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Conjugated polymers for visible-light-driven photocatalysis

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Conjugated polymers have recently been under active investigation as promising alternatives to traditional inorganic semiconductors for photocatalysis. This is due to their unique advantages of low cost, high chemical stability, and molecularly tunable optoelectronic properties. This critical review summarizes the recent advancements in π -conjugated polymers for visible-light-driven photocatalytic applications including water splitting, CO₂ reduction, organic transformation and degradation of organic dyes. Special emphasis is placed on how the changes in the polymer structure could influence their physicochemical properties and photocatalytic activities. This structure–activity relationship analysis should guide rational molecular design of conjugated polymers for improved photocatalytic activity.

Broader context

The exploitation of efficient photocatalysts to directly convert the radiant sunlight into chemical energy is of great importance to address the current energy and environmental challenges. Conjugated polymers (CPs) consisting of photoactive π -systems represent an attractive platform for solar energy utilization. They have been intensively studied for a variety of photocatalytic applications and many exciting performances were reported through facile molecular design. A comprehensive review is thus timely to summarize the progress of the field. This review article systematically presents the recent advances in conjugated polymers for visible-light-driven photocatalysis, including water splitting, CO₂ reduction, organic transformation and degradation of organic dyes. The synthesis and design principles of conjugated polymers in these photocatalytic applications are illustrated, with an emphasis on the correlation between polymer structures and their photocatalytic activities. We expect that the systematic discussion in this review will not only provide a general overview of the field, but also promote the development of conjugated polymers with fascinating properties for photocatalysis.

1. Introduction

The sun is known as a super “energy warehouse” to irradiate solar light continuously to the Earth. On average, the solar energy reaching the Earth’s surface per hour is enough to meet

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the current human energy demand in a whole year.^{1,2} Despite the advantages of inexpensive, abundant and clean features of sunlight, it is difficult to utilize solar energy very efficiently because of its low energy density as well as diffuse and intermittent nature. Therefore, it is desirable to convert sunlight into chemical energy, which represents an appealing solution to address several challenges facing the world today. For example, clean hydrogen fuel production by photocatalytic water splitting holds great promise to alleviate the global energy crisis, which promises to be clean, renewable and eco-friendly.^{3–9} Enabling photocatalytic technology will also allow the reduction of CO₂ into value-added chemicals (CO, CH₄, and CH₃OH, *etc.*), thereby creating huge benefits by simultaneously solving the current energy shortage and environmental issues caused by overloaded CO₂ release.^{10–14} Moreover, light-driven organocatalysis is considered as an attractive strategy for direct chemical bond functionalization with a broad substrate scope under mild conditions, which is a vitally important contribution to organic synthetic methodology.^{15–20} Last but not least, photocatalytic degradation of organic dyes in wastewater is a powerful technology for environmental remediation.^{21–24}

In a photocatalytic process, the photon energy required for photoexcitation is dependent on the optical gap of the photocatalyst. As compared to the UV region that comprises only 5% of solar light, the visible light (400–800 nm) accounts for about 53% of the incident solar energy (Fig. 1).^{4,25,26} Therefore, it is crucial to develop photocatalysts with optical gap below 3.0 eV to increase the absorption of solar light with a focus on the visible light region.

Over the past few decades, research efforts have been intensively devoted to pursuing suitable photocatalysts for practical applications. Among them, organic π -conjugated polymers have recently garnered significant attention as promising candidates for various photocatalytic applications. Many interesting features allow π -conjugated polymers to stand out in the field of photocatalysis over inorganic semiconductors. Firstly, they are environmentally friendly semiconductor materials,

consisting of earth-abundant elements, and they possess tunable energy levels for oxidation and reduction reactions. Secondly, they could be facilely prepared under mild conditions and have excellent chemical stability against photobleaching. Thirdly, their molecular structures can be fine-tuned to better utilize visible light, while many traditional inorganic photocatalysts are only UV-active. Last but not least, the π -conjugation along the polymer backbone enables them with unique photogenerated charge carrier separation and transport properties, which is critical to trigger photoredox reactions.

Upon diverse modification of conjugated polymers (CPs) at the molecular level, many exciting achievements in photocatalysis have recently been reported, which clearly justify a comprehensive review on this research topic. So far, most CP based photocatalysis reviews have been based on the family of graphitic carbon nitride (g-C₃N₄) materials with a focus on one or two photocatalytic applications.^{18,27–35} There is an urgent need to provide a review focused on π -conjugated polymers in photocatalysis from a broad perspective to present an overview of the recent developments on this research topic.

This review presents the recent advances of CPs in applications of photocatalytic water splitting, CO₂ photoreduction, light-driven organic transformation and photocatalytic degradation of organic dyes (Fig. 2). We focus on an in-depth understanding of how the CP structure modification could influence the optical properties, photogenerated charge separation and transport, and photocatalytic activity. The review starts from a brief discussion of general photocatalytic processes and the synthesis of CP photocatalysts. This is followed by the illustration of CPs for each photocatalytic application. Finally, perspectives toward exploring new CPs for photocatalytic applications are proposed, which is expected to stimulate the future development of more exciting CPs for photocatalysis.

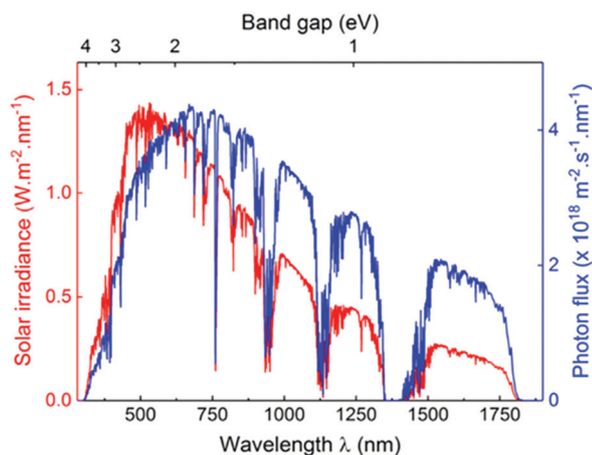


Fig. 1 AM1.5 solar light radiation spectrum (red) and photon flux curve (blue) as functions of light wavelength and material band gap. Reprinted with permission from ref. 25. Copyright 2017 Wiley-VCH.

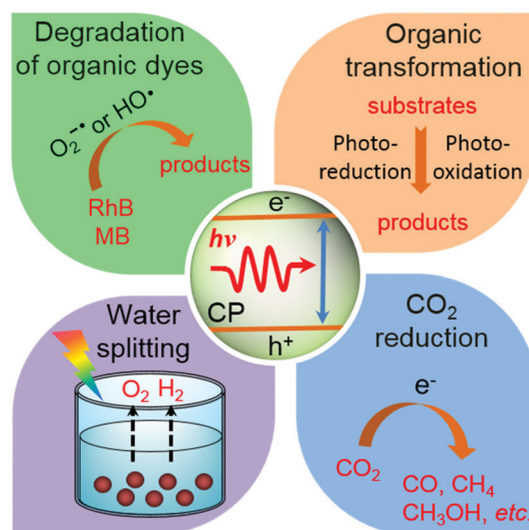


Fig. 2 Overview of π -conjugated polymers for various photocatalytic applications, which are summarized in this review.

2. Basic aspects of π -conjugated polymers for photocatalytic applications

2.1 Fundamentals of heterogeneous photocatalysis

The main processes of polymeric semiconductors in specific light-driven redox reactions are described in many reviews^{3,11,14,27,28,36–39} and the general pathway is briefly illustrated in Fig. 3a. Upon light irradiation, the semiconductor harvests incident photons and charges (electron–hole pairs) are formed inside the photocatalyst. The photogenerated charges subsequently separate and migrate to the semiconductor surface where redox reactions take place. On the other hand, most of the photogenerated electrons and holes could undergo bulk or surface recombination, which are competitive processes to the surface reactions.

According to the proposed mechanism of photocatalytic redox reactions, an efficient CP photocatalyst should at least have the following properties: (a) a strong and broad absorption in the visible region to effectively capture solar light; (b) high charge separation and migration efficiency for surface redox reactions; (c) appropriate energy level alignments to enable efficient charge separation as well as the reduction or oxidation half reaction. Therefore, light harvesting, energy levels as well as photogenerated charge separation and migration are three key factors affecting the photocatalytic activities of these CPs (Fig. 3b). This requires rational design and synthesis of CPs for effective photocatalysis.

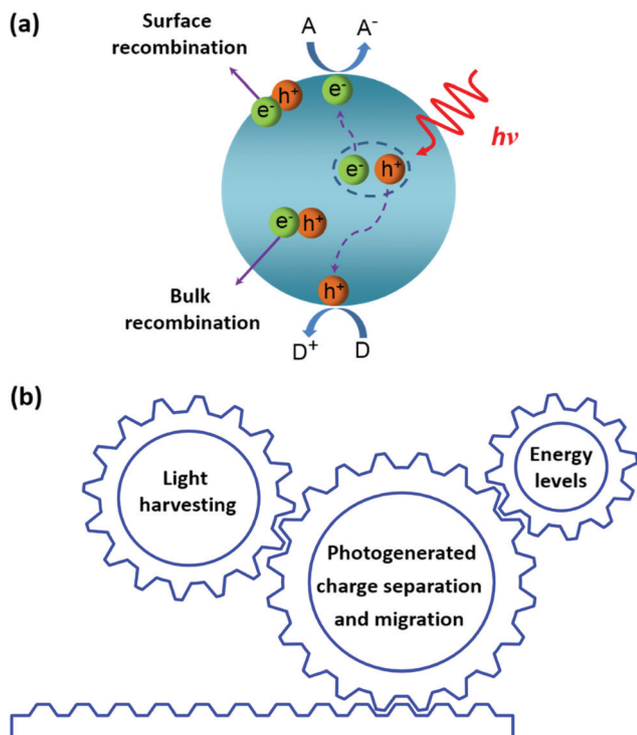


Fig. 3 (a) The main process of polymeric semiconductors for photocatalysis (D: electron donor; A: electron acceptor) and (b) critical factors affecting their photocatalytic performances.

2.2 Synthesis of conjugated polymers for heterogeneous photocatalysis

The extensively available conjugated organic monomers and diverse synthetic methods enable facile synthesis of a broad range of CPs. Moreover, the polymer structures could be precisely controlled for specific photocatalytic applications. Among these methods, Pd-catalyzed Suzuki–Miyaura coupling polymerization between halogenated aromatics and diboron ester has been widely used to prepare linear copolymers and conjugated porous polymers (Fig. 4). The reaction is typically carried out in the presence of a base (always K_2CO_3) under a N_2 or Ar atmosphere. Another commonly used method for preparing CPs by C–C coupling is the Sonogashira coupling reactions of aromatic halides and terminal alkynes, in which $Pd(PPh_3)_2Cl_2$ was employed as the catalyst with an amine as the base, and CuI as the cocatalyst. Furthermore, Ni(0)-promoted Yamamoto dehalogenation coupling reaction is a convenient pathway to synthesize CPs from dibromo-aromatic monomers. In this reaction, $Ni(COD)_2$ and 2,2'-bipyridyl are used as the catalyst and promoter for polymerization, respectively. Oxidative coupling reactions are exploited as a versatile methodology for performing self-polymerization of arylethynylene monomers, in which CuBr is used as the catalyst and a base is used as the binding agent.

Interestingly, conjugated polymer nanomaterials could be readily prepared under appropriate reaction conditions. For example, conjugated microporous polymer nanosheets could be obtained by one-step oxidative coupling of aromatic alkynes⁴⁰ or a condensation reaction.^{41,42} Conjugated microporous polymer nanoparticles could be directly prepared by polymerization in high internal phase emulsion, which is concentrated emulsion with a dispersed phase volume fraction over 74.05%.^{43,44} The morphologies of nanoparticles can be varied by introducing monomers with different electron-donating/withdrawing properties.⁴⁵

Additionally, the condensation reaction between amino and aldehyde groups based on Schiff-chemistry is widely performed to prepare crystalline covalent organic frameworks (COFs) for photocatalysis.⁵¹ The reaction is typically performed under harsh experimental conditions with carefully controlled temperature (usually 120 °C), pressure and reaction time (usually 3–5 days). Acetic acid is employed as the catalyst in the reaction. It is believed that an error-correction process occurs during the conversion of amorphous intermediates to crystalline products. The reaction solvent is crucial for the assembly of framework units. In a typical condensation reaction, mesitylene/dioxane and *n*-butanol/*o*-dichlorobenzene are commonly used reaction solvents. Besides, a suitable building block is also important to obtain a highly crystalline COF. Rigid and planar π -conjugated units are preferred to prepare 2D COFs.

Covalent triazine frameworks (CTFs) could be prepared by a trimerization reaction of aromatic nitriles in the presence of a Brønsted or Lewis acid catalyst.⁵³ In a typical ionothermal preparation, $ZnCl_2$ is chosen as the catalyst and high temperature (>400 °C) is always required for the reversible reaction. By contrast, a much lower temperature (<100 °C) could be employed

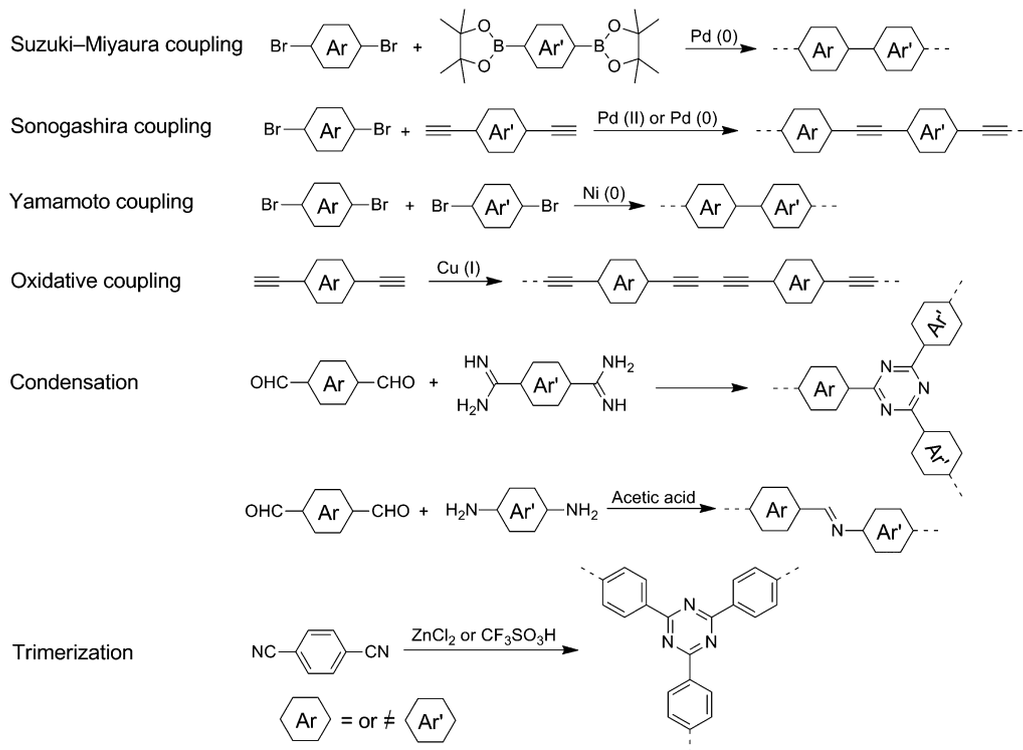


Fig. 4 Typical polymerization methods used to synthesize conjugated polymers for photocatalysis.

when $\text{CF}_3\text{SO}_3\text{H}$ is used as the catalyst for the condensation reaction. Alternatively, CTFs could be synthesized by condensation of an aldehyde and an amidine dihydrochloride, which involves the generation of a Schiff base followed by Michael addition. The reaction is carried out in DMSO with Cs_2CO_3 as a base.

Graphitic carbon nitride ($\text{g-C}_3\text{N}_4$) consisting of *s*-triazine or tri-*s*-triazine (heptazine) units has been synthesized by thermal polycondensation of nitrogen-rich precursors, such as melamine, thiourea, urea, dicyanadiamide and cyanamide, *etc.*^{18,29,31} For more detailed preparation of the above-mentioned CPs, please refer to other reviews^{46–53} and books.^{54,55}

3. Conjugated polymers for photocatalytic water splitting

Photocatalytic water splitting is of particular interest as the technology provides an effective pathway for clean hydrogen production using water and sunlight, both of which are abundant in nature.

In a light-driven water splitting process, photogenerated electrons and holes were used for the reduction and oxidation of water, respectively. The photocatalytic reaction is a thermodynamically uphill reaction and a Gibbs free energy (ΔG_0) of $+237.2 \text{ kJ mol}^{-1}$ is required.³ In principle, to drive the redox half reaction, the energy level alignments of the polymer photocatalyst and the redox potential of water must be well matched. This means that the lowest unoccupied molecular orbital (LUMO) of the polymer photocatalyst should be more

negative than the reduction potential of H^+/H_2 (0 eV vs. normal hydrogen electrode (NHE), pH = 0), while its highest occupied molecular orbital (HOMO) must be more positive than the oxidation potential of $\text{O}_2/\text{H}_2\text{O}$ (1.23 eV vs. NHE, pH = 0). Therefore, the band gap (E_g) of the polymer should be over 1.23 eV to trigger water splitting. In this part, we summarize the state-of-the-art advancements in photocatalytic H_2 evolution using various types of organic CPs, and their applications in photocatalytic O_2 evolution and overall water splitting are also briefly described.

3.1 Photocatalytic H_2 evolution

Hydrogen is an ideal fuel with gravimetric energy density up to 142 MJ kg^{-1} , which is almost three times larger than that of gasoline.⁵⁶ More importantly, H_2 combustion produces water with no pollution to the environment. Photocatalytic hydrogen evolution using semiconductors is a sustainable and economical technology to convert solar energy into hydrogen energy. Since Honda and Fujishima reported water splitting on TiO_2 photoelectrodes in 1972, a huge number of semiconductors have been studied for hydrogen production.^{3,34,57–64}

As promising photocatalysts, CPs have been employed for hydrogen production in the early 1980s. Unfortunately, they haven't attracted much attention until three-dimensional (3D) poly(azomethine) networks were demonstrated to be efficient photocatalysts for hydrogen production from water in 2010.⁶⁵ Since then, various types of organic CPs have been developed to yield greatly improved H_2 -production efficiency. This was achieved through the optimization of CP photophysical properties,

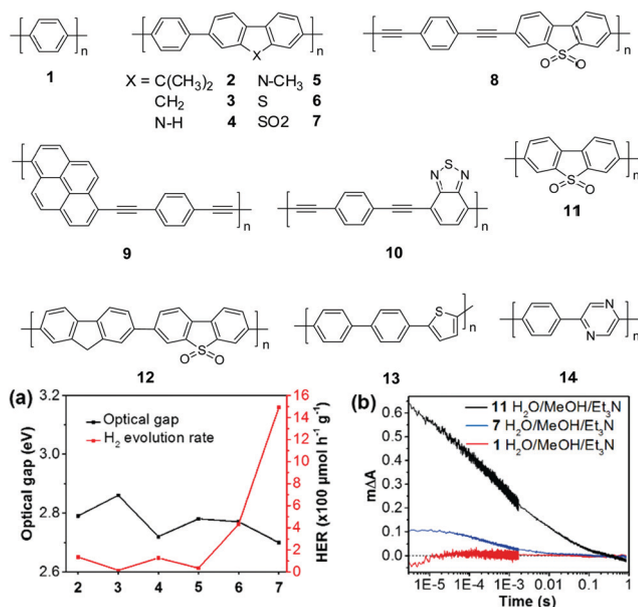


Fig. 5 Chemical structures of polymers 1–14, (a) the correlation between the photocatalytic H₂ evolution and their optical gaps of polymers 2–7 (>420 nm); (b) transient kinetics probed at 630 nm for polymers 1, 7, and 11 in the Et₃N/CH₃OH/H₂O mixture (excited at 355 nm). Reprinted with permission from ref. 72. Copyright 2018 Nature Publishing Group.

mainly light harvesting in the visible region, band gap alignment, as well as photogenerated charge generation and transport, etc.

3.1.1 Linear conjugated polymers. The early report of linear organic CPs as the photocatalyst for water splitting could be dated back to 1985.⁶⁶ Yanagida *et al.* prepared poly(*p*-phenylene) (PPP) (1, Fig. 5) by the polycondensation of 1,4-dibromobenzene. Polymer 1 shows a band gap of 2.9 eV and a hydrogen evolution rate (HER) of 207.5 μmol h⁻¹ g⁻¹ under broad spectrum irradiation (>290 nm). However, this polymer is almost inactive under visible light (>400 nm). Following this work, they further reported some analogues of 1, but very low activities were achieved even with the assistance of Ru nanoparticles.^{67–69}

In 2016, Cooper and co-workers demonstrated the new breakthrough in H₂ evolution by incorporating extended planarized units into the main chain of 1, which is expected to facilitate the photogenerated charge dissociation and thus promote the proton reduction reaction.⁷⁰ Polymers 2–7 show band gaps ranging from 2.70 to 2.86 eV (Fig. 5a). Among them, polymers 2 and 4 exhibited an almost equal HER of ~128 μmol h⁻¹ g⁻¹, which was 3-fold improvement over 1. More excitingly, polymer 7 with the lowest band gap of 2.70 eV exhibited an impressive HER of 1492 μmol h⁻¹ g⁻¹, which was 46-fold higher than that of 1 (Table 1).

Encouraged by the good activity of polymer 7, Chen *et al.* further modified this polymer by introducing an ethynyl group into the main chain, which led to a reduced band gap and accelerated charge separation of the resulting polymers. Polymer 8 has a band gap of 1.89 eV and showed a HER of up to 6023 μmol h⁻¹ g⁻¹ when triethanolamine (TEOA) was used as the sacrificial agent (>420 nm).⁷¹ Similarly, compared to their counterparts without ethynyl insertion, polymers 9 and 10

showed large red shifts of 167 and 154 nm in their light absorption edges, respectively. The HER was enhanced to 4270 μmol h⁻¹ g⁻¹ for 9 and 3187 μmol h⁻¹ g⁻¹ for 10.

Notably, dibenzo[*b,d*]thiophene sulfone has been recognized as one of the most attractive building blocks for the synthesis of H₂-production polymer photocatalysts. As shown above, many polymers with incorporated sulfone units show outstanding photocatalytic performances. To study the critical role of the sulfone unit in H₂ evolution, Cooper and co-workers investigated the photophysical properties of polymers 1, 7 and 11.⁷² Compared with 1 and 7, the homopolymer of sulfone 11 showed a much higher HER of 3260 μmol h⁻¹ g⁻¹. The decay kinetics at 630 nm exhibited a significantly larger signal for 11 than 7, while almost no signal was observed for 1 (Fig. 5b). This result indicated a higher polaron concentration of 11 and 7, which is in line with the H₂ evolution activities of the three polymers. Recently, they further prepared nanoparticles of about 160 nm using polymer 11 by mini-emulsion polymerization.⁷³ With a polymer concentration of 0.1 mg mL⁻¹, the nanoparticle suspension yielded a HER of 14.52 mmol h⁻¹ g⁻¹, while a significantly lower HER of 6.13 mmol h⁻¹ g⁻¹ was measured for the bulk material of 11, evidently suggesting a size-dependent H₂ evolution of the polymer. Moreover, when the polymer concentration was decreased to 13 μg mL⁻¹, the suspension gave a superior average HER of 60.6 mmol h⁻¹ g⁻¹ over 5 h, which is the highest value among the reported organic polymers so far.

