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Biscoumarin-containing acenes as stable organic semiconductors for photocatalytic oxygen reduction to hydrogen peroxide†

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Conversion of solar energy into chemical energy in the form of hydrogen peroxide and other reactive oxygen species has been predicted to be an efficient strategy, yet few organic materials systems support these types of photochemical conversion reactions. Herein we report a simple synthetic route to yield biscoumarin-containing acenes, semiconducting small molecules with exceptional stability and tunable electrochemical and electrical properties. We find that these semiconductors are photo(electro)catalysts capable of reducing oxygen to hydrogen peroxide. Visible light irradiation of thin films on insulating substrates in pure water results in H_2O_2 photogeneration with water as the sacrificial electron donor. Thin films on conducting substrates are robust catalytic photocathodes for producing H_2O_2 . These semiconductor photoelectrodes retain their catalytic properties in a pH range from 2–13. Photocatalytic or photoelectrocatalytic deployment of biscoumarin-containing acenes does not lead to measurable degradation. This work demonstrates a strategy to synthesize stable organic semiconductors not only suitable for thin-film electronic devices but also next-generation photocatalytic concepts.

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1. Introduction

Demand for robust small molecule semiconductors with improved and novel functionality has encouraged materials chemists to search for solutions in many branches of organic chemistry. One of the most fruitful approaches is to tap into the extensive literature on dyes and pigments, which has led to a present renaissance of many old and forgotten molecular structures. Many classical chromophores such as indigos, quinacridones, diketopyrrolopyrroles, epindolindiones, such as indigos, and indanthrones are purposed as functional chromophores due to their exceptional photophysical properties and potential applications in organic

electronics. As organic light-emitting diode technology has achieved large-scale commercial success and organic photovoltaic technologies edge closer to technological maturity, finding next generation applications for organic semiconductors becomes pertinent. Two such emerging fields for organic semiconductors are bioelectronics^{13,14} and (photo) catalysis. For both of these application directions, stability in oxygenated and aqueous environments is critical. Many of the well-known organic semiconductors are unfortunately unstable under these conditions.

In our work, we have focused on catalytic oxygen reduction to H_2O_2 . Photochemical generation of H_2O_2 , a high energy-density compound, has been suggested as an alternative solar-to-fuel concept to compete with better-known hydrogen evolution or CO2 reduction concepts.18-21 A number of semiconductor photocatalysts, most notably ZnO, are known to photochemically reduce O₂ to H₂O₂ while oxidizing a range of sacrificial electron donors, such as phenol, alcohols, etc. 22-24 On the other hand it was recently reported that an archetypical carbonyl pigment, perylenetetracarboxylic diimide, can achieve the oxygen evolution reaction in the presence of some sacrificial electron acceptors.25 A structurally-related naphthalene diimide polymer was also shown to function as a photoanode for water oxidation.26 Several works, on the other hand have focused on researching sustainable photocatalytic systems for H₂O₂ production via oxygen reduction. 27,28 H₂O₂ can be photochemically generated to some extent by TiO2 via water oxidation.29,30

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[†] Electronic supplementary information (ESI) available. CCDC 1531580, 1531582, 1531581, 1531579, 1531584 and 1531583 contain the supplementary crystallographic data for compounds **8**, **9** and **10** at 100 K and 298 K. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c7ta05882a

The overall photocatalytic process of oxygen reduction to H₂O₂ accompanied by oxidation of water has been recently observed for the first time by Hirai et al. using a graphitic carbon nitride (g-C₃N₄) semiconductor, 31,32 and recently found also to be catalyzed by graphene oxide.²⁸ In 2016, we showed that acridone pigments like epindolidione and quinacridone can photocathodically reduce O₂ to H₂O₂, with the results implicating carbonyl groups as the critical catalytic site.33 This motivated us to evaluate carbonyl pigments as potential catalysts: while searching for better photocatalysts we turned our attention to coumarins.³⁴ Coumarins have been extensively studied in the past as colorants, and have found application notably in dye lasers.35 Recently we discovered an oxidative ring closure reaction of biscoumarins that can lead to large fused-ring acene-like structures.36 Preliminary observations have suggested that these compounds possess high stability under various conditions. Their general structural formula suggests that strong π - π stacking in the solid state could lead to effective charge transport. The aim of this study was to synthesize such fused biscoumarins and to study their performance in photochemical generation of H₂O₂. Herein we would like to present the results of this study.

