; Fedeli, S.; Caprasecca, S.; Mennucci, B. [Combined](https://doi.org/10.1021/jp505957q?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as) [Experimental](https://doi.org/10.1021/jp505957q?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as) and Theoretical Study of Efficient and Ultrafast Energy Transfer in a [Molecular](https://doi.org/10.1021/jp505957q?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as) Dyad. *J. Phys. Chem. C* 2014, *118*, 23476− 23486.

(31) Iagatti, A.; Patrizi, B.; Basagni, A.; Marcelli, A.; Alessi, A.; Zanardi, S.; Fusco, R.; Salvalaggio, M.; Bussotti, L.; Foggi, P. [Photophysical](https://doi.org/10.1039/C6CP08721C) Properties and Excited State Dynamics of 4,7-Dithien-[2-yl-2,1,3-Benzothiadiazole.](https://doi.org/10.1039/C6CP08721C) *Phys. Chem. Chem. Phys.* 2017, *19*, 13604−13613.

(32) Patrizi, B.; Iagatti, A.; Abbondanza, L.; Bussotti, L.; Zanardi, S.; Salvalaggio, M.; Fusco, R.; Foggi, P. Ultrafast [Intramolecular](https://doi.org/10.1021/acs.jpcc.8b11191?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as) and Solvation Dynamics in [4,7-Bis\(4,5-Dibutylbenzo\[1,2-](https://doi.org/10.1021/acs.jpcc.8b11191?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as)*b*:4,3-*b*′]- Bisthiophene[1,2-*b*:4,3-*b*′[\]Bisthiophen-2-yl\)-2,1,3-Benzothiadiazole.](https://doi.org/10.1021/acs.jpcc.8b11191?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as) *J. Phys. Chem. C* 2019, *123*, 5840−5852.

(33) van Stokkum, I. H. M.; Larsen, D. S.; van Grondelle, R. [Global](https://doi.org/10.1016/j.bbabio.2004.04.011) and Target Analysis of [Time-Resolved](https://doi.org/10.1016/j.bbabio.2004.04.011) Spectra. *Biochim. Biophys. Acta, Bioenerg.* 2004, *1657*, 82−104.

(34) Snellenburg, J. J.; Laptenok, S. P.; Seger, R.; Mullen, K. M.; van Stokkum, I. H. M. Glotaran: A *Java*-Based [Graphical](https://doi.org/10.18637/jss.v049.i03) User Interface for the *R* [Package](https://doi.org/10.18637/jss.v049.i03) TIMP. *J. Stat. Software* 2012, *49*, 1−22.

(35) Debije, M. G.; Evans, R. C.; Griffini, G. [Laboratory](https://doi.org/10.1039/D0EE02967J) Protocols for Measuring and Reporting the Performance of [Luminescent](https://doi.org/10.1039/D0EE02967J) Solar [Concentrators.](https://doi.org/10.1039/D0EE02967J) *Energy Environ. Sci.* 2021, *14*, 293−301.

(36) Yang, C.; Atwater, H. A.; Baldo, M. A.; Baran, D.; Barile, C. J.; Barr, M. C.; Bates, M.; Bawendi, M. G.; Bergren, M. R.; Borhan, B.; Brabec, C. J.; Brovelli, S.; Bulović, V.; Ceroni, P.; Debije, M. G.; Delgado-Sanchez, J.-M.; Dong, W.-J.; Duxbury, P. M.; Evans, R. C.; Forrest, S. R.; Gamelin, D. R.; Giebink, N. C.; Gong, X.; Griffini, G.; Guo, F.; Herrera, C. K.; Ho-Baillie, A. W. Y.; Holmes, R. J.; Hong, S.- K.; Kirchartz, T.; Levine, B. G.; Li, H.; Li, Y.; Liu, D.; Loi, M. A.; Luscombe, C. K.; Makarov, N. S.; Mateen, F.; Mazzaro, R.; McDaniel, H.; McGehee, M. D.; Meinardi, F.; Menéndez-Velázquez, A.; Min, J.; Mitzi, D. B.; Moemeni, M.; Moon, J. H.; Nattestad, A.; Nazeeruddin, M. K.; Nogueira, A. F.; Paetzold, U. W.; Patrick, D. L.; Pucci, A.; Rand, B. P.; Reichmanis, E.; Richards, B. S.; Roncali, J.; Rosei, F.; Schmidt, T. W.; So, F.; Tu, C.-C.; Vahdani, A.; van Sark, W. G. J. H. M.; Verduzco, R.; Vomiero, A.; Wong, W. W. H.; Wu, K.; Yip, H.-L.; Zhang, X.; Zhao, H.; Lunt, R. R. Consensus Statement: [Standardized](https://doi.org/10.1016/j.joule.2021.12.004) Reporting of [Power-Producing](https://doi.org/10.1016/j.joule.2021.12.004) Luminescent Solar Concentrator [Performance.](https://doi.org/10.1016/j.joule.2021.12.004) *Joule* 2022, *6*, 8−15.

