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YAL SOCIETY CHEMISTRY

Journal Name

ARTICLE

Received 00th January 20xx, Accepted 00th January 20xx

DOI: 10.1039/x0xx00000x

www.rsc.org/

Electron Donor-free Photoredox Catalysis via Electron Transfer Cascade by Cooperative Organic Photocatalysts

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Electron-donating sacrificial reagents are highly important for certain photo-redox reactions. However, the use of usually excessive amount of sacrificial reagents, mostly amines, often leads to undesired side products and is especially troublesome for product purification. Herein, we take the light-induced electron transfer cascade process of the nature photosystems as roll model and assemble organic photocatalysts into cooperative photocatalyst couples. The cooperative photocatalyst couples could undergo intermolecular electron transfer to ease the charge separation process and overcome the need of an extra electron donor. Time-resolved photoluminescence spectroscopy was conducted to precisely characterize the photo-excited dynamics within the cooperative photocatalyst couples. As model photoredox reaction, the carbon-carbon formation reaction between heteroarenes and malonates, which usually requires electron-donating sacrificial reagents such as amines, was conducted to demonstrate the feasibility of the cooperative photocatalyst couples under visible light irradiation. A significant reaction conversion improvement from trace conversion for single photocatalyst system to over 90% by cooperative photocatalyst couples was achieved.

Introduction

Electron-donating sacrificial reagents are of great importance for photocatalytic redox reactions including chemical transformation and water decomposition into hydrogen and oxygen.¹⁻³ Especially for semiconductor-catalyzed reactions, the catalytic efficiency can be largely improved while the photo-generated holes can be scavenged by the electrondonating reagent molecules and the charge recombination can be reduced.² Examples of organic photoredox reactions without the employment of sacrificial reagent were conducted using reactive substrates such as aryl diazonium or trifluoromethanesulfonyl salts⁵, as well as when the solvent molecule could also act as electron donor.⁶ However, for specific photoredox reactions involving less reactive substrates, the use of electron donating reagents is necessary. And the use of usually excessive amount of the donor reagents during the catalytic cycle, mostly amines,⁷⁻¹⁰ often leads to undesired oxidized side products and is especially troublesome for product purification.¹¹ To overcome the above-mentioned challenge and maintain reduced charge recombination within the semiconductor-based photocatalyst, an enhanced photogenerated electron transfer pathway should be developed.

Max Planck Institute for Polymer Research, Ackermannweg 10, 55128 Mainz, Germany. E-mail: kai.zhang@mpip-mainz.mpg.de Nature photosystems offer a promising model to overcome the charge recombination issue, where a series of stepwise electron transfer processes is involved to enhance the photogenerated charge separation and transfer process.¹² So far, chemists have designed different photocatalytic systems for enhancing the photocatalytic efficiency. For example, bridged transition metal complexes¹³⁻¹⁵ have been reported to greatly promote the photogenerated charge separation and transfer.

Organic semiconductor (OS)-containing photocatalysts have emerged as a promising alternative to the traditional metal-containing photocatalysts due to their highly tunable electronical properties via flexible structural design. Recent research activities have demonstrated a vast number of structural design methods of small molecular^{8, 10, 16-20} or macromolecular²¹⁻³⁰ OS systems as efficient photocatalysts for visible light-promoted photoredox reactions. Nevertheless, most developed OS photocatalysts, similar to traditional transition metal complexes,³¹⁻³⁸ are single photocatalytic systems. As illustrated in Fig. 1a, the photocatalytic ability of the OS system originates from the photo-generated electron/hole pair, which can function as the reductive and oxidative species. To maintain the efficient electron transfer from the ground to the exited state of the OS, extra sacrificial agents are usually needed due to direct recombination of the electron/hole pair. To overcome the necessary use of electrondonating sacrificial reagents, the single photocatalyst system is not sufficient; a new concept of catalyst design is needed.

Electronic Supplementary Information (ESI) available: Experiment details and characterization, 1H and 13C NMR spectra, fluorescence quenching experiments and control experiments are described. See DOI: 10.1039/x0xx00000x

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Fig. 1. Difference between (a) conventional single organic photocatalyst system with direct electron/hole recombination, sacrificial reagent needed; (b) cooperative photocatalyst system with enhanced electron/hole separation via electron transfer cascade, no sacrificial reagent needed. OS: organic semiconductor; SA: sacrificial electron donor reagent; S: substrate.

Herein, we aim to take the mechanism of the light-induced electron transfer cascade process of nature photosystems as roll model and assemble organic photocatalysts into cooperative photocatalyst couples. As illustrated in Fig. 1b, the two molecular OS photocatalysts within the cooperative couple consisted of different organic photocatalysts possessing different highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) levels. By coupling the OSs, an extra intermolecular electron transfer could occur between the OSs, leading to enhanced photogenerated electron/hole separation under light irradiation and thereby more stable reductive and oxidative species. The intermolecular electron transfer within the OS couple can decay the direct recombination of the electron/hole pair and overcome the mandatory requirement of extra electron donors during the photoredox cycle. As model reaction, the photocatalytic carbon-carbon formation reaction between electron-rich heteroarenes and malonates, which usually required electron-donating sacrificial reagents,^{9, 39} was chosen to demonstrate the feasibility and photocatalytic activity of the cooperative photocatalyst couple. No extra sacrificial reagents were needed. The light-induced electron transfer between the OSs within the cooperative photocatalyst couple and the substrates could be revealed by mechanistic studies.