We have adapted this synthetic scheme (Fig. 1a) to yield π extended structures 8-10 (Fig. 1b). X-ray crystal structures reveal that 8 and 9 are relatively planar extended π systems, while the larger 10 is a propeller-shaped molecule (Fig. 1c). In this work, we find that these large acene-like molecular semiconductors that are intrinsically stable and display impressive semiconductor catalytic properties in a wide pH range: 2-13. The two electron-withdrawing coumarin units flank the carbocyclic core

of the molecule, resulting in effective stability against oxidative degradation. The electron-poor coumarin substituents also lead to favourable n-type conduction properties, and quasi-reversible reduction in electrochemistry, with irreversible oxidation. Increasing the size of the molecular core from a pentacene to a dibenzo [a,f] heptacene core raises the HOMO level and yields p-type behavior as well, leading to overall ambipolar transport and both quasi-reversible reduction and oxidation. We have discovered that visible light irradiation of thin films of materials 8-10 in pure water leads to H2O2 photosynthesis. Based on photoelectrochemical measurements we identified that H₂O₂ evolution proceeds via photoreduction of dissolved O2, accompanied by water oxidation. In our work we have found that our biscoumarin compounds are the first carbonyl pigments to achieve this photocatalytic transformation, only g-C₃N₄ derivatives were previously reported to give this catalytic functionality.31,32,37 The catalytic efficiency afforded by 8-10 in terms of quantity of H₂O₂ evolution rate per amount catalyst exceeds g-C₃N₄ by at least a factor of 2-4×. Aside from pure photocatalysis, materials 9 and 10 are suitable for fabrication of semiconductor photoelectrodes capable of anodic or cathodic photoelectrocatalysis. This way electrical power can be harvested alongside photosynthesis of H2O2. Depending on pH and applied potential, either photoanodic or photocathodic modes are accessible, and operation under continuous illumination for at least twelve days in a two-electrode photochemical cell is possible. From the point of view of H₂O₂ photosynthesis, 8-10 have a significant advantage over the known inorganic catalysts such as ZnO, namely the latter can only function at neutral pH.

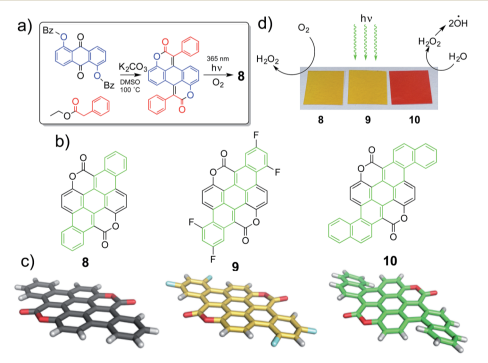


Fig. 1 (a) Synthesis scheme of π -extended biscoumarins. (b) Molecular structures of the three pigments evaluated in this work, with the acene π system highlighted in green. (c) Molecular structures in the solid state as measured by XRD. (d) Thin-films of the three pigments deposited on plastic foils photocatalytically reduce oxygen to hydrogen peroxide while water is oxidized to hydrogen peroxide, which apparently is further converted to hydroxyl radicals.

These findings of novel catalytic and electrochemical functionality in aqueous conditions is enabling both for application at the interface with biology as well as catalysis, both new frontiers for small molecule semiconductors.

2. Results and discussion

2.1. Synthesis and solid-state structure

Following the previously developed general strategy³⁶ we carried out double Knoevenagel condensation of compound **1** with esters **2**, **3** and **4** to obtain biscoumarins **5–7** in good yields (Scheme 1). Subsequently, all three biscoumarins were subjected to UV