To further extend the conjugation degree of polymer 7, we introduced large planar units (fluorene and pyrene) into the main chain.⁷⁴ Polymer 12 exhibited a smaller band gap of 2.07 eV in comparison with 7 (2.70 eV) and a good HER of 5040 μmol h⁻¹ g⁻¹ was achieved. Very recently, Wang *et al.* reported an analogue of polymer 12 by replacing the fluorene in the main chain with electron-rich dibenzothiophene to construct a local D–A heterojunction, which led to a reduced exciton binding energy from 91 to 88 meV and 1.3-fold HER as compared to 12 in the presence of TEOA.⁷⁵

Another modification of PPP (1) is the copolymerization of 2,5-phenylene with heterocycles, which results in optimized band gaps and energy levels of the as-prepared polymers. A maximum HER enhancement of 6.5-fold compared to PPP was achieved by polymer 13 with 33% thiophene fraction and a band gap of 2.42 eV.⁷⁶ Polymer 14 bearing a 2,5-pyrazine attained a 15-fold higher HER than PPP.⁷⁷

Compared to those insoluble bulk polymers, the development of soluble CPs provides a good choice for easy processing of polymers into a desirable pattern for photocatalytic H₂ production. By virtue of its inherent hydrophobicity, CP could be self-assembled into polymer dots (Pdots) in water.^{78–80} Pdots have much smaller size relative to the bulk materials; hence, the distance covered by photogenerated charges migrating to the surface of photocatalyst particles becomes short, and the recombination probability of electrons and holes decreases to improve the photocatalytic efficiency.⁸¹ In 2016, Tian *et al.* prepared a Pdot suspension by the nano-precipitation method using polymer 15 and a matrix polymer PS-PEG-COOH.⁸² In the preparation, the mixture of polymer and matrix in THF

Table 1 Summary of representative photocatalytic H₂ evolution using CPs under visible light

Polymer	Optical gap (eV)	S _{BET} (m ² g ⁻¹)	Cocatalyst	Sacrificial agent	Light source ^a	HER (μmol h ⁻¹ g ⁻¹)	AQY ^b (%)	Ref.
1	2.9	—	—	DEA	> 400 nm (Xe)	Trace	—	66
2	2.79	—	—	TEA	> 420 nm (Xe)	136	—	70
3	2.86	—	—	TEA	> 420 nm (Xe)	16	—	70
4	2.72	—	—	TEA	> 420 nm (Xe)	128	—	70
5	2.78	—	—	TEA	> 420 nm (Xe)	36	—	70
6	2.77	—	—	TEA	> 420 nm (Xe)	432	1.1	70
7	2.70	—	—	TEA	> 420 nm (Xe)	1492	2.3	70
8	1.89	—	—	TEOA	> 420 nm (Xe)	6023	4.2	71
9	1.64	—	—	TEOA	> 420 nm (Xe)	4270	0.85	71
10	1.94	—	—	TEOA	> 420 nm (Xe)	3187	—	71
11	2.62	—	—	TEA	> 420 nm (Xe)	3260	11.6	72
12	2.07	—	—	TEA	> 420 nm (Xe)	5040	2.13	74
13	2.42	17	—	TEA	> 420 nm (Xe)	420	—	76
14	2.45	69	—	TEA	> 420 nm (Xe)	960	—	77
15	2.38	—	—	AA	> 420 nm (LED)	8300	~0.73	82
16	1.98	—	—	AA	> 420 nm (LED)	50 000	~0.88	83
17	2.46	—	—	AA	> 420 nm (LED)	NA	—	83
18	2.03	—	—	DEA	> 420 nm (LED)	12 700	—	84
19	2.11	—	—	TEA	> 420 nm (Xe)	512	—	89
20	—	—	Pt	AA	> 420 nm (Xe)	12 800	0.13	90
21	2.33	995	—	DEA	> 420 nm (Xe)	174	—	96
22a	1.81	39	—	TEA	> 400 nm (Xe)	17	—	105
22b	1.95	59	—	TEA	> 400 nm (Xe)	140	—	105
23a	2.89	228	—	TEA	> 400 nm (Xe)	397	—	105
23b	2.45	279	—	TEA	> 400 nm (Xe)	1460	—	105
24a	2.31	121	—	TEA	> 400 nm (Xe)	260	—	105
24b	2.19	99	—	TEA	> 400 nm (Xe)	786	—	105
25a	2.33	80	—	TEA	> 400 nm (Xe)	414	—	105
25b	2.06	104	—	TEA	> 400 nm (Xe)	1900	—	105
26	1.88	—	—	TEA	> 400 nm (Xe)	2590	—	106
27	2.44	280	Pt	TEOA	> 420 nm (Xe)	400	—	107
28	2.18	58	—	TEOA	≥ 420 nm (Xe)	9600	1.8	110
29	2.19	499	Pt	TEOA	> 420 nm (Xe)	30	—	111
30	2.21	669	Pt	TEOA	> 420 nm (Xe)	134	—	112
31	2.28	750	Pt	TEOA	> 420 nm (Xe)	598	—	112
32	2.14	564	Pt	TEOA	> 420 nm (Xe)	908	2.0	112
33	2.37	834	Pt	TEOA	> 420 nm (Xe)	620	—	112
34	2.53	811	—	TEOA	> 420 nm (Xe)	2460	—	113
35	2.62	915	—	TEOA	> 420 nm (Xe)	188	—	113
36	2.64	447	—	TEOA	> 420 nm (Xe)	116	—	113
37	2.84	895	—	TEA	> 420 nm (Xe)	120	—	114
38	2.56	431	—	TEA	> 420 nm (Xe)	3106	13.2	115
39	2.37	723	—	TEOA	> 420 nm (Xe)	5697	—	116
40	2.11	692.5	Pt	TEOA	> 420 nm (Xe)	2103.2	6.4	117
41	2.8	1603	Pt	TEOA	> 420 nm (Xe)	1970	—	119
42	2.67	702	Pt	TEOA	420–900 nm (Xe)	22.5	—	120
43	2.68	326	Pt	TEOA	420–900 nm (Xe)	90	0.08	120
44	2.62	1537	Pt	TEOA	420–900 nm (Xe)	438	0.19	120
45	2.65	1046	Pt	TEOA	420–900 nm (Xe)	1703	0.44	120
46	2.10	985	Pt	AA	> 420 nm (Xe)	4440	—	121
47	1.85	1288	Pt	AA	> 420 nm (Xe)	10 100	—	121
48	2.28	919	Pt	AA	> 420 nm (Xe)	1600	—	121
49	2.34	523	Pt	TEOA	≥ 395 nm (Xe)	30	—	122
50	2.31	758	Pt	TEOA	≥ 395 nm (Xe)	324	1.8	122
51	2.09	541	Pt	TEOA	> 420 nm (Xe)	8300	—	123
52	2.06	348	Pt	TEOA	> 420 nm (Xe)	1560	—	123
53	1.92	162	Pt	TEOA	> 420 nm (Xe)	220	—	123
54	2.03	602	Pt	TEOA	> 420 nm (Xe)	1360	—	124
55	2.94	19	Pt	TEOA	≥ 420 nm (Xe)	200	—	128
56	2.38	52	Pt	TEOA	> 420 nm (Xe)	6600	7.3	130
57	2.22	807	Pt	TEOA	> 420 nm (Xe)	1238	—	132
58	2.13	764	Pt	TEOA	> 420 nm (Xe)	1582	—	132
59	2.33	663	Pt	TEOA	> 420 nm (Xe)	1460	—	132
60	2.42	757	Pt	TEOA	> 420 nm (Xe)	2647	—	132
61	2.17	569	Pt	TEOA	> 420 nm (Xe)	10 760	4.07	133
62	2.47	533	Pt	TEOA	> 420 nm (Xe)	5320	4.11	133
63	2.67	650	Pt	TEOA	> 420 nm (Xe)	1440	2.10	133
64	—	599	Pt	TEOA	> 420 nm (Xe)	5100	—	138
65	—	520	Pt	TEOA	> 420 nm (Xe)	2400	—	138
66	—	598	Pt	TEOA	> 420 nm (Xe)	650	—	138

Table 1 (continued)

Polymer	Optical gap (eV)	S_{BET} ($\text{m}^2 \text{g}^{-1}$)	Cocatalyst	Sacrificial agent	Light source ^a	HER ($\mu\text{mol h}^{-1} \text{g}^{-1}$)	AQY ^b (%)	Ref.
CN-F	2.63	—	Pt	TEOA	> 420 nm (Xe)	~130	—	142
CN-N	2.65	9.21	Pt	TEOA	> 400 nm (Xe)	553.5	—	143
CN-C/P	2.98	141.1	Pt	TEOA	> 420 nm (Xe)	1493.3	2.14	144
CN-BA	2.57	22	Pt	TEOA	> 420 nm (Xe)	294	—	145
CN-P15	—	—	—	TEOA	> 420 nm (Xe)	722.3	—	152
CN-P17	2.24	—	—	TEOA	> 400 nm (Xe)	929.3	5.7	153
CN-PyP	2.46	120	—	TEOA	≥ 420 nm (Xe)	600	—	154
CN-COF	—	85.2	Pt	TEOA	> 420 nm (Xe)	10 100	—	155
CN-CTF	—	—	Pt	TEOA	≥ 420 nm (Xe)	850	—	156

^a Xe: xenon lamp. ^b At 420 nm. NA: no activity; AQY: apparent quantum yield; HER: hydrogen evolution rate; DEA: diethylamine; TEA: trimethylamine; TEOA: triethanolamine; AA: ascorbic acid.

was injected into pure water under sonication, and a clear Pdot suspension was obtained after THF evaporation (Fig. 6). The as-prepared Pdots displayed an impressive initial rate of $8300 \mu\text{mol h}^{-1} \text{g}^{-1}$. Compared to polymer 15, Pdots of polymer

16 with extended absorption renders an excellent initial HER of up to $50 \text{ mmol h}^{-1} \text{g}^{-1}$.⁸³ The theoretical calculations suggested that N atoms in the 2,1,3-benzothiadiazole (BT) units provided the reactive sites in the formation of hydrogen, polymer 17 without a BT unit did not show any activity although its optical gap and reduction potential are appropriate for proton reduction. Using the nano-precipitation method, Chou *et al.* prepared cycloplatinated Pdots using polymer 18, which provided an impressive HER of $12.7 \text{ mmol h}^{-1} \text{g}^{-1}$, while only 1.3 and $5.46 \text{ mmol h}^{-1} \text{g}^{-1}$ were obtained for Pdots without a Pt complex and their Pt-complex-blended counterpart under identical conditions, respectively.⁸⁴ This was likely attributed to the improved charge transfer after grafting the Pt complex into the polymer backbone.

Conjugated polyelectrolytes (CPEs) are a subclass of interesting water-soluble CPs with wide applications in photovoltaic devices, biomedical imaging and therapy, as well as biosensing, *etc.*^{85–88} Their rigid conjugated structures endow them with excellent optoelectronic properties, while the hydrophilic side chains with charged groups allow for unique dispersibility of CPEs in polar solvents. We prepared cationic CPE 19 by one-step quaternization of the precursor polymer attaching a hexyl bromide side chain.⁸⁹ In the mixture of $\text{Et}_3\text{N}/\text{CH}_3\text{OH}/\text{H}_2\text{O}$, polymer 19 could self-assemble into homogeneous nanoparticles of ~180 nm. The suspension gives a HER of $512 \mu\text{mol h}^{-1} \text{g}^{-1}$, which is 3-fold that of its counterpart without the side chain. The enhanced HER was attributed to the improved light absorption and charge separation of the nanoparticles.

Different from the cationic CPEs, attaching oligoethylene glycol (OEG) side chains endows polymer 20 with more interesting functions.⁹⁰ The O atom in the OEG side chain favours the interface contact of the polymer backbone with water, which resulted in a much smaller water contact angle of 20 (27.5°) than that of an alkyl-functionalized polymer (101.2°). Moreover, the side chains could interact with Pt, resulting in more efficient charge transport from the polymers to Pt. Polymer 20 loaded with 3 wt% Pt gave a HER of $12.8 \text{ mmol h}^{-1} \text{g}^{-1}$, which is much higher than its alkyl-attached counterpart.

Besides, soluble CPs could also be processed into films for photocatalytic H_2 production. By drop-casting, Cooper's group prepared a film of polymer 4 attaching a 2-ethylhexyl side chain. The film demonstrated a HER of $72 \mu\text{mol h}^{-1} \text{g}^{-1}$ when immersed in the $\text{Et}_3\text{N}/\text{CH}_3\text{OH}/\text{H}_2\text{O}$ mixture ($> 420 \text{ nm}$).⁹¹

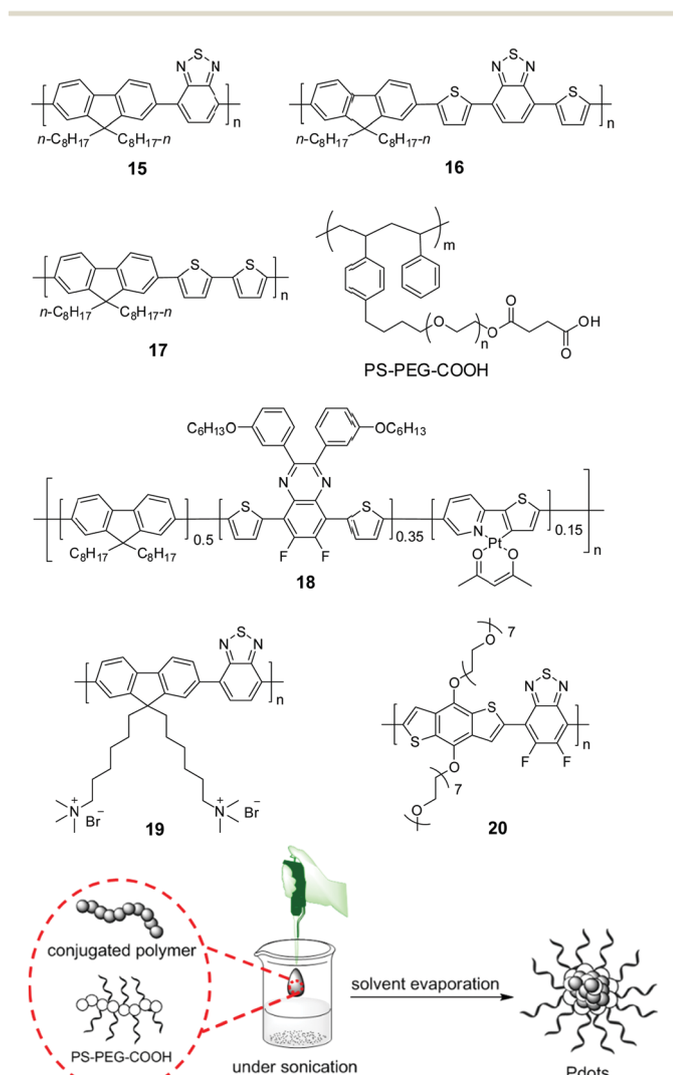


Fig. 6 Chemical structures of polymers 15–20 and preparation of polymer dots (Pdots) by the nano-precipitation method for photocatalytic H_2 production. The scheme is adapted with permission from ref. 82, Copyright 2016 Wiley-VCH.

3.1.2 Conjugated porous polymers. Crosslinked CPs with a three-dimensional porous feature have gained ever-increasing interest because of their large electron delocalized network structure, excellent stability and tunable optoelectronic properties. These advantages have stimulated their wide applications in energy storage, chemical sensing, gas absorption and storage, heterogeneous catalysis, *etc.*^{50,92–95} In recent years, conjugated porous polymers (CPPs) have attracted good research interest in photocatalytic H₂ evolution. Their inherently porous structures can enhance the interaction between the polymers and water molecules and promote the charge transfer in different dimensions, which is expected to boost the photocatalytic reactions.^{33,95}

The statistical copolymerization of aromatic monomers was reported as an important strategy to enhance the catalytic activity of CPPs by improving their physical properties such as band gap, porosity and charge mobility, *etc.* For instance, Cooper and co-workers synthesized fifteen conjugated microporous polymers (CMPs) by adjusting the feed ratio of the four comonomers shown in Fig. 7.⁹⁶ The CMPs have high surface areas of 597–1710 m² g⁻¹ and their band gaps gradually decrease from 2.95 to 1.94 eV with the increase of pyrene content in the polymer. Without adding a metal cocatalyst, the polymers showed gradually enhanced HER when the band gap decreased from 2.95 to 2.33 eV. With the ratio of 1,4-benzene diboric acid and 1,2,4,5-tetrabromobenzene of 2 : 1, polymer **21** exhibited the highest HER of 174 μmol h⁻¹ g⁻¹. However, further narrowing the band gap of polymer led to a decrease in the H₂ production. This was likely ascribed to the increased nonradiative recombination of electron and hole in the pyrene-rich polymers.

Similarly, Jiang *et al.* synthesized D-π-A CPPs using pyrene as a donor, benzothiadiazole as an acceptor, and benzene or biphenyl as a π-crosslinker unit.⁹⁷ Compared to D-A polymers (without π unit), the extended π-conjugation endows the D-π-A polymers with more efficient charge migration to the surface of

polymer particles. In addition, the electron push-pull interaction in the D-π-A polymer backbone could facilitate charge separation in the photocatalytic process. Thereby, D-π-A polymers exhibited better hydrogen production than the D-A and D-π-D polymers (without acceptor). When the ratio of pyrene to benzothiadiazole was 9 : 2, a maximum HER of 296 μmol h⁻¹ g⁻¹ was obtained (>420 nm). Xiang *et al.* developed a series of CPPs by copolymerizing 1,3,6,8-tetrabromopyrene and 3,8-dibromophenanthroline. A remarkable HER of 4200 μmol h⁻¹ g⁻¹ was achieved by the polymer having a 3 : 1 ratio for 1,3,6,8-tetrabromopyrene and 3,8-dibromophenanthroline.⁹⁸ In addition, Wang *et al.* further modified polymer **21** by gradually replacing the benzene units in the polymer network with dibenzo[*b,d*]thiophene sulfone units.⁹⁹ They claimed that the sulfone unit could act as an electron-output “tentacle” to trap photogenerated electrons from polymers, which is beneficial for the proton reduction reaction. The HERs of the resulting polymers were significantly enhanced with increasing content of sulfone unit in the polymer. This molecular modification provided a maximum HER of 400 μmol h⁻¹.

The donor-acceptor combination approach is one of the most prevailing methods used in polymer solar cells to enhance the power conversion efficiency.^{100–104} The alternating electron-rich donor (D) and electron-deficient acceptor (A) in the main chain could induce intramolecular charge transfer (ICT) interactions between D and A units, resulting in a narrowing of the band gap and molecular energy level modulation. Owing to this, the D-A approach has attracted a great deal of interest in designing CPs for photocatalytic H₂ evolution.

In 2016, Yu *et al.* reported CPPs by the combination of various chromophores (**M1–M4**) with biphenyl and bipyridyl, respectively.¹⁰⁵ With the chromophores from strong acceptors (**M1**) to strong donors (**M4**), the band gaps could be fine-tuned from 1.81 to 2.89 eV for **22a–25a**. The replacement of biphenyl (weak donor) with bipyridyl (weak acceptor) led to lower band gaps of 2.06–2.45 eV for **23b–25b** (Fig. 8a). The overall H₂ evolution of bipyridyl containing polymers is better than that of biphenyl containing polymers (Fig. 8b). Polymer **22b** gave the highest HER of 1900 μmol h⁻¹ g⁻¹ (>400 nm) as a result of the improved light absorption, wettability and charge separation. Later, a higher HER of 2590 μmol h⁻¹ g⁻¹ was obtained by polymer **26** through further optimizing nitrogen-containing acceptors in **M4** based polymers.¹⁰⁶

2,1,3-Benzothiadiazole (BT) is a strong acceptor unit used in designing D-A copolymers. Wang and co-workers demonstrated that the substitution position of BT on the central phenyl unit has a profound effect on the porosity and morphology of the resulting polymers.¹⁰⁷ CPPs with 1,2,4,5 and 1,2,4-substitution exhibited a fused particle-like shape and a fibre shape was observed for their counterpart with 1,3,5-substitution, while the BET surface areas of the three polymers ranged from 40 to 280 m² g⁻¹ with optical gaps of 2.25–2.44 eV. Consequently, polymer **27** loaded with 3 wt% Pt yielded a HER of 400 μmol h⁻¹ g⁻¹, while 8 μmol h⁻¹ g⁻¹ and 26 μmol h⁻¹ g⁻¹ were observed for the polymers with BT at the 1,2,4 and 1,3,5-positions of the central phenyl ring, respectively. Interestingly, compared to those 3D CPPs, a 1D linear

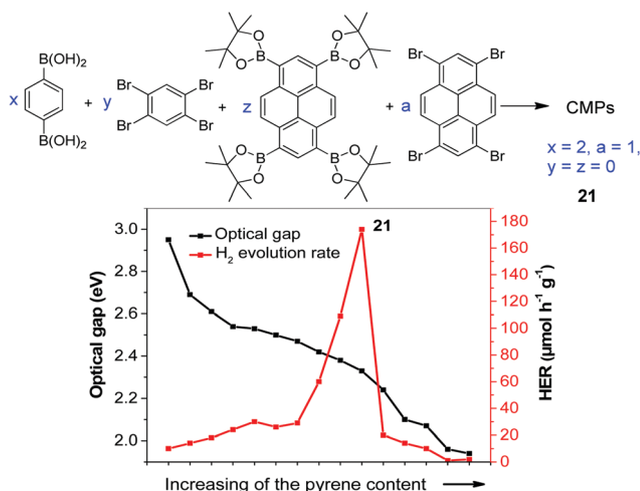


Fig. 7 Synthesis of conjugated microporous polymers with varying ratios of benzene and pyrene as well as the correlation of optical gap (red line) and hydrogen production (black line) as a function of increasing pyrene content in the polymer backbone.

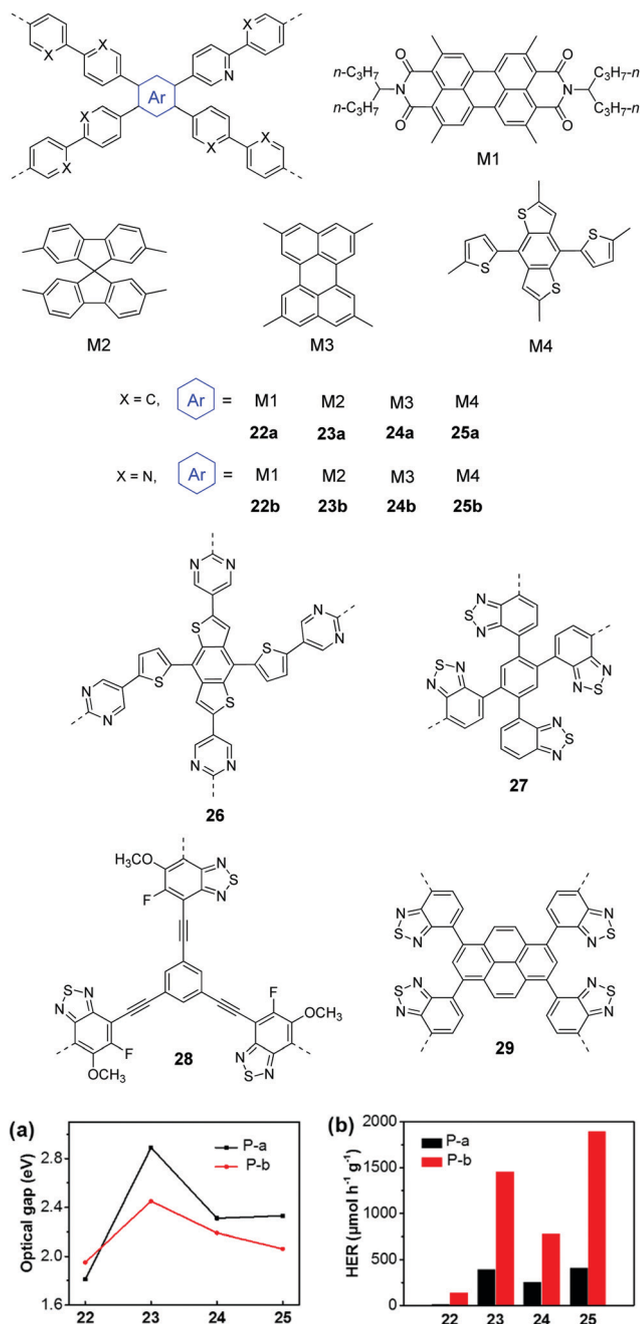


Fig. 8 Chemical structures of D-A type CPPs and (a) the optical gaps as well as (b) photocatalytic H₂ production of polymers 22–25 (P-a: 22a–25a; P-b: 22b–25b). Reprinted with permission from ref. 105. Copyright 2016 American Chemical Society.

polymer consisting of alternating phenyl and benzothiadiazole units showed a much higher HER of $2320 \mu\text{mol h}^{-1} \text{g}^{-1}$, which was ascribed to its narrowest band gap (2.17 eV) and improved charge transport in the polymer.