(37) Wang, Y.-K.; Wu, S.-F.; Li, S.-H.; Yuan, Y.; Wu, F.-P.; Kumar, S.; Jiang, Z.-Q.; Fung, M.-K.; Liao, L.-S. [D-A-A-Type](https://doi.org/10.1002/adom.201700566) Emitter Featuring Benzo[*c*] [\[1,2,5\]Thiadiazole](https://doi.org/10.1002/adom.201700566) and Polar C≡N Bond as Tandem Acceptor for [High-Performance](https://doi.org/10.1002/adom.201700566) Near-Infrared Organic [Light-Emitting](https://doi.org/10.1002/adom.201700566) Diodes. *Adv. Opt. Mater.* 2017, *5*, 1700566.

(38) Wałęsa-Chorab, M.; Yao, C.; Tuner, G.; Skene, W. G. Electrochemical and Solvent-Mediated [Visible-to-Near-Infrared](https://doi.org/10.1002/chem.201903291) Spectroscopic Switching of [Benzoselenadiazole](https://doi.org/10.1002/chem.201903291) Fluorophores. *Chem.*� *Eur. J.* 2020, *26*, 17416−17427.

(39) De Nisi, F.; Francischello, R.; Battisti, A.; Panniello, A.; Fanizza, E.; Striccoli, M.; Gu, X.; Leung, N. L. C.; Tang, B. Z.; Pucci, A. [Red-](https://doi.org/10.1039/c7qm00008a)Emitting AIEgen for Luminescent Solar [Concentrators.](https://doi.org/10.1039/c7qm00008a) *Mater. Chem. Front.* 2017, *1*, 1406−1412.

(40) Ma, W.; Li, W.; Cao, M.; Liu, R.; Zhao, X.; Gong, X. [Large](https://doi.org/10.1016/j.orgel.2019.06.017) Stokes-Shift AIE Fluorescent Materials for [High-Performance](https://doi.org/10.1016/j.orgel.2019.06.017) Luminescent Solar [Concentrators.](https://doi.org/10.1016/j.orgel.2019.06.017) *Org. Electron.* 2019, *73*, 226−230.

(41) Gao, S.; Balan, B.; Yoosaf, K.; Monti, F.; Bandini, E.; Barbieri, A.; Armaroli, N. Highly Efficient Luminescent Solar [Concentrators](https://doi.org/10.1002/chem.202001210) Based on [Benzoheterodiazole](https://doi.org/10.1002/chem.202001210) Dyes with Large Stokes Shifts. *Chem.*� *Eur. J.* 2020, *26*, 11013−11023.

(42) Mateen, F.; Meti, P.; Hwang, D.-Y.; Swelm, W.; Algarni, H.; Al-Sehemi, A. G.; Kim, Y.-C.; Gong, Y.-D.; Hong, S.-K. Highly [Efficient](https://doi.org/10.1016/j.dyepig.2022.110563) [Indoor/Outdoor](https://doi.org/10.1016/j.dyepig.2022.110563) Light Harvesting Luminescent Solar Concentrator Employing [Aggregation-Induced](https://doi.org/10.1016/j.dyepig.2022.110563) Emissive Fluorophore. *Dyes Pigm.* 2022, *205*, 110563.