Scheme 1 Synthesis of biscoumarins 8-10

 6π -electrocyclization³⁸ irradiation, which double causes combined with concomitant oxidation of the intermediates to aromatic compounds 8-10. This two-step strategy allowed us to prepare three structurally unique heterocyclic pigments 8-10 in overall yields in the range 29-38% (Scheme 1). It is noteworthy that in the case of biscoumarin 7 full regioselectivity was observed in the final cyclization step and reaction occurred exclusively at position 1 of the naphthalene substituent. Very poor solubility of compounds 8-10 precluded performing NMR studies and the identity and purity were established based on high-resolution mass spectrometry and combustional analysis. Compounds 8-10 were purified by repeated temperature gradient sublimation in a vacuum of $<1 \times 10^{-6}$ mbar. The main impurities were found to be uncyclized precursor or singly-cyclized derivatives. As expected, these impurities were more volatile than the desired fully-cyclized molecules and traveled to lower temperature zones of the tube. Single crystals were grown at a pressure of 1 atm by sublimation of purified material in a stream of N₂. The crystal growth zone had a temperature of 120-150 °C. Experimental and crystallographic details can be found in the ESI.† Crystal packing shows pillars held together by π - π stacking interactions (distance between mean planes of packed molecules are comparable: 3.35(5) Å in 8, 3.38(9) Å in 9 and 3.80(1) Å in 10). 8 and 9 pillars are built of alternating molecules that are flipped with respect to the mean molecular plane. Neighboring columns are kept packed through weak dipole-dipole contacts, all with the same orientation in 8 and 10 or with a herringbone layered pattern in 2. The functionalization with fluorine atoms changes packing symmetry due to a different superimposition area between molecules stacked on the same pillar and a significant tilt angle change between neighboring molecular columns (less than 1° in 8 and almost 18° in 9). In contrast with highly planar 8 and 9, peripheral benzene rings in 10 break planarity due to steric repulsion of proximal hydrogens and slightly bend the polyaromatic scaffold of the molecule. This leads to a longer π - π packing distance in 10 compared with the more planar molecules, though as it will be shown later this apparently does not have a detrimental effect on charge transport.

2.2. Optical and electrochemical properties

UV-Vis absorption and photoluminescence spectra were recorded for 8-10 in dilute solution in toluene as well as for sublimated thin films (Fig. 2). All three materials in thin films demonstrated bathochromic shifts in absorption of roughly 50 nm, accompanied with broadening of the absorption features. Nevertheless a clear vibronic structure is apparent in both solution and solid-state spectra. While there is a very small Stokes shift in solution, in solid-state the luminescence is markedly red-shifted, broad, and weak, signaling that luminescence may primarily originate from defect states in the polycrystalline films rather than molecular excitons. The optical band gaps for each material, estimated from absorption onset, were very similar, around 2.1-2.3 eV (Table 1). The stronger absorption in the green region for 10 results in the apparent red colour of the pigment, in contrast to 1 and 2, which appear yellowish-orange. While the optical properties of the three π -

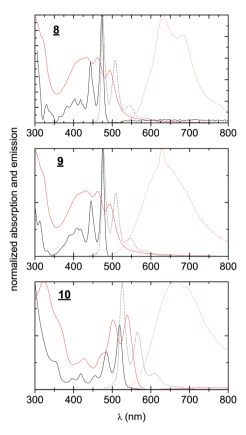


Fig. 2 UV-visible absorption and photoluminescence spectra of 8-10. Black continuous line: absorption of solutions, 0.1 mM in toluene. Black dotted line: photoluminescence of the same solutions. Red continuous line: absorption of evaporated thin films. Red dotted line: photoluminescence of the same films.

Table 1 Hydrogen peroxide photosynthesis in pure water. Literature data on best to-date results for H₂O₂ production using graphitic carbon nitride derivatives is shown for comparison

Material	Optical band gap (eV)	H_2O_2 evolution rate (µg H_2O_2 /mg catalyst/hour)
$g-C_3N_4 + PDI^{25,26}$ 8 9 10	2.4 2.2 2.3 2.1	$0.7 ext{-}0.8 \ 1.5 \pm 0.4 \ 2.5 \pm 0.2 \ 3.3 \pm 0.2$

extended biscoumarins are not remarkably different from each other, their electrochemical behaviour varies considerably (Fig. 3). We measured cyclic voltammetry (CV) of thin films evaporated on indium tin oxide substrates in acetonitrile electrolyte under inert (N_2) conditions. 8 and 9 demonstrated two quasi-reversible reduction peaks, with irreversible oxidation. The tetrafluorinated 9, as can be expected from the electronegativity of the fluorine atoms, affords a reduction onset 500 mV more positive than 8. In addition to this substantial lowering of the LUMO level, the four fluorine atoms in 9 also lower the HOMO level by 200 mV with respect to 8. Compound 10, in contrast, gave not only quasi-reversible reduction but also quasi-reversible 2-electron oxidation. This can be rationalized

