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Organic photoelectrode engineering: accelerating photocurrent generation *via* donor—acceptor interactions and surface-assisted synthetic approach†

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Conventional photoelectrocatalysts composed of precious metals and inorganic elements have limited synthetic design, hence, hampered modularity of their photophysical properties. Here, we demonstrate a scalable, one-pot synthetic approach to grow organic polymer films on the surface of the conventional copper plate under mild conditions. Molecular precursors, containing electron-rich thiophene and electron-deficient triazine-rings, were combined into a donor-acceptor π -conjugated polymer with a broad visible light adsorption range due to a narrow bandgap of 1.42 eV. The strong charge push-pull effect enabled the fabricated donor-acceptor material to have a marked activity as an electrode in a photoelectrochemical cell, reaching anodic photocurrent density of 6.8 μ A cm⁻² (at 0.6 V vs. Ag/AgCl, pH 7). This value is 3 times higher than that of the model donor-donor thiophene-only-based polymer and twice as high as that of the analogue synthesized in bulk using the heterogenous CuCl catalyst. In addition, the fabricated photoanode showed a 2-fold increase in the photoelectrocatalytic oxygen evolution from water upon simulated sunlight irradiation with the photocurrent density up to 4.8 mA cm⁻² (at 1.0 V vs. Ag/AgCl, pH 14). The proposed engineering strategy opens new pathways toward the fabrication of efficient organic "green" materials for photoelectrocatalytic solar energy conversion.

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Introduction

Artificial photosynthesis is a key technology for clean, sustainable, fossil-fuels-free energy generation. The conversion of abundant solar energy into a useful feedstock or chemical fuel using a semiconducting photocatalyst has attracted considerable attention in the scientific community. In the light of that, photoelectrochemical (PEC) water splitting into oxygen and

hydrogen offers a great potential for clean and secure energy

production without dependence on finite and non-renewable resources.² In this process the harvested light excites charge

carriers (electrons and holes) in the semiconducting material,

There is a considerable number of reports about efficient PEC systems based on transition metal oxides, ⁵⁻⁷ metal chalcogenides, and dichalcogenides. ⁸⁻¹¹ Despite their good performance, the majority of these photoactive materials are still suffering from low modularity, narrow light absorption range and require complicated and costly synthetic approaches, such as atomic layer deposition, solvothermal synthesis or solution-based deposition with subsequent high-temperature annealing. ^{12,13} In that respect, organic conjugated semiconducting polymers emerged as a promising alternative for the fabrication of the materials for solar energy harvesting. ¹⁴⁻¹⁶ The metal-free fully organic molecular backbone enables precise tuneability

which subsequently promotes the hydrogen evolution (HER) and oxygen evolution (OER) half-reactions at the catalyst–electrolyte interface.³ Therefore, the ideal photocatalyst should possess a broad light absorption range, facilitate fast interfacial charge transfer without rapid charge recombination and be stable under operating conditions.⁴ Moreover, to meet industrial requirements the active material has to be fabricated and easily scaled up from low-cost and readily available precursors.

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of photophysical properties, such as bandgap and charge transport, thus, empowering these materials to be utilised as photocatalysts in light-driven H₂ production.¹⁷ In 2008 graphitic carbon nitride (g-C₃N₄) became the first example of metal-free synthetic photocatalyst, which was capable to operate under visible light irradiation.18 The wide optical bandgap of 2.7 eV, however, severely limits the light absorption range of g-C₃N₄ (up to 460 nm), therefore, numerous attempts to modify this material have been made to improve its photocatalytic activity.19-21

Along with g-C₃N₄, several other classes of organic porous materials, such as covalent organic frameworks (COFs), covalent triazine-based frameworks (CTFs), and conjugated microporous polymers (CMPs) have emerged as promising materials for photocatalytic water splitting reaction.²²⁻²⁴ On one hand, typical synthetic protocols of COFs and CTFs often involve hightemperature/high-pressure conditions, such as solvothermal reaction in pressurised vessels, or ionothermal/superacidcatalysed cyclotrimerization.^{25,26} Not every molecular precursor can withstand such harsh synthetic conditions. On the other hand, the relatively mild solution-processed bulk polymerisation of CMPs leads to materials with intertwisted molecular backbone struts and, as a consequence, numerous structural defects, which can inhibit the efficient transfer and separation of the photoexcited charge carriers.²⁷ Therefore, it is essential to find a reliable, easy scalable synthetic approach for the fabrication of semiconducting conjugated polymer photocatalyst with improved photophysical properties.