Fluorine (F) substitution on the CP backbones is an effective way to engineer their optoelectronic properties.¹⁰⁸ The van der Waals radii of F atom is only 1.35 Å, which is close to that of the hydrogen atom (1.2 Å).¹⁰⁹ Besides, with a Pauling electronegativity value of up to 4.0, the F atom tends to impair the

electron density in the D-A type CPs. As a consequence, the introduction of F does not cause a significant steric hindrance, but has a profound influence on the molecular orbits of the polymers.

Zhu and co-workers studied the F substitution effect in the case of BT incorporated linear CPs and conjugated porous polymers on their photocatalytic activities.¹¹⁰ It was found that F substitution had a weak influence on the optical gaps of polymers but the charge transfer efficiency was improved significantly. With the increase of F atom number in BT from 0 to 2, the HOMO and LUMO levels of polymers decrease simultaneously. Polymer 28 incorporating CH₃O and F substituted BT units showed a HER of $9.6 \text{ mmol h}^{-1} \text{g}^{-1}$, which was 28.8-fold as large as that of its counterpart without fluorination. The F substitution was also examined by Wang and co-workers.¹¹¹ They found that the F substitution on BT led to a more negative LUMO level, which could increase the driving force for proton reduction. In this work, polymer 29 showed a 5-fold enhancement of HER compared to its non-fluorinated counterpart.

In general, extending the π -conjugation along the skeleton had multiple impacts on the morphologies, porous properties and photophysical properties of the resulting polymers, which have a synergistic effect on the photocatalytic performances of the CPs. For example, Zhang and co-workers examined the π -extended conjugation effect of polymers 30–33 on their H₂ production performances.¹¹² The spacer change led to a profound effect on the porosity of the polymers. Polymer 30 with a small *para*-phenylene spacer has micropores, while both micro- and mesopores were found for 31 and 33 with a longer spacer. By contrast, polymer 32 showed a much broader pore size distribution, which was due to the longest polyphenylene spacer in the polymer network. As a result, the BET surface areas were 669, 750, 564, and 834 $\text{m}^2 \text{g}^{-1}$ for 31, 32, 33, and 34, respectively. In addition, the four polymers showed different energy level alignments due to the donor (arene units) and acceptor (tricyanomesitylene) interactions (Fig. 9a). The HERs were found to be 134, 598, 908, and 620 $\mu\text{mol h}^{-1} \text{g}^{-1}$ for 30, 31, 32, and 33, respectively. Despite the similar band gap, the highest photocatalytic activity of 33 as compared to 30 and 31 was ascribed to its nanoparticle morphology and the largest surface area.

Similarly, Jiang *et al.* developed polymers 34–36 by changing the central π -linker from benzene to *p*-terphenyl.¹¹³ Because of the higher steric hindrance of the biphenyl than phenyl, 34 showed a smaller BET surface area of $811 \text{ m}^2 \text{g}^{-1}$ than 35 ($915 \text{ m}^2 \text{g}^{-1}$), while 36 with the longest linker showed the lowest surface area of $447 \text{ m}^2 \text{g}^{-1}$ (Fig. 9b). Moreover, the introduction of the extended linker seemed to favour the formation of larger pores. The three polymers showed a similar pore distribution but more mesopores or macropores were found for 35 and 36 (Fig. 9c). In addition, polymer 34 has a smaller band gap of 2.53 eV than 35 (2.62 eV) and 36 (2.64 eV). Polymer 34 exhibits better charge separation efficiency as evidenced by a much lower emission than that of 35 and 36 (Fig. 9d). Accordingly, polymer 34 yielded the highest HER of $2460 \mu\text{mol h}^{-1} \text{g}^{-1}$, while HERs of 188 and $116 \mu\text{mol h}^{-1} \text{g}^{-1}$ were obtained by 35 and 36, respectively.

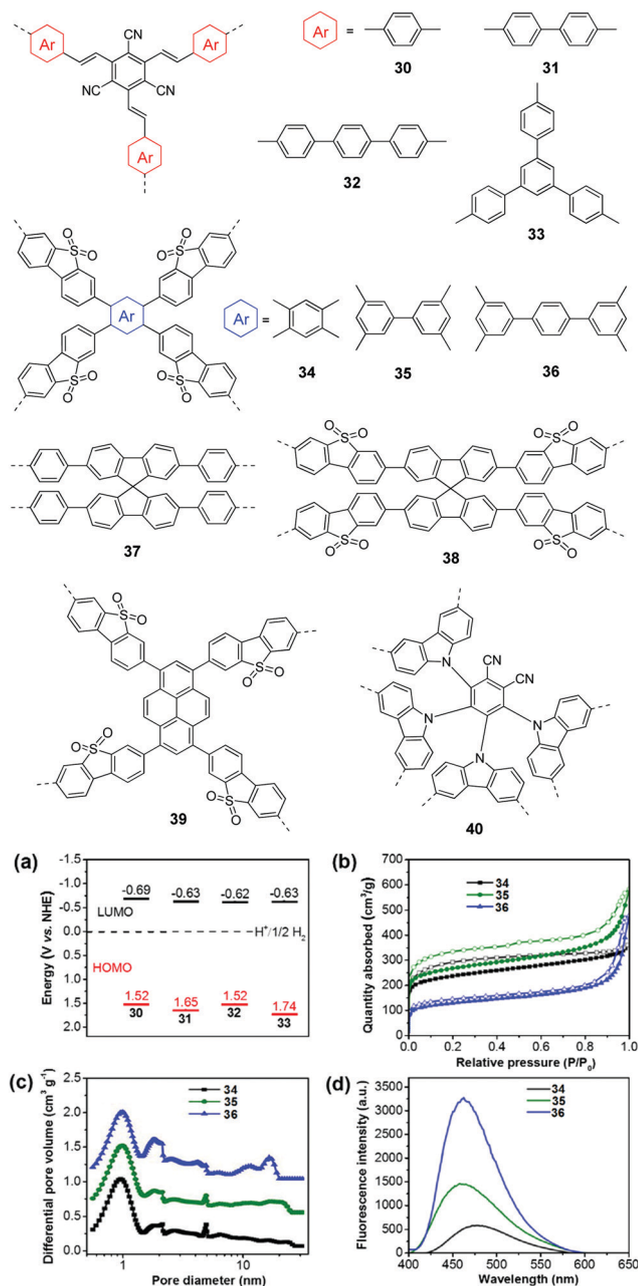


Fig. 9 Chemical structures of polymers **30–40**. (a) HOMO and LUMO band positions of polymers **30–33**. (b) Nitrogen sorption isotherms, (c) photoluminescence spectra, and (d) hydrogen evolution of polymers **34–36**. Reprinted with permission from ref. 112 and 113. Copyright 2017 Wiley-VCH and 2016 American Chemical Society.

Cooper *et al.* found that the optical gaps of the CMPs could be tuned by changing the monomer linker length and type of linkage, which led to different H₂ evolution activities.¹¹⁴ With spirobifluorene as the core, polymer **37** exhibited a 15-fold HER as compared to that of the analogue with phenyl as the core. In addition, by extending the central core from phenyl to spirobifluorene, polymer **38** was reported with an improved HER of 3106 $\mu\text{mol h}^{-1} \text{g}^{-1}$.¹¹⁵

Jiang *et al.* demonstrated that the substitution position of dibenzo[*b,d*]thiophene sulfone had a significant influence on

the photocatalytic performances.¹¹⁶ They prepared two CMPs by linking pyrene at the 3,7 and 2,8 positions of sulfone, respectively. Polymer **39** with the 3,7-linking pattern showed a narrower optical gap and lower emission intensity than the other polymer, which indicated better photogenerated charge migration and thus led to a much higher HER of 8523 $\mu\text{mol h}^{-1} \text{g}^{-1}$ than that of the 2,8-linked polymer (2650 $\mu\text{mol h}^{-1} \text{g}^{-1}$). Similarly, Su and co-workers found that substitution positions of CN groups and carbazole on the central phenyl ring could affect spectral absorption and charge separation of the resulting polymers. With an optimized structure, polymer **40** showed an excellent HER of 2103.2 $\mu\text{mol h}^{-1} \text{g}^{-1}$ and 6.4% apparent quantum yield at 420 nm.¹¹⁷

3.1.3 Covalent organic frameworks. Covalent organic frameworks (COFs) are an emerging class of 2D or 3D crystalline polymers formed by linking organic building units into extended structures.^{47,49,51,118} The high degree of crystallinity favours charge separation and transport, which endows COFs with promising catalytic activity. Moreover, the well-defined pores, excellent stability and fine-tuned physicochemical properties of COFs make them attractive photocatalysts for H₂ production.

In an early contribution, Lotsch *et al.* synthesized COF **41** (TFPT-COF) by a condensation reaction of 2,5-diethoxyterephthalohydrazide and 1,3,5-tris(4-formylphenyl)triazine in a 1,4-dioxane/mesitylene mixture.¹¹⁹ COF **41** showed a honeycomb-type layered structure with mesopores of 3.8 nm and its BET surface area was up to 1603 $\text{m}^2 \text{g}^{-1}$. Pt-loaded COF **41** could produce H₂ from water at a constant rate of 1970 $\mu\text{mol h}^{-1} \text{g}^{-1}$. Subsequently, they reported a series of azine-linked COFs **42–45** (N_x-COFs, x = 0–3) with different N numbers in the central aryl ring (Fig. 10).¹²⁰ The molecular design is based on the varied dihedral angles between the central aryl ring and peripheral phenyl rings by replacing the H atom (green dots) with a N atom (Fig. 11a), which led to different planarity of the platform and thus varied crystallinity and porosity of the resulting COFs. As displayed in Fig. 11b, compared to **42** (N₀-COF) and **43** (N₁-COF), **44** (N₂-COF) and **45** (N₄-COF) have better crystallinity as revealed by their sharper PXRD peaks, which is beneficial for charge migration in the photocatalytic reaction. Besides, the four COFs showed close band gaps of 2.6–2.7 eV, but different BET areas of 326 to 1536 $\text{m}^2 \text{g}^{-1}$. The HERs of Pt-modified **42**, **43**, **44** and **45** were 23, 90, 438 and 1703 $\mu\text{mol h}^{-1} \text{g}^{-1}$, respectively. The progressively enhanced HER was likely attributed to the increased surface area and improved charge migration when the number of N atoms was increased in the frameworks.

The facile choice of building blocks did permit a good modulation of the porosity, wettability and optoelectronic properties of β -ketoenamine linked COFs, which led to enhanced photocatalytic performance. When different building blocks were introduced, the corresponding band gaps were 2.10, 1.85 and 2.28 eV for **46** (F-COF), **47** (FS-COF) and **48** (TP-COF), respectively (Fig. 11c).¹²¹ Moreover, **46** and **47** containing a sulfone unit exhibited better wettability than **48** as revealed by the water contact angle measurement, which facilitates dispersion of COF particles in water and the interaction between COF and water in the photocatalytic reaction. Loaded with 3 wt% Pt,

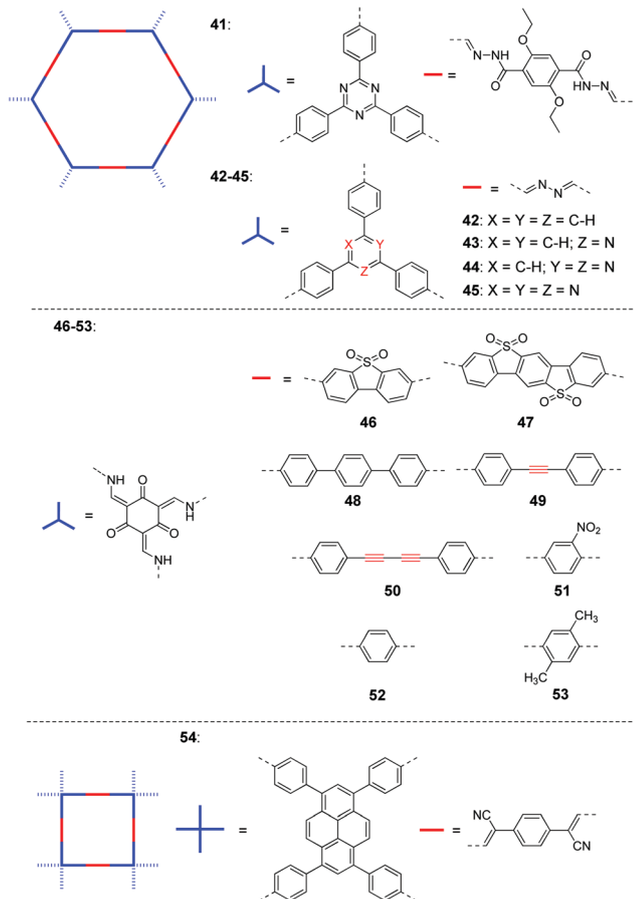


Fig. 10 Chemical structures of COFs 41–54 for photocatalytic H₂ production under visible light.

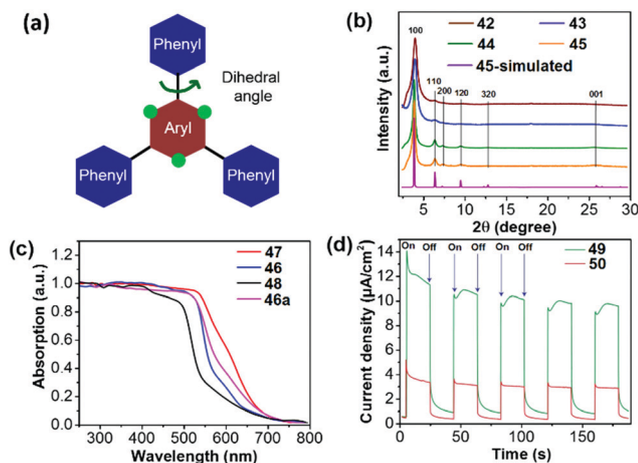


Fig. 11 (a) Design principle and (b) PXRD patterns of COFs 42–45. (c) Diffuse reflectance spectra of COFs 46–48 (46a is an amorphous analogue of 46). (d) Transient photocurrent for COFs 49 and 50 measured in 0.5 M KCl solution under visible light irradiation. Reprinted with permission from ref. 120–122. Copyright 2015 Nature Publishing Group, 2018 Nature Publishing Group and 2018 American Chemical Society.

the HER of 46 was up to 4.44 mmol h⁻¹ g⁻¹, while 10.1 and 1.6 mmol h⁻¹ g⁻¹ were reached for 47 and 48, respectively.

The best H₂ evolution of 47 was ascribed to its lowest band gap, good wettability, highest BET area (985 m² g⁻¹ for 46, 1288 m² g⁻¹ for 47 and 919 m² g⁻¹ for 48).

In addition, the photocatalytic activity of COFs is susceptible to linking bond length between comonomers. Thomas and co-workers reported that the catalytic activity of the diacetylene-based COF 49 (TP-EDDA) was much better than the acetylene-based COF 50 (TP-BDDA).¹²² Despite the almost identical band gaps (2.34 eV for 49 and 2.31 eV for 50), COF 50 has an HER of 324 μmol h⁻¹ g⁻¹, whereas a relatively low HER of 30 μmol h⁻¹ g⁻¹ was achieved for 49. The significantly enhanced HER was attributed to the higher charge carrier generation of 50 under visible light, which could be revealed by the larger photocurrent response of 50 as shown in Fig. 11d.

Substituent modification in the COF could also bring good photocatalytic performances. Sun *et al.* prepared three ketoenamine-based COFs (51–53) and studied the effect of different functional groups on their photocatalytic activities.¹²³ The photocurrent density of COF 53 (TpPa-COF-(CH₃)₂) is about 2.3-fold and 6.5-fold higher than that of 52 (TpPa-COF-H) and 51 (TpPa-COF-NO₂), respectively. Moreover, the electron transfer resistance value decreased gradually when the electron-donating ability of the substituent groups became stronger (from NO₂ to H, then CH₃). These results indicated that the charge separation of COFs was improved by increasing the electron-donating properties of the substituents. With an optical gap of 2.06 eV and the best charge separation efficiency, COF 52 showed a HER of 8.33 mmol h⁻¹ g⁻¹, which was much higher than that of 51 (1.56 mmol h⁻¹ g⁻¹) and 53 (0.22 mmol h⁻¹ g⁻¹).

In comparison with those reported COFs based on imine-, hydrazone-, or azine linkers, fully π-conjugated COFs are expected to allow for more efficient exciton separation and migration over the framework, which is good for H₂ production. Very recently, Jiang and co-workers reported a 2D sp² carbon-COF 54 (sp²c-COF) as the H₂-producing photocatalyst, into which electron-withdrawing cyano groups were incorporated to impart a conjugated electron donor-acceptor structure and the phenylenevinylene further improved π conjugation.¹²⁴ The BET surface area of COF 54 was 602 m² g⁻¹ and its optical gap was 2.05 eV. A stable HER of 1360 μmol h⁻¹ g⁻¹ was obtained under visible light. The HER could be enhanced to 2120 μmol h⁻¹ g⁻¹ by attaching 3-ethylrhodanine as an end-capping group to strengthen the push-pull effect.

3.1.4 Covalent triazine frameworks. As another class of organic framework materials, covalent triazine frameworks (CTFs) are featured by unique porosity and high nitrogen content, which could be obtained by the choice of suitable organic linkers and synthetic conditions. Different from COFs, most of CTFs reported so far are amorphous or semicrystalline. Recently, CTFs have been employed for a range of applications, such as CO₂ capture and storage, photocatalysis and energy storage.^{53,125–127}

In 2015, Wu and co-workers prepared CTF 55 (CTF-T1) by a trimerization reaction of terephthalonitrile using CF₃SO₃H as the catalyst at room temperature (Fig. 12).¹²⁸ The CTF exhibits a band gap of 2.94 eV and a low HER of 200 μmol h⁻¹ g⁻¹ after loading 3 wt% Pt.

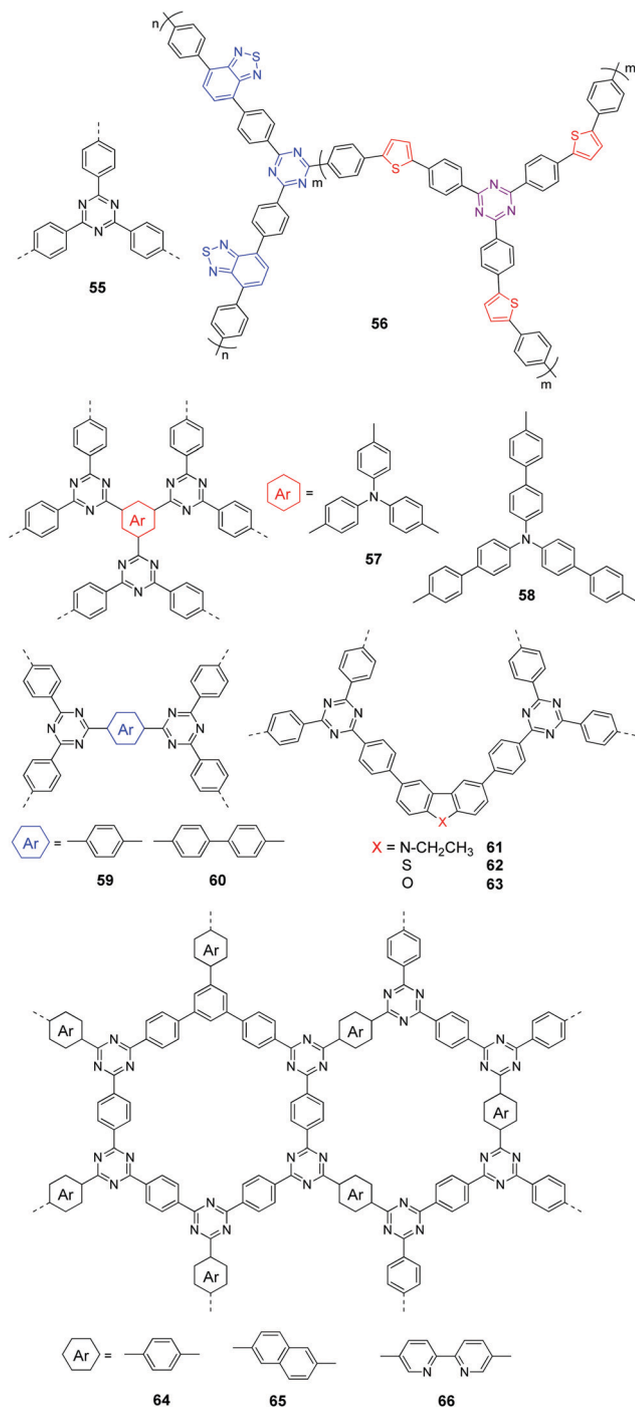


Fig. 12 Chemical structures of polymers 55–66.

Afterward, Thomas *et al.* showed that a CTF with a low HER could be converted into a more efficient photocatalyst by further mixing the pre-synthesized CTF in $\text{CHCl}_3/\text{CF}_3\text{SO}_3\text{H}$ with ZnCl_2 salt for 10 minutes. The HER could increase from 186 to $1072 \mu\text{mol h}^{-1} \text{g}^{-1}$ due to the decreased optical gap (2.42 eV) and improved crystallinity.¹²⁹ Alternatively, Wang *et al.* further processed the pre-synthesized CTF 55 through thermal annealing at 300 °C, which induced structural reorganization of 55 and yielded new conjugated heterocycles in the polymer.

With improved light absorption and charge mobility, 3 times enhancement of HER was achieved under visible light as a result of the thermal treatment of 55.¹³⁰

The fast recombination of photogenerated electrons and holes of CTFs is another major disadvantage preventing them from achieving higher efficiencies. In this context, Li *et al.* developed a molecular heterostructure 56 (CTF-Th) via a two-step synthetic strategy.¹³¹ CTF 56 exhibited higher photocurrent and charge carrier density than its single-component counterparts, which suggested a largely improved charge separation efficiency. As a result, a HER of $6.6 \text{ mmol h}^{-1} \text{g}^{-1}$ was obtained for 56 under visible light.

Compared to the commonly used ionothermal method (usually at 400 °C), polycondensation under relatively gentle conditions is desirable for the preparation of CTFs with tunable functions and geometries. Tan and co-workers developed CTFs 57–60 by condensation between 1,4-benzenedicarboximidamide and different aldehydes in the presence of Cs_2CO_3 .¹³² The structural variations have an obvious impact on the porosity and light absorption of the resulting CTFs. The BET surfaces areas are 807, 764, 663, and $757 \text{ m}^2 \text{g}^{-1}$ for 57 (CTF-HUST-3), 58 (CTF-HUST-3), 59 (CTF-HUST-1) and 60 (CTF-HUST-2), respectively (Fig. 13a). CTFs 57 and 58 showed broader absorption compared to 59 and 60 (Fig. 13b). Under visible light, CTF 60 showed the highest HER of $2647 \mu\text{mol h}^{-1} \text{g}^{-1}$, while the HERs were 1238, 1582 and $1460 \mu\text{mol h}^{-1} \text{g}^{-1}$ for 57, 58 and 59, respectively. Using a similar synthetic method, Tan *et al.* prepared donor–acceptor CTFs 61–63.¹³³ Upon incorporating a strong electron-donating carbazole unit, 61 (CTF-N) showed the lowest band gap of 2.17 eV compared to 62 (CTF-S, 2.47 eV) and 63 (CTF-O, 2.67 eV) (Fig. 13c). Meanwhile, the enhanced charge separation/transfer efficiency of 61 was studied by employing fluorescence and photoelectrochemical measurements (Fig. 13d). Therefore, CTF 61 decorated with 3 wt% Pt exhibits the best HER of $10760 \mu\text{mol h}^{-1} \text{g}^{-1}$.