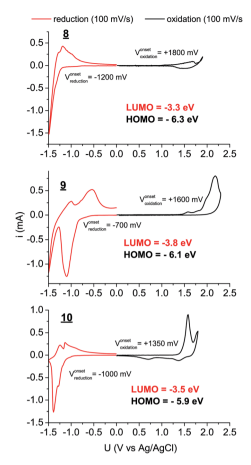


Fig. 3 Cyclic voltammograms of 8-10 deposited on ITO, functioning as the working electrode. Platinum and an Ag/AgCl wire functioned as the counter and quasi-reference electrodes, respectively. 0.1 M tetrabutylammonium hexafluorophosphate in CH₃CN was used as the electrolyte solution

by the presence of the larger extended electron-rich π -system in this compound. The Frontier molecular orbital energies are estimated (values shown in Fig. 3) assuming an NHE electrode value of -4.75 eV on the Fermi scale.39

Semiconducting properties

In order to evaluate the electrical properties of 8-10, we fabricated field-effect transistors (FETs) with the device structure shown in Fig. 4a. Compounds 8 and 9 demonstrated n-type transport, with electron mobility of 0.06 and 0.004 cm² V⁻¹ s⁻¹, respectively (Fig. 4b and c). Hole transport was not measurable with these two materials under any conditions. In contrast, 10 afforded well-balanced n- and p-type mobility. With aluminium source-drain contacts, enhanced n-type behaviour was observed (Fig. 4d), while higher workfunction gold source-drain electrodes allowed the fabrication of a device showing ambipolar behavior (Fig. 4e). The observed transport polarity for the three materials corresponds logically to the electrochemical results: 8 and 9 show only reversible reduction, and thus only n-type behavior, while 10, with both reduction and oxidation accessible electrochemically, supports

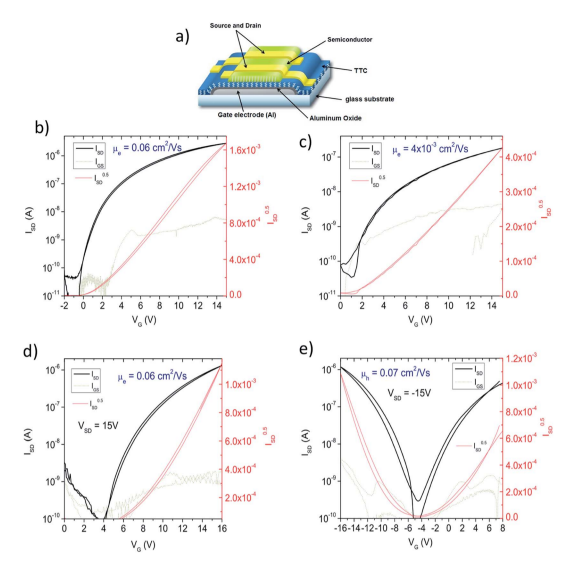


Fig. 4 Thin film field-effect transistors. (a) Schematic of the device configuration used. Aluminum with anodically-grown aluminum oxide was used (oxide thickness: 32 nm), which was passivated with a 20 nm thick layer of tetratetracontane (TTC). Materials 8-10 were evaporated to form 80 nm thick films. Source-drain contacts were then evaporated through a shadow mask to give W/L=2 mm/60 μ m. (b) nFET with material 8, with aluminum source-drain contacts. (c) nFET with material 9, using aluminum source-drain contacts. (e) Ambipolar FET with material 10, using gold source-drain contacts.

ambipolar transport. The LUMO level of each material is too high-energy to observe air-stable electron transport⁴⁰ in OFETs, however p-type transport in 3 was found to be stable over 150 days of measurements, where the devices were stored and measured in ambient conditions. Mobility declined to $0.04~{\rm cm^2~V^{-1}~s^{-1}}$ after a few days and remained stable at this value (ESI S2†).