In 2010, Li and co-workers reported a copper-foil catalysed synthesis of a carbon-rich diacetylene-containing network graphdiyne - a structural analogue to the theoretically predicted graphyne,28 which is believed to exhibit unique optical, electronic and mechanical properties.29 Recently, Schwarz et al. employed a similar synthetic strategy to fabricate photocatalytically active triazine-based analogue of graphdiyne triazine-based graphdiyne (TzG) framework.30 Here, the copper foil is playing a role of both a catalyst and a surface template, that allows the growing of large-scale and easy-processable polymer films. The inclusion of nitrogen heteroatoms creates an additional charge push-pull effect, thus leading to a comparatively narrow bandgap of 1.91 eV and improved electron transfer processes with the ability to evolve H₂ gas from water under light irradiation without the need for a noble metal cocatalyst. However, the use of acetylenic organic polymer frameworks in PEC systems, which combines the benefits of solar energy harvesting and electrochemical water splitting, remains largely unexplored. A recent example from Zhang et al. shows the preparation of poly(1,3,5-triethynylbenzene) (PTEB) with incorporation of thieno[3,2-b]thiophene unit via coppersurface-mediated polycondensation.31 Reported materials have notable activity as photoelectrodes in PEC devices and show promise in light-driven hydrogen evolution reaction.

Herein, we present a fully organic π -conjugated triazinethiophene (Tz-Th)-based polymer (denoted as Tz-Th@Cu). To synthesise this material, we employed the following engineering strategy: (1) we use a copper foil as both a catalyst and a template to grow a polymer film in a basic solution under mild

conditions (60 $^{\circ}$ C, ambient pressure); (2) we predesign the C_3 symmetrical molecular precursor that is able to form a 3D polymer network with an extended π -conjugated backbone; (3) we incorporate into the molecular building block both electronaccepting triazine (C3N3) and electron-donating thiophene repeating units that give the resulting material an additional charge push-pull effect that narrows down the optical band gap to 1.42 eV, and extends the light absorption range up to 845 nm. The D-A polymer concept is believed to improve not only the optical properties, such as the bandgap, but also benefit toward efficient transfer and separation of photoexcited charge carriers, thus leading to better photocatalytic performance of the D-A materials.32 To demonstrate the benefits of D-A polymer architecture, we also synthesised a framework composed of only repeating electron-rich benzotrithiophene core as a material for comparison (denoted as BTT@Cu). As expected, the D-A Tz-Th@Cu network showed better overall performance as a photoelectrode in the PEC cell with marked saturated photocurrent density up to 6.8 μA cm⁻² (at 0.6 V vs. Ag/AgCl reference electrode). Given the highest photoelectrochemical activity of the fabricated materials at a potential range between 0-0.8 V (vs. Ag/AgCl, pH = 7), the electrodes can potentially be employed as photoelectrocatalysts for OER. As a proof-of-concept, we demonstrated that D-A Tz-Th@Cu shows a 2-fold increase in the photocurrent density under simulated sunlight illumination (up to 4.8 mA cm⁻² at 1.0 V vs. Ag/AgCl), which is more than 10-times higher than that of D-D BTT@Cu (0.46 mA cm⁻² at 1.0 V vs. Ag/AgCl). Our results demonstrate that organic conjugated polymers built by the D-A approach and via a robust and easy-scalable Cu-surface-catalysed route can be used to fabricate highly active photoelectrodes for PEC devices.

Results and discussion

Materials synthesis

Molecular precursors 2,4,6-tris(5-ethynylthiophen-2-yl)-1,3,5triazine (Tz-Th) and 2,5,8-triethynylbenzo[1,2-b:3,4-b':5,6-b"]trithiophene (BTT) were prepared by similar reaction protocols (Fig. 1a, Schemes S1 and S2†). In brief, the respective brominated synthon was reacted with trimethylsilyl-acetylene under Negishi cross-coupling reaction conditions. Subsequently, the obtained intermediate was subjected to the deprotection reaction in the presence of a base (aqueous NaOH in the case of Tz-Th, and K₂CO₃ in the case of BTT, more details can be found in the Experimental section of the ESI†).

We employed "wet" chemistry approach to fabricate polymeric materials by using a conventional copper plate that acts both as a surface template and a catalyst for polymeric network formation. The polymer growth reaction is based on the homocoupling of tris-ethynyl-substituted monomer and can be governed by two possible reaction pathways: (1) Glaser-type (Glaser) polycondensation of terminal alkynes; (2) [2 + 2 + 2]cyclotrimerization (cyclo) of carbon-carbon triple bonds (Fig. 1a).30 At the initial point the synthons have to get absorbed at the catalytically active copper surface, and, due to their increased local concentration at the solid-liquid interface, should form the statistically more preferred [2 + 2 + 2] product.

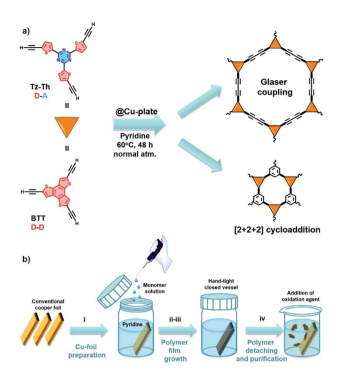


Fig. 1 (a) Synthetic pathways toward the preparation of Tz-Th- and BTT-based polymers on the surface of a conventional copper plate (@Cu-plate). There are two possible mechanisms that can occur during the reaction – Glaser-type coupling or [2+2+2] cyclotrimerization of terminal alkenes; (b) schematic representation of the synthetic procedure that includes: (i) preparation of the Cu-foil, (ii) submerging of the Cu-foil in pyridine and addition of the monomer, (iii) reaction in the hand-tight closed vessel at 60 °C for 48 h; (iv) detaching of the polymer film from Cu-foil with subsequent purification and drying.