(43) Hohenberg, P.; Kohn, W. [Inhomogeneous](https://doi.org/10.1103/PhysRev.136.B864) Electron Gas. *Phys. Rev.* 1964, *136*, B864−B871.

(44) Kohn, W.; Sham, L. J. [Self-Consistent](https://doi.org/10.1103/PhysRev.140.A1133) Equations Including Exchange and [Correlation](https://doi.org/10.1103/PhysRev.140.A1133) Effects. *Phys. Rev.* 1965, *140*, A1133− A1138.

(45) Adamo, C.; Jacquemin, D. The Calculations of [Excited-State](https://doi.org/10.1039/C2CS35394F) Properties with [Time-Dependent](https://doi.org/10.1039/C2CS35394F) Density Functional Theory. *Chem. Soc. Rev.* 2013, *42*, 845−856.

(46) Laurent, A. D.; Adamo, C.; Jacquemin, D. Dye [Chemistry](https://doi.org/10.1039/C3CP55336A) with [Time-Dependent](https://doi.org/10.1039/C3CP55336A) Density Functional Theory. *Phys. Chem. Chem. Phys.* 2014, *16*, 14334−14356.

(47) Frisch, M. J.; Trucks, G. W.; Schlegel, H. B.; Scuseria, G. E.; Robb, M. A.; Cheeseman, J. R.; Scalmani, G.; Barone, V.; Petersson, G. A.; Nakatsuji, H.; Li, X.; Caricato, M.; Marenich, A. V.; Bloino, J.; Janesko, B. G.; Gomperts, R.; Mennucci, B.; Hratchian, H. P.; Ortiz, J. V.; Izmaylov, A. F.; Sonnenberg, J. L.; Williams-Young, D.; Ding, F.; Lipparini, F.; Egidi, F.; Goings, J.; Peng, B.; Petrone, A.; Henderson, T.; Ranasinghe, D.; Zakrzewski, V. G.; Gao, J.; Rega, N.; Zheng, G.; Liang, W.; Hada, M.; Ehara, M.; Toyota, K.; Fukuda, R.; Hasegawa, J.; Ishida, M.; Nakajima, T.; Honda, Y.; Kitao, O.; Nakai, H.; Vreven, T.; Throssell, K.; Montgomery, J. A., Jr.; Peralta, J. E.; Ogliaro, F.; Bearpark, M. J.; Heyd, J. J.; Brothers, E. N.; Kudin, K. N.; Staroverov, V. N.; Keith, T. A.; Kobayashi, R.; Normand, J.; Raghavachari, K.; Rendell, A. P.; Burant, J. C.; Iyengar, S. S.; Tomasi, J.; Cossi, M.; Millam, J. M.; Klene, M.; Adamo, C.; Cammi, R.; Ochterski, J. W.; Martin, R. L.; Morokuma, K.; Farkas, O.; Foresman, J. B.; Fox, D. J. *Gaussian 16*, Revision C.01; Gaussian, Inc.: Wallingford, CT, 2016.

(48) Lee, C.; Yang, W.; Parr, R. G. [Development](https://doi.org/10.1103/PhysRevB.37.785) of the Colle-Salvetti [Correlation-Energy](https://doi.org/10.1103/PhysRevB.37.785) Formula into a Functional of the Electron [Density.](https://doi.org/10.1103/PhysRevB.37.785) *Phys. Rev. B: Condens. Matter Mater. Phys.* 1988, *37*, 785− 789.

(49) Becke, A. D. [Density-functional](https://doi.org/10.1063/1.464913) Thermochemistry. III. The Role of Exact [Exchange.](https://doi.org/10.1063/1.464913) *J. Chem. Phys.* 1993, *98*, 5648−5652.

(50) Yanai, T.; Tew, D. P.; Handy, N. C. A New Hybrid [Exchange](https://doi.org/10.1016/j.cplett.2004.06.011)− Correlation Functional Using the [Coulomb-Attenuating](https://doi.org/10.1016/j.cplett.2004.06.011) Method [\(CAM-B3LYP\).](https://doi.org/10.1016/j.cplett.2004.06.011) *Chem. Phys. Lett.* 2004, *393*, 51−57.