2.4. Aqueous photo(electro)catalysis

Applications of organic semiconductors have often been plagued with stability problems, and considerable attention has been devoted to producing materials that are stable with respect to oxygen and water. ³⁴ It is therefore not surprising that organic semiconductors are not often explored for aqueous catalytic applications, in stark contrast to inorganic semiconductors, which are a major focus of renewable energy research. ^{41,42} We observed that thin films of **8–10** in pure 18 M Ω water under

ambient conditions, irradiated with a commercial white LED (30 mW cm⁻²), produced H_2O_2 , with evolution rates of 1-3 µg H₂O₂ per mg of compound per hour (Table 1). The amount of H₂O₂ was quantified using the tetramethylbenzidine/ horseradish peroxidase assay (TMB/HRP). Since the only available reagents are water and dissolved atmospheric oxygen, the apparent reaction is 2-electron reduction of O2 to H2O2, with simultaneous oxidation of water. The oxidation of the semiconductor itself, as a sacrificial donor, can be excluded on the basis that the amount of peroxide produced exceeds by at least a factor of ten the total molar quantity of semiconducting molecules in the film. The only precedent from the literature is a g-C₃N₄ derivative that was recently reported to achieve this catalytic transformation.31 In terms of catalytic performance, 8-10 are all at least twice as efficient as this previous report, which is shown for comparison in Table 1. In these reports and

our experiments only pure water is used, though in our case we perform the experiment in ambient air instead of under 1 atm of pure oxygen, further underscoring the better performance of the biscoumarin molecules. The question of the identity of the anodic process of water oxidation will be elucidated by photoelectrochemical measurements discussed in the following. After verifying the photocatalytic potential for 8-10 and determining that 9 and 10 had the best performance in terms of H₂O₂ photosynthesis, we moved on to measure the photoelectrocatalytic properties of these two materials (Fig. 5). To fabricate photoelectrodes, we evaporated 220 nm thick films of 9 or 10 on fluorine-doped tin oxide (FTO). Measuring linear sweep voltammetry with modulated on/off illumination (Fig. 5a) reveals clearly enhanced photoanodic performance for 9 versus a better photocathodic performance for 10. At low pH, both materials give photocathodic currents at voltages ≤0 V vs. Ag/ AgCl. Increasing pH leads to larger photoanodic currents. 10 gives the highest photocathodic currents, cyclic voltammetry in the dark and light at different pH shows the crossover from photocathodic to photoanodic regimes as the pH is increased (Fig. 5b). Photocathodic current is easily correlated to oxygen reduction to H₂O₂, based on the evidence that complete removal of O2 by purging with argon leads to elimination of photocathodic current, and H2O2 presence is quantified using the TMB/HRP assay. The fact that H2O2 photosynthesis occurs in pure water, as discussed above, indicates that H2O is the electron-donor in the photoanodic process, and three scenarios are in principle possible to explain the photoanodic process, given in order of increasing electrochemical potential: 4-electron oxidation of H₂O to yield O₂, two-electron oxidation of H₂O to H₂O₂, and finally single-electron oxidation of hydroxyl ions, formed from water autodissociation, to hydroxyl radicals. Photoelectrochemistry of thin films on electrodes allow elucidation of the mechanism. The oxygen evolution reaction we excluded by measuring (under N2 atmosphere) dissolved oxygen concentration during photoanodic electrolysis of a previously deoxygenated electrolyte using a Clark electrode. This leaves the possibility of 2-electron oxidation to H₂O₂. However, we find that photoanodic current in both oxygenated and deoxygenated conditions does not lead to measurable H₂O₂ generation, even after several hours of photoelectrolysis. The direct generation of hydroxyl radicals by single-electron photooxidation is not possible thermodynamically, however we conclude that the sustained photoanodic current with no measureable O2 evolution and no H₂O₂ means that peroxide is indeed produced but then rapidly degraded. It can be further oxidized to short-lived hydroxyl radical species, or these form by homolytic cleavage. We thus speculate that the dominant photoanodic process present in 8-10 must be peroxide formation followed by hydroxyl radical generation. The photoanodic behaviour is clearly complex and should be a topic of more detailed future studies.