Further during the reaction Cu-ions are leached into the solution due to passivation and solvation effects, and therefore will promote the diffusion-controlled coupling of the terminal alkyne groups via the Glaser mechanism.33 For both Tz-Th@Cu and BTT@Cu we used pyridine that simultaneously acts as a solvent and a base and the reaction mixture was kept under an ambient atmosphere in a hand-tight-closed vessel at 60 °C for 48 h (Fig. 1b). After completion of the reaction, the polymer film can be easily released from the copper surface via the addition of 0.5 M aqueous solution of (NH₄)₂S₂O₈ as a mild oxidative agent (Fig. S1†). An average polymer film thickness of $\sim 1 \mu m$ for Tz-Th@Cu and \sim 1.5 μ m for BTT@Cu was obtained after 48 h, which is evidenced by scanning electron microscopy (SEM, Fig. S2 and S3†). Using atomic force microscopy (AFM) we established that the polymer film growth almost stopped after 24 h of reaction time, which can be explained by the limited diffusion path length of the leached copper species (Fig. S4 and S5†). The reaction was carried out for an additional 24 h to achieve a homogeneous film structure.

Structural characterisation

We probed the chemical make-up of **Tz-Th-** and **BTT-**based networks by performing ¹³C cross-polarization magic-angle-

spinning (CP-MAS) solid-state NMR (ssNMR) spectroscopy (Fig. 2a). In the case of **Tz-T@Cu** we observed characteristic triazine-ring carbon signals at \sim 170 ppm. In addition, two distinguishable signals at \sim 140 and 130 ppm confirm the presence of other structural motifs – thiophene and phenyl rings. The ssNMR spectrum of **BTT@Cu** shows a broad feature between 130–150 ppm, which is indicative of the fused benzotrithiophene core.³⁴ Moreover, both materials have a weak signal at \sim 80 ppm corresponding to the carbon–carbon triple bonds. This phenomenon points toward the twinned growth of Glaser-type and [2 + 2 + 2] cyclotrimerization heterostructures in both polymers, which was also observed before in the case of TzF/TzG system.³⁰

We analysed the elemental composition of the synthesised polymers by combustion elemental analysis (EA). All materials have close to ideal C/N and C/S ratios, which proves the high degree of polymerisation reaction (Tables S1 and S2†). The inductively coupled plasma optical emission spectrometry (ICP-OES) and energy dispersive X-ray spectroscopy (EDS) showed a residual content of P(0.35-0.38 wt%) and Cu (0.3-2.49 wt%), that might arise from the preparation of molecular precursors and Cu-catalysed polycondensation reaction (Tables S1-S4†). We corroborate these findings by performing the thermogravimetric analysis (TGA) under the oxidative atmosphere, which revealed the incombustible inorganic residues from 0.66 to 5.71 wt% (Fig. S6a†). Noteworthy, all prepared materials have excellent structural integrity and, therefore, high thermal stability, which is proved by TGA performed under inert conditions - more than 40 wt% of polymers are remaining after heating up to 1000 °C (Fig. S6b†).

We further analysed the chemical states of the main composition elements (C, N, and S) using high-resolution photoelectron spectroscopy (XPS, Fig. S7 and S8†). Deconvolution of the C 1s region for Tz-Th@Cu resulted in three peaks at 282.6 eV, 284.5 eV, and 286.9 eV, which can be attributed to the sp² hybridized carbons of the phenyl rings (C-C_{phenyl}), thiophene ring (C-S=C)/adventitious (Cadv) carbons and triazine ring carbons (C-N=C) (Fig. S7b†). 35,36 Deconvolution of the N 1s region identified one peak with the binding energy of 397.4 eV, which can be assigned to nitrogen atoms in C_3N_3 ring (C-N=C) (Fig. S7c†). In the S 2p region, we observed two dominant peaks at 162.4 eV and 163.6 eV, which are characteristic for the S 2p_{3/2} and S 2p_{1/2} spin-orbit components of sulfur in thiophene units (C-S=C) (Fig. S7d†).36 The XPS spectrum of BTT@Cu confirmed the presence of carbon and sulfur as main structural elements with 3 carbon signals in the C 1s region at 282.4 eV, 284.9 eV, and 288.4 eV (C-C_{phenyl}, C-S=C/C_{adv}, and C-O/C=O) and 2 main peaks in the S 2p region at 164.04 and 165.2 eV (2p_{3/2} and S $2p_{1/2}$) (Fig. S8b and c†).