CTFs could also be fabricated by a Pd-catalyzed coupling reaction. In 2017, Cooper *et al.* synthesized two series of CTFs with similar structures by either Suzuki coupling of 2,4,6-tris-(4-bromophenyl)-1,3,5-triazine and aromatic monomers or trimerization of dicyanobenzene derivatives.¹³⁴ In this case, Pt loaded CTFs from trimerization exhibited higher activity than their counterparts from Suzuki coupling. Moreover, the length of the *para*-phenylene spacer has a profound effect on the band gaps and porosity for both series of CTFs and thus affects their photocatalytic activities. Based on the D–A interaction between thiophene derivatives (donor) and triazine units (acceptor), Bojdyś *et al.* designed eight CTFs with fine-tuned band gaps (2.06–2.70 eV) and charge transfer in the polymer.¹³⁵ As a result, a maximum HER of $3158 \mu\text{mol h}^{-1} \text{g}^{-1}$ was achieved by the CTF consisting of benzotrithiophene and triazine.

Instead of introducing different building blocks into CTFs, heteroatom doping is another efficient method to boost the HERs of CTFs. Su *et al.* prepared S-doped CTF samples (CTFS₁₀) by annealing treatment of CTFs with sulfur.¹³⁶ The S concentration in the framework was 0.5–0.52 atom% when the CTF was mixed with different amounts of S (5–30 wt%). The characteristic peak of C–S at 286.5 eV revealed by XPS analysis showed that the

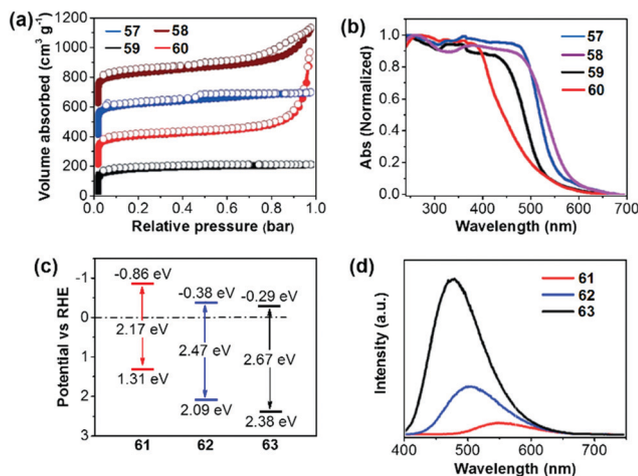


Fig. 13 (a) Nitrogen sorption curves and (b) UV-vis absorption spectra of CTFs **57–60**. (c) Energy diagram and (d) fluorescence spectra of CTFs **61–63**. Reproduced with permission from ref. 132 and 133. Copyright 2017 Wiley-VCH and 2018 Royal Society of Chemistry.

S atom is covalently bound to the CTF backbone. After depositing 3 wt% Pt, the CTFs mixed with 10 wt% S showed the best HER of 2000 $\mu\text{mol h}^{-1} \text{g}^{-1}$, which is 5-fold enhanced compared to that of the pristine CTFs. This is due to the sharply decreased band gap (from 2.87 eV to 1.87 eV) and improved charge transport after S doping. Alternatively, the photocatalytic activity of CTFs could be enhanced by phosphorus doping due to the narrower band gap and more efficient charge transfer.¹³⁷

Besides the above-mentioned amorphous CTFs, Tan and co-workers reported the successful preparation of crystalline CTFs **64–66** by *in situ* oxidizing alcohols into aldehyde monomers and studied their H₂ production activities.¹³⁸ It was claimed that high crystallinity of CTFs could be achieved by slowing down their nucleation process through controlling aldehyde monomer generation in the polycondensation reaction. All CTFs showed good crystallinity with BET surface areas of 520–599 $\text{m}^2 \text{g}^{-1}$. The HERs of **64** (CTF-HUST-C1), **65** (CTF-HUST-C5) and **66** (CTF-HUST-C6) were 5100, 2400 and 650 $\mu\text{mol h}^{-1} \text{g}^{-1}$, respectively. In addition, Zou and co-workers prepared crystalline CTFs by condensation of melamine (MA) and pyromellitic dianhydride (PMDA) under thermal conditions. The obtained CTF displayed a band gap of 2.7 eV and HER of 7 $\mu\text{mol h}^{-1}$.¹³⁹

3.1.5 Carbon nitrides. Pure g-C₃N₄ materials have an optical gap of about 2.7 eV and suitable CB/VB positions for water reduction and oxidation. g-C₃N₄ has captured broad research interest toward photocatalytic H₂ evolution since the pioneering report by Wang and co-workers in 2009.¹⁴⁰ The photocatalytic activity of g-C₃N₄ has subsequently been improved by morphology modulation, crystallinity control, doping and composite construction, *etc.*^{29–35,141} Given the fact that these developments have been summarized in previous reviews,^{29–35,141} this section briefly presents the representative research on molecular modification of g-C₃N₄ photocatalysts for H₂ production from water.

The elemental doping strategy has been intensively developed to boost the photocatalytic performance of g-C₃N₄ by tuning its band structure, light absorption and charge carrier transport.^{27–32}

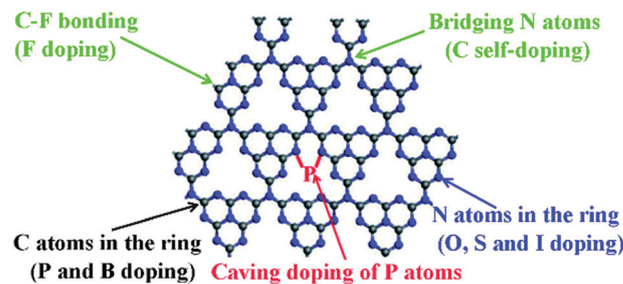


Fig. 14 Possible elemental doping sites in the single layer of g-C₃N₄. Reproduced with permission from ref. 141. Copyright 2017 Royal Society of Chemistry.

Various elements such as F, Br, I, N, P, O, S and B have been doped in the skeleton of tri-s-triazines based g-C₃N₄, as shown in Fig. 14.¹⁴¹ For instance, Wang *et al.* prepared F doped g-C₃N₄ (CN-F) by adding NH₄F in the polymerization of dicyanamide. The formation of C–F bonds in the polymer led to a slightly red-shifted optical absorption and 2.7 times higher HER ($\sim 130 \mu\text{mol h}^{-1} \text{g}^{-1}$) than that of the unmodified sample ($> 420 \text{ nm}$).¹⁴² Nitrogen-doped g-C₃N₄ (CN-N) could be synthesized using pre-treated melamine with hydrazine hydrate.¹⁴³ This modification gave rise to improved charge transfer and a slightly narrowed band gap from 2.72 eV to 2.65 eV, as a result, CN-N showed a 1.8-fold higher HER of 553.5 $\mu\text{mol h}^{-1} \text{g}^{-1}$ than that of pristine g-C₃N₄ ($> 400 \text{ nm}$). On the other hand, the doping of multiple elements in g-C₃N₄ provided more variable properties.¹⁴¹ Dai *et al.* reported carbon and phosphorus co-doped g-C₃N₄ (CN-C/P), which showed a significantly larger surface area of 141.1 $\text{m}^2 \text{g}^{-1}$ than pristine g-C₃N₄ (24.9 $\text{m}^2 \text{g}^{-1}$).¹⁴⁴ Moreover, this modulation also led to extended optical absorption and promoted charge transfer in the polymer. Benefiting from this, CN-C/P exhibited a HER of 1493.3 $\mu\text{mol h}^{-1} \text{g}^{-1}$, which was 9.7-fold as compared to that of pristine g-C₃N₄.

Organic small molecules are also capable of doping into g-C₃N₄ by copolymerization to adjust its π -electronic system and band structure. In 2010, Wang and co-workers prepared modified g-C₃N₄ (CN-BA) using dicyandiamide and barbituric acid as the precursor and dopant, respectively.¹⁴⁵ With an optimization of amount of dopants, CN-BA exhibited a maximum HER of 294 $\mu\text{mol h}^{-1} \text{g}^{-1}$ mainly due to the extended light absorption. After that, many other small molecules such as pyrimidine,¹⁴⁶ quinoline,¹⁴⁷ and thiophene¹⁴⁸ were incorporated into g-C₃N₄ by several groups.

Structure defect engineering is another efficient approach for improving the photocatalytic performance of g-C₃N₄. The introduction of defects into the g-C₃N₄ backbone can not only modify the band structure by generating mid-gap states to extend the light absorption, but also act as trapping sites for charge carriers to inhibit the recombination of electrons and holes. In the presence of KOH, Zhang *et al.* introduced nitrogen vacancies and cyano groups into g-C₃N₄, which resulted in a reduced band gap from 2.68 to 2.36 eV and enhanced charge carrier separation.¹⁴⁹ Therefore, the defect-rich g-C₃N₄ showed a HER of 6.9 $\text{mmol h}^{-1} \text{g}^{-1}$, which was nearly twice that of pristine g-C₃N₄. In a very recent study, Wang *et al.* introduced

defects into $g\text{-C}_3\text{N}_4$ by NaBH_4 reduction of a crystalline pristine polymer under an inert atmosphere, which achieved nearly 8 times higher HER than the pristine polymer.¹⁵⁰ Apart from those modification strategies, the development of analogous polymers of $g\text{-C}_3\text{N}_4$ such as $\text{C}_3\text{N}_3\text{S}_3$ ¹⁵¹ offered new opportunities for exciting findings in photocatalytic H_2 production.

In some cases, the construction of polymer heterojunctions (PHJs) can efficiently modulate the band structure and charge transport of $g\text{-C}_3\text{N}_4$ through inter/intramolecular $\pi\text{-}\pi$ interactions between $g\text{-C}_3\text{N}_4$ and the aromatic structures. Shen *et al.* prepared three PHJs by coating different polyfluorene polymers on the surface of $g\text{-C}_3\text{N}_4$, which resulted in extended optical absorption and promoted charge transfer within the PHJs.¹⁵² Under visible light, all three pure CPs have no photocatalytic activity and pure $g\text{-C}_3\text{N}_4$ showed a very low HER of $87.4 \mu\text{mol h}^{-1} \text{g}^{-1}$. Among the three PHJs, 5% polymer **15** coated $g\text{-C}_3\text{N}_4$ (CN-P15) exhibited a maximum HER of $722.2 \mu\text{mol h}^{-1} \text{g}^{-1}$. Using a similar method, Chen *et al.* deposited Pdots on the surface of $g\text{-C}_3\text{N}_4$ nanosheets, which yielded an enhanced HER *via* improved charge transfer from Pdots to $g\text{-C}_3\text{N}_4$ nanosheets.¹⁵³ In this case, 5 wt% polymer **17** deposited PHJ (CN-P17) exhibited a HER of $929.3 \mu\text{mol h}^{-1} \text{g}^{-1}$, which is higher than that of its counterpart coated with polymer **15** ($439.6 \mu\text{mol h}^{-1} \text{g}^{-1}$). Besides, Wang *et al.* fabricated type II heterojunctions by anchoring pyrene-based polymers onto CN to improve light absorption and interface charge transfer.¹⁵⁴ By optimizing the amount and molecular structures of polymers, CN-PyP hybrids attained a maximum HER of $600 \mu\text{mol h}^{-1} \text{g}^{-1}$ without Pt deposition. PHJs based on $g\text{-C}_3\text{N}_4$ could also be prepared by linking another component through a chemical bond. Yan *et al.* prepared heterojunctions (CN-COF) by *in situ* formation of a COF on the surface of pre-synthesized $g\text{-C}_3\text{N}_4$ through condensation of amino group in $g\text{-C}_3\text{N}_4$ and aldehyde group in the COF.¹⁵⁵ CN-COF attained a dramatically enhanced HER of $10.1 \text{ mmol h}^{-1} \text{g}^{-1}$, which is much higher than those of the pure COF ($0.16 \text{ mmol h}^{-1} \text{g}^{-1}$), $g\text{-C}_3\text{N}_4$ ($1.05 \text{ mmol h}^{-1} \text{g}^{-1}$) and their mixture ($0.93 \text{ mmol h}^{-1} \text{g}^{-1}$). Zou and co-workers fabricated heterostructured photocatalysts (CN-CTF) by linking $g\text{-C}_3\text{N}_4$ and CTFs with amide bonds, which demonstrated a good HER of $850 \mu\text{mol h}^{-1} \text{g}^{-1}$ and high stability compared to their counterpart (CN and COFs connected *via* van der Waals forces).¹⁵⁶ These investigations offer more opportunities for the design of H_2 -production polymer photocatalysts. The photocatalytic activity and stability of these materials remain a great concern.

3.1.6 The role of the Pt cocatalyst and residual Pd on photocatalytic H_2 production of CPs. As shown above, noble metals were commonly deposited as cocatalysts on the surface of polymer materials to promote their H_2 production activities. The deposition of noble metal cocatalysts can effectively lower the overpotential or activation energy for proton reduction reactions. Moreover, the cocatalysts could act as charge trap centre to extract more electrons from the polymer photocatalysts, which favours the electron-hole separation at the cocatalyst/polymer interface.^{157–159} Notably, as early as 1990s, Yanagida *et al.* have found that the catalytic activity of linear CPs could be improved by loading Ru nanoparticles.^{67–69}

Among the commonly used metal cocatalysts such as Ru, Pd, Rh, Au, and Ag, Pt is a versatile cocatalyst mainly because of its largest work function for trapping photogenerated electrons and lowest overpotential for photocatalytic H_2 evolution.¹⁵⁸ In general, Pt is deposited on CPs by photoreduction of H_2PtCl_6 in the reaction system. Under light illumination, $\text{Pt}(\text{vi})$ is reduced by the photo-induced electrons of the polymer to produce $\text{Pt}(0)$ nanoparticles with sizes of 3–5 nm *in situ* on the polymer surface. Compared to the pristine polymer, the Pt-polymer composite exhibited greatly enhanced H_2 evolution. H_2PtCl_6 with a Pt mass concentration of 3 wt% was commonly used in the photodeposition experiment, because too low Pt concentration provided only a few active sites for proton reduction, which resulted in a low enhancement of catalytic activity. Excessive amount of Pt deposited on the surface of the polymer photocatalyst could prevent light absorption by shielding the incident light and stimulate the aggregation of Pt nanoparticles, which is detrimental to the activity of Pt.

Pd-catalyzed cross-coupling polymerization reactions are widely used to synthesize CPs for photocatalytic water splitting. Residual Pd inevitably existed in the polymer and may play a vital role in its photocatalytic process.¹⁶⁰ Yu *et al.* have shown that the Pd content in the porous polymer greatly affected the H_2 evolution.¹⁰⁵ By changing the initial feeding ratio of $\text{Pd}(\text{PPh}_3)_4$ and monomer from 0.005 to 0.40, the Pd content was increased from 0.04% to 1.88% as measured by inductively coupled plasma mass spectrometry (ICP-MS). As shown in Fig. 15a, the HER of polymer **25b** was sharply improved with increasing Pd content from 0.04% to 0.46%, which became saturated at a higher residual Pd content. In addition, upon loading with 2 wt% Pt, a ~ 5 -fold enhancement of the AQY at 350 nm was measured for **25b** (Fig. 15b), which verified the importance of the Pt cocatalyst in the enhancement of photoactivity.

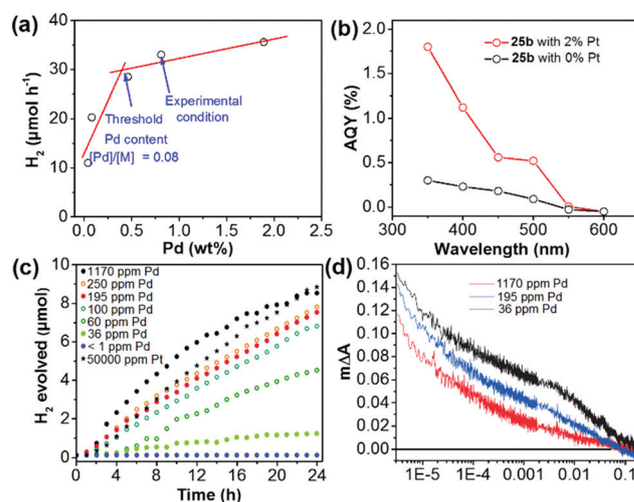


Fig. 15 (a) H_2 evolution of polymer **25b** with different residual Pd contents. (b) Apparent quantum yield (AQY) of polymer **25b** with 2% and without Pt loading at different wavelengths. (c) H_2 evolution of nanoparticles of polymer **15** with different amounts of Pd or Pt. (d) Transient kinetics of nanoparticles of polymer **15** with different Pd concentrations in a diethylamine/water mixture. Reprinted with permission from ref. 105 and 161. Copyright 2016 American Chemical Society and 2018 Wiley-VCH.

To study the role of Pd residues in determining the H₂ evolution activity, Jan Kosco and co-workers prepared colloidal polymer **15** nanoparticles with different Pd contents.¹⁶¹ In the sample preparation, they first purified the as-prepared polymer **15** by gel permeation chromatography (GPC), and then washed the polymer with diethyldithiocarbamate. After the treatment, the Pd content in polymer **15** was below 1 ppm and the content could be gradually adjusted to 1170 ppm by adding Pd₂(dba)₃ solution. Fig. 15c displays that the H₂ evolution of nanoparticles of polymer **15** is gradually enhanced with increasing Pd content from <1 ppm to ~100 ppm, at which point the evolved H₂ begins to saturate. Transient absorption spectroscopy (TAS) suggested that Pd could mediate electron transfer from the nanoparticles of polymer **15** to protons in the reaction media (Fig. 15d). It was found that photogenerated electrons were increasingly accumulated with decreasing Pd content in nanoparticles of polymer **15**. This indicates that the Pd cocatalyst could effectively suppress the accumulation of long-lived electrons and thereby accelerate photocatalytic H₂ generation of polymer **15**.

3.2 Conjugated polymers for photocatalytic O₂ evolution

The photocatalytic O₂ evolution is a critical step in water splitting because the four-electron transfer is an uphill process with large overpotentials and slow kinetics, which makes it very challenging to develop robust photocatalysts for water oxidation.^{162,163}

To date, most of the O₂ evolution photocatalysts have been based on nitrogen-rich CPs. In 2017, Xu *et al.* reported polymer **67**, which has a nanosheet morphology and a very low band gap of ~1.22 eV.⁴¹ With AgNO₃ as the electron scavenger, **67** exhibits an oxygen evolution rate (OER) of 1.0 μmol h⁻¹, which could be enhanced to 14.3 μmol h⁻¹ after loading Co(OH)₂ as the cocatalyst. More interestingly, polymer **67** can catalyze water oxidation under near-infrared light (>800 nm) and the OER is ~0.4 μmol h⁻¹ while g-C₃N₄ is not active.

Inspired by the facile tunability of CTFs, Wang *et al.* synthesized polymers **68–70** with different numbers of phenyl units in the polymer backbone.¹⁶⁴ The band gaps of **68–70** decreased from 2.98 eV to 2.36 eV with increasing number of phenyl units (Fig. 16a). Importantly, their HOMO/LUMO levels straddle the potential of water splitting, suggesting sufficient driving force for both photocatalytic water reduction and oxidation. Furthermore, both the HOMO and LUMO levels decreased with the increase of phenyl number in the polymer. After loading with 3 wt% Co(NO₃)₂ as a cocatalyst, polymer **69** showed the highest OER of 3 μmol h⁻¹ (>300 nm) compared to **68** (~1 μmol h⁻¹) and **70** (~0.4 μmol h⁻¹) in the presence of AgNO₃, which was attributed to the synergistic effect of improved light absorption and charge transfer. In addition, the catalytic activity of **69** is also much better than that of C₃N₄ (Fig. 16b) due to the lower LUMO position of **69**. Notably, polymer **69** could produce 7.2 μmol of O₂ after 8 h visible-light irradiation. Considering that high crystallinity of CTF materials is desirable for exciton separation and thus enhancing the photocatalytic activity, Tang and co-workers reported highly ordered CTFs made by microwave-assisted polymerization.¹⁶⁵ The crystallinity of the resulting CTFs became higher with the increase of microwave

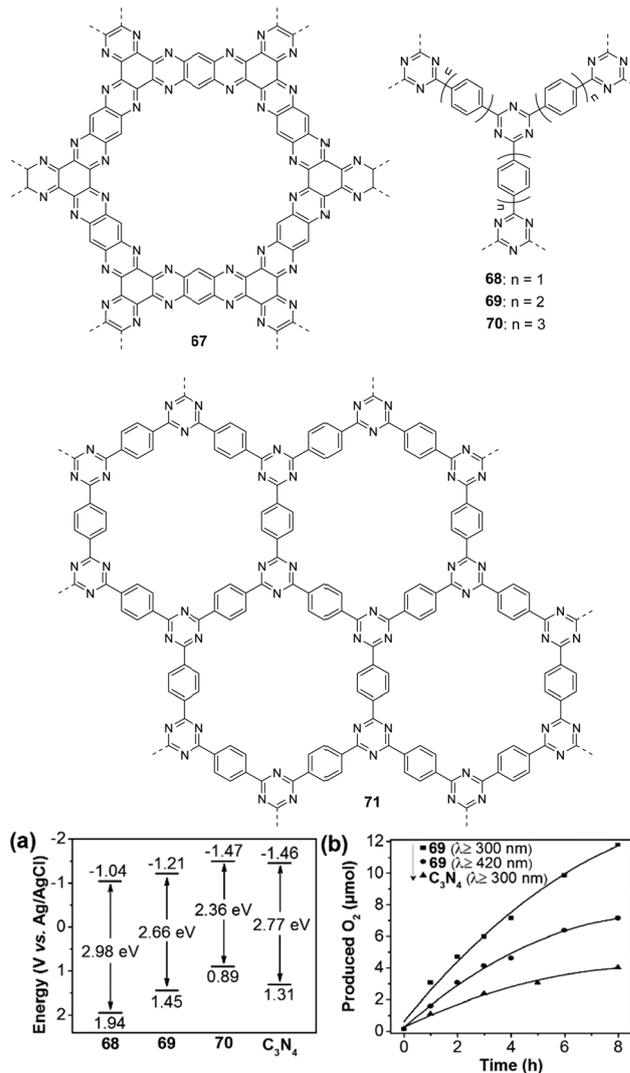


Fig. 16 Chemical structures of polymers **67–71**; (a) band structure diagram and (b) time course of the O₂ production for polymer **69** and C₃N₄. Reprinted with permission from ref. 164. Copyright 2018 Wiley-VCH.

power from 20 to 100 W. However, a further increase in microwave power can destroy the hexagonal units. At a power of 100 W, polymer **71** has a well-defined planar structure as evidenced by a G⁺ peak at 1613 cm⁻¹ from the Raman spectra analysis of CTFs. In addition, polymer **71** has conduction band (CB)/valence band (VB) levels of -0.74/+2.18 eV (vs. NHE, pH = 0), which is suitable for the photocatalytic water oxidation and reduction reaction. Polymer **71** gave an O₂ production rate of 140 μmol h⁻¹ g⁻¹ upon loading with 3 wt% RuOx in the presence of AgNO₃, which is higher than those of its counterparts prepared under different powers.

3.3 Conjugated polymers for photocatalytic overall water splitting

In comparison with the aforementioned photocatalytic water reduction or water oxidation, stoichiometric splitting of pure water into hydrogen and oxygen is cost-saving because both

photogenerated electrons and holes are utilized without using a sacrificial agent to suppress one of the half-reactions.^{8,166–168} Until now, the reports of efficient CPs for photocatalytic overall water splitting remains rare.