Results and discussion



Scheme 1. Structure and HOMO/LUMO levels (vs. SCE) of the OS molecules in both designed photocatalyst couples.

In this study, three molecular organic semiconductors (OS), in particular, 4,7-di(thiophen-2-yl) benzo[c][1,2,5]thiadiazole (**Th-BT-Th**), 2-butyl-4,7-di(thiophen-2-yl)-2H-benzo[d][1,2,3] triazole (**Th-BTz-Th**), and 4,7-bis(1-butyl-1H-1,2,3-triazol-4-yl)benzo[c] [1,2,5]thiadiazole (**TA-BT-TA**) were designed. The structures and energy band positions of the OSs are shown in Scheme 1. The synthetic and characteristic details of the OSs are described in the Experimental Section or Electronic Supplementary Information (ESI).



Fig. 2. (a) UV/Vis absorption and (b) fluorescence spectra of the three OS molecules. (c), (d) PL decay times of the pristine materials and the blend of the cooperative couples with exciton lifetimes collected at the fluorescence maximum. (e), (f) Jablonski diagram illustrates energy and charge transfer in the photocatalyst couples, respectively. Excitation wavelength: 400 nm. EET: excitation energy transfer; ET: electron transfer.

The three OSs possessed different HOMO and LUMO band positions, which could be determined via cyclic voltammetry (Fig. S1). As shown in Scheme 1, Th-BTz-Th exhibited a significantly high LUMO level at -1.68 V vs. saturated calomel electrode (SCE), followed by TA-BT-TA with -1.20 V vs. SCE and Th-BT-Th with -1.15 V vs. SCE. In comparison, TA-BT-TA exhibited the lowest HOMO level at +1.54 V vs. SCE, followed by Th-BT-Th with +1.12 V vs. SCE and Th-BTz-Th with +0.95 V vs. SCE. In addition, the excited state redox potentials are calculated to be +1.16 V and -1.01 V vs. SCE for Th-BT-Th, +0.62 V and -1.35 V vs. SCE for Th-BTz-Th, and +1.14 V and - 0.80 V vs. SCE for TA-BT-TA, respectively. The date are listed in Table S1 in the ESI using the calculation described in previous report.⁴⁰ Given the different energy band positions, we assembled the three OSs in two photocatalyst couples for this

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study, particularly, Th-BTz-Th/Th-BT-Th for cooperative photocatalyst couple 1 and Th-BTz-Th/TA-BT-TA for couple 2.

Steady-state fluorescence quenching experiments showed that by only exciting Th-BT-Th (at 500 nm) or TA-BT-TA (at 450 nm) in both couples, the fluorescence intensity of couple 2 exhibited a larger decrease in PL intensity than that of couple 1, indicating an enhanced electron transfer within couple 2, which was likely caused by the larger distance between the two HOMO levels (Fig. S2).

To further study the photo-induced charge transfer within . couples, time-resolved photoluminescence (TRPL) the spectroscopy was conducted to characterize the photo-excited dynamics within the OS couples. The PL measurements of the pristine materials and a 1:1 (mol-%) blend were all performed in DMF solution with excitation at 400 nm (Fig. 2c and 2d). The selected PL decays correspond to the emission maximum for each component OS (Fig. 2c). The fluorescence features of the different component materials overlap with each other. Therefore, decay associated spectra (DAS) were derived by a global fitting analysis⁴¹ of the TRPL data (Fig. S2). This allows us to characterize the spectral distribution and lifetime of each emitting species, particularly for the measurements of couple 1 and couple 2. In couple 1 (Fig. S2a), one can see that DAS 1, centered at 460 nm, matches exactly with the fluorescence spectrum of Th-BTz-Th and decays in 0.6 ns, which is quenched with respect to the pristine donor (2.6 ns). DAS 2, which is centered at 590 nm, matches the fluorescence spectrum of Th-BT-Th and decays with the same ~12 ns lifetime as the pristine Th-BT-Th sample, exhibiting no quenching. This indicates that in couple 1 excitation energy transfer (EET) from Th-BTz-Th to Th-BT-Th likely dominates the excited state dynamics, as illustrated in Fig. 2e. In couple 2 (Fig. S2b), the DAS 1, centered at 460 nm, is again originating from Th-BTz-Th and decays in 0.97 ns, which is guenched with respect to the pristine Th-BTz-Th (2.6 ns). DAS 2, centered at 530 nm, is assigned to the emission of TA-BT-TA and has a decay time of 10.6 ns. Here, the fluorescence of both Th-BTz-Th and TA-BT-TA are guenched with respect to the pristine materials. This indicates an electron transfer from Th-BTz-Th to TA-BT-TA and an enhanced charge transfer within couple 2 (Fig. 2f).