The high degree of network-forming reactions was confirmed via Fourier-transform infrared (FT-IR) spectroscopy – the sharp peak of C \equiv C–H at 3270–3280 cm $^{-1}$ for both **BTT** and **Tz-Th** precursors is absent in the spectra of the respective polymers (Fig. 2b and c). Moreover, there are no strong features which can point toward free OH- and NH-groups of some oxidized or hydrolysed fragments in the region between 3000 and 3600 cm $^{-1}$. The sharp signals at 1500 cm $^{-1}$ and 805 cm $^{-1}$

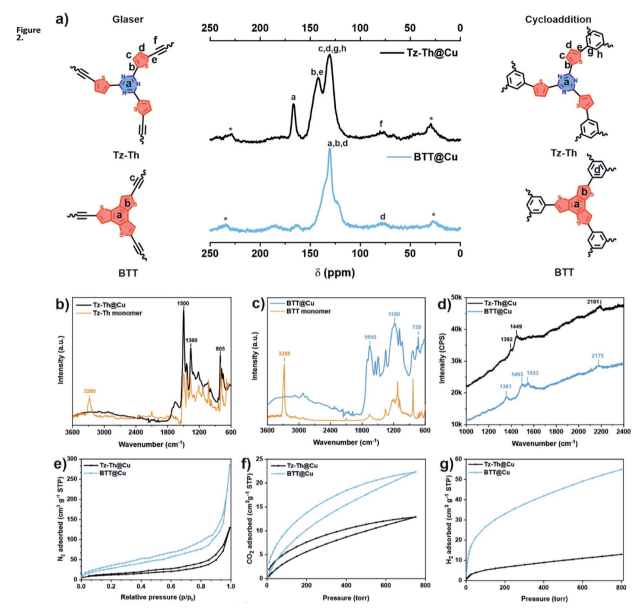


Fig. 2 Structural characterisation of the obtained materials. (a) ¹³C CP-MAS solid-state NMR spectra of Tz-Th@Cu and BTT@Cu. Signals of carbon atoms are labelled in accordance with 2 possible structures - products of Glaser-coupling and cycloaddition reaction (schematic synthons on the left and right of ssNMR spectra. Spinning side bands are marked with an asterisk (*); (b) and (c) FTIR spectra of Tz-Th@Cu and BTT@Cu respectively, showing the comparison with the starting monomers; (d) Raman spectra of both Th-Th@Cu (black) and BTT@Cu (blue); (e) N₂ ad-/desorption isotherms measured at 77 K; (f) CO₂ ad-/desorption isotherms measured at 298 K and 1 bar; (d) H₂ adsorption isotherms recorded at 77 K and 1 bar.

for Tz-Th@Cu can be attributed to C₃N₃-ring vibration and breathing modes.37 Typical C-S bond symmetric and asymmetric vibrations in the thiophene ring between 830-730 cm⁻¹ and phenyl ring vibrations in the region from 1400 to 1650 ${
m cm}^{-1}$ can be observed for both Tz-Th@Cu and BTT@Cu.38 Complementary Raman spectra show the presence of C≡C stretching at 2191 cm⁻¹ for **Tz-Th@Cu** and at 2175 cm⁻¹ for **BTT@Cu**, which corroborate the formation of conjugated divene linkages (Fig. 2d).³⁹ The peaks at $\sim 1360-1390$ cm⁻¹ and $\sim 1450-1390$ 1550 cm⁻¹ can be assigned to the aromatic (phenyl, thiophene, triazine) rings breathing and stretching modes, respectively.40

Morphology and porosity investigation

Both Tz-Th and BTT-based polymers have predominantly amorphous morphology in the bulk, which is evidenced by powder X-ray diffraction (PXRD) patterns (Fig. S9†). Broad peaks in the range from 20° to 30° 2θ can be attributed to the stacking distances of individual layers (2.8-3.8 Å), which were also observed for other porous organic networks, such as COFs and CTFs.41 More atomistic structural information for both Tz-Th@Cu and BTT@Cu, however, cannot be deduced from PXRD patterns due to the low scattering contrast of 2D-3D intertwined heterostructures and additional free rotation along the C-C

bonds between repeating units, which leads to amorphization of the polymer structures in bulk.

The flake-like morphological growth of Tz-Th@Cu and BTT@Cu films is particularly interesting for nanoscale investigation using transmission electron microscopy (TEM). Accordingly, we performed high-resolution TEM imaging of both polymers and observed pronounced moiré fringes with hexagonal repeating patterns, that can be attributed to the crystalline domains formed within the polymeric network (Fig. S10 and S12†).42 The corresponding selected area diffraction (SAED) pattern made from Tz-Th@Cu confirms the hexagonal bonding motif within 2D crystalline lattice of obtained polymer, which is typical for structural analogues, such as CTFs (Fig. S11†).42 The SAED for BTT@Cu shows visible concentric rings, that correspond to regular distances within the principal unit-cells and indicate that this material features polycrystalline morphology (Fig. S13†).43 Both recorded SAED patterns are in a reasonable agreement with simulated monolayer patterns (Fig. S11 and S13†).