In 2016, Wang *et al.* demonstrated that Pt modified g-C₃N₄ was an efficient photocatalyst for overall water splitting and the morphology of g-C₃N₄ significantly affected the catalytic activity.¹⁶⁹ Among the three types of g-C₃N₄, the urea-derived g-C₃N₄ exhibits a 2D nanosheet structure and compact layer-like morphology, which showed better charge mobility as compared to the other two samples made from dicyandiamide and ammonium thiocyanate as the precursor. All three types of pure g-C₃N₄ have no photocatalytic activity in the absence of Pt. After depositing Pt on the three samples, g-C₃N₄ nanosheets exhibited the best activity with H₂ and O₂ evolution rates of 1.2 and 0.6 μmol h⁻¹, respectively (>420 nm). Very recently, Zhang and co-workers prepared 3D g-C₃N₄ with high crystallinity, displayed a larger surface area (130 m² g⁻¹) than those of bulk g-C₃N₄ (10.83 m² g⁻¹) and g-C₃N₄ nanosheets (93.84 m² g⁻¹).¹⁷⁰ Surprisingly, after loading 1 wt% Pt and 3 wt% IrO₂, 3D g-C₃N₄ exhibited H₂ and O₂ generation rates of 101.4 and 49.1 μmol h⁻¹ g⁻¹, respectively, which are about 11.8 and 5.1-fold higher than those of bulk g-C₃N₄ and g-C₃N₄ nanosheets. These examples demonstrated the critical role of the cocatalyst and surface morphology control of g-C₃N₄ in photocatalytic overall water splitting performance.

In 2017, Xu and co-workers prepared 1,3-diyne-linked CMPs 72–73 by Glaser oxidative coupling of terminal alkynes.⁴⁰ TEM measurements revealed that both polymers have sheet-like structures (Fig. 17a and b), which favours the separation and transport of photogenerated excitons to the surface of polymer nanosheets for water redox reactions. The CB/VB levels were determined to be -3.55/-6.5 eV for 72 and -2.9/-5.77 eV for 73, indicating that both polymers have appropriate electronic band structures for overall water splitting under the given conditions. As a result, both H₂ and O₂ are evolved for 73 with expected 2:1 stoichiometry upon light irradiation (Fig. 17c). The HER is 218 μmol h⁻¹ g⁻¹ for 73, which is higher than that of 72 (102 μmol h⁻¹ g⁻¹). This could be attributed to the lower

band gap of 73 (2.85 eV) compared to 72 (2.94 eV). This work demonstrates the great potential of polymer nanosheets as unique photocatalysts for overall water splitting.

4. Conjugated polymers for photocatalytic CO₂ reduction

4.1 Photocatalytic CO₂ reduction

The massive consumption of fossil fuels has caused the CO₂ levels in the atmosphere to increase over the years, which has raised serious environmental and safety concerns. One of the best solutions to address this challenge is to convert CO₂ into hydrocarbon fuels using solar light as an energy source.^{10,11,13,171–174} The technology could create huge economic and environmental benefits by simultaneously reducing the greenhouse effect and solving the energy shortage issue. However, CO₂ is a very stable linear molecule. Compared with that of C–C (336 kJ mol⁻¹), C–H (411 kJ mol⁻¹), and C–O (327 kJ mol⁻¹), an energy input of ~750 kJ mol⁻¹ is required for the cleavage of a C=O bond,¹³ which makes the conversion of CO₂ very challenging. In a typical CO₂ photoreduction process, water is required to provide a hydrogen source to achieve CO₂ transformation. The adsorption and activation of CO₂ by photocatalysts was a prerequisite for this transformation. Importantly, the redox potential alignment of the photocatalyst must be sufficient to drive the reaction. The potentials of CO₂ reduction in water are given in Table 2 (pH = 7).¹³ Reduction of CO₂ by one-electron transfer to generate CO₂^{•-} is not feasible due to the large negative reduction potential of -1.85 V (*vs.* NHE). However, relatively lower reduction potentials of -0.61, -0.53, -0.48, -0.38 and -0.24 V are required for the generation of HCOOH, CO, HCHO, CH₃OH, and CH₄, respectively. A favourable potential of -0.42 V is required for water reduction to generate H₂, which is a strong competitive reaction during the CO₂ photoreduction process. Therefore, depending on the reaction pathways taken and the number of photogenerated electrons transferred, many different products are possible for CO₂ photoreduction, making it difficult to achieve both good selectivity and high efficiency by a given photocatalyst. The selectivity (Φ) of the reacted electrons for CO₂ reduction could be calculated using the following equation:

$$\Phi(\%) = \frac{\text{Number of reacted electrons for CO}_2 \text{ reduction}}{\text{Number of reacted electrons in all reduction reactions}} \times 100\%$$

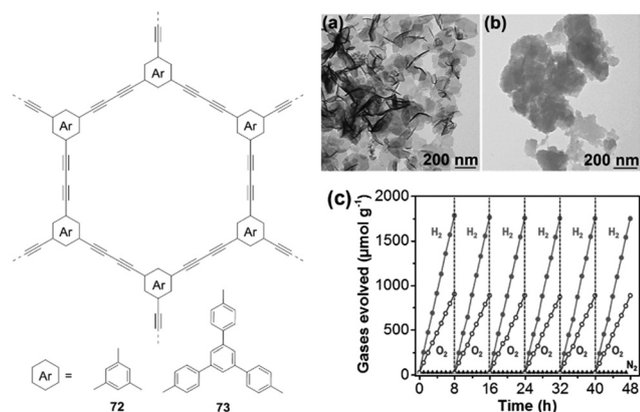


Fig. 17 Chemical structures of polymers 72 and 73; TEM pictures of (a) 72 and (b) 73 as well as (c) time course of H₂ and O₂ production of polymer 73. Reprinted with permission from ref. 40. Copyright 2017 Wiley-VCH.

Table 2 The reduction potentials *versus* NHE for CO₂ reduction in aqueous media (pH = 7). Adapted with permission from ref. 13. Copyright 2018 Wiley-VCH

Entry	Reaction	E ₀ <i>vs.</i> NHE (V)
1	CO ₂ + e ⁻ → CO ₂ ^{•-}	-1.85
2	CO ₂ + 2H ⁺ + 2e ⁻ → HCOOH	-0.61
3	CO ₂ + 2H ⁺ + 2e ⁻ → CO + H ₂ O	-0.53
4	CO ₂ + 2H ⁺ + 4e ⁻ → HCHO + H ₂ O	-0.48
5	CO ₂ + 6H ⁺ + 6e ⁻ → CH ₃ OH + 6H ₂ O	-0.38
6	CO ₂ + 8H ⁺ + 8e ⁻ → CH ₄ + 2H ₂ O	-0.24
7	2H ⁺ + 2e ⁻ → H ₂	-0.42

Heterogeneous CO₂ photoreduction over porous polymers was performed in either liquid–solid or gas–solid systems. In a liquid–solid system, the polymer particles were suspended in the CO₂-saturated aqueous solution. Under light irradiation, the CO₂ reduction reactions take place at the interface of the liquid–solid phase. Organic sacrificial agents such as triethanolamine are needed to consume undesirable holes. In contrast, CO₂ photoreduction in a gas–solid system is much simpler and only water vapour is used as the electron donor. In this case, photocatalysts were uniformly dispersed at the bottom of the reactor. Water vapour and CO₂ were chemisorbed and activated by a porous polymer followed by the reduction of CO₂ under light irradiation.

Most of the photocatalysts for CO₂ reduction are inorganic compounds such as metal oxides or sulphides, *e.g.* TiO₂, CdS, BiVO₄, *etc.*^{163,171} Organic polymers have been relatively less explored so far. For the development of highly active porous CPs for CO₂ photoreduction, some critical factors such as band gaps, surface area, CO₂ absorption, energy level alignments and photogenerated charge separation of the polymers are needed to be taken into account for pursuing good photocatalytic activities and selectivity. In addition, the external environment and operating conditions, such as catalyst quantity, solution pH, CO₂ pressure, temperature, *etc.* will also likely influence the activity and selectivity of the reactions. For the liquid–solid system, the very low CO₂ solubility in the reaction media is one major hurdle to achieve high CO₂ reduction efficiency for the photocatalyst. To address this issue to some extent, Liu *et al.* used ionic liquids to capture more CO₂ from air for the reduction reaction.¹⁷⁵ It was estimated that ~33 mg of CO₂ was captured by 1 g of ionic liquid. By introducing different functional monomers, the CBs of polymers 74–78 (Fig. 18) are varied from –0.78 and –1.52 eV, indicating sufficient driving force for the reduction of CO₂ to CO. The CO production rate follows the order: 78 > 76 > 77 > 75 > 74. The result was largely determined by the CB values of the CPs. Polymer 78 shows the highest CO production rate of 47.37 μmol h⁻¹ g⁻¹ with an excellent selectivity of up to 98.3%, which is attributed to its most negative CB value and the largest photocurrent.

g-C₃N₄ also has a favourable CB position to drive the reduction of CO₂ into hydrocarbons, which makes them potential candidates in this field.^{30,176} For example, Huang *et al.* reported that fluorine doped boron carbon nitride (BCN-F) could efficiently catalyse the reduction of CO₂ into CO in the CH₃CN/H₂O/TEOA mixture, which gave 3-times higher CO generation rate (7.75 μmol h⁻¹) than pristine BCN.¹⁷⁷ Wang *et al.* synthesized barbituric acid modified g-C₃N₄ nanosheets (CN-BAN) through copolymerization between urea and barbituric acid (BA).¹⁷⁸ With an optimal amount of BA, CN-BAN could produce CO and H₂ with a rate of ~14 μmol h⁻¹ and ~3.1 μmol h⁻¹, respectively. In addition, Guo and co-workers prepared g-C₃N₄ nanosheets with nitrogen vacancies (CN-NV) by polymerization of dicyandiamide in the presence of tartaric acid.¹⁷⁹ It was found that the introduction of defects could not only prolong the lifetimes of charge carriers, but also improve the visible light absorption and charge transfer; thereby, CN-NV exhibited

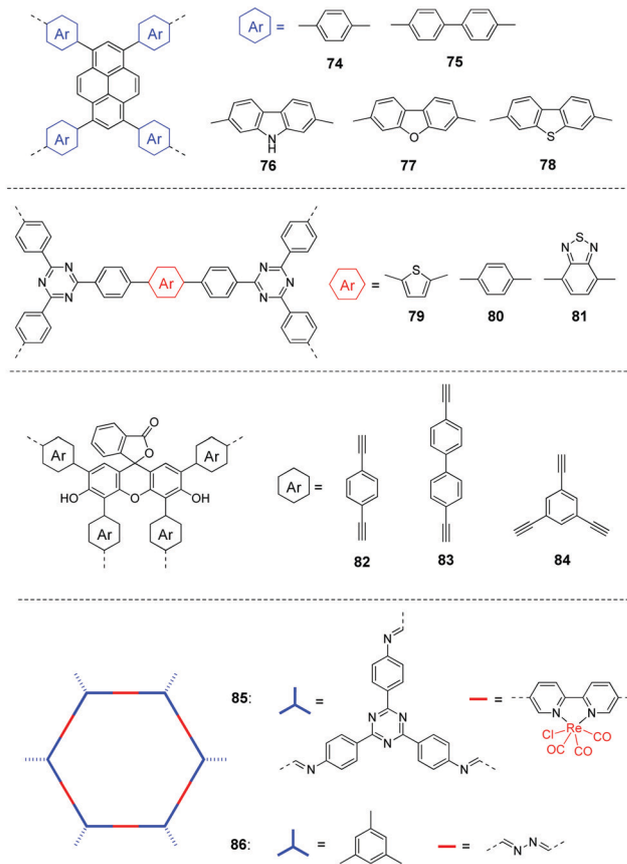


Fig. 18 Chemical structures of polymers 74–86 for photocatalytic CO₂ reduction.

a CO formation rate of 56.9 μmol h⁻¹ g⁻¹, which is about 8 times higher than that of pristine g-C₃N₄.

Apart from g-C₃N₄, CTFs with high nitrogen content are expected to have good CO₂-philic ability and thus facilitate the reduction of CO₂. Wang and co-workers developed triazine-based conjugated polymers 79–81,¹⁸⁰ which showed tunable band gaps of 2.24–2.93 eV due to the D–A interaction between the triazine unit and comonomers. The BET surface areas were 409, 52, and 37 m² g⁻¹ for 79, 80, and 81, respectively. After loading CoCl₂ and dipyriddy as cocatalysts, polymer 81 gave the highest CO generation rate of 18.2 μmol h⁻¹ with 81.6% selectivity, which is due to the enhanced charge separation by incorporating electron-withdrawing BT units into the polymer.

Considering the advantages of relatively simple and inexpensive gas–solid systems, Liu *et al.* reported polymers 82–84 by introducing different arylethynylenes.¹⁸¹ The BET surface areas were 445, 131, and 610 m² g⁻¹ for 82, 83, and 84, respectively. The highest BET surface area of polymer 84 has thus led to a CO₂ adsorption capacity of 62 mg g⁻¹ at 1.0 bar, while lower capacities of 39 and 24 mg g⁻¹ were measured for 82 and 83, respectively. In addition, the band gaps are 2.42–2.53 eV for the three polymers. The combination of electrochemical impedance spectroscopy (EIS) and transient photocurrent response measurements demonstrated that 82 had the best charge transfer

efficiency. Therefore, polymer **82** exhibited the highest CO production rate of $33 \mu\text{mol g}^{-1} \text{h}^{-1}$ with 92% selectivity (Table 3).

COFs have inherent features of crystalline porous structures, which facilitate more efficient charge separation. Thus, this kind of material is used for CO₂ photoreduction. Huang and co-workers developed COF **85**, in which the photoactive COF served as the photosensitizer and Re complex (Re(bpy)(CO)₃Cl) was incorporated as a CO₂ reduction catalyst.¹⁸² Transient absorption spectroscopy showed a lifetime of 171 ps for the excited state of **85**, which was much longer than that of the pristine COF, indicating that the incorporation of the Re moiety inhibits charge recombination in **85**. COF **85** could produce $\sim 15 \text{ mmol CO/g}$ with 98% selectivity in the TEOA/water mixture after visible light illumination for 22 h.

g-C₃N₄ could also be modified for CO₂ photoreduction in the gas phase. For instance, Yu *et al.* prepared O-doped g-C₃N₄ (CN-O) with a tube-like morphology by a two-step process, which involves thermal exfoliation of the bulk g-C₃N₄ into a nanosheet and curling-condensation of the nanosheet into a nanotube.¹⁸³ Compared with bulk g-C₃N₄, the as-prepared hierarchical nanotube exhibited a narrower band gap, higher CO₂ uptake and better charge separation. Therefore, CN-O showed a methanol production rate of $0.88 \mu\text{mol h}^{-1} \text{g}^{-1}$, which is 5 times larger than that of bulk g-C₃N₄. Wang and co-workers prepared modified g-C₃N₄ with carbon vacancies (CN-CV) by heat treatment of pre-synthesized g-C₃N₄ under an NH₃ atmosphere.¹⁸⁴ This modification improves light absorption and CO₂ adsorption/activation for CN-CV, as compared to pristine g-C₃N₄. Moreover, the charge generation and

separation of g-C₃N₄ are also enhanced. As a result, CN-CV displayed a CO production rate of $4.18 \mu\text{mol h}^{-1} \text{g}^{-1}$, nearly 3.3 times higher than that of pristine g-C₃N₄.

COFs are also efficient in a gas-solid system; azine-linked COFs **45** and **86** were reported for the photoconversion of CO₂ with gaseous H₂O into methanol under visible light.¹⁸⁵ COF **45** showed a higher BET surface area of $1412 \text{ m}^2 \text{g}^{-1}$, much higher than **86** ($1053 \text{ m}^2 \text{g}^{-1}$), which is expected to absorb more CO₂ for the reduction reaction. With a smaller optical gap of 2.57 eV than **86** (2.67 eV) and better photogenerated charge transfer, COF **45** produced $13.7 \mu\text{mol g}^{-1}$ of CH₃OH after 24 h. The amount is much higher than that of **86** ($8.6 \mu\text{mol g}^{-1}$) and C₃N₄ ($4.8 \mu\text{mol g}^{-1}$) under the same conditions.

5. Conjugated polymers for photocatalytic organic transformation

Light-initiated organic transformation is a fascinating tool for the synthesis of many important compounds under mild conditions.^{15–17} In recent years, CPs have been intensively studied for organic transformation and different types of photochemical reactions were carried out in high yields with good selectivity under visible light (Fig. 19).^{18,19,157–160} As compared to those homogeneous systems for organic transformation, heterogeneous polymer photocatalysts have distinct advantages in terms of structural robustness, non-toxicity, low cost and good recyclability. In general, a photocatalytic organic transformation is initiated by excitation of the polymer photocatalyst to generate electrons

Table 3 Summary of photocatalytic CO₂ reduction using porous CPs under visible light

Polymer	Optical gap (eV)	HOMO/LUMO	S _{BET} (m ² g ⁻¹)	Cocatalyst	Sacrificial agent	Main products and highest yield	Selectivity (%)	Ref.
74	2.48	1.70/−0.78 (eV)	27.6	—	TEOA	14.53 (CO)/5.90 (H ₂) (μmol h ⁻¹ g ⁻¹)	71.1	175 ^a
75	2.17	1.26/−0.91 (eV)	14.0	—	TEOA	27.45 (CO)/1.16 (H ₂) (μmol h ⁻¹ g ⁻¹)	95.9	175 ^a
76	2.86	1.40/−1.46 (eV)	442.8	—	TEOA	41.44 (CO)/1.20 (H ₂) (μmol h ⁻¹ g ⁻¹)	97.2	175 ^a
77	2.76	1.61/−1.15 (eV)	4.5	—	TEOA	31.76 (CO)/1.16 (H ₂) (μmol h ⁻¹ g ⁻¹)	96.5	175 ^a
78	2.66	1.14/−1.52 (eV)	23.9	—	TEOA	47.37 (CO)/0.81 (H ₂) (μmol h ⁻¹ g ⁻¹)	98.3	175 ^a
79	2.24	1.55/−0.69 (V vs. NHE)	52	CoCl ₂ and dipyritydyl	TEOA	4 (CO)/2 (H ₂) (μmol h ⁻¹)	66.7	180 ^b
80	2.93	1.66/−0.77 (V vs. NHE)	409	CoCl ₂ and dipyritydyl	TEOA	10 (CO)/3 (H ₂) (μmol h ⁻¹)	76.9	180 ^b
81	2.43	2.08/−0.85 (V vs. NHE)	37	CoCl ₂ and dipyritydyl	TEOA	18.2 (CO)/4.1 (H ₂) (μmol h ⁻¹)	81.6	180 ^b
82	2.45	1.50/−0.95 (eV)	445	—	—	33 (CO)/2.8 (H ₂) (μmol h ⁻¹)	92	181 ^c
83	2.53	1.57/−0.96 (eV)	131	—	—	~26 (CO)/~3.0 (H ₂) (μmol h ⁻¹)	90	181 ^c
84	2.42	1.48/−0.94 (eV)	610	—	—	~12 (CO)/~29 (H ₂) (μmol h ⁻¹)	29	181 ^c
85	—	—	—	—	TEOA	~900 (CO)/~18 (H ₂) (μmol h ⁻¹ g ⁻¹)	98	182 ^d
44	2.57	—	1412	—	—	37 (CH ₃ OH) (μmol h ⁻¹ g ⁻¹)	~100	185 ^e
86	2.69	—	1053	—	—	57 (CH ₃ OH) (μmol h ⁻¹ g ⁻¹)	~100	185 ^e
BCN-F	2.81	1.93/−0.88 (eV) (V vs. NHE)	—	CoCl ₂ and dipyritydyl	TEOA	7.75 (CO) (μmol h ⁻¹)	~100	177 ^f
CN-BAN	—	—	—	CoCl ₂ and dipyritydyl	TEOA	~14 (CO)/~3.1 (H ₂) μmol h ⁻¹	81.9	178 ^g
CN-NV	2.49	1.66/−0.83 (V vs. NHE)	29.5	CoCl ₂ and dipyritydyl	TEOA	56.9 (CO)/10.3 (H ₂) μmol h ⁻¹	84.7	179 ^h
CN-O	2.61	1.73/−0.88 (V vs. Ag/AgCl)	36	—	—	0.88 (CH ₃ OH) μmol h ⁻¹ g ⁻¹	100	183 ⁱ
CN-CV	2.79	1.60/−1.19 (V vs. NHE)	135.66	—	—	4.18 (CO) μmol h ⁻¹ g ⁻¹	100	184 ^j

Reaction conditions: ^a 50 mg of polymer, 10 g of CO₂-saturated ionic liquid, 1 g of TEOA, 220 W xenon lamp (> 420 nm). ^b Polymer (15 mg), CoCl₂ (1 mmol), dipyritydyl (5 mmol), TEOA (1 mL), solvent (5 mL, acetonitrile/water = 4 : 1), CO₂ (1 atm), white light (> 420 nm), 30 °C. ^c 10 mg of polymer powder, 300 W Xe lamp (> 420 nm), CO₂ and H₂O vapour. ^d 0.9 mg of polymer, 3 mL of CH₃CN, and 0.2 mL of TEOA, 225 W Xe lamp (> 420 nm), irradiated for 10 h. ^e 10 mg of polymer, 500 W Xe lamp (800 nm ≥ λ ≥ 420 nm), CO₂ (0.4 MPa) and H₂O vapour, the reactor temperature was kept at 80 °C. ^f 50 mg of catalyst, dipyritydyl (20 mg), CoCl₂·6H₂O (1 mmol), solvent (5 mL, acetonitrile : H₂O = 3 : 2), TEOA (1 mL), CO₂ (1 atm), 300 W xenon lamp (≥ 420 nm). ^g 30 mg of catalyst, 1 μmol CoCl₂, 15 mg of bipyridine, 5 mL of solvent containing 1 mL of TEOA, CO₂ (1 atm), 300 W xenon lamp (> 420 nm). ^h 5 mg of catalyst, 4 mL of CH₃CN, 1 mL of TEOA, bipyridine (10 mmol L⁻¹), as well as 25 μL of 20 mmol L⁻¹ CoCl₂ aqueous solution, 300 W xenon lamp (> 400 nm). ⁱ 50 mg of sample, CO₂ and H₂O vapour, 300 W xenon lamp (> 420 nm). ^j 30 mg of polymer, CO₂ and H₂O vapour, 300 W xenon lamp (≥ 420 nm).