2.5. H₂O₂-evolving photoelectrochemical cell

We found that over the course of repeated measurements, photoelectrode samples of **9** and **10** on FTO did not degrade or change in performance. Encouraged by the apparent

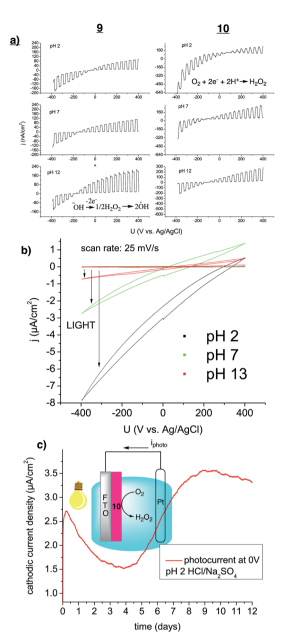


Fig. 5 (a) Photoelectrodes comprising FTO coated with **9** or **10** measured with linear sweep voltammetry with a scan rate of 25 mV s $^{-1}$ and a white LED (10 mW cm $^{-2}$) modulated at 0.125 Hz. Both photocathodic and photoanodic regimes are present in each material, the former corresponding to oxygen reduction to hydrogen peroxide, the latter to oxidation of hydroxyl ions to peroxide followed by further oxidation to hydroxyl radicals. (b) Cyclic voltammograms of **10** on FTO measured in the dark and with 60 mW cm $^{-2}$ illumination. (c) **10** on FTO measured in a two-electrode photoelectrochemical cell at pH 2 over 12 days of continuous illumination. The primary anodic reaction on Pt is chloride oxidation to Cl₂.

photoelectrochemical stability of the materials, we fabricated a simple photoelectrochemical cell comprising 220 nm film of 3 on FTO and a platinum counter electrode (Fig. 5c). The device was measured in pH 2 electrolyte using a source meter while being continuously illuminated at 200 mW cm⁻² over 12 days. In this configuration, a consistent closed-circuit photocurrent of \sim 3 μ A cm⁻² was generated. The open circuit potential of this

Fig. 6 Proposed mechanism for selective oxygen reduction to H₂O₂.

photocell was 200 mV. The anodic reaction on Pt is chloride oxidation to Cl₂, which bubbles out of solution. Fresh HCl is replenished to the cell every 24 h to ensure continuous operation. This arrangement yields a photoelectrochemical cell that produces photovoltaic power while accumulating hydrogen peroxide and chlorine as chemical products. Not only did the device function without any apparent degradation, the photocathodic current was actually found to increase over the course of 4–5 days before levelling off. This behaviour we attribute to exposure of more catalytically-active sites at the surface, a phenomenon found to occur recently for acridone pigments.³³

3. Conclusion

Based on combining observations herein with evidence from previous works, we believe we can make some conclusions

concerning organic catalysts for H2O2 production: it appears that the presence of duel carbonyl functional groups is the crucial catalytic site that allows the selective 2e⁻/2H⁺ reduction of oxygen to H2O2. In our previous findings27 on acridones (epindolidione and quinacridone) as H₂O₂-evolving photoelectrocatalysts, which contain two aromatic amines and two carbonyl functions, we postulated that the two-electron twoproton photoreduction of the pigment generated a reduced leuco species at the surface, which initiates nucleophilic attack on O2 in the same manner as is known for the industrial anthraquinone synthesis43 of H2O2. We found that N-methylation of epindolidione did not preclude O2 reduction to H2O2, suggestion that the NH group is not critical for this reaction, but rather only the carbonyl groups. In the work outlined here, we found that biscoumarin compounds containing two carbonyl functions are photocatalysts for producing H₂O₂. In the work of Hirai et al., the authors found that only when g-C₃N₄ was functionalized with pyromellitic diimide, a unit with 4 carbonyl groups, was H₂O₂ evolution observed - other g-C₃N₄ materials did not accomplish this reaction.25 This strongly suggests that the well-known quinone-like 2e⁻/2H⁺ electrochemistry of carbonyl pigments is at the heart of all these observations of H₂O₂ photosynthesis. Based on this, we outline our proposed mechanism, illustrated for compound 10, in Fig. 6. In conclusion, we have reported a synthetic scheme to a family of organic small molecule semiconductors with competitive semiconductor benchmark performance and novel catalytic ability. The application of organic semiconductors as catalytic materials in their own right is a nascent field where the body of knowledge in dye and pigment chemistry can be used to achieve next generation catalytic technologies. We believe the work we have presented will be inspiring to move this concept forward.