We further probed the pore structure of the synthesised polymers using different gas adsorption analyses. The guest accessible surface area of the obtained materials was determined by N2 ad-/desorption experiments at 77 K (Fig. 2e). Both Tz-Th@Cu and BTT@Cu showed moderate nitrogen gas uptake, with respective specific surface areas of 42 and 116 m² g⁻¹, calculated according to the Brunauer-Emmett-Teller (BET) model (Table 1). The N2 adsorption isotherms show almost no discernible hysteresis loops, which is pointing to the high rigidity of prepared networks. Obtained materials are predominantly mesoporous with the largest pore fraction between 5 and 25 nm, which is proved by pore size distribution (PSD) analysis, performed according to Horvath-Kawazoe (HK) and Barrett-Joyner-Halenda (BJH) models (Fig. S14†) The CO₂ adsorption isotherms show steep linear gas uptake without any discernible hysteresis, which is typical for micro-and mesoporous polymers (Fig. 2f).44 In addition, we investigated the hydrogen gas storage capacity of synthesised networks (Fig. 2g). The CO₂ and H₂ uptake of Tz-Th@Cu have close values of 0.57 and 0.58 mmol g^{-1} , whereas in the case BTT@Cu there is an almost 3-fold difference – 0.99 mmol g⁻¹ for CO₂ and 2.45 mmol g⁻¹ for H₂ adsorption (Table 1). Higher BET surface area together with the results of CO2 and H2 sorption analyses point to the formation of more uniform pore channels in the case of BTT@Cu, which is the consequence of the preferential AA-stacking structure.

Structural arrangement and theoretical modelling

For a better understanding of the polymer structural arrangement, we performed additional density functional theory (DFT) calculations. In the case of [2 + 2 + 2] cycloaddition network, the formation of both AA and AB-stacking structures has similar interaction energies (E_{int}) , hence, similar stability. However, in the case of Glaser-type polymer, the AA-arrangements are more stable than AB-arrangements (Fig. S15-S17 and Table S5†). Considering this, it is difficult to determine the dominant interlayer stacking in the resulting materials solely depending on E_{int} . To rationalize the difference in BET surface area observed for the prepared materials, we considered the solvation effect of pyridine to explain their structural stability and preferential interlayer stacking. Pyridine as a solvent is also an electron-acceptor and can alter the polarity of the solvated material, thus leading to sliding of the individual polymer layers, and therefore to the formation of a staircase-like ABstacking arrangement with blocked pore channels and decreased accessible surface area. 45 On the one hand, this effect is more pronounced in the case of polar D-A Tz-Th@Cu polymer, where the AB-stacking with the reduced accessible surface area will be favoured. On the other hand, the electronwithdrawing effect of pyridine will have limited influence on the relatively apolar D-D BTT@Cu, therefore causing less interlayer sliding and resulting in the AA-stacked structure with a higher BET surface area.

Optoelectronic properties

We further investigated the optical properties of the prepared polymers *via* diffuse-reflectance UV-Vis performed in solid-state. The UV-Vis spectra confirm that all materials absorb light in the whole visible part of the electromagnetic spectrum with discernible light absorption edges starting around 740 nm for **BTT**@Cu and 845 nm for **Tz-Th**@Cu (Fig. 3a). We calculated the optical bandgaps from the UV-Vis adsorption edges by plotting the respective Tauc plots using Kubelka–Munk theory and found them to be equal to 1.42 eV for **Tz-Th**@Cu and 1.85 eV **BTT**@Cu (Table 1 and Fig. 3b). ⁴⁶ Theoretical modelling predicts the bandgaps to be in the range from 1.04 to 1.93 eV and 1.30 to 1.95 eV for **Tz-Th**@Cu and **BTT**@Cu respectively,

Table 1 Porous and optical properties of the prepared materials

Material	$S_{\mathrm{BET}}^{a} (\mathrm{m}^2 \mathrm{g}^{-1})$	CO ₂ uptake ^b (mmol g ⁻¹)	H_2 uptake ^c (mmol g ⁻¹)	$E_{\mathrm{g\ dir.}}^{d}\left(\mathrm{eV}\right)$	J^e ($\mu A \text{ cm}^{-2}$)
Tz-Th@Cu	42	0.57	0.58	1.42	6.8
BTT@Cu	116	0.99	2.45	1.85	2.3
Tz-Th _b	868	1.04	3.42	1.48	3.7
BTT_b	563	1.38	4.08	1.58	2.6

^a Surface area calculated from N₂ adsorption isotherm using BET equation. ^b CO₂ gas uptake measured at 298 K and 1 bar. ^c H₂ gas uptake measured at 77 K and 1 bar. ^d Optical bandgaps calculated *via* the Kubelka–Munk function from the diffuse-reflectance UV-Vis adsorption edge. ^e Saturated photocurrent density obtained at an applied potential of 0.6 V (vs. Ag/AgCl) under simulated sunlight irradiation in 0.01 M Na₂SO₄ aqueous solution.