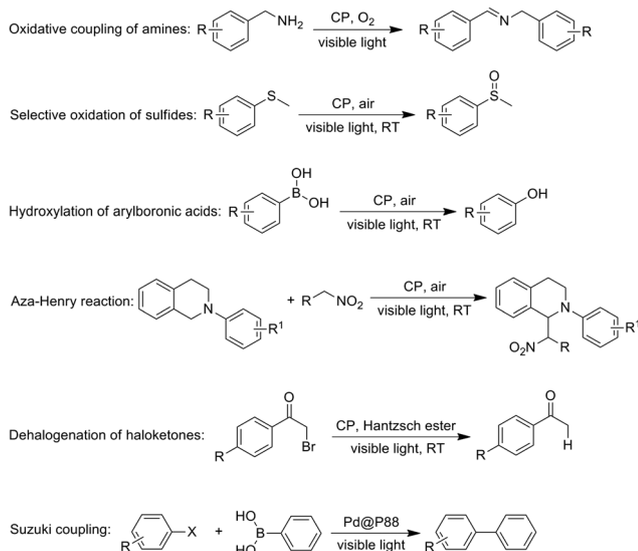


Fig. 19 Typical organic transformation reactions using CPs as the photocatalysts.

and holes, which react with organic substrates to form radical intermediates. These active intermediate species are further converted into final products through rearrangements or reactions with other species at the surface or in bulk solution.^{18–20}

5.1 Photocatalytic oxidative coupling of amine

Under light irradiation, polymer semiconductors could generate singlet oxygen ($^1\text{O}_2$), which promotes amine oxidative coupling with a broad range of substrates. In 2012, Lin *et al.* synthesized phosphorescent porous polymer **87** with incorporated $[\text{Ru}(\text{bpy})_3]^{2+}$ by oxidative homocoupling (Fig. 20).¹⁸⁶ Polymer **87** possesses $[\text{Ru}(\text{bpy})_3]^{2+}$ loadings of up to 91 wt%, which not only improves the light absorption but also favours energy migration of the triplet excited states in the polymer network to promote oxidative reactions. Under visible light and at 60 °C, stoichiometric conversions (99%) were achieved for all three substituted benzylamines. Subsequently, Zhang *et al.* developed three CMPs **88–90** by changing the heteroatoms in the benzoxalchogenadiazole unit from oxygen to sulfur to selenium.¹⁸⁷ All three polymers can catalyze the oxidative coupling of benzylamine derivatives at room temperature and polymer **89** exhibits the best activity compared to **88** and **90**, which was rationalized by the synergistic effect of the highest BET area, optimized VB and CB band levels and the highest efficiency of electron–hole pair generation. With polymer **89** as the photocatalyst, the reaction illustrates a broad substrate scope with conversions of 5–74%. In parallel, they showed that substitution positions of the benzoxadiazole unit at the centred phenyl unit has a significant impact on the CB/VB of the polymers and their photocatalytic activities. Compared to the other two 3D polymers connecting benzoxadiazole on the 1,2,4- and 1,2,4,5-positions of the phenyl unit, polymer **88** with the 1,3,5-substitution position of the centred phenyl ring demonstrated superior photocatalytic activity.¹⁸⁸ With a high specific surface area of 1137 m² g^{−1} and a broad light absorption in the visible region, the excited

polymer **91** could reduce the molecular O₂ to generate superoxide radicals ($E_{\text{red}} = -0.86$ V vs. SCE), which can oxidize the primary amine to imine products with 94–98% yields.¹⁸⁹ Directly using natural sunlight, polymer **92** exhibited good conversion ranging from 77 to 99% for oxidative homocoupling of benzylamine, which provided a cost-effective way for large-scale photochemical applications in the future.¹⁹⁰

5.2 Photocatalytic selective sulfide oxidation

In recent years, significant progress has been made by applying CPPs as photocatalysts for selective sulfide oxidation.^{191–193} In 2014, Zhang *et al.* presented the photooxidation of thioanisole using polymer **93** with high porosity.¹⁹⁴ By introducing a BT unit as the strong acceptor into the conjugated backbone, polymer **93** shows a lower band gap of 2.47 eV compared to its counterpart polymer incorporating a carbazole unit. A number of thioanisole derivatives could be photooxidized by polymer **93** with 33–98% conversion and selectivity up to 88–99% after 24 h irradiation of visible light. Upon incorporation of BODIPY dyes into the polymer network, polymer **94** exhibited a bright red emission in the solid state with efficient singlet oxygen generation, which contributed to trigger the oxidation of sulfides with excellent conversion and selectivity.¹⁹⁵ With superior surface area and good activity to reduce O₂ into the superoxide radical, polymer **91** could oxidize thioanisole to the corresponding sulfoxide with excellent selectivity (>93%) and 90–99% yields.¹⁸⁹ Notably, the reactions were completed in a short reaction time of 4.5–12 h, which was strongly influenced by the electron-donating/withdrawing substituents on various substrates.

5.3 Hydroxylation of arylboronic acid

As versatile intermediates for the synthesis of important natural products and pharmaceutical compounds, phenols could be directly synthesized by the hydroxylation of arylboronic acids in air under light illumination. Zhang *et al.* designed polymer **95** using biphenyl as the core and the polymer had a BET area of up to 2065 m² g^{−1}.¹⁹⁶ Moreover, polymer **95** exhibited a half wave potential at +1.38 V (vs. SEC), indicating its strong oxidation ability. With Et₃N as a sacrificial reductant, this polymer can catalyze the hydroxylation of arylboronic acids bearing various substituents with excellent yields (71–97%). Subsequently, Wang *et al.* reported the conversion of arylboronic acid derivatives to the corresponding phenols using benzoxazole-based COF **103** as the metal-free photocatalyst.¹⁹⁷ Prepared by reversible/irreversible cascade reactions, COF **103** exhibited super stability against light, strong acids and bases. The COF has an optical gap of 2.02 eV with a BET area of up to 1035 m² g^{−1}. Under air, polymer **103** displayed 55–99% yields for a range of substrates and the polymer could be recycled for at least 20 catalytic runs.

5.4 Aza-Henry reaction

The aerobic cross dehydrogenative coupling reaction offers a powerful method for the conversion of C–H bonds adjacent to nitrogen atoms into C–C bonds under mild conditions. This reaction plays an increasingly critical role in obtaining natural products and biologically active molecules. Over the

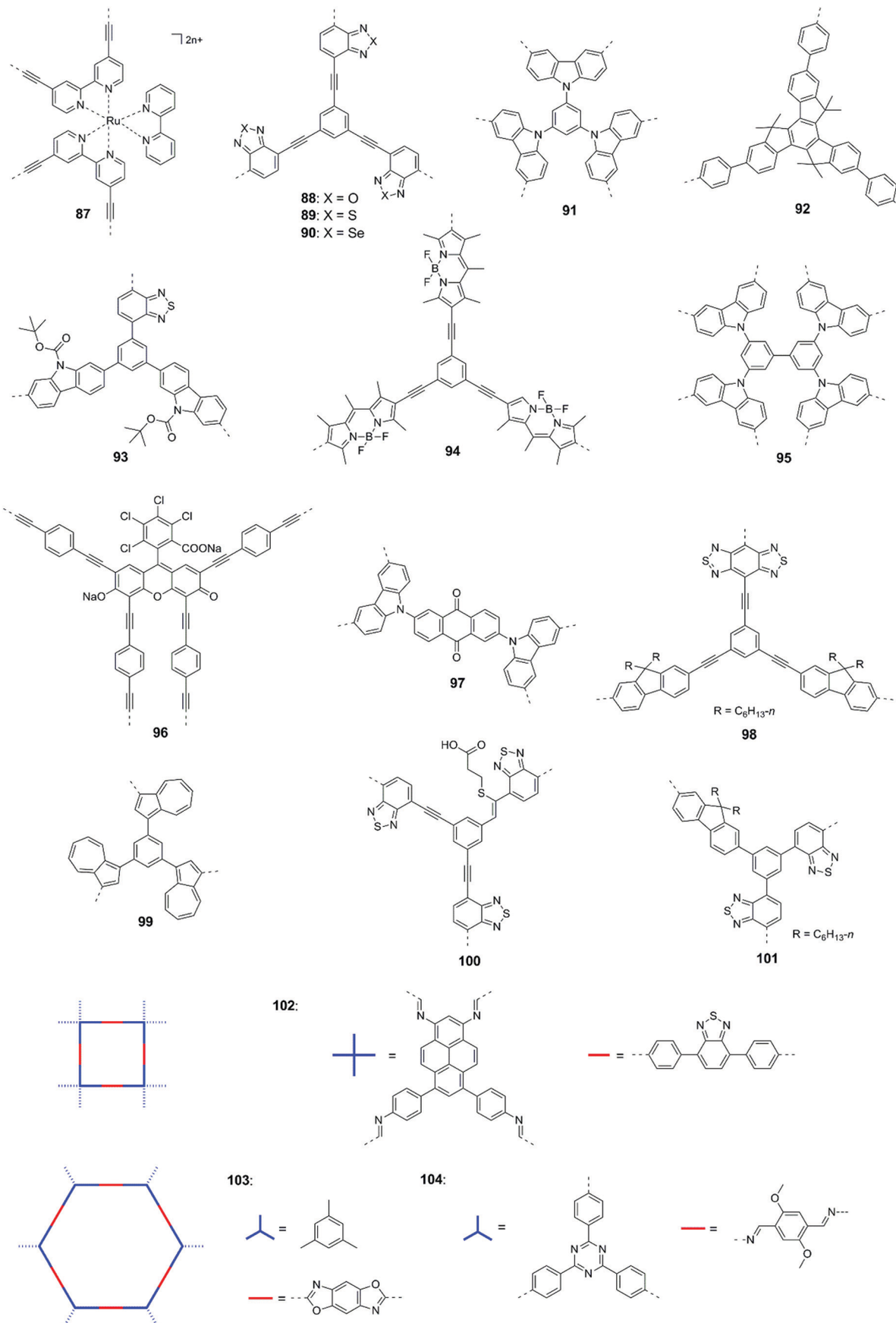


Fig. 20 Typical polymerization methods used to synthesize conjugated polymers for photocatalysis.

last decade, there has been increasing interest in developing CP photocatalysts for this reaction. As a commonly used organic dye, rose bengal was incorporated into polymer **96**, which shows a broad absorption from 350 to 700 nm and a BET surface area of $833 \text{ m}^2 \text{ g}^{-1}$.¹⁹⁸ 23 substrates were transformed into desirable products with 80–97% isolated yields using a common fluorescent lamp as the light source at room temperature. With good crystallinity and stability, 2D COF **104** exhibited excellent performance for the C–C/C–P coupling reaction with 45–92% yields and a reaction time of 6–11 h without loading any cocatalysts.¹⁹⁹ In this case, the interaction between the donor (2,5-dimethoxybenzene) and acceptor (2,4,6-triphenyl-1,3,5-triazine) in the polymer was expected to improve the photoelectric properties of the resulting COF. Based on the donor–acceptor strategy, they recently synthesized three CMPs by alternating the central core with different electron-deficient units.²⁰⁰ Compared to the CMPs with incorporated benzene (2.385 eV) and 4,7-diphenyl-2,1,3-benzothiadiazole (2.199 eV), polymer **97** exhibited the smallest optical gap of 1.994 eV. Under visible light irradiation, polymer **97** could catalyse the conversion of 1,2,3,4-tetrahydroisoquinoline derivatives to the desired products with 82–94% yields in 1 h.

5.5 Reductive dehalogenation of haloketones

Besides photo-oxidation reactions, CPs are also very efficient in catalysing some reduction reactions. Photocatalytic reductive dehalogenation of haloketones was one of the most widely investigated reactions. In 2014, Zhang *et al.* reported that polymer **98**, prepared by Sonogashira-coupling in a high internal phase emulsion, exhibited excellent activity for dehalogenation of haloketones with 15–97% yields in 4–5 h at room temperature.⁴³ The incorporation of benzobisthiadiazole (BBT) unit as a strong electron acceptor into the polymer backbone led to a much lower band gap of 1.5 eV and significantly higher electron paramagnetic resonance (EPR) signals for polymer **98** as compared to its counterpart without the BBT unit. With promising photoredox properties and high porosity, polymer **95** can catalyze the transformation of phenacyl bromide derivatives into acetophenones in 79–98% yields after 24 h light irradiation.²⁰¹ Polymer **102** exhibits good crystallinity with a strong diffraction peak at 2.67° and forms irregular block nanocrystals with a size of 60–80 nm.²⁰² With a BET area of $945 \text{ m}^2 \text{ g}^{-1}$ and a band gap of 2.08 eV, polymer **102** could catalyse many α -bromoacetophenones to the corresponding acetophenone compounds in 69–88 yields.

5.6 C–C coupling reactions

Cross-coupling reactions, such as Pd(0)-catalyzed Suzuki or Stille coupling reactions, are one of the most important methods for the construction of C–C bonds in organic synthesis.^{203,204} In general, these coupling reactions are carried out in homogeneous systems under thermal conditions. However, the Pd(0) catalysts are not stable enough and tend to undergo rapid deactivation during the reaction, which greatly affects the reaction efficiency. Therefore, the synthesis of these targets by stable and robust photocatalysts has attracted increasing attention.^{205,206}

Zhang *et al.* reported Pd decorated CMP **88** (Pd@P88) for C–C coupling reactions at ambient temperature.²⁰⁷ TEM images demonstrated that Pd NPs with sizes of 5–10 nm were distributed in the polymer network after treating the polymer with Pd(OAc)₂ at 90 °C for 1 h. The Pd content in Pd@P88 was determined to be 3% by ICP measurements. After immobilization with Pd NPs, the band gap of polymer **88** was slightly decreased from 2.43 eV to 2.38 eV. Pd@P88 has a BET surface area of $176 \text{ m}^2 \text{ g}^{-1}$ and VB/CB of +1.43/–0.95 eV (*vs.* SCE). 75–98% conversion was achieved for the coupling between aryl halides and arylboronic acid with a range of substrates. CPs could also be used for the photocatalytic Stille coupling reaction. Zhang *et al.* disclosed that an azulene-containing CMP **99** could catalyse the Stille coupling reaction between aryl iodides and aryl stannanes under visible light at room temperature.²⁰⁸ Polymer **99** exhibits a band gap of 2.03 eV with HOMO/LUMO of 1.14/–1.10 V (*vs.* SCE). Notably, only electron-withdrawing-group substituted aryl iodides could be coupled with aryl stannanes due to their lower activated energies than the LUMO of **99** (–1.10 V *vs.* SCE), while electron-donating-group substituted or unsubstituted phenyl iodides are not applicable.

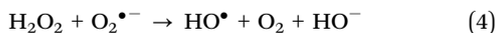
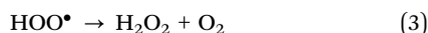
Besides those critical advances in photoredox organic transformations, several new reactions were recently exploited using CPs as the photocatalysts. For instance, polymer **89** was used for photocatalytic C–C double bond cleavage in aqueous media.²⁰⁹ Various styrene derivatives could be cleaved to afford aldehydes with 27–65% conversion and 92–99% selectivity. Yu and co-workers reported three conjugated microporous polymers for C-3 functionalization of indoles;²¹⁰ morphology and oxidation of polymers played an important role in determining the catalytic efficiency.

6. Conjugated polymers for photocatalytic degradation of organic dyes

As one of the most common contaminants in wastewater from textile industries, commercial organic dyes such as methylene blue (MB), rhodamine B (RhB), methyl orange (MO) are chemically stable and could not be removed by microbial processes.²¹¹ Other techniques such as adsorption, ozonation and flocculation have been proven to be insufficient for treatment of these effluents. Photocatalytic degradation of organic dyes into less hazardous compounds offers an attractive method for environmental remediation.^{22,23,36,39,212,213} The degradation rate of organic dyes could be facilely monitored using UV-vis absorption spectroscopy.

The degradation process of organic dyes using semiconductors is complicated and the detailed mechanism is not well revealed in the literature. A combination of electron paramagnetic resonance (ESR) analyses and radical species trapping experiments revealed that singlet oxygen (¹O₂), superoxide radical ([•]O₂[–]), photo-generated hole (h⁺), and HO[•] are the main reactive species in

the process, and the generation of these species could be proposed as follows:



The photogenerated e^- could react with adsorbed O_2 in the reaction media to generate non-selective $O_2^{\bullet-}$ radicals (eqn (1)) because of the low negative redox potential of $O_2^{\bullet-}/O_2$ (-0.13 eV vs. NHE, pH = 7).³⁶ $O_2^{\bullet-}$ could react with H^+ to give HO_2^{\bullet} (eqn (2)), which could decompose to H_2O_2 and O_2 (eqn (3)). Besides, HO^{\bullet} could be produced by the reaction of H_2O_2 with $O_2^{\bullet-}$ or h^+ , as well as its decomposition (eqn (4)–(6)). Moreover, the ROS generation from O_2 by trapping e^- also contributes to inhibit the recombination of electrons and holes. During photocatalysis, these reactive species promote the degradation of organic dyes into CO_2 , H_2O and other small molecular products.

Linear CPs with unique nanostructures show great promise for organic dye degradation. Remita *et al.* prepared nanofibers of poly(diphenylbutadiyne) (**105**) by photopolymerization using a soft templating approach (Fig. 21).²¹⁴ The nanofibers are a few micrometres long with a diameter of about 19 nm. Without additional sacrificial reagents or cocatalysts, 75% MO were decomposed in water after 240 min irradiation, whereas only 17% MO degradation was achieved by Ag-modified TiO_2 . The photocatalytic performance of the nanofibers was almost unchanged even after 15 repeated cycles. Using the same method, they further prepared polymer **106** with vesicle and spindle morphologies, respectively.²¹⁵ The spindles are 40 nm thick and the length is several hundred nanometres, while the vesicles are spherical hollow capsules with a diameter of around 1 μm and the thickness of the wall is around 40 nm. A clear shape-dependent photocatalytic activity was demonstrated as shown in Fig. 22a. The nanospindles displayed 100% degradation of MO after 180 min visible light irradiation; by contrast, the vesicles remained totally inactive under identical conditions. Besides, the photocatalytic activity of the polymer **107** nanospindles is much better than that of the polymer **106** nanofibers.

With good photoactivity and excellent water dispersibility, conjugated polyelectrolytes (CPES) were reported by Zhang and co-workers for organic dye degradation.²¹⁶ They prepared polymer **108** by attaching 1-alkyl-3-vinylimidazolium bromide onto the side chains of the poly(benzothiadiazolyfluorene) backbone, which was conducted through a self-initiated radical polymerization process under visible light. With a band gap of 2.11 eV, polymer **108** could almost quantitatively decompose MB and RhB after 70 and 90 min, respectively (Fig. 22b). The good singlet oxygen generation ability of polymer **107** under light excitation is likely responsible for the organic dye photodegradation. In addition, RhB was degraded almost quantitatively in

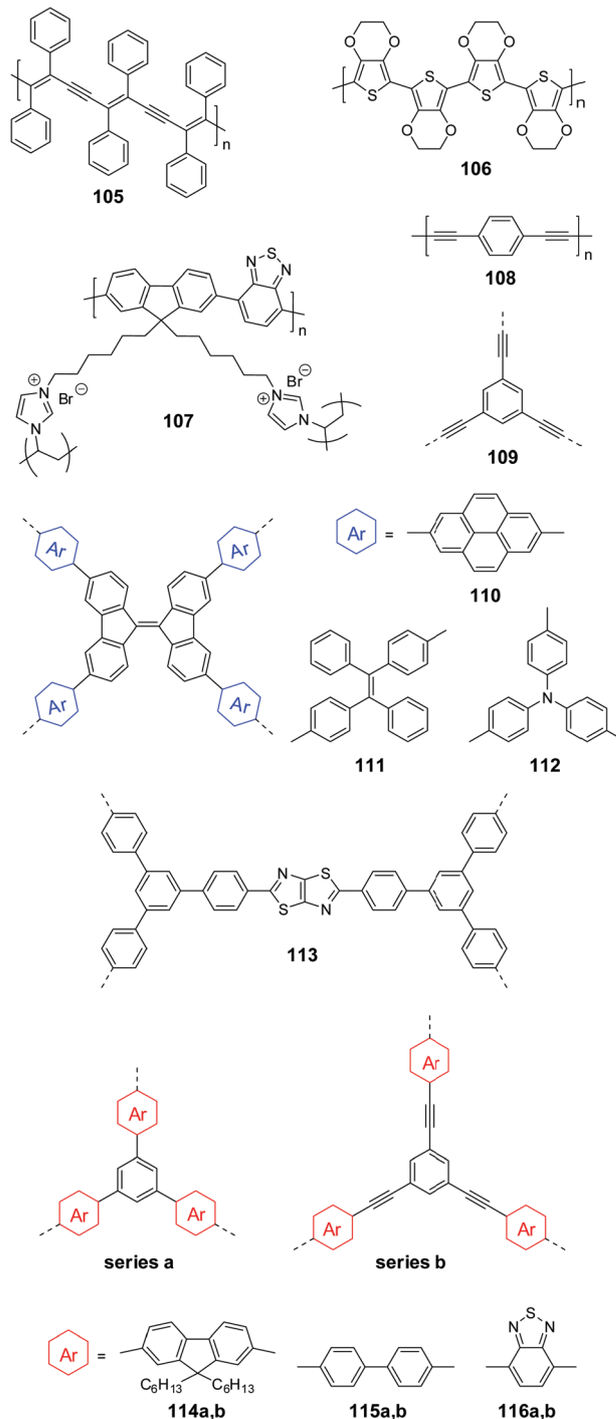


Fig. 21 Representative conjugated polymers for photocatalytic degradation of organic dyes.

repeated 10 runs, demonstrating excellent stability and reusability of **108**.

Recently, Ouyang and co-workers synthesized three polymers with similar backbone structures and studied the effect of structural variation on their photocatalytic activities.²¹⁷ Among them, polymer **109** had a 3D network structure, while the linear polymers **108a** and **108b** were obtained by varying the concentrations of the feeding monomers.

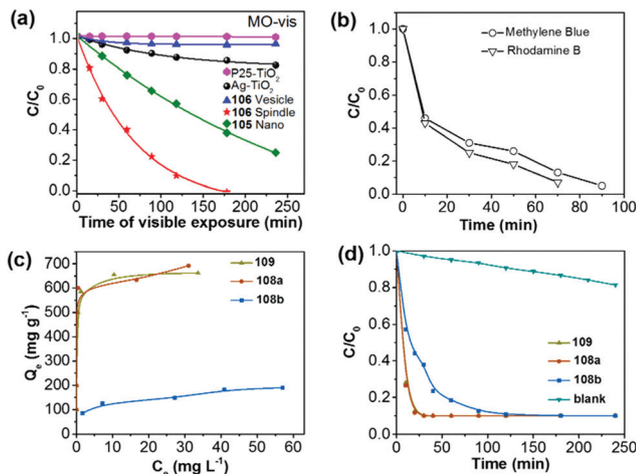


Fig. 22 (a) Comparative photocatalytic degradation of MO of commercial TiO₂, Ag-TiO₂, polymer nanofibers of **105**, vesicles and nanospindles of polymer **106** under visible light (P25-TiO₂ is TiO₂ with a diameter of 25 nm). (b) Photocatalytic degradation of RhB and MB using polymer **107**. (c) The adsorption isotherms of RhB for polymers **108a**, **108b** and **109** at 25 °C. (d) Removal of RhB with polymers **108a**, **108b** and **109** in the supernatant. Reprinted with permission from ref. 215–217. Copyright 2015 Nature Publishing Group, 2015 Wiley-VCH and 2015 Royal Society of Chemistry.

The BET areas were 1204, 571, 1265 m² g⁻¹ for **109a**, **109b** and **109**, respectively. As shown in Fig. 22c, with a high surface area, the adsorption capacities of **110** (661 mg g⁻¹) and **108a** (688 mg g⁻¹) toward RhB were much higher than that of **108b** (189 mg g⁻¹). Polymers **108a** and **108b** exhibited a very close band gap (1.95 eV for **108a** and 1.97 eV for **108b**), which was narrower than that of **109** (2.28 eV). By the synergistic adsorption-photocatalysis, RhB was totally decomposed in 30 min for both **109** and **108a**, while 120 min was needed for **108b** to achieve the same performance (Fig. 22d). These results implied that the BET surface area of the polymer played an important role in photocatalytic activity.

The adsorption-photocatalysis synergy was also reported by Chen *et al.*²¹⁸ They prepared 9,9'-bifluorenylidene-based porous polymers **110–112** by introducing different comonomers such as pyrene, tetraphenylethene and triphenyl. Polymer **110** with incorporated pyrene units exhibited the lowest band gap of 1.55 eV and the largest BET surface area of 1306 m² g⁻¹, while lower surface areas of 777 and 590 m² g⁻¹ were determined for **111** and **112**, respectively. All three polymers demonstrated fast uptake of RhB in an aqueous solution. After 45 min, the absorbed percentages of RhB were 97%, 96% and 89% for **110**, **111** and **112**, respectively. As shown in Fig. 23a, polymer **110** exhibited a remarkably high adsorption capacity of 1905 mg g⁻¹ of RhB, which is much higher than those of **111** (1024 mg g⁻¹) and **112** (926 mg g⁻¹). As shown in Fig. 23b, more than 81% of RhB was decomposed in the presence of polymer **110** after 30 min visible light irradiation (>450 nm).