We chose the photocatalytic carbon-carbon formation reaction between electron-rich heteroarenes and malonates as model reaction.9, 10 Previous studies demonstrated the mandatory use of sacrificial reagents, mostly amines, which ought not to only act as electron donor, but also as electron mediator between the radical intermediate and the photogenerated hole of the photocatalyst (Fig. S4). We first tested the three designed OSs as single photocatalyst system without using sacrificial reagent. As expected, by using 3methylbenzofuran and diethyl bromomalonate in the model reaction, no product was obtained using Th-BT-Th and TA-BT-TA as photocatalysts (entries 1-2 in Table 1). Interestingly, using Th-BTz-Th as photocatalyst led to a slight conversion of 9%. This indicates that a minimal electron transfer could still occur between the photocatalyst and the substrate, which corresponds with the highest LUMO level of Th-BTz-Th among

the three OSs and thereby the largest overpotential for the reduction of diethyl bromomalonate.

| able 1. Photocatalytic C-C bond formation between diethyl bromomalonate and | |
|---|--|
| neteroarenes using different photocatalyst systems. ^a | |
| | |

| | K + Br- | COOEt photoca | MF, RT | | : |
|-----------------|----------------------------------|---------------|----------------|--------------------------|---------------------------|
| Entry | Catalyst | Substrate | Product | Conv [%] ^b | Yield [%] ^c |
| 1 | Th-BT-Th | | | trace | - |
| 2 | TA-BT-TA | | | 0 | - |
| 3 | Th-BTz-Th | | | 9 | - |
| 4 | Couple 1 | | | 43 | 39 |
| 5 | Couple 2 | | | 93 | 87 |
| 6 ^d | Couple 2 | | | 98 | 85 |
| 7 ^e | Couple 2 | | | 78 | 75 |
| 8 ^e | Couple 2 | | | 92 | 85 |
| 9 ^e | Couple 2 | COOCH3 | EtOOC COOEt | 85 | 80 |
| 10 ^f | Th-BTz- Th/TA-BT- TA (2:1) | | | 72 | 60 |
| 11 ^f | Connected cat. | | | 31 | 22 |
| 12 ^g | - | | | 0 | - |
| 13 ^h | Couple 2 | | | 0 | - |

^aReaction conditions: 1 equiv. (0.38 mmol) heteroarene, 2 equiv. diethyl bromomalonate, 0.1 equiv. photocatalyst (for the coupled system, the ratio between the two OSs was 1:1) in 2.5 mL DMF, white LED lamp (0.07 Wcm⁻²), 24 h. ^bDetermined by GC-MS. ^cIsolated yield. ^dblue LED (450 nm, 0.1 Wcm⁻²). ^e48h. ^f0.05 equiv. photocatalyst, 24h. ^gno catalyst, under light. ^hno light.

In contrast, by using both cooperative photocatalyst couples, the product could be obtained in significantly higher conversions, in particular, 43% for couple 1 and 93% for couple 2, respectively (entries 4-5). Both the apparent quantum yield (Φ_{AQY}) and catalytic turnover number (TON) for the model reaction (entry 5) using coupling 2 as photocatalyst were calculated to be 0.04% and 8.7, respectively (ESI). A kinetic study confirmed the superior catalytic activity of couple 2

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DOI: 10.1039/C8CY01072B Journal Name

compared to couple 1 via fluorescence quenching experiment by the substrates (Fig. S6). This is in agreement with the photophysical study (Fig. 2), indicating that the enhanced charge transfer between Th-BTz-Th and TA-BT-TA in couple 2 led to a better catalytic efficiency than that of couple 1 containing Th-BTz-Th and Th-BT-Th. The reason could lie on the larger HOMO energy gap ($\Delta E_{\rm HOMO} = 0.59$ V) between Th-BTz-Th and TA-BT-TA than that ($\Delta E_{\rm HOMO} = 0.17$ V) in couple 1 between Th-BTz-Th and Th-BT-Th.^{42, 43} By using a single wavelength light source (blue LED, 450 nm), where only TA-BT-TA of couple 2 could be excited, the model reaction exhibited a high conversion of 98%. This indicates a hole transfer from TA-BT-TA to Th-BTz-Th, leading a similar effect of an electron transfer from Th-BTz-Th back to TA-BT-TA, as illustrated in Fig. 2f.

In comparison to the use of Th-BT-Th as single photocatalyst with the employment of an extra sacrificial electron donor, in particular, triphenylamine from our previous report for the model reaction,⁴⁴ where the apparent quantum yield was calculated to be 0.96%, the apparent quantum yield of the couple 2 was lower due to the longer reaction time. However, the avoidance of the extra sacrificial electron donor using the molecular photocatalyst couples is still advantageous. It can efficiently prevent the postpurification procedures caused by the addition of usually excess amount of the amine donors.