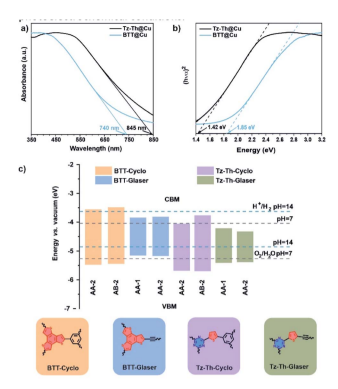


Fig. 3 Optical properties and bandgap characterisation of the prepared polymer networks. (a) Diffuse reflectance solid-state UV-Vis spectra ND (b) Tauc plots showing an estimation of optical bandgaps of Tz-Th@Cu (black) and BTT@Cu (blue); (c) relative locations of valence band maxima (VBM) and conduction band minima (CBM) for both Tz-Th and BTT-based polymers derived from DFT calculations for different AA and AB-stacking arrangements. The redox potential of water is indicated at pH = 7 and pH = 14. On the bottom – schematic structures of repeating subunits of Glaser- and cycloaddition-types of synthesised frameworks.

depending on the structure and interlayer stacking arrangement (Fig. 3c, S18, S19 and Table S5†). Additionally, the positive slope of the Mott-Schottky curves shows that both prepared materials are n-type semiconductors, and the relative electronic band positions are in broad agreement with theoretical predictions (Fig. S20†).47

According to DFT calculations, the Glaser-type products always have relatively low band gaps, while [2 + 2 + 2] cycloaddition products should have significantly higher energy gaps. Therefore, the high band gap experimentally measured for BTT@Cu (1.85 eV) can be mostly attributed to the [2 + 2 + 2]cycloaddition product, whereas the lower band gap obtained for Tz-Th@Cu (1.42 eV) should be related to the Glaser-type product. However, we cannot completely rule out the existence of [2 + 2 + 2] cycloaddition product in the structure of Tz-Th@Cu. The narrower bandgap value of Tz-Th@Cu polymer is a consequence of its D-A composition - the combination of tectons with different electron affinity (electron-poor triazine and electron-rich thiophene rings) results in a material with a smaller energy gap. We observed a similar phenomenon for Sand N-containing and S-only containing conjugated microporous polymers.34 Moreover, the highly polarised triazine-thiophene couple can also withdraw electron density from electron-

rich 1,3-butadyine linkages of the Glaser-coupling component, thus creating an additional charge push-pull effect within the Tz-Th@Cu network. This in turn, can lead to increased mobility of the generated charge carriers under photoelectrochemical conditions.48

Photoelectrodes fabrication and PEC characterisations

Narrow optical bandgaps (1.42-1.85 eV), micro-and mesoporosity, and improved transport of photogenerated electrons and holes make a promise for the synthesised materials to be applied in PEC devices. Accordingly, we conducted photoelectrochemical experiments with Tz-Th@Cu and BTT@Cu polymers spray-coated on an indium tin oxide (ITO) substrate as photoelectrodes in a 0.01 M Na_2SO_4 solution (pH = 7). The electrochemical measurements were performed in a 3-electrode setup with a Pt-disk counter electrode and applied bias voltage relative to the Ag/AgCl reference electrode. Photoanodes with the area of ~0.36 cm² fabricated from both Tz-Th-and BTTbased polymers showed a clear on/off response at an applied bias under intermitted light irradiation (100 mW cm⁻²) (Fig. 4a). To find the optimal equilibrium between the light adsorption ability and charge-transfer properties, we prepared a series of electrodes with different thicknesses.31 The highest photocurrent for Tz-Th@Cu was achieved for a medium-thick (\sim 0.9 μ m) coating, whereas thinner and thicker polymer films resulted in a lower photocurrent (Fig. S21†). In the case of BTT(a)Cu the highest photocurrent was achieved at \sim 1 µm and attempts to prepare a thicker coating resulted in unstable polymer films, which were easily detached from the ITO substrate when exposed to the electrolyte solution (Fig. S22†).

Both materials showed stable switching behaviour at 0.6 V vs. Ag/AgCl with Tz-Th@Cu reaching the saturated photocurrent density of \sim 6.8 μA cm⁻² and **BTT@Cu** reaching \sim 2.3 μA cm⁻² (after subtracting the dark current, Fig. 4b). The appearance of anodic photocurrent confirms that the prepared materials are n-type semiconductors,49 and the obtained values of the photocurrent densities are on par with the highest hitherto reported for organic polymeric photoelectrodes (Table S6†). The benefit of D-A composition of Tz-Th@Cu results in an almost 3fold increase in photocurrent density, clearly pointing toward improved excited charge transport and separation. The decreased arc radii of the Nyquist plots under light illumination for both Tz-Th@Cu and BTT@Cu suggest the reduced charge transport resistance in comparison to the dark conditions (Fig. 4c and d). This phenomenon confirms the photogeneration of electrons and holes upon simulated sunlight irradiation. Moreover, we compared the charge transfer resistance of both polymers under illuminated conditions, which is significantly lower for Tz-Th@Cu in contrast to BTT@Cu (~440 $k\Omega \nu s. > 800 k\Omega$, Fig. S23†). This fact points toward much better transport of photogenerated charge carriers through the more polarised D-A Tz-Th@Cu network.50 Noteworthy, both synthesised materials show good stability with negligible change in the photocurrent density under constant illumination for more than 2 h, which is imperative for real device fabrication (Fig. S24†).