(51) Lynch, B. J.; Fast, P. L.; Harris, M.; Truhlar, D. G. [Adiabatic](https://doi.org/10.1021/jp000497z?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as) [Connection](https://doi.org/10.1021/jp000497z?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as) for Kinetics. *J. Phys. Chem. A* 2000, *104*, 4811−4815.

<span id="page-18-0"></span>(52) Tomasi, J.; Mennucci, B.; Cammi, R. Quantum [Mechanical](https://doi.org/10.1021/cr9904009?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as) [Continuum](https://doi.org/10.1021/cr9904009?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as) Solvation Models. *Chem. Rev.* 2005, *105*, 2999−3094.

(53) Bernini, C.; Zani, L.; Calamante, M.; Reginato, G.; Mordini, A.; Taddei, M.; Basosi, R.; Sinicropi, A. Excited State [Geometries](https://doi.org/10.1021/ct500328t?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as) and Vertical Emission Energies of Solvated Dyes for DSSC: A [PCM/TD-](https://doi.org/10.1021/ct500328t?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as)DFT [Benchmark](https://doi.org/10.1021/ct500328t?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as) Study. *J. Chem. Theory Comput.* 2014, *10*, 3925− 3933.

(54) Khambhati, D. P.; Sachinthani, K. A. N.; Rheingold, A. L.; Nelson, T. L. Regioselective [Copper-Catalyzed](https://doi.org/10.1039/C7CC01781B) Direct Arylation of [Benzodithiophene-S,S-Tetraoxide.](https://doi.org/10.1039/C7CC01781B) *Chem. Commun.* 2017, *53*, 5107− 5109.

(55) Punzi, A.; Capozzi, M. A. M.; Di Noja, S.; Ragni, R.; Zappimbulso, N.; Farinola, G. M. Solvent-Free [Pd-Catalyzed](https://doi.org/10.1021/acs.joc.8b01284?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as) [Heteroaryl](https://doi.org/10.1021/acs.joc.8b01284?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as)−Aryl Coupling via C−H Bond Activation for the Synthesis of Extended [Heteroaromatic](https://doi.org/10.1021/acs.joc.8b01284?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as) Conjugated Molecules. *J. Org. Chem.* 2018, *83*, 9312−9321.

(56) Dessì, A.; Calamante, M.; Sinicropi, A.; Parisi, M. L.; Vesce, L.; Mariani, P.; Taheri, B.; Ciocca, M.; Di Carlo, A.; Zani, L.; Mordini, A.; Reginato, G. Thiazolo[5,4-*d*[\]Thiazole-Based](https://doi.org/10.1039/D0SE00124D) Organic Sensitizers with Improved Spectral Properties for Application in [Greenhouse-](https://doi.org/10.1039/D0SE00124D)Integrated [Dye-Sensitized](https://doi.org/10.1039/D0SE00124D) Solar Cells. *Sustainable Energy Fuels* 2020, *4*, 2309−2321.

(57) Dessì, A.; Chalkias, D. A.; Bilancia, S.; Sinicropi, A.; Calamante, M.; Mordini, A.; Karavioti, A.; Stathatos, E.; Zani, L.; Reginato, G. D−A−*π*−A organic dyes with tailored green light [absorption](https://doi.org/10.1039/d0se01610a) for potential application in [greenhouse-integrated](https://doi.org/10.1039/d0se01610a) dye-sensitized solar [cells.](https://doi.org/10.1039/d0se01610a) *Sustainable Energy Fuels* 2021, *5*, 1171−1183.

(58) Goti, G.; Calamante, M.; Coppola, C.; Dessì, A.; Franchi, D.; Mordini, A.; Sinicropi, A.; Zani, L.; Reginato, G. [Donor-Acceptor-](https://doi.org/10.1002/ejoc.202100199)Donor [Thienopyrazine-Based](https://doi.org/10.1002/ejoc.202100199) Dyes as NIR-Emitting AIEgens. *Eur. J. Org. Chem.* 2021, *2021*, 2655−2664.