4. Experimental section

4.1 Synthesis

1,7-Bis(3,5-difluorophenyl)benzo[1,2,3-de:4,5,6-d'e']dichro mene-2,8-dione (6). To a mixture of 1,5-di(benzoyloxy)anthraquinone (896 mg, 2 mmol) and K2CO3 (2.76 g, 10 mmol) in 20 ml of DMSO, and methyl 2-(3,5-difluorophenyl)acetate (3, 2.2 g, 6 mmol) was added under an argon atmosphere and the resulting suspension was stirred at 100 °C for 1 hour. The reaction mixture was poured into an aqueous solution of acetic acid (0.2%, 300 ml) and the suspension was cooled down. The precipitate was filtered off and washed with ethanol. Recrystallization from CHCl₃/EtOH gave the pure biscoumarin as yellow solid (694 mg, 68%). 1 H NMR (CDCl₃, 500 MHz): δ 7.42– 7.46 (m, 2H, Ar), 7.33 (t, J = 8.2 Hz, 2H, Ar), 7.15–7.19 (m, 2H, Ar), 6.93–7.02 (m, 6H, Ar) 13 C NMR (CDCl₃, 151 MHz): δ 162.7– 165.0 (m), 160.3, 152.0, 139.2, 138.5 (t, J = 10.1 Hz), 130.7, 127.0, 123.9, 128.5, 119.1, 118.2, 113.3–113.5 (m), 105.0 (t, J = 25.2 Hz). HRMS (EI): m/z calculated for $C_{30}H_{12}O_4F_4$ [M⁺] = 512.0672; found: 512.0676 mp > 250 °C. Rf (hexane/DCM, 1:4): 0.54.

1,7-Di(naphthalen-2-yl)benzo[1,2,3-de:4,5,6-d'e']dichromene-2,8-dione (7). To a mixture of 1,5-di(benzoyloxy)anthraquinone (448 mg, 1 mmol) and K_2CO_3 (1.38 g, 10 mmol) in 10 ml of DMSO and t-butyl 2-(naphthalen-2-yl)acetate (4, 1.45 g, 6 mmol)

was added under an argon atmosphere and the resulting suspension was stirred at 100 $^{\circ}\text{C}$ for 1 hour. The reaction mixture was then poured into an aqueous solution of 0.2% acetic acid (300 ml), followed by addition of 200 ml of DCM. The resulting mixture was separated and the water phase was extracted 3 times with DCM. Next the organic extracts were combined and dried with anhydrous Na2SO4. After drying with anhydrous Na₂SO₄ the organic phase was filtered through a short DCVC column and eluted with DCM. Fractions containing biscoumarin were combined and after concentration, ethanol (30-40 ml) was added. After evaporating DCM the remaining organic residue was cooled down. The precipitate was filtered off and washed with ethanol yielding 7 as a yellow solid (293 mg, 54%). ¹H NMR (CDCl₃, 500 MHz): δ 7.97 (d, J = 8.6 Hz, 2H, Ar), 7.92 (d, J = 8.1 Hz, 2H, Ar), 7.86 (d, J = 8.1 Hz, 2H, Ar), 7.52-7.60 (m, 4H, Ar), 7.49 (d, J = 8.1 Hz, 2H, Ar), 7.33 (dd, J =8.0 Hz, J = 0.8 Hz, 2H, Ar), 7.15 (dd, J = 7.9, J = 0.8 Hz, 2H, Ar), 7.08 (t, J = 8.1 2H, Ar). ¹³C NMR (CDCl₃, 151 MHz): δ 161.5, 151.8, 138.8, 133.8, 133.4, 132.9, 130.0, 129.8, 129.3, 128.5, 127.9, 127.7, 127.5, 127.2, 127.1, 126.5, 125.9, 118.8, 118.3. HRMS (EI): m/z calculated for $C_{38}H_{20}O_4$ [M⁺] = 540.1362; found: 540.1356. Mp > 250 °C. Rf (hexane/DCM, 1:4): 0.58.

General photocyclization of biscoumarins. 2 mmol of biscoumarin was dissolved in THF (1 L) and photoirradiated (365 nm) in photoreactor (specially assembled for these reactions, see ESI† for details) for 24 h at room temperature. The reaction volume was reduced to around 200 ml and resulting precipitate was filtered off and washed with THF to give fused derivatives.