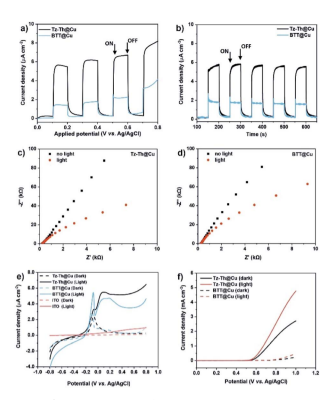


Fig. 4 PEC characterization of the Tz-Th- and BTT-based photo-anodes. (a) Current density–potential curves vs. applied potential of Tz-Th@Cu (black) and BTT@Cu (blue). Under intermittent irradiation at 1 mV s⁻¹ scan rate; (b) transient current density vs. time at 0.6 V under intermittent light irradiation recorded for both materials; EIS Nyquist plots at applied voltage of 0.6 V under dark (black) and light irradiation (red) of Tz-Th@Cu (c) and BTT@Cu (d); (e) linear sweep voltammetry recorded for obtained polymers at 1 mV s⁻¹ scan rate and applied potential from -0.8 to 0.8 V (a–e measured in 0.01 M Na $_2$ SO $_4$, pH = 7 vs. Ag/AgCl reference electrode); (f) photocurrent responses of Tz-Th@Cu (solid lines) and BTT@Cu (dashed lines) under dark (black) and illuminated (red) conditions measured in 1 M NaOH solution (pH = 14) vs. Ag/AgCl reference electrode.

Unlike with the majority of organic π -conjugated photoelectrodes, both Tz-Th@Cu and BTT@Cu show a more pronounced photocurrent response in the positive potential window from 0 to 0.8 V vs. Ag/AgCl reference electrode (Fig. 4e).51-53 Additionally, calculated band positions for both materials are suitable for water oxidation at the high pH value of the OER electrolyte (1 M NaOH, pH = 14) (Fig. 3c), therefore, as a proof-of-concept, we tested the prepared photoelectrodes in photoelectrochemical OER. The photocurrent density, and hence, the OER activity of Tz-Th@Cu increased almost 2 times under simulated solar irradiation – from 2.6 to 4.8 mA cm⁻² at 1.0 V vs. Ag/AgCl reference electrode without the addition of any transition-metal-cocatalyst (Fig. 4f). For BTT(a)Cu we observed similar behaviour under dark and illuminated conditions a roughly 2-fold increase in the photocurrent density from 0.26 to 0.46 mA cm⁻², which is, however, significantly lower than that of D-A Tz-Th@Cu material (Fig. 4f).

Both synthesised materials have a similar guest-accessible area with a predominance of mesopores, therefore the external surface wettability may play a crucial role in PEC OER

(Table 1).54 We estimated the contact angle with water to be slightly less for Tz-Th@Cu compared to BTT@Cu (51.3 vs. 56.5 deg., Fig. S25†). The negligible difference in the surface polarity clearly demonstrates that the affinity to water should not be a detrimental factor for the better OER performance of Tz-Th@Cu. The D-A composition creates a strong charge pushpull effect and improves the photophysical properties of Tz-Thbased network, i.e., narrows down the bandgap and enhances the charge transport in the photoexcited state. We corroborate these findings with GC-MS measurements that show 5.7 increase in the amount of evolved O2 under light illumination for Tz-Th@Cu vs. 4.3 increase in the case of BTT@Cu (Fig. S26 and Table S7†). Moreover, the Tz-Th-based material demonstrated overall better OER efficiency with the ability to produce 2-times and 1.5-times more oxygen with and without light irradiation, respectively. These results are pointing to the accelerated generation and better separation of photoexcited holes that can reach the surface of the photocatalyst and promote the oxygen evolution reaction.

The residual Cu content left in both networks after the synthesis, may influence the performance in electrochemical OER. The EDS showed a similar nonstochiometric amount of Cu left in both materials (2.49 and 2.48 wt% for Tz-Th@Cu and BTT@Cu respectively, Table S2†). Since D-A Tz-Th@Cu performed better in the photoelectrochemical OER, there is no direct correlation between the OER activity and residual Cu content. Moreover, we observed no catalytic activity when conventional Cu foil was used as a working electrode in the OER (Fig. S27†). In addition, we confirmed that increase in the photocurrent is not due to the oxidation of active materials, since their structure remains intact during the OER process. No additional bands appear in the FTIR spectra recorded after the OER measurements under illuminated conditions (Fig. S28†).