(59) Zani, L.; Dessì, A.; Franchi, D.; Calamante, M.; Reginato, G.; Mordini, A. Transition [Metal-Catalyzed](https://doi.org/10.1016/j.ccr.2019.04.007) Cross-Coupling Methodologies for the Engineering of Small Molecules with [Applications](https://doi.org/10.1016/j.ccr.2019.04.007) in Organic Electronics and [Photovoltaics.](https://doi.org/10.1016/j.ccr.2019.04.007) *Coord. Chem. Rev.* 2019, *392*, 177−236.

(60) Wilson, L. R.; Richards, B. S. [Measurement](https://doi.org/10.1364/AO.48.000212) Method for [Photoluminescent](https://doi.org/10.1364/AO.48.000212) Quantum Yields of Fluorescent Organic Dyes in Polymethyl Methacrylate for Luminescent Solar [Concentrators.](https://doi.org/10.1364/AO.48.000212) *Appl. Opt.* 2009, *48*, 212−220.

(61) Li, Y.; Zhang, X.; Zhang, Y.; Dong, R.; Luscombe, C. K. [Review](https://doi.org/10.1002/pola.29192) on the Role of Polymers in Luminescent Solar [Concentrators.](https://doi.org/10.1002/pola.29192) *J. Polym. Sci., Part A: Polym. Chem.* 2019, *57*, 201−215.

(62) Sperling, L. H. *Introduction to Physical Polymer Science*; Wiley, 2005. DOI: [10.1002/0471757128](https://doi.org/10.1002/0471757128?urlappend=%3Fref%3DPDF&jav=VoR&rel=cite-as).

(63) Griffini, G.; Levi, M.; Turri, S. Thin-Film [Luminescent](https://doi.org/10.1016/j.renene.2015.01.009) Solar [Concentrators:](https://doi.org/10.1016/j.renene.2015.01.009) A Device Study towards Rational Design. *Renewable Energy* 2015, *78*, 288−294.

(64) Haines, C.; Chen, M.; Ghiggino, K. P. The Effect of [Perylene](https://doi.org/10.1016/j.solmat.2012.06.030) Diimide [Aggregation](https://doi.org/10.1016/j.solmat.2012.06.030) on the Light Collection Efficiency of Luminescent [Concentrators.](https://doi.org/10.1016/j.solmat.2012.06.030) *Sol. Energy Mater. Sol. Cells* 2012, *105*, 287−292.

(65) Ceriani, C.; Corsini, F.; Mattioli, G.; Mattiello, S.; Testa, D.; Po, R.; Botta, C.; Griffini, G.; Beverina, L. [Sustainable](https://doi.org/10.1039/d1tc03536c) by Design, Large Stokes Shift [Benzothiadiazole](https://doi.org/10.1039/d1tc03536c) Derivatives for Efficient Luminescent Solar [Concentrators.](https://doi.org/10.1039/d1tc03536c) *J. Mater. Chem. C* 2021, *9*, 14815−14826.

(66) Micheletti, C.; Wang, Q.; Ventura, F.; Turelli, M.; Ciofini, I.; Adamo, C.; Pucci, A. Red-emitting [Tetraphenylethylene](https://doi.org/10.1002/agt2.188) Derivative with [Aggregation-induced](https://doi.org/10.1002/agt2.188) Enhanced Emission for Luminescent Solar [Concentrators:](https://doi.org/10.1002/agt2.188) A Combined Experimental and Density Functional [Theory](https://doi.org/10.1002/agt2.188) Study. *Aggregate* 2022, *3*, No. e188.

(67) Corsini, F.; Nitti, A.; Tatsi, E.; Mattioli, G.; Botta, C.; Pasini, D.; Griffini, G. Large-Area [Semi-Transparent](https://doi.org/10.1002/adom.202100182) Luminescent Solar Concentrators Based on Large Stokes Shift [Aggregation-Induced](https://doi.org/10.1002/adom.202100182) [Fluorinated](https://doi.org/10.1002/adom.202100182) Emitters Obtained Through a Sustainable Synthetic [Approach.](https://doi.org/10.1002/adom.202100182) *Adv. Opt. Mater.* 2021, *9*, 2100182.