Screening experiments using different substrates confirmed the feasibility of the OS couple as photocatalyst (entries 7-9). Other control experiments either without the photocatalyst or without light led to no product formation (entries 12 and 13), indicating the mandatory roles of both components.



Fig. 3. Study of the electron transfer pathway using comparative substrates with different reduction potentials. Light source: white LED lamp (0.07 Wcm⁻², λ > 420 nm). *Interm.*: intermediate.

To gain more mechanistic insight of the interaction between the photocatalyst and substrates, we conducted

further fluorescence quenching experiments. It was shown that the fluorescence of the OSs as single photocatalyst could be only minimal quenched by adding diethyl bromomalonate (Fig. S5). As a contrast, the fluorescence of both photocatalyst couples could be gradually quenching by adding diethyl bromomalonate (Fig. S6), confirming the possible electron transfer between the photocatalyst couple and the substrate.



Fig. 4. Control experiments: (a) Study of the impact of the OS ratio on the reaction conversion; (b) Comparison between cooperative photocatalyst couple 2 with the C6-connected molecule Th-BTz-Th@TA-BT-TA; (c) Structure of Th-BTz-Th@TA-BT-TA. Reaction conditions: 1 equiv. (0.38 mmol) 3-methylbenzofuran, 2 equiv. diethyl bromomalonate, 0.05 equiv. photocatalyst in 2.5 mL DMF, white LED (0.07 Wcm⁻², λ > 420 nm), 24 h, conversion determined by GC-MS.

To precisely study the concrete electron transfer process within the catalytic cycle, we then employed ethyl bromoacetate as substrate, which possessed a reduction potential of -1.43 V vs. SCE (Fig. 3). By using couple 2 as photocatalyst, it could be determined that only traces of the final product could be obtained (Fig. S11). This reveals that the photo-generated electron could indeed only transferred from the LUMO of Th-BTz-Th (-1.68 V vs. SCE) to the LUMO of TA-BT-TA (-1.20 V vs. SCE) in the first step, from which the further electron transfer to the higher reduction level of ethyl bromoacetate was not possible. Additionally, a direct electron transfer from Th-BTz-Th to ethyl bromoacetate could not occur either. In comparison, diethyl bromomalonate ($E_{red.}$ =-1.00 V vs. SCE) could be reduced by via electron transfer originated from the LUMO of TA-BT-TA, leading to the formation of the which further reacts malonate radical. with 3methylbenzofuran to form the final in the following step of the catalytic cycle. An additional model reaction using both Th-BTz-Th as the single photocatalyst system and triphenylamine as sacrificial reagent led to the formation of the final product, confirming that a direct electron transfer from Th-BTz-Th to ethyl bromoacetate was only possible in the absence of TA-BT-TA (Fig. S12). It is also worth to note that further scope

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reactions with other substrates led to high product yields using couple 2 as phototocatalyst, demonstrating a general feasibility of the cooperative photocatalyst design. It is worth to note that a possible radical chain mechanism as introduced by Yoon *et al.*⁴⁵ could be clearly ruled out due to our previous study on the influence of the oxidation potential of the photocatalysts on the reaction rate.¹⁰

The reason for the lower efficiency using the cooperative photocatalyts couple in comparison to the one catalyzed in the presence of sacrificial reagents might be caused by different factors. The multiple step electron transfer within the photocatalyst couple and further to the substrate might not be efficient due to the low concentration of the catalysts. Another important factor might be the electron transfer steps between the intermediates and the photogenerated hole. For this point, an additional study was conducted as described below.

The photo-induced electron transfer within the cooperative photocatalyst couple usually occurs based on the rather random contact between the two OSs in the liquid reaction medium. To create a non-random contact and thereby a more defined electron transfer process, we then connected Th-BTz-Th and TA-BT-TA via C6 alkyl chain as molecular bridge as a control photocatalyst. As displayed in Fig. 4c, a large molecule of Th-BTz-Th@TA-BT-TA containing both OSs in couple 2 was synthesized. Remarkably, the fluorescence intensity of Th-BTz-Th/TA-BT-TA strongly decreased after connection compared to their unconnected form as couple 2 (Fig. S8). This result suggests a strong interaction between two OSs inside the large molecule. A study on the impact of Th-BTz-Th/TA-BT-TA in couple 2 on their photocatalytic efficiency showed that the highest reaction rate was achieved by a ratio of 2:1 for of Th-BTz-Th/TA-BT-TA (Fig. 4a), which in fact corresponds to the connected molecule of Th-BTz-Th@TA-BT-TA. However, when comparing the catalytic efficiency between the connected catalyst of Th-BTz-Th@TA-BT-TA and unconnected couple 2 with the same ratio between both OSs, only a decreased conversion was observed for the connected catalyst (Fig. 4b). In spite of the better electron transfer occurred within connected system, the reduced catalytic efficiency could be likely caused by the sterical hindrance within the large molecule or the competing charge recombination process, which limits the contact between the substrate and the individual OSs in the large molecule and efficient electron transfer between the catalyst and substrate. To overcome this limitation, new molecular bridges other than the C6 alkyl chain could be needed.