Preparation and characterisation of bulk materials

To further demonstrate the benefit of surface-assisted polymerisation for the fabrication of organic photoelectrodes, we prepared Tz-Th- and BTT-based frameworks in bulk using copper(1) chloride as a catalyst (denoted as Tz-Th_b and BTT_b). We confirmed the similarity of their chemical composition to the Tz-Th@Cu and BTT@Cu by 13C CP-MAS ssNMR (Fig. S29 and S30†), elemental analysis and EDS (Tables S8-S11†), and XPS (Fig. S31 and S32†). Moreover, the completion of the network forming reactions was also evidenced by TGA (Fig. S33†) and FTIR (Fig. S34†). Opposite to the pronounced flake-like morphology of on-copper grown frameworks, Tz-Thb and BTT_b show random 'cauliflower' aggregation due to kinetically controlled nucleation growth, which is typical for transition-metal-catalysed bulk reactions (Fig. S35†).55 Similarly to their thin-film analogues, bulk Tz-Thb and BTTb have an overall amorphous structure according to PXRD patterns (Fig. S36†). The N₂ gas sorption analyses resulted in BET surface areas of 868 and 563 m² g⁻¹ for Tz- Th_b and BTT_b , respectively (Fig. S14a†), with the majority of micro- and mesopores having a diameter in the range from 2 to 5 nm according to PSD analysis (Fig. S14b†). The synthetic approach directly influences

the morphology, hence the porosity of the prepared materials. Bulk-prepared networks have a higher accessible surface area due to random interdigitation of polymeric struts that formed predominantly under kinetic control. Additionally, Tz-Thb showed CO2 and H2 adsorption capacities of 1.04 and 3.42 mmol g⁻¹, whereas BTT_b showed higher capacities of 1.38 and 4.08 mmol g^{-1} despite lower S_{BET} (Table 1, Fig. S14c and d†). This can be attributed to the larger S-content (Tables S8-S11†), which is believed to be beneficial for CO₂ gas capturing.⁵⁶ Noteworthy, the obtained values for CO2 and H2 adsorption capacities are comparable to the highest of the hitherto reported values for benchmark CMPs and COFs.57,58

We affirmed the similarity of optical properties (bandgap and photoluminescence) of bulk materials with their onsurface-grown analogues (Table 1, Fig. S37 and S38†). The PEC measurements showed an almost 2-fold drop in the saturated photocurrent density of Tz-Thb in comparison to the Tz-**Th@Cu** – from 6.8 to 3.7 μ A cm⁻², whereas there is almost no difference between BTT@Cu and BTT_b - 2.3 and 2.6 μA cm⁻² respectively (Table 1 and Fig. S39†). These results indicate that not only the synthetic conditions but also the molecular composition will determine the efficiency of the resulting organic photoelectrode. Here, the strong D-A interactions dramatically improve the intra- and interlayer charge transport within Tz-Th@Cu network, leading to enhanced PEC performance in comparison with the bulk synthesised material.⁵⁹ In the case of BTT-based frameworks, however, weak D-D interactions do not enable efficient mobility of photoexcited charge carriers, and, consequently, there is no improvement in the photocurrent generation of layered BTT@Cu in comparison to the bulk BTT_b.

Conclusions

In conclusion, we demonstrated herein a rational approach toward the design and fabrication of highly efficient photoelectrodes composed of fully organic triazine- and thiophenebased porous polymers. First, the facile and easy-scalable surface-assisted polymerisation using conventional copper foil and mild "wet chemistry" reaction conditions allowed us to produce polymeric heterostructures with pronounced layered morphology and permanent micro- and mesoporosity. Second, the selection of modular molecular precursors enables finetuning of the photophysical properties (bandgap, photoluminescence) of the resulting materials. The combination of building blocks with different electron affinity enables strong donor-acceptor interactions, which in turn narrow the bandgap and improve the photoexcited charge transfer within the obtained network. As a result, the photoanode, fabricated from the donor-acceptor Tz-Th@Cu polymer showed a 3-fold increase in PEC performance in comparison to donor-donor BTT@Cu, reaching an exceptional photocurrent density of 6.8 μA cm⁻², which is also 3-times higher than that of the bulk synthesised Tz-Th_b material (at 0.6 V vs. Ag/AgCl). Moreover, the donoracceptor Tz-Th@Cu network showed a double increase in activity under simulated sunlight irradiation as OER photoelectrocatalyst, and performed as twice as better in comparison

to donor-donor BTT@Cu. The presented strategy, which combines robust and simple copper-surface-assisted synthesis and modular terminal alkyne coupling, may lead to the fabrication of photoelectrochemically active materials for efficient and sustainable utilisation of solar energy.

Author contributions

M. P. and Y. S. K. devised and designed the project. Y. S. K. carried out the materials synthesis, SEM, EDS, TEM, TGA, PXRD, FTIR, Raman, AFM, solid-state UV-Vis, and PL measurements, analysed the data and wrote the paper. S. M. B. M. designed and performed the PEC experiments, analysed the data. B. K. performed photoelectrochemical O₂ evolution experiments and GC analysis. P. L. performed DFT calculations and analysis the data. M. J. B. contributed reagents and materials. M. P. conceived and designed the project, contributed reagents/materials/analysis tools, and wrote the paper.

Conflicts of interest

There are no conflicts to declare.

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