(68) Rosadoni, E.; Bellina, F.; Lessi, M.; Micheletti, C.; Ventura, F.; Pucci, A. Y-Shaped [Alkynylimidazoles](https://doi.org/10.1016/j.dyepig.2022.110262) as Effective Push-Pull

Fluorescent Dyes for Luminescent Solar [Concentrators](https://doi.org/10.1016/j.dyepig.2022.110262) (LSCs). *Dyes Pigm.* 2022, *201*, 110262.

(69) Meti, P.; Mateen, F.; Hwang, D. Y.; Lee, Y.-E.; Hong, S.-K.; Gong, Y.-D. Luminescent Solar [Concentrator](https://doi.org/10.1016/j.dyepig.2022.110221) Based on Large-Stokes Shift [Tetraphenylpyrazine](https://doi.org/10.1016/j.dyepig.2022.110221) Fluorophore Combining Aggregation-Induced Emission and [Intramolecular](https://doi.org/10.1016/j.dyepig.2022.110221) Charge Transfer Features. *Dyes Pigm.* 2022, *202*, 110221.

(70) Seybold, G.; Wagenblast, G. New Perylene and [Violanthrone](https://doi.org/10.1016/0143-7208(89)85048-X) Dyestuffs for [Fluorescent](https://doi.org/10.1016/0143-7208(89)85048-X) Collectors. *Dyes Pigm.* 1989, *11*, 303−317.

# **Recommended by ACS**

**[Green](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[-](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[Light](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Wavelength](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[-](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[Selective](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Organic](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Solar](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Cells](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Based](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [on](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Poly](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[\(](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[3](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[-](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[hexylthiophene](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[\)](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) **[and](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Naphthobisthiadiazole](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[-](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[Containing](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Acceptors](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [toward](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Agrivoltaics](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**

Seihou Jinnai, Yutaka Ie, et al. JANUARY 12, 2023 ACS SUSTAINABLE CHEMISTRY & ENGINEERING [READ](http://pubs.acs.org/doi/10.1021/acssuschemeng.2c06426?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) READ

**[Multifaceted](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Sulfone](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[–](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[Carbazole](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[-](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[Based](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [D](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[–](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[A](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[–](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[D](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Materials](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[:](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) **[A](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Blue](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Fluorescent](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Emitter](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [as](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [a](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Host](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [for](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Phosphorescent](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [OLEDs](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [and](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Triplet](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[–](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[Triplet](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Annihilation](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Up](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[-](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[Conversion](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[...](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)

Tsz Chung Yiu, Yuan Jay Chang, et al. DECEMBER 28, 2022 ACS APPLIED MATERIALS & INTERFACES [READ](http://pubs.acs.org/doi/10.1021/acsami.2c21294?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) CONTROL AND THE READ CONTROL AND READ CONTROL AT A READ CONTROL AND A READ CONTR

**[Photovoltaic](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Effects](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [of](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Dye](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[-](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[Sensitized](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Solar](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Cells](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Using](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Double](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[-](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[Layered](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [TiO](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)[2](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Photoelectrodes](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [and](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Pyrazine](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[-](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[Based](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Photosensitizers](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**

Mi-Ra Kim, Songyi Lee, et al. APRIL 12, 2023 ACS OMEGA [READ](http://pubs.acs.org/doi/10.1021/acsomega.3c00707?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) 

#### **[Outdoor](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Stability](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [of](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Triplet](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Generating](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [and](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Triplet](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Harvesting](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Materials](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[;](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) **[Pentacene](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [and](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Tetracene](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Electron](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Donors](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Merged](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [with](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Boron](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Subphthalocyanines](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [as](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) [Electr](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)**[...](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362)

Adam R. Tetreault, Timothy P. Bender, et al.

FEBRUARY 24, 2023 ACS APPLIED ENERGY MATERIALS **[READ](http://pubs.acs.org/doi/10.1021/acsaem.2c04185?utm_campaign=RRCC_aaemcq&utm_source=RRCC&utm_medium=pdf_stamp&originated=1684158038&referrer_DOI=10.1021%2Facsaem.3c00362) LARGE SERVICE** 

**[Get More Suggestions >](https://preferences.acs.org/ai_alert?follow=1)**