Very recently, Jin and co-workers prepared three CMPs with thiazolo[5,4-*d*]thiazole as the linker for photodegradation of organic dyes.²¹⁹ Interestingly, the polymers showed nanosheet morphologies and their porous and photophysical properties

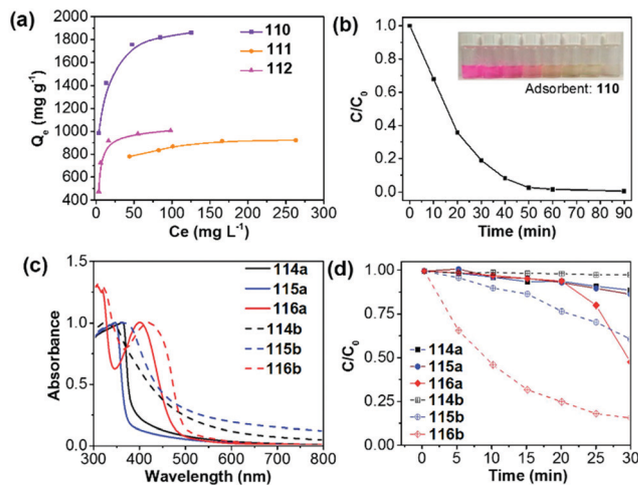


Fig. 23 (a) The adsorption isotherms for RhB on polymers **110–112**. (b) Photocatalytic degradation kinetics of RhB using polymer **110**. (c) UV-vis absorption spectra of NPs of polymers **114–116** in a water dispersion. (d) Photocatalytic degradation of RhB in the presence of NPs of polymers **114–116** in water. Reprinted with permission from ref. 218–220. Copyright 2018 American Chemical Society, 2019 American Chemical Society and 2015 Royal Society of Chemistry.

could be fine-tuned by introducing diphenyl, triphenylbenzene and tetraphenylbenzene. With the highest surface area of 673 m² g⁻¹ among the three polymers, polymer **113** could absorb 57% of RhB, while the amount of RhB absorbed by the other two polymers is lower than 15%. Polymer **113** can completely degrade RhB in 30 min and 60 min is needed for the degradation of 80% MO.

Conjugated microporous polymer (CMP) nanoparticles offer unique advantages of high porosity and excellent solution dispersibility compared to bulk catalysts, which is desirable for organic dye photodegradation. Inspired by this, Zhang *et al.* reported the direct preparation of CMP nanoparticles *via* palladium-catalysed polycondensation reactions in a miniemulsion.²²⁰ Interestingly, the difference in the comonomers and linkages has a clear impact on the morphologies and optical properties of the resulting polymers. CMP nanoparticles obtained *via* Sonogashira-coupling are larger than the ones obtained *via* Suzuki-coupling. For BT-containing polymers, polymer **116a** showed a rod-like shape, while polymer **116b** formed nanorings in the reaction media. As shown in Fig. 23c, the introduction of BT into the backbone has led to a narrower band gap of the resulting polymers. Over 80% of RhB was degraded in the presence of **116b** after 25 min, while only about 50% of RhB was decomposed for **116a** (band gap: 1.96 eV) after the same period (Fig. 23d). This result is in accordance with the lowest optical band gap of **116b** (1.76 eV).

7. Summary and outlook

In this Review, we have summarized the recent advances in CPs for various photocatalytic applications, including water splitting, CO₂ reduction, organic transformation and organic dye degradation. Since g-C₃N₄ was first demonstrated for water

splitting under visible light in 2009,¹⁴⁰ CPs have drawn intensive interest and represent a new platform in the field of photocatalysis on account of their inherent advantages of earth-abundant nature, tunable band structure and high stability in the photocatalytic process. We discussed all classes of CPs including *g*-C₃N₄, linear CPs, porous CPs, water-soluble conjugated polyelectrolytes (CPEs), and organic framework materials (COFs and CTFs) in this review article and presented how polymer design could contribute to different photocatalytic reactions. Structurally, linear CPs consist of alternative conjugated units, while CPPs have 3D porous structures, which are expected to facilitate charge transfer in multiple dimensions. Compared to CPPs and linear CPs, crystalline COFs possess a stacked 2D structure, which is beneficial for photogenerated charge transfer, thus boosting the photocatalytic activity. On the other hand, to date, COFs used for heterogeneous photocatalysis still suffer from limited structural variations. Moreover, a Pt cocatalyst is commonly needed for COFs in photocatalytic H₂ production while CPs prepared by Pd-catalyzed coupling reactions could be directly used without adding any cocatalysts. Besides those noble metal cocatalysts (Pt and Pd), the development of earth-abundant and low-cost cocatalysts has recently attracted attention. For instance, nickel-thiolate cluster was found to be a very efficient H₂ production cocatalyst on a COF photoabsorber.²²¹ This system demonstrated long-term hydrogen evolution activity with a turnover number of up to 103 and the maximum HER reached 941 μmol h⁻¹ g⁻¹ in the TEOA/H₂O mixture. To improve the water-wettability of the polymers, a family of polymer photocatalysts enriched with CPEs with conjugated backbones and charged side-chains have also been developed, which offered new potential to this field.

The molecular design has undoubtedly played a vital role in achieving the preferred properties of the resulting polymers, such as strong and wide visible light absorption, appropriate band structure, high efficiency of charge separation and transport, as well as good wettability and surface reaction efficiency. These synergy improvements in principle could result in remarkably enhanced catalytic activity. Among the reported strategies, elemental and molecular doping and the introduction of structural defects into the skeleton of *g*-C₃N₄ have been well developed for achieving enhanced photocatalytic water splitting and CO₂ reduction. In addition, copolymerization of well-designed donors and acceptors has been widely employed to design CPs for photocatalysis, which have brought extended light absorption and improved charge mobility of polymers. The statistical copolymerization for the synthesis of CPPs offered additional opportunity to adjust the band gaps and porosity of polymers.

Despite the great progress made so far, the investigations in this field are in its infancy. The reported CPs suffer from low photocatalytic efficiency, which are far from the practical requirements. Many issues need to be resolved which may include the following:

(1) The development of photocatalytic H₂ production using CPs has evolved rapidly in the past few years. A recorded H₂ generation rate of 60.6 mmol h⁻¹ g⁻¹ has been achieved,

which is very competitive to inorganic counterparts. It is highly expected that more exciting performance could be achieved in the future by designing new polymers, and the combination of computation and experiment appears to be a promising approach for high-throughput screening of polymer libraries. In a very recent report, Cooper and co-workers computationally studied photocatalytic H₂ production using 6354 potential alternating co-polymers, among which more than 170 co-polymers have been synthesized and examined for their H₂ evolution activities.²²² This work well explored the structure–property–performance relationships of candidate co-polymers and successfully discovered new polymers with HERs greater than 6 mmol h⁻¹ g⁻¹. Apart from photocatalytic water reduction or oxidation, photocatalytic overall water splitting has been reported by conjugated microporous polymer nanosheets.⁴⁰ However, the fundamentals of the photocatalytic process and structure–activity relationship of the polymer are not well understood. Besides, loading metals as cocatalyst could considerably increase the photocatalytic H₂ production activity, but knowledge about the interaction mechanism between the polymer and metal is still not clear.

(2) Until now, relatively few CPs were exploited for photocatalytic CO₂ reduction and most of them suffer from low activities. Likewise, knowledge about reaction pathways and fundamental enhancements for CO₂ reduction is scarce. In order to increase the catalytic performance of the reaction, more work is in urgent demand to optimize the photoactivity and porosity of polymers.

(3) The use of CPs as heterogeneous photocatalysts is rapidly expanding the toolbox of chemical transformations. They have demonstrated good catalytic activities and recyclability for a variety of photochemical reactions. In particular, some reactions were performed directly under natural sunlight. However, the applied reaction types and substrate scope are still limited as compared to those of traditional transition-metal complexes. It is highly desirable to explore new organic molecule transformations, such as more challenging C-heteroatom or C–C functionalization, for heterogeneous polymer photocatalysts.

(4) Photodegradation of organic dyes using CPs presented an effective technology for environmental remediation. Interestingly, nanostructured CPs with good water dispersion have demonstrated excellent degradation ability. Another important finding is that CPPs could decompose organic dyes very efficiently by the adsorption–photocatalysis synergy. In order to achieve better catalytic activity, advanced research should be devoted to optimize the photophysical properties of conjugated polymers to improve the reactive species generation efficiency.

As CPs have the inherent advantages of diverse molecular structures and readily tunable physicochemical properties, there are unlimited opportunities and challenges in developing high-efficiency CPs for various photocatalytic applications. We expect that this review will be a good reference for the next decade of research in this exciting field. With more advances in polymer chemistry and materials science, it is anticipated that critical innovations in photocatalytic technology using CPs could play an increasingly important role in practical applications.

Conflicts of interest

There are no conflicts of interest to declare.

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References

- J. Barber, *Chem. Soc. Rev.*, 2009, **38**, 185–196.
- A. J. Carrillo, J. González-Aguilar, M. Romero and J. M. Coronado, *Chem. Rev.*, 2019, **119**, 4777–4816.
- X. Chen, S. Shen, L. Guo and S. S. Mao, *Chem. Rev.*, 2010, **110**, 6503–6570.
- T. Hisatomi, J. Kubota and K. Domen, *Chem. Soc. Rev.*, 2014, **43**, 7520–7535.
- T. Wang and J. Gong, *Angew. Chem., Int. Ed.*, 2015, **54**, 10718–10732.
- J. Willkomm, K. L. Orchard, A. Reynal, E. Pastor, J. R. Durrant and E. Reisner, *Chem. Soc. Rev.*, 2016, **45**, 9–23.
- Y. Wang, H. Suzuki, J. Xie, O. Tomita, D. J. Martin, M. Higashi, D. Kong, R. Abe and J. Tang, *Chem. Rev.*, 2018, **118**, 5201–5241.
- Z. Wang, C. Li and K. Domen, *Chem. Soc. Rev.*, 2019, **48**, 2109–2125.
- T. Hisatomi and K. Domen, *Nat. Catal.*, 2019, **2**, 387–399.
- S. N. Habisreutinger, L. Schmidt-Mende and J. K. Stolarczyk, *Angew. Chem., Int. Ed.*, 2013, **52**, 7372–7408.
- X. Liu, S. Inagaki and J. Gong, *Angew. Chem., Int. Ed.*, 2016, **55**, 14924–14950.
- R. Shi, G. I. N. Waterhouse and T. Zhang, *Sol. RRL*, 2017, **1**, 1700126.
- Z. Sun, N. Talreja, H. Tao, J. Texter, M. Muhler, J. Strunk and J. Chen, *Angew. Chem., Int. Ed.*, 2018, **57**, 7610–7627.
- C. Wang, Z. Sun, Y. Zheng and Y. H. Hu, *J. Mater. Chem. A*, 2019, **7**, 865–887.
- L. Shi and W. Xia, *Chem. Soc. Rev.*, 2012, **41**, 7687–7697.
- J. W. Beatty and C. R. Stephenson, *Acc. Chem. Res.*, 2015, **48**, 1474–1484.
- I. Ghosh, L. Marzo, A. Das, R. Shaikh and B. König, *Acc. Chem. Res.*, 2016, **49**, 1566–1577.
- A. Savateev, I. Ghosh, B. König and M. Antonietti, *Angew. Chem., Int. Ed.*, 2018, **57**, 15936–15947.
- R. Li, J. Byun, W. Huang, C. Ayed, L. Wang and K. A. I. Zhang, *ACS Catal.*, 2018, **8**, 4735–4750.
- A. Savateev and M. Antonietti, *ACS Catal.*, 2018, **8**, 9790–9808.
- U. G. Akpan and B. H. Hameed, *J. Hazard. Mater.*, 2009, **170**, 520–529.
- L. G. Devi and R. Kavitha, *Appl. Catal., B*, 2013, **140–141**, 559–587.
- C.-C. Wang, J.-R. Li, X.-L. Lv, Y.-Q. Zhang and G. Guo, *Energy Environ. Sci.*, 2014, **7**, 2831–2867.
- L. Nie and Q. Zhang, *Inorg. Chem. Front.*, 2017, **4**, 1953–1962.
- L. Bucher, N. Desbois, P. D. Harvey, G. D. Sharma and C. P. Gros, *Sol. RRL*, 2017, **1**, 1700127.
- S. Y. Tee, K. Y. Win, W. S. Teo, L.-D. Koh, S. Liu, C. P. Teng and M.-Y. Han, *Adv. Sci.*, 2017, **4**, 1600337.
- X. Wang, S. Blechert and M. Antonietti, *ACS Catal.*, 2012, **2**, 1596–1606.
- Y. Zheng, L. Lin, B. Wang and X. Wang, *Angew. Chem., Int. Ed.*, 2015, **54**, 12868–12884.
- W.-J. Ong, L.-L. Tan, Y. H. Ng, S.-T. Yong and S.-P. Chai, *Chem. Rev.*, 2016, **116**, 7159–7329.
- J. Fu, J. Yu, C. Jiang and B. Cheng, *Adv. Energy Mater.*, 2018, **8**, 1701503.
- Y. Zheng, J. Liu, J. Liang, M. Jaroniec and S. Z. Qiao, *Energy Environ. Sci.*, 2012, **5**, 6717–6731.
- J. Zhang, Y. Chen and X. Wang, *Energy Environ. Sci.*, 2015, **8**, 3092–3108.
- V. S. Vyas, V. W.-h. Lau and B. V. Lotsch, *Chem. Mater.*, 2016, **28**, 5191–5204.
- G. Zhang, Z. A. Lan and X. Wang, *Angew. Chem., Int. Ed.*, 2016, **55**, 15712–15727.
- L. Lin, Z. Yu and X. Wang, *Angew. Chem., Int. Ed.*, 2019, **58**, 6164–6175.
- C. Chen, W. Ma and J. Zhao, *Chem. Soc. Rev.*, 2010, **39**, 4206–4219.
- M. Marszewski, S. Cao, J. Yu and M. Jaroniec, *Mater. Horiz.*, 2015, **2**, 261–278.
- J. L. White, M. F. Baruch, J. E. Pander III, Y. Hu, I. C. Fortmeyer, J. E. Park, T. Zhang, K. Liao, J. Gu, Y. Yan, T. W. Shaw, E. Abelev and A. B. Bocarsly, *Chem. Rev.*, 2015, **115**, 12888–12935.
- L. V. Bora and R. K. Mewada, *Renewable Sustainable Energy Rev.*, 2017, **76**, 1393–1421.
- L. Wang, Y. Wan, Y. Ding, S. Wu, Y. Zhang, X. Zhang, G. Zhang, Y. Xiong, X. Wu, J. Yang and H. Xu, *Adv. Mater.*, 2017, **29**, 1702428.
- L. Wang, Y. Wan, Y. Ding, Y. Niu, Y. Xiong, X. Wu and H. Xu, *Nanoscale*, 2017, **9**, 4090–4096.
- L. Wang, X. Zheng, L. Chen, Y. Xiong and H. Xu, *Angew. Chem., Int. Ed.*, 2018, **57**, 3454–3458.
- Z. J. Wang, S. Ghasimi, K. Landfester and K. A. I. Zhang, *J. Mater. Chem. A*, 2014, **2**, 18720–18724.
- Z. J. Wang, S. Ghasimi, K. Landfester and K. A. I. Zhang, *Chem. Commun.*, 2014, **50**, 8177–8180.
- B. C. Ma, S. Ghasimi, K. Landfester, F. Vilela and K. A. I. Zhang, *J. Mater. Chem. A*, 2015, **3**, 16064–16071.
- Y.-J. Cheng, S.-H. Yang and C.-S. Hsu, *Chem. Rev.*, 2009, **109**, 5868–5923.
- X. Feng, X. Ding and D. Jiang, *Chem. Soc. Rev.*, 2012, **41**, 6010–6022.
- C. Zhu, L. Liu, Q. Yang, F. Lv and S. Wang, *Chem. Rev.*, 2012, **112**, 4687–4735.
- S.-Y. Ding and W. Wang, *Chem. Soc. Rev.*, 2013, **42**, 548–568.
- Y. Xu, S. Jin, H. Xu, A. Nagai and D. Jiang, *Chem. Soc. Rev.*, 2013, **42**, 8012–8031.
- J. L. Segura, M. J. Mancheno and F. Zamora, *Chem. Soc. Rev.*, 2016, **45**, 5635–5671.

- 52 J.-R. Pouliot, F. Grenier, J. T. Blaskovits, S. Beaupré and M. Leclerc, *Chem. Rev.*, 2016, **116**, 14225–14274.
- 53 M. Liu, L. Guo, S. Jin and B. Tan, *J. Mater. Chem. A*, 2019, **7**, 5153–5172.
- 54 M. Leclerc and J.-F. Morin, *Design and synthesis of conjugated polymers*, John Wiley & Sons, 2010.
- 55 Y. Chujo, *Conjugated polymer synthesis: methods and reactions*, John Wiley & Sons, 2011.
- 56 M. Karayilan, W. P. Brezinski, K. E. Clary, D. L. Lichtenberger, R. S. Glass and J. Pyun, *Angew. Chem., Int. Ed.*, 2019, **58**, 7537–7550.
- 57 Y. Ma, X. Wang, Y. Jia, X. Chen, H. Han and C. Li, *Chem. Rev.*, 2014, **114**, 9987–10043.
- 58 S. Cao and J. Yu, *J. Phys. Chem. Lett.*, 2014, **5**, 2101–2107.
- 59 L.-Z. Wu, B. Chen, Z.-J. Li and C.-H. Tung, *Acc. Chem. Res.*, 2014, **47**, 2177–2185.
- 60 S. Kauffhold, L. Petermann, R. Staehle and S. Rau, *Coord. Chem. Rev.*, 2015, **304**, 73–87.
- 61 X. Zhang, T. Peng and S. Song, *J. Mater. Chem. A*, 2016, **4**, 2365–2402.
- 62 M. Wen, K. Mori, Y. Kuwahara, T. An and H. Yamashita, *Appl. Catal., B*, 2017, **218**, 555–569.
- 63 Y. Huang and B. Zhang, *Angew. Chem., Int. Ed.*, 2017, **56**, 14804–14806.
- 64 Y.-J. Yuan, D. Chen, Z.-T. Yu and Z.-G. Zou, *J. Mater. Chem. A*, 2018, **6**, 11606–11630.
- 65 M. G. Schwab, M. Hamburger, X. Feng, J. Shu, H. W. Spiess, X. Wang, M. Antonietti and K. Müllen, *Chem. Commun.*, 2010, **46**, 8932–8934.
- 66 S. Yanagida, A. Kabumoto, K. Mizumoto, C. Pac and K. Yoshino, *J. Chem. Soc., Chem. Commun.*, 1985, **8**, 474–475.
- 67 T. Shibata, A. Kabumoto, T. Shiragami, O. Ishitani, C. Pac and S. Yanagida, *J. Phys. Chem.*, 1990, **94**, 2068–2076.
- 68 S. Matsuoka, T. Kohzuki, A. Nakamura, C. Pac and S. Yanagida, *J. Chem. Soc., Chem. Commun.*, 1991, **8**, 580–581.
- 69 S. Matsuoka, H. Fujii, T. Yamada, C. Pac, A. Ishida, S. Takamuku, M. Kusaba, N. Nakashima and S. Yanagida, *J. Phys. Chem.*, 1991, **95**, 5802–5808.
- 70 R. S. Sprick, B. Bonillo, R. Clowes, P. Guiglion, N. J. Brownbill, B. J. Slater, F. Blanc, M. A. Zwiijnenburg, D. J. Adams and A. I. Cooper, *Angew. Chem., Int. Ed.*, 2018, **57**, 2520.
- 71 X.-H. Zhang, X.-P. Wang, J. Xiao, S.-Y. Wang, D.-K. Huang, X. Ding, Y.-G. Xiang and H. Chen, *J. Catal.*, 2017, **350**, 64–71.
- 72 M. Sachs, R. S. Sprick, D. Pearce, S. J. Hillman, A. Monti, A. A. Y. Guilbert, N. J. Brownbill, S. Dimitrov, X. Shi, F. Blanc, M. A. Zwiijnenburg, J. Nelson, J. R. Durrant and A. I. Cooper, *Nat. Commun.*, 2018, **9**, 4968.
- 73 C. M. Aitchison, R. S. Sprick and A. I. Cooper, *J. Mater. Chem. A*, 2019, **7**, 2490–2496.
- 74 C. Dai, S. Xu, W. Liu, X. Gong, M. Panahandeh-Fard, Z. Liu, D. Zhang, C. Xue, K. P. Loh and B. Liu, *Small*, 2018, **14**, 1801839.
- 75 Z.-A. Lan, G. Zhang, X. Chen, Y. Zhang, K. A. I. Zhang and X. Wang, *Angew. Chem., Int. Ed.*, 2019, **58**, 10236–10240.
- 76 R. S. Sprick, C. M. Aitchison, E. Berardo, L. Turcani, L. Wilbraham, B. M. Alston, K. E. Jelfs, M. A. Zwiijnenburg and A. I. Cooper, *J. Mater. Chem. A*, 2018, **6**, 11994–12003.
- 77 R. S. Sprick, L. Wilbraham, Y. Bai, P. Guiglion, A. Monti, R. Clowes, A. I. Cooper and M. A. Zwiijnenburg, *Chem. Mater.*, 2018, **30**, 5733–5742.
- 78 J. Pecher and S. Mecking, *Chem. Rev.*, 2010, **110**, 6260–6279.
- 79 K. Li and B. Liu, *J. Mater. Chem.*, 2012, **22**, 1257–1264.
- 80 L. Feng, C. Zhu, H. Yuan, L. Liu, F. Lv and S. Wang, *Chem. Soc. Rev.*, 2013, **42**, 6620–6633.
- 81 S. Kundu and A. Patra, *Chem. Rev.*, 2016, **117**, 712–757.
- 82 L. Wang, R. Fernández-Terán, L. Zhang, D. L. Fernandes, L. Tian, H. Chen and H. Tian, *Angew. Chem., Int. Ed.*, 2016, **55**, 12306–12310.
- 83 P. B. Pati, G. Damas, L. Tian, D. L. Fernandes, L. Zhang, I. B. Pehlivan, T. Edvinsson, C. M. Araujo and H. Tian, *Energy Environ. Sci.*, 2017, **10**, 1372–1376.
- 84 P.-J. Tseng, C.-L. Chang, Y.-H. Chan, L.-Y. Ting, P.-Y. Chen, C.-H. Liao, M.-L. Tsai and H.-H. Chou, *ACS Catal.*, 2018, **8**, 7766–7772.
- 85 C. V. Hoven, A. Garcia, G. C. Bazan and T. Q. Nguyen, *Adv. Mater.*, 2008, **20**, 3793–3810.
- 86 H. Jiang, P. Taranekar, J. R. Reynolds and K. S. Schanze, *Angew. Chem., Int. Ed.*, 2009, **48**, 4300–4316.
- 87 A. Duarte, K.-Y. Pu, B. Liu and G. C. Bazan, *Chem. Mater.*, 2010, **23**, 501–515.
- 88 Q. Cui and G. C. Bazan, *Acc. Chem. Res.*, 2017, **51**, 202–211.
- 89 C. Dai, M. Panahandeh-Fard, X. Gong, C. Xue and B. Liu, *Sol. RRL*, 2019, **3**, 1800255.
- 90 Z. Hu, Z. Wang, X. Zhang, H. Tang, X. Liu, F. Huang and Y. Cao, *iScience*, 2019, **13**, 33–42.
- 91 D. J. Woods, R. S. Sprick, C. L. Smith, A. J. Cowan and A. I. Cooper, *Adv. Energy Mater.*, 2017, **7**, 1700479.
- 92 A. I. Cooper, *Adv. Mater.*, 2009, **21**, 1291–1295.
- 93 R. Dawson, A. I. Cooper and D. J. Adams, *Prog. Polym. Sci.*, 2012, **37**, 530–563.
- 94 J. Zhu, C. Yang, C. Lu, F. Zhang, Z. Yuan and X. Zhuang, *Acc. Chem. Res.*, 2018, **51**, 3191–3202.
- 95 F. Vilela, K. Zhang and M. Antonietti, *Energy Environ. Sci.*, 2012, **5**, 7819–7832.
- 96 R. S. Sprick, J.-X. Jiang, B. Bonillo, S. Ren, T. Ratvijitvech, P. Guiglion, M. A. Zwiijnenburg, D. J. Adams and A. I. Cooper, *J. Am. Chem. Soc.*, 2015, **137**, 3265–3270.
- 97 Y. Xu, N. Mao, C. Zhang, X. Wang, J. Zeng, Y. Chen, F. Wang and J.-X. Jiang, *Appl. Catal., B*, 2018, **228**, 1–9.
- 98 Y. Liu, Z. Liao, X. Ma and Z. Xiang, *ACS Appl. Mater. Interfaces*, 2018, **10**, 30698–30705.
- 99 Z.-A. Lan, W. Ren, X. Chen, Y. Zhang and X. Wang, *Appl. Catal., B*, 2019, **245**, 596–603.
- 100 Y. Li, *Acc. Chem. Res.*, 2012, **45**, 723–733.
- 101 H. Zhou, L. Yang and W. You, *Macromolecules*, 2012, **45**, 607–632.
- 102 C. Gao, L. Wang, X. Li and H. Wang, *Polym. Chem.*, 2014, **5**, 5200–5210.
- 103 H. Yao, L. Ye, H. Zhang, S. Li, S. Zhang and J. Hou, *Chem. Rev.*, 2016, **116**, 7397–7457.