Conclusions

In conclusion, we have taken the light-induced electron transfer cascade process in nature photosystems as role model and developed a new sacrificial reagent-free photoredox pathway by employing molecular organic semiconductors as cooperative photocatalyst couples. The photocatalyst couples consists of two molecular organic semiconductors with complementary HOMO/LUMO band positions, leading to enhanced photo-induced intermolecular charge transfer between the organic photocatalysts within the couple. By the cooperative photocatalyst design, no extra electron-donating sacrificial reagents are needed and a new paradigm in the photoredox reaction without undesired side reaction under the oxidation of the sacrificial reagents can be established. We believe that the cooperative photocatalyst design ought not to be limited in chemical transformation reactions; it can also be applied in the field of photocatalytic water splitting or CO_2 reduction.

Experimental

Materials and Methods

Chemicals were purchased from commercial sources and used as received without further purification. 4.7dibromobenzo[c][1,2,5]thiadiazole (97%) was bought from Combi-Blocks. Sodium tert-butoxide (97%), tert-Butanol (99.5%), Sodium borohydride (96%), 1-Bromobutane (99%), 2-(Tributylstannyl)thiophene (97%), Ethynyltrimethylsilane (98%), Sodium ascorbate (98%), Pd(PPh₃)Cl₂ (98%), diethyl bromomalonate (92%), 3-Methylbenzofuran (97%), 1-Methylindole (97%) and 3-Methylindole (98%) were bought from Sigma-Aldrich. Other common used solvents were bought from Fisher Scientific and VWR International. Glassware was dried under heating by a heating gun at 600 C and cooled under vacuum prior to use if necessary. All photocatalyst reactions were conducted using common dry, inert atmosphere techniques via a Schlenk tube. White LED (Silent Air System White OL M-018) is a product of OSA Opto Light Gmbh with an intensity of 0.07 Wcm⁻² from a distance of 5 cm. UV-Vis absorption and emission spectra were recorded on a Perkin Elmer Lambda 100 spectrophotometer and J&M TIDAS spectrofluorometer at ambient temperature, respectively. Cyclic voltammetry measurement was performed with Autolab PGSTAT204 potentiostat/galvanostat of Metrohm (a three electrode cell system): glassy carbon electrode as the working electrode, Hg/HgCl₂ electrode as the reference electrode, platinum wire as the counter electrode, and Bu₄NPF₆ (0.1 M Acetonitrile) as supporting electrolyte with a scan rate of 100 mV/s in the range of -2 eV to 2.5 eV. $^1\!H$ and $^{13}\!C$ spectra were recorded on a Bruker Avance 300 spectrometer, and d-CHCl₃ was used as solvent unless otherwise noted. GC-MS spectra were recorded on a SHIMADZU GCMS-QP2010 gas chromatograph mass spectrometer. Time-resolved photoluminescence (TRPL) on a nanosecond timescale was taken with a Streak Camera System (Hamamatsu C4742) in slow sweep mode. The excitation wavelength of 400 nm was provided using the frequency-doubled output of a commercial titanium:sapphire amplifier (Coherent LIBRA-HE, 3.5 mJ, 1 kHz, 100 fs).

Synthesis of the molecular organic semiconductor molecules

Synthesis of Th-BT-Th⁴⁴

A 100 ml Schlenk tube equipped with a stirring bar and stopper was heated under vacuum then cooling three times before 4,7-

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DOI: 10.1039/C8CY01072B Journal Name

dibromo-2,1,3-benzothiadiazole (1.76 g, 6.00 mmol), 2-(tributylstannyl)thiophene (4.76 ml, 15 mmol) and Pd(PPh₃)₂Cl₂ (201.60 mg, 5 mol%) were added. 30 ml anhydrous THF was added via syringe and the reactor was degassed under vacuum and then back filled with nitrogen three times. It was heated under 90 °C overnight, then poured into 200 ml water and extracted with CH₂Cl₂ three times. The organic layer was washed with water twice, dried by anhydrous MgSO₄, solvent was removed under vacuum. Purified with a short column then crystalized from methanol give a needle like reddish solid (1.14 g, 63%). ¹H NMR (300 MHz, CDCl₃) δ : 8.04 (dd, *J* = 3.7, 1.2 Hz, 2H), 7.88 (s, 2H), 7.39 (dd, *J* = 5.1, 1.2 Hz, 2H), 7.15 (dd, *J* = 5.2, 3.7 Hz, 2H). ¹³C NMR (300 MHz, CDCl₃) δ : 152.66, 139.36, 128.02, 127.52, 126.81, 126.02, 125.79.