Pentacene[5,6,7-*cdef*:12,13,14-*c'd'e'f'*]dichromene-2,10-dione (8). Orange solid (454 mg, 52%). Anal. calcd for $C_{30}H_{12}O_4$: C, 82.56; H, 2.77; found: C, 82.81; H, 2.65. HRMS (EI): m/z calculated for $C_{30}H_{12}O_4$ [M⁺] = 436.0736; found: 436.0732 mp > 250 °C.

4,6,12,14-Tetrafluoropentacene[5,6,7-*cdef*:12,13,14-*c'd'e'f'*] **dichromene-2,10-dione** (9). Orange solid (438 mg, 43%). Anal. calcd for $C_{30}H_8F_4O_4$: C, 70.88; H, 1.59; F, 14.95 found: C, 71.05; H, 1.39; F, 14.80. HRMS (EI): m/z calculated for $C_{30}H_8O_4F_4$ [M^{+}] = 508.0359; found: 508.0364 mp > 250 °C.

 $\begin{aligned} \textbf{Dibenzo}[3,4:10,11] & \textbf{pentacene}[5,6,7\text{-}cdef\text{:}12,13,14\text{-}c'd'e'f'] \\ \textbf{dichromene-10,20-dione} & \textbf{(10)}. \text{ Red solid (727 mg, 68\%). Anal.} \\ & \textbf{calcd for } C_{38}H_{16}O_4\text{: C, 85.07; H, 3.01; found: C, 85.18; H, 2.96.} \\ & \textbf{HRMS (EI): } \textit{m/z } \textbf{calculated for } C_{38}H_{16}O_4 \left[\textbf{M}^+ \right] = 536.1049; found: 536.1053. \text{ Mp} > 250 \, ^{\circ}\text{C.} \end{aligned}$

4.2 Thin film device fabrication

8–10 were processed by sublimation in vacuum ($\leq 1 \times 10^{-6}$ mbar) in a custom-built organic thin film evaporator from resistively-heated alumina crucibles. Rate was controlled by quartz crystal microbalance. For thin film transistor devices, the semiconductor material was evaporated at a rate of 0.2–0.3 Å s⁻¹ to a total thickness of 80 nm. The gate structures comprised aluminum with an anodically-grown 32 nm thick alumina layer passivated with a 20 nm thick tetratetracontane layer prepared as reported previously. Aluminium or gold source-drain electrodes were evaporated at a rate of 5–10 Å s⁻¹ through a stainless steel shadow mask to give W/L=2 mm/60 μ m.

4.3 Photo(electro)catalysis

Photocatalysis in pure H_2O (18 $M\Omega$) was carried out by evaporating 220 nm of 1-3 with a fast rate (2-6 Å s^{-1}) onto PET foil. Following coating, 1×1 cm² squares were cut out of the foil and placed into a polystyrene 24-well culture plate. Each cell was then filled with a 2 ml volume of water, and the 24-well plate was placed onto an array of white LEDs, with one LED illuminating each well with a light intensity of 30 mW cm⁻². H₂O₂ concentration was measured using the tetramethylbenzidine/ horseradish peroxidase assay, according to the exact procedure we have previously reported.33 Photoelectrocatalysis experiments were performed using conductive fluorine-doped tin oxide (FTO) as an optically transparent substrate, with 220 nm thick evaporated films prepared in the identical way as for photocatalytic experiments. An IPS Elektroniklabor GmbH PGU 10 V - 1 A potentiostat was used for all three-electrode measurements. The FTO/pigment served always as the working electrode, while a graphite rod and Ag/AgCl wire were used as counter and quasireference electrodes, respectively. In two-electrode photoelectrochemical cell experiments, the counter electrode was replaced with a platinum foil and a Keithley 2400 source-meter was used to record current at short circuit conditions (0 V applied). In all aqueous experiments ionic strength of electrolytes was kept constant at 0.1 M by using Na₂SO₄ with suitable addition of HCl or NaOH to adjust pH. As indicated in the text, three types of light sources were used for different experiments: white LEDs as described before, a tungsten halogen lamp with an irradiance of 60 mW cm⁻² and finally for the long-term twoelectrode photoelectrochemical cell we used a 500 W Xe lamp, with a substantial UV component in order to subject our samples to harsh conditions to test their resistance to degradation.

Conflicts of interest

The authors declare no conflicts of interest.

Acknowledgements

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