- 104 C. Liu, K. Wang, X. Gong and A. J. Heeger, *Chem. Soc. Rev.*, 2016, **45**, 4825–4846.
- 105 L. Li, Z. Cai, Q. Wu, W.-Y. Lo, N. Zhang, L. X. Chen and L. Yu, *J. Am. Chem. Soc.*, 2016, **138**, 7681–7686.
- 106 L. Li, W.-y. Lo, Z. Cai, N. Zhang and L. Yu, *Macromolecules*, 2016, **49**, 6903–6909.
- 107 C. Yang, B. C. Ma, L. Zhang, S. Lin, S. Ghasimi, K. Landfester, K. A. I. Zhang and X. Wang, *Angew. Chem., Int. Ed.*, 2016, **55**, 9202–9206.
- 108 Q. Zhang, M. A. Kelly, N. Bauer and W. You, *Acc. Chem. Res.*, 2017, **50**, 2401–2409.
- 109 W. Zhong, J. Liang, S. Hu, X.-F. Jiang, L. Ying, F. Huang, W. Yang and Y. Cao, *Macromolecules*, 2016, **49**, 5806–5816.
- 110 Y. Xiang, X. Wang, L. Rao, P. Wang, D. Huang, X. Ding, X. Zhang, S. Wang, H. Chen and Y. Zhu, *ACS Energy Lett.*, 2018, **3**, 2544–2549.
- 111 C. Cheng, X. Wang, Y. Lin, L. He, J.-X. Jiang, Y. Xu and F. Wang, *Polym. Chem.*, 2018, **9**, 4468–4475.
- 112 S. Bi, Z. A. Lan, S. Paasch, W. Zhang, Y. He, C. Zhang, F. Liu, D. Wu, X. Zhuang and E. Brunner, *Adv. Funct. Mater.*, 2017, **27**, 1703146.
- 113 Z. Wang, X. Yang, T. Yang, Y. Zhao, F. Wang, Y. Chen, J. H. Zeng, C. Yan, F. Huang and J.-X. Jiang, *ACS Catal.*, 2018, **8**, 8590–8596.
- 114 R. S. Sprick, B. Bonillo, M. Sachs, R. Clowes, J. R. Durrant, D. J. Adams and A. I. Cooper, *Chem. Commun.*, 2016, **52**, 10008–10011.
- 115 R. S. Sprick, Y. Bai, A. A. Guilbert, M. Zbiri, C. M. Aitchison, L. Wilbraham, Y. Yan, D. J. Woods, M. A. Zwijnenburg and A. I. Cooper, *Chem. Mater.*, 2019, **31**, 305–313.
- 116 Y. Zhao, W. Ma, Y. Xu, C. Zhang, Q. Wang, T. Yang, X. Gao, F. Wang, C. Yan and J.-X. Jiang, *Macromolecules*, 2018, **51**, 9502–9508.
- 117 G. Zhang, W. Ou, J. Wang, Y. Xu, D. Xu, T. Sun, S. Xiao, M. Wang, H. Li, W. Chen and C. Su, *Appl. Catal., B*, 2019, **245**, 114–121.
- 118 F. Beuerle and B. Gole, *Angew. Chem., Int. Ed.*, 2018, **57**, 4850–4878.
- 119 L. Stegbauer, K. Schwinghammer and B. V. Lotsch, *Chem. Sci.*, 2014, **5**, 2789–2793.
- 120 V. S. Vyas, F. Haase, L. Stegbauer, G. Savasci, F. Podjaski, C. Ochsenfeld and B. V. Lotsch, *Nat. Commun.*, 2015, **6**, 8508.
- 121 X. Wang, L. Chen, S. Y. Chong, M. A. Little, Y. Wu, W.-H. Zhu, R. Clowes, Y. Yan, M. A. Zwijnenburg and R. S. Sprick, *Nat. Chem.*, 2018, **10**, 1180.
- 122 P. Pachfule, A. Acharjya, J. r. m. Roeser, T. Langenhahn, M. Schwarze, R. Schomäcker, A. Thomas and J. Schmidt, *J. Am. Chem. Soc.*, 2018, **140**, 1423–1427.
- 123 J. L. Sheng, H. Dong, X. B. Meng, H. L. Tang, Y. H. Yao, D. Q. Liu, L. L. Bai, F. M. Zhang, J. Z. Wei and X. J. Sun, *ChemCatChem*, 2019, **11**, 2313–2319.
- 124 E. Jin, Z. Lan, Q. Jiang, K. Geng, G. Li, X. Wang and D. Jiang, *Chem*, 2019, **5**, 1632–1647.
- 125 A. Thomas, *Angew. Chem., Int. Ed.*, 2010, **49**, 8328–8344.
- 126 K. Sakaushi and M. Antonietti, *Acc. Chem. Res.*, 2015, **48**, 1591–1600.
- 127 P. Puthiaraj, Y.-R. Lee, S. Zhang and W.-S. Ahn, *J. Mater. Chem. A*, 2016, **4**, 16288–16311.
- 128 J. Bi, W. Fang, L. Li, J. Wang, S. Liang, Y. He, M. Liu and L. Wu, *Macromol. Rapid Commun.*, 2015, **36**, 1799–1805.
- 129 S. Kuecken, A. Acharjya, L. Zhi, M. Schwarze, R. Schomäcker and A. Thomas, *Chem. Commun.*, 2017, **53**, 5854–5857.
- 130 Z.-A. Lan, Y. Fang, X. Chen and X. Wang, *Chem. Commun.*, 2019, **55**, 7756–7759.
- 131 W. Huang, Q. He, Y. Hu and Y. Li, *Angew. Chem., Int. Ed.*, 2019, **131**, 1–6.
- 132 K. Wang, L. M. Yang, X. Wang, L. Guo, G. Cheng, C. Zhang, S. Jin, B. Tan and A. I. Cooper, *Angew. Chem., Int. Ed.*, 2017, **56**, 14149–14153.
- 133 L. Guo, Y. Niu, H. Xu, Q. Li, S. Razzaque, Q. Huang, S. Jin and B. Tan, *J. Mater. Chem. A*, 2018, **6**, 19775–19781.
- 134 C. B. Meier, R. S. Sprick, A. Monti, P. Guiglion, J.-S. M. Lee, M. A. Zwijnenburg and A. I. Cooper, *Polymer*, 2017, **126**, 283–290.
- 135 Y. S. Kochergin, D. Schwarz, A. Acharjya, A. Ichangi, R. Kulkarni, P. Eliášová, J. Vacek, J. Schmidt, A. Thomas and M. J. Bojdys, *Angew. Chem., Int. Ed.*, 2018, **57**, 14188–14192.
- 136 L. Li, W. Fang, P. Zhang, J. Bi, Y. He, J. Wang and W. Su, *J. Mater. Chem. A*, 2016, **4**, 12402–12406.
- 137 Z. Cheng, W. Fang, T. Zhao, S. Fang, J. Bi, S. Liang, L. Li, Y. Yu and L. Wu, *ACS Appl. Mater. Interfaces*, 2018, **10**, 41415–41421.
- 138 M. Liu, Q. Huang, S. Wang, Z. Li, B. Li, S. Jin and B. Tan, *Angew. Chem.*, 2018, **130**, 12144–12148.
- 139 S. Chu, Y. Wang, Y. Guo, P. Zhou, H. Yu, L. Luo, F. Kong and Z. Zou, *J. Mater. Chem.*, 2012, **22**, 15519–15521.
- 140 X. Wang, K. Maeda, A. Thomas, K. Takanebe, G. Xin, J. M. Carlsson, K. Domen and M. Antonietti, *Nat. Mater.*, 2009, **8**, 76.
- 141 A. Naseri, M. Samadi, A. Pourjavadi, A. Z. Moshfegh and S. Ramakrishna, *J. Mater. Chem. A*, 2017, **5**, 23406–23433.
- 142 Y. Wang, Y. Di, M. Antonietti, H. Li, X. Chen and X. Wang, *Chem. Mater.*, 2010, **22**, 5119–5121.
- 143 J. Fang, H. Fan, M. Li and C. Long, *J. Mater. Chem. A*, 2015, **3**, 13819–13826.
- 144 H. Wang, B. Wang, Y. Bian and L. Dai, *ACS Appl. Mater. Interfaces*, 2017, **9**, 21730–21737.
- 145 J. Zhang, X. Chen, K. Takanebe, K. Maeda, K. Domen, J. D. Epping, X. Fu, M. Antonietti and X. Wang, *Angew. Chem., Int. Ed.*, 2010, **49**, 441–444.
- 146 M. K. Bhunia, K. Yamauchi and K. Takanebe, *Angew. Chem., Int. Ed.*, 2014, **53**, 11001–11005.
- 147 X. Fan, L. Zhang, R. Cheng, M. Wang, M. Li, Y. Zhou and J. Shi, *ACS Catal.*, 2015, **5**, 5008–5015.
- 148 J. Zhang, M. Zhang, S. Lin, X. Fu and X. Wang, *J. Catal.*, 2014, **310**, 24–30.
- 149 H. Yu, R. Shi, Y. Zhao, T. Bian, Y. Zhao, C. Zhou, G. I. Waterhouse, L. Z. Wu, C. H. Tung and T. Zhang, *Adv. Mater.*, 2017, **29**, 1605148.
- 150 W. Ren, J. Cheng, H. Ou, C. Huang, M. M. Titirici and X. Wang, *ChemSusChem*, 2019, **12**, 3257–3262.

- 151 Z. Zhang, J. Long, L. Yang, W. Chen, W. Dai, X. Fu and X. Wang, *Chem. Sci.*, 2011, **2**, 1826–1830.
- 152 J. Chen, C. L. Dong, D. Zhao, Y. C. Huang, X. Wang, L. Samad, L. Dang, M. Shearer, S. Shen and L. Guo, *Adv. Mater.*, 2017, **29**, 1606198.
- 153 W. Zhou, T. Jia, H. Shi, D. Yu, W. Hong and X. Chen, *J. Mater. Chem. A*, 2019, **7**, 303–311.
- 154 S. Zang, G. Zhang, P. Yang, D. Zheng and X. Wang, *Chem. – Eur. J.*, 2019, **25**, 6102–6107.
- 155 M. Luo, Q. Yang, K. Liu, H. Cao and H. Yan, *Chem. Commun.*, 2019, **55**, 5829–5832.
- 156 G. Zhou, L.-L. Zheng, D. Wang, Q.-J. Xing, F. Li, P. Ye, X. Xiao, Y. Li and J.-P. Zou, *Chem. Commun.*, 2019, **55**, 4150–4153.
- 157 J. Yang, D. Wang, H. Han and C. Li, *Acc. Chem. Res.*, 2013, **46**, 1900–1909.
- 158 J. Ran, J. Zhang, J. Yu, M. Jaroniec and S. Z. Qiao, *Chem. Soc. Rev.*, 2014, **43**, 7787–7812.
- 159 X. Li, J. Yu, M. Jaroniec and X. Chen, *Chem. Rev.*, 2019, **119**, 3962–4179.
- 160 J. Kosco and I. McCulloch, *ACS Energy Lett.*, 2018, **3**, 2846–2850.
- 161 J. Kosco, M. Sachs, R. Godin, M. Kirkus, L. Francas, M. Bidwell, M. Qureshi, D. Anjum, J. R. Durrant and I. McCulloch, *Adv. Energy Mater.*, 2018, **8**, 1802181.
- 162 K. L. Materna, R. H. Crabtree and G. W. Brudvig, *Chem. Soc. Rev.*, 2017, **46**, 6099–6110.
- 163 D. Kong, Y. Zheng, M. Kobielski, Y. Wang, Z. Bai, W. Macyk, X. Wang and J. Tang, *Mater. Today*, 2018, **21**, 897–924.
- 164 Z. A. Lan, Y. Fang, Y. Zhang and X. Wang, *Angew. Chem., Int. Ed.*, 2018, **57**, 470–474.
- 165 J. Xie, S. A. Shevlin, Q. Ruan, S. J. Moniz, Y. Liu, X. Liu, Y. Li, C. C. Lau, Z. X. Guo and J. Tang, *Energy Environ. Sci.*, 2018, **11**, 1617–1624.
- 166 S. Chen, T. Takata and K. Domen, *Nat. Rev. Mater.*, 2017, **2**, 17050.
- 167 G. Zhang, Z.-A. Lan and X. Wang, *Chem. Sci.*, 2017, **8**, 5261–5274.
- 168 L. Wang, Y. Zhang, L. Chen, H. Xu and Y. Xiong, *Adv. Mater.*, 2018, **30**, 1801955.
- 169 G. Zhang, Z.-A. Lan, L. Lin, S. Lin and X. Wang, *Chem. Sci.*, 2016, **7**, 3062–3066.
- 170 X. Chen, R. Shi, Q. Chen, Z. Zhang, W. Jiang, Y. Zhu and T. Zhang, *Nano Energy*, 2019, **59**, 644–650.
- 171 A. Dhakshinamoorthy, S. Navalon, A. Corma and H. Garcia, *Energy Environ. Sci.*, 2012, **5**, 9217–9233.
- 172 J. Mao, K. Li and T. Peng, *Catal. Sci. Technol.*, 2013, **3**, 2481–2498.
- 173 W. Tu, Y. Zhou and Z. Zou, *Adv. Mater.*, 2014, **26**, 4607–4626.
- 174 J. Ran, M. Jaroniec and S. Z. Qiao, *Adv. Mater.*, 2018, **30**, 1704649.
- 175 Y. Chen, G. Ji, S. Guo, B. Yu, Y. Zhao, Y. Wu, H. Zhang, Z. Liu, B. Han and Z. Liu, *Green Chem.*, 2017, **19**, 5777–5781.
- 176 Y. Fang and X. Wang, *Chem. Commun.*, 2018, **54**, 5674–5687.
- 177 F. Xing, Q. Liu, M. Song and C. Huang, *ChemCatChem*, 2018, **10**, 5270–5279.
- 178 J. Qin, S. Wang, H. Ren, Y. Hou and X. Wang, *Appl. Catal., B*, 2015, **179**, 1–8.
- 179 H. Shi, S. Long, J. Hou, L. Ye, Y. Sun, W. Ni, C. Song, K. Li, G. G. Gurzadyan and X. Guo, *Chem. – Eur. J.*, 2019, **25**, 5028–5035.
- 180 C. Yang, W. Huang, L. C. da Silva, K. A. I. Zhang and X. Wang, *Chem. – Eur. J.*, 2018, **24**, 17454–17458.
- 181 X. Yu, Z. Yang, B. Qiu, S. Guo, P. Yang, B. Yu, H. Zhang, Y. Zhao, X. Yang, B. Han and Z. Liu, *Angew. Chem., Int. Ed.*, 2019, **58**, 632–636.
- 182 S. Yang, W. Hu, X. Zhang, P. He, B. Pattengale, C. Liu, M. Cendejas, I. Hermans, X. Zhang and J. Zhang, *J. Am. Chem. Soc.*, 2018, **140**, 14614–14618.
- 183 J. Fu, B. Zhu, C. Jiang, B. Cheng, W. You and J. Yu, *Small*, 2017, **13**, 1603938.
- 184 M. Shen, L. Zhang, M. Wang, J. Tian, X. Jin, L. Guo, L. Wang and J. Shi, *J. Mater. Chem. A*, 2019, **7**, 1556–1563.
- 185 Y. Fu, X. Zhu, L. Huang, X. Zhang, F. Zhang and W. Zhu, *Appl. Catal., B*, 2018, **239**, 46–51.
- 186 J.-L. Wang, C. Wang, K. E. deKrafft and W. Lin, *ACS Catal.*, 2012, **2**, 417–424.
- 187 Z. J. Wang, K. Garth, S. Ghasimi, K. Landfester and K. A. I. Zhang, *ChemSusChem*, 2015, **8**, 3459–3464.
- 188 Z. J. Wang, S. Ghasimi, K. Landfester and K. A. I. Zhang, *Adv. Mater.*, 2015, **27**, 6265–6270.
- 189 C. Su, R. Tandiana, B. Tian, A. Sengupta, W. Tang, J. Su and K. P. Loh, *ACS Catal.*, 2016, **6**, 3594–3599.
- 190 V. R. Battula, H. Singh, S. Kumar, I. Bala, S. K. Pal and K. Kailasam, *ACS Catal.*, 2018, **8**, 6751–6759.
- 191 C. Yang, Q. Jin, H. Zhang, J. Liao, J. Zhu, B. Yu and J. Deng, *Green Chem.*, 2009, **11**, 1401–1405.
- 192 R. D. Chakravarthy, V. Ramkumar and D. K. Chand, *Green Chem.*, 2014, **16**, 2190–2196.
- 193 B. Zhang, J. Li, B. Zhang, R. Chong, R. Li, B. Yuan, S.-M. Lu and C. Li, *J. Catal.*, 2015, **332**, 95–100.
- 194 Z. J. Wang, S. Ghasimi, K. Landfester and K. A. I. Zhang, *Chem. Commun.*, 2014, **50**, 8177–8180.
- 195 M. Liras, M. Iglesias and F. L. Sánchez, *Macromolecules*, 2016, **49**, 1666–1673.
- 196 J. Luo, X. Zhang and J. Zhang, *ACS Catal.*, 2015, **5**, 2250–2254.
- 197 P.-F. Wei, M.-Z. Qi, Z.-P. Wang, S.-Y. Ding, W. Yu, Q. Liu, L.-K. Wang, H.-Z. Wang, W.-K. An and W. Wang, *J. Am. Chem. Soc.*, 2018, **140**, 4623–4631.
- 198 J.-X. Jiang, Y. Li, X. Wu, J. Xiao, D. J. Adams and A. I. Cooper, *Macromolecules*, 2013, **46**, 8779–8783.
- 199 Y. Zhi, Z. Li, X. Feng, H. Xia, Y. Zhang, Z. Shi, Y. Mu and X. Liu, *J. Mater. Chem. A*, 2017, **5**, 22933–22938.
- 200 Y. Zhi, S. Ma, H. Xia, Y. Zhang, Z. Shi, Y. Mu and X. Liu, *Appl. Catal., B*, 2019, **244**, 36–44.
- 201 J. Luo, X. Zhang and J. Zhang, *ACS Catal.*, 2015, **5**, 2250–2254.
- 202 Z. Lia, Y. Zhi, P. Shao, H. Xia, G. Li, X. Feng and X. Chen, *Appl. Catal., B*, 2019, **245**, 334–342.

- 203 V. Polshettiwar, A. Decottignies, C. Len and A. Fihri, *ChemSusChem*, 2010, **3**, 502–522.
- 204 I. P. Beletskaya, F. Alonso and V. Tyurin, *Coord. Chem. Rev.*, 2019, **385**, 137–173.
- 205 Q. Xiao, S. Sarina, A. Bo, J. Jia, H. Liu, D. P. Arnold, Y. Huang, H. Wu and H. Zhu, *ACS Catal.*, 2014, **4**, 1725–1734.
- 206 F. Wang, C. Li, H. Chen, R. Jiang, L.-D. Sun, Q. Li, J. Wang, J. C. Yu and C.-H. Yan, *J. Am. Chem. Soc.*, 2013, **135**, 5588–5601.
- 207 Z. J. Wang, S. Ghasimi, K. Landfester and K. A. I. Zhang, *Chem. Mater.*, 2015, **27**, 1921–1924.
- 208 S. Ghasimi, S. A. Bretschneider, W. Huang, K. Landfester and K. A. I. Zhang, *Adv. Sci.*, 2017, **4**, 1700101.
- 209 C. Ayed, L. C. da Silva, D. Wang and K. A. I. Zhang, *J. Mater. Chem. A*, 2018, **6**, 22145–22151.
- 210 W. Zhang, J. Tang, W. Yu, Q. Huang, Y. Fu, G. Kuang, C. Pan and G. Yu, *ACS Catal.*, 2018, **8**, 8084–8091.
- 211 T. Robinson, G. McMullan, R. Marchant and P. Nigam, *Bioresour. Technol.*, 2001, **77**, 247–255.
- 212 I. K. Konstantinou and T. A. Albanis, *Appl. Catal., B*, 2004, **49**, 1–14.
- 213 X. Liu, J. Iocozzia, Y. Wang, X. Cui, Y. Chen, S. Zhao, Z. Li and Z. Lin, *Energy Environ. Sci.*, 2017, **10**, 402–434.
- 214 S. Ghosh, N. A. Kouamé, L. Ramos, S. Remita, A. Dazzi, A. Deniset-Besseau, P. Beaunier, F. Goubard, P.-H. Aubert and H. Remita, *Nat. Mater.*, 2015, **14**, 505.
- 215 S. Ghosh, N. A. Kouame, S. Remita, L. Ramos, F. Goubard, P.-H. Aubert, A. Dazzi, A. Deniset-Besseau and H. Remita, *Sci. Rep.*, 2015, **5**, 18002.
- 216 S. Ghasimi, S. Prescher, Z. J. Wang, K. Landfester, J. Yuan and K. A. I. Zhang, *Angew. Chem., Int. Ed.*, 2015, **54**, 14549–14553.
- 217 J. Wang, H. Yang, L. Jiang, S. Liu, Z. Hao, J. Cheng and G. Ouyang, *Catal. Sci. Technol.*, 2018, **8**, 5024–5033.
- 218 B. Wang, Z. Xie, Y. Li, Z. Yang and L. Chen, *Macromolecules*, 2018, **51**, 3443–3449.
- 219 Q. Huang, L. Guo, N. Wang, X. Zhu, S. Jin and B. Tan, *ACS Appl. Mater. Interfaces*, 2019, **11**, 15861–15868.
- 220 B. C. Ma, S. Ghasimi, K. Landfester, F. Vilela and K. A. Zhang, *J. Mater. Chem. A*, 2015, **3**, 16064–16071.
- 221 B. P. Biswal, H. A. Vignolo-González, T. Banerjee, L. Grunenberg, G. Savasci, K. Gottschling, J. Nuss, C. Ochsenfeld and B. V. Lotsch, *J. Am. Chem. Soc.*, 2019, **141**, 11082–11092.
- 222 Y. Bai, L. Wilbraham, B. J. Slater, M. A. Zwijnenburg, R. S. Sprick and A. I. Cooper, *J. Am. Chem. Soc.*, 2019, **141**, 9063–9071.