Synthesis of Th-BTz-Th



Synthesis of **1a**: The suspension of 4,7-dibromobenzo[c]-1,2,5thiadiazole (2.94 g, 10 mmol) in 100 ml methanol was previously cooled to -5 $^{\circ}$ C in water-ice bath. Then sodium borohydride (8 g, 21.7 mmol) was added portion wise to the suspension. The mixture was warmed up to room temperature and stirred overnight. Then, 500 ml water was added, the mixture was extracted with CH₂Cl₂, the organic layer was washed with water twice, dried by anhydrous MgSO₄, solvent was removed under vacuum to give the final product as a pale yellow solid in 80% yield. The H-NMR was consistent with the literature and used without further purification.⁴⁷

Synthesis of **1b**: Dibromobenzene-1,2-diamine (2 g, 7.5 mmol) (**1a**) was added to a flask containing 15 ml glacial acetic acid. Sodium nitrite (1.03 g, 15 mmol) was dissolved in 10 ml water and added dropwise to the solution. The mixture was further reacted for another 3 hours before it was cooled by ice to 0 °C. The suspension was filtrated, washed with water until PH reaches 7. The crude product was dried in oven at 70 °C under vacuum overnight. The H-NMR was consistent with the literature¹ and used without further purification.

Synthesis of **1c**: To a 100 ml schlenk tube was added 4,7dibromo-2H-benzo[d][1,2,3]triazole (2.4 g, 8.6 mmol)(1b) and Nat-OBu (1.24 g, 12.9 mmol), then 20 ml methanol and 1bromobutyl(4.3 ml, 40 mmol) was added via syringe. The mixture was degassed under vacuum slightly then back filled with N₂ three times. After stirred under 90 °C for 24h, the mixture was poured into 200 ml water and extracted with CH₂Cl₂ three times. The organic layer was washed with water twice, dried by anhydrous MgSO₄, solvent was removed under vacuum. Purified with a column using CH₂Cl₂/Hexane = 1 : 1 as eluent give the final product as white solid (1.28 g, 45%). ¹H NMR (300 MHz, CDCl₃) δ : 7.37 (s, 2H), 4.72 (t, *J* = 7.4 Hz, 2H), 2.06 (m, 2H), 1.41-1.29 (m, 2H), 0.91 (t, *J* = 7.4 Hz, 3H). ¹³C NMR (300 MHz, CDCl₃) δ : 143.71, 129.51, 109.97, 57.22, 32.17, 19.82, 13.51.

Synthesis of Th-BTz-Th: A 100ml Schlenk tube equipped with a stirring bar and stopper was heated under vacuum then cooling three times before 4,7-dibromo-2-(n-butyl)-2Hbenzo[d]-[1,2,3]triazole (1 g, 3.00 mmol) (1c), 2-(tributylstannyl)thiophene (2.38 ml, 7.5 mmol) and Pd(PPh₃)₂Cl₂ (100.8 mg, 5 mol%) were added. 15 ml anhydrous THF was added via syringe and the reactor was degassed under vacuum and then back filled with nitrogen three times. It was heated under 90 °C overnight, then poured into 200 ml water and extracted with CH₂Cl₂ three times. The organic layer was washed with water twice, dried by anhydrous MgSO₄, solvent was removed under vacuum. Purified with a column using CH_2Cl_2 /Hexane = 1 : 1 as eluent give the final product as light red solid (713 mg, 70%). ¹H NMR (300 MHz, CDCl₃) δ : 8.03 (dd, J = 3.7, 1.2 Hz, 2H), 7.56 (s, 2H), 7.31 (dd, J = 5.1, 1.2 Hz, 2H), 7.12 (dd, J = 5.1, 3.7 Hz, 2H), 4.76 (t, J = 7.4 Hz, 2H), 2.16-2.07 (m, 2H), 1.44-1.34 (m, 2H), 0.94 (t, J = 7.4 Hz, 3H). ¹³C NMR (75 MHz, CDCl₃) δ : 142.10, 139.97, 128.12, 126.97, 125.55, 123.59, 122.78, 56.61, 32.08, 19.89, 13.57.

Synthesis of TA-BT-TA



Synthesis of **2a**: A 100ml Schlenk tube equipped with a stirring bar and stopper was heated under vacuum then cooling three times before 4,7-dibromobenzothiadiazole (3 g, 10.2 mmol), Pd(PPh₃)₂Cl₂ (143.2 mg, 2%), Cul (77 mg, 4%) were added. 30 ml THF and 10ml NEt₃ were added via syringe. The mixture was cooled to freeze in liquid N₂, then trimethylsilyl acetylene (4 ml, 30.6 mmol) was added. The reactor was degassed via the freeze-pump-thaw method and reacted under 50 °C overnight. Then it was poured into 100 ml water and extracted with CH₂Cl₂ three times. The organic layer was washed with water twice, dried by anhydrous MgSO₄, solvent was removed under vacuum. Purified with a column using CH₂Cl₂/Hexane = 1 : 1 as eluent give the final product as light yellow solid (2.78 g, 83%). The H-NMR was consistent with the literature reported.

Synthesis of **2b**: In a 100 ml two neck flask was added KOH methanol solution (1 mol L^{-1} , 50 ml) and 4,7-bis[2-(trimethylsilyl)ethynyl]benzothiadiazole (1 g, 3 mmol)(2a). The solution was stirred under room temperature for 2 h, and then

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it was poured into 300 ml water and extracted with CH_2CI_2 three times. The organic layer was washed with water twice, dried by anhydrous MgSO₄, solvent was removed under vacuum. Purified with a column using CH_2CI_2 /Hexane = 1 : 1 as eluent give the final product as light brown solid (420 mg, 876%). The H-NMR was consistent with the literature reported.⁴⁸

TA-BT-TA: 4,7-Diethynylbenzothiadiazole (400 mg, 2.17 mmol)(2b), 1-azidobutane (synthesis 1-azidobutane: a mixture of 10 mmol 1-bromobutane, 20 mmol sodium azide in 100 ml DMSO was stirred over night at RT, then the reaction mixture was poured into 500ml water and extracted with CH₂Cl₂ then dried by anhydrous MgSO₄, solvent was removed under vacuum used as synthesized without column purification. Caution: the excessive amount of sodium azide in water phase muss be slowly added to a solution of NaOCI before deposition.), CuSO₄ (51.4 mg, 0.32 mmol), Na-ascorbate (87 mg, 0.44 mmol), 20 ml t-BuOH, 2 ml H₂O were added into a 100 ml two nech flask. The reactor was degassed under vacuum and then back filled with nitrogen three times. The mixture was heated under 60 °C overnight, then poured into 200 ml water and extracted with CH₂Cl₂ three times. The organic layer was washed with water twice, dried by anhydrous MgSO₄, solvent was removed under vacuum. Purified with a column using $CH_2Cl_2/Acetone = 10:1$ as eluent then recrystallized from methanol give the final product as a sheet-like yellow solid (747 mg, 90%). ¹H NMR (300 MHz, $CDCl_3$) δ : 8.67 (s, 2H), 8.61 (s, 2H), 4.44 (t, J = 7.4 Hz, 4H), 2.00-1.90 (m, 4H), 1.43-1.32 (m, 4H), 0.94 (t, J = 7.4 Hz, 6H). ¹³C NMR (75 MHz, CDCl₃) δ : 152.29, 143.10, 126.08, 123.74, 122.66, 50.27, 32.37, 19.79, 13.52.

Synthesis of the connected catalyst Th-BT-Th@TA-BT-TA



Synthesis of **3a**: To a 100 ml schlenk tube was added 4,7dibromo-2H-benzo[d][1,2,3]triazole (3 g, 10.8 mmol) and Nat-OBu (1.56 g, 16.2 mmol), then 40 ml methanol and 1,6dibromohexane (8.3 ml, 54 mmol) was added via syringe. The mixture was degassed under vacuum slightly then back filled with N₂ three times. After stirred under 90 °C for 24 h, the mixture was poured poured into 200 ml water and extracted with CH₂Cl₂ three times. The organic layer was washed with water twice, dried by anhydrous MgSO₄, solvent was removed under vacuum. Purified with a column using CH₂Cl₂/Hexane = 1 : 1 as eluent give the final product as a white solid (1.82 g, 41%). ¹H NMR (300 MHz, CDCl₃) δ : 7.38 (s, 2H), 4.73 (t, *J* = 7.4 Hz, 2H), 3.33 (t, *J* = 7.4 Hz, 2H), 2.16-2.06 (m, 2H), 1.84-1.75 (multi, 2H), 1.50-1.39 (multi, 2H), 1.34-1.29 (multi, 2H). ¹³C NMR (75 MHz, CDCl₃) δ : 143.76, 129.64, 110.00, 57.24, 33.58, 32.39, 29.99, 27.54, 25.70.

Synthesis of **3b**: 4,7-Dibromo-2-(bromohexane)benzo[d][1,2,3]triazole (1.8 g, 2.27 mmol)(**3a**), sodium azide (295 mg, 4.54 mmol) were dissolved in 20 ml DMSO and stirred under room temperature overnight. Then the mixture was poured into 200 ml water and extracted with CH_2Cl_2 three times. The organic layer was washed with water twice, dried by anhydrous MgSO₄, solvent was removed under vacuum. Product was used directly for the next step without any purification.

Synthesis of 3c: In a 100 ml two neck flask was added 4,7diethynylbenzothiadiazole (184 mg, 1 mmol)(3b), 4,7-dibromo-2-(bromohexane)-benzo[d][1,2,3]triazole, CuSO₄ (24 mg, 0.15 mmol), Na-ascorbate (41 mg, 0.21 mmol), 10 ml t-BuOH, 1 ml H₂O. The reactor was degassed under vacuum and then back filled with nitrogen three times. The suspention was stirred under heating at 60 °C for two days, then it was poured into 100 ml water and extracted with CH₂Cl₂ three times. The organic layer was washed with water twice, dried by anhydrous MgSO₄, solvent was removed under vacuum. Purified with a column using $CH_2Cl_2/Aceton = 20:1$ as eluent give the final product as a yellow solid (750 mg, 76%). ¹H NMR (300 MHz, CDCl₃) δ : 8.66 (s, 2H), 8.59 (s, 2H), 7.36 (s, 4H), 4.72 (t, J = 7.4 Hz, 4H), 4.42 (t, J = 7.4 Hz, 4H), 2.16-2.07 (m, 4H), 2.00-1.94 (m, 4H), 1.42-1.39 (m, 8H). ¹³C NMR (75 MHz, CDCl₃) δ : 152.26, 143.74, 143.14, 129.65, 126.15, 123.82, 122.58, 109.98, 57.12, 50.30, 30.11, 29.86, 25.94.

Th-BT-Th@TA-BT-TA: In a capped vial, 3c (750 mg, 0.76 mmol), 2-(tributylstannyl)thiophene (1.64 ml, 4.56 mmol) and Pd(PPh₃)₂Cl₂ (53 mg, 10 mol%) were added. 10 ml anhydrous THF was added via syringe and the reactor was degassed via bubbling nitrogen for 5 min. The vial was put in a microwave reactor (120 °C, 0.8 T) for 1 h, then poured into 100 ml water and extracted with CH₂Cl₂ three times. The organic layer was washed with water twice, dried by anhydrous MgSO₄, solvent was removed under vacuum. Purified with a column using $CH_2Cl_2/Acetone = 20:1$ as eluent give the final product as a yellow-red solid (403 mg, 53%). ¹H NMR (300 MHz, CDCl₃) δ : 8.59 (s, 2H), 8.55 (s, 2H), 7.98 (dd, J = 3.7, 1.2 Hz, 4H), 7.52 (s, 4H), 7.29-7.27 (m, 4H), 7.08 (dd, J = 5.1, 3.7 Hz, 4H), 4.74 (t, J = 7.4 Hz, 4H), 4.39 (t, J = 7.4 Hz, 4H), 2.16-2.12 (m, 4H), 1.98-1.93 (m, 4H), 1.45-1.42 (m, 8H). ^{13}C NMR (75 MHz, CDCl_3) δ : 152.21, 143.15, 142.09, 139.84, 128.10, 126.92, 126.04, 125.63, 123.72, 123.55, 122.82, 122.59, 56.52, 50.30, 30.15, 29.70, 26.03, 25.97.

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Catalysis Science & Technology Accepted Manuscript

DOI: 10.1039/C8CY01072B

Journal Name

General procedure for the electron-donating sacrificial reagentfree and photocatalytic C-C bond formation reactions using cooperative photocatalyst systems

A 25 ml Schlenk tube equipped with a stirring bar and stopper was heated under vacuum then cooling for one time and then back filled with argon before the electron-rich heteroaromates (3-methyl benzofuran et al.) (0.38 mmol, 1 equiv), cooperative photocatalyst couple system (declared amount and ratio as described in Table 2), diethyl bromomalonate (0.76 mmol, 2.0 equiv), 2.5 ml DMF were added. The reactor was degassed via the freeze-pump-thaw method and irradiated under a white LED lamp (0.07 Wcm⁻²) in a water bathe (see photograph in ESI) for 24 h. The conversion was determined via GC-MS and the pure product was obtained via chromatography.

Comparison control experiment for the C-C bond formation reactions between 3-methylbenzofuran and ethyl bromoacetate using Th-BTz-Th as catalyst and 4methoxyltriphenylamine as electron-donating sacrificial reagent

A 25 ml Schlenk tube equipped with a stirring bar and stopper was heated under vacuum then cooling for one time and then back filled with argon before 3-mehtylbenzofuran (0.38 mmol, 1 equiv), 4-methoxyltriphenylamine (0.76 mmol, 2 equiv), ethyl bromomalonate (0.76 mmol, 2.0 equiv), Th-BTz-Th (1 mol%), 2.5 ml DMF were added. The reactor was degassed via the freeze-pump-thaw method and irradiated under a white LED. The reaction time was determined by GC-MS when the signal of starting material totally disappeared. After the reaction was completed, the mixture was poured into 20 ml water and extracted by CH₂Cl₂, the organic layer was dried by anhydrous MgSO₄, and the solvent removed under vacuum. The crude product was purified on silica gel using the indicated solvent system to offer the desired product. ¹H NMR (300 MHz, CDCl₃) δ: 7.40-7.32 (m, 2H), 7.20-7.11 (m, 2H), 4.11 (q, J = 7.4 Hz, 2H), 3.69 (s, 3H), 2.13 (s, 3H), 1.19 (t, *J* = 7.4 Hz, 3H). ¹³C NMR (75 MHz, CDCl_3) $\delta:$ 169.14, 154.15, 146.12, 129.91, 123.92, 122.23, 119.13, 112.81, 110.95, 61.35, 32.84, 14.19, 7.97. Molecular weight C₁₃H₁₄O₃: calculated 218.09, found 218.10.

Acknowledgements

The authors thank the Max Planck Society for financial support. L.W. thanks the China Scholarship Council (CSC) for scholarship. The work of I. R. is part of the research program of the Dutch Polymer Institute (project #763).

Conflict of interest

The authors declare no conflict of interest.

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