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## Advanced functional polymer materials

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The research on advanced functional polymers is being driven by the fast-growing demand for new functional materials that can be used in revolutionary technologies. Polymers can be endowed with functions by using certain special preparation methods or by introducing functional groups or fillers into materials. These functions are either intrinsically possessed by materials or actuated by external stimuli. In this review, we present an overview of the recent developments made in the research hotspots of functional polymers, encompassing polymerization methodologies, luminescent polymers, photovoltaic polymers, other electronic and optical polymers (including low-*k* polyimides and second-order nonlinear optical polymers), biorelated polymers (particularly those for biomedical applications), supramolecular polymers, stimuli-responsive polymers, shape-memory polymers, separation polymer membranes, energy storage polymers, and covalent organic framework polymers. The concepts, design strategies at the molecular level, preparation methods, classifications, properties, potential applications, and recent progress made in such polymers are summarized. Challenges and future perspectives of each type of functional polymers are also addressed, including research efforts regarding the design and fabrication of functional polymers for serving the increasing demand for new materials.

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## 1. Introduction

Conventional polymers refer to materials that are used in our daily lives, such as plastics, artificial fibers, rubbers, and paint coatings. With the development of modern society, however, special functions are required in certain fields, where conventional polymers cannot serve the full needs, thereby promoting the development of novel polymers with specific functions. Such polymers are called advanced functional polymers that have potential applications in many fields such as energy harvesting and storage, wireless communication industry, biomedicine, oil/water or gas separation, and intelligent and bionic industries.<sup>1,2</sup> Functional polymer science, developed in the 1960s, is an interdisciplinary field encompassing polymer chemistry, polymer physics, materials science, biology, energy, nanoscience, and environment.<sup>3</sup> Besides the mechanical

properties of conventional polymeric materials, advanced functional polymeric materials can also possess chemical reactivity, photosensitivity, electrical conductivity, catalytic properties, biocompatibility, biological activity, pharmacological properties, selective separation, and energy conversion. They have become an important cornerstone of the modern industry and advanced technology. A considerable amount of attention has been paid toward the development of new functional polymers to meet the major demand necessary for the development of modern society.

Advanced functional polymers follow their own characteristics and laws. The first is that the syntheses of monomers are usually complex and diverse, which involve organic chemistry and metal catalysis. Second, new polymerization reactions, methods, mechanisms, and catalysts are needed to be developed because of the introduction of functional monomers. Third, their functions are not only related to the chemical structures of the

polymer chains but also closely related to their aggregate states. Therefore, it is necessary to combine the synthesis, processing, and morphology of polymers to obtain excellent functions. The cutting-edge development of other disciplines can be used to create new materials and improve the performance of the existing polymeric materials because the field of functional polymeric materials is an interdisciplinary subject.

Advanced functional polymers have drawn much attention from researchers in many fields. As a result, a number of review papers have been published with different emphases on this particular subject. Almost all of them involve specific aspects of advanced functional polymers, such as conductive polymers,<sup>4</sup> polymer micelles or vesicles for drug and gene delivery,<sup>5,6</sup> polymer gels,<sup>7,8</sup> polymer acceptors,<sup>9–11</sup> or/and donors<sup>12</sup> for use in organic solar cells (OSCs), biomedical polymers,<sup>13,14</sup> shape-memory polymers (SMPs),<sup>15–17</sup> stimuli-responsive polymers,<sup>18,19</sup> separation polymer membranes,<sup>20,21</sup> luminescent polymers,<sup>22,23</sup>

covalent organic framework (COF) polymers,<sup>24</sup> and supramolecular polymers (SPs).<sup>25</sup> All the above reviews only focused on one specific aspect of functional polymers. To the best of our knowledge, no reviews have summarized and reported all the research hotspots of advanced functional polymers. For a better understanding of advanced functional polymers in a comprehensive manner, a review of functional polymeric materials is necessary. Owing to advanced functional polymers covering too many topics, we mainly present research directions studied by the authors' own groups in this review. We start by discussing polymerization methodologies of advanced functional polymers and then focus on several research directions involving functional polymers, such as luminescent polymers, photovoltaic polymers, other electronic and optical polymers, biorelated polymers, SPs, stimuli-responsive polymers, SMPs, separation polymer membranes, energy storage polymers, and COF polymers.

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## 2. Polymerization methodologies

The key element of a functional polymer is its preparation methodology, which can facilitate the creation of new functional polymers and improve their performance. Much effort has been devoted toward the development of new polymerizations, which are not only owing to research interests but also because of various potential applications of polymers. Theoretically, polymerizations can be classified into chain-growth and step-growth polymerizations. Chain-growth polymerization, *i.e.*, addition polymerization, requires the initiation of a monomer to begin the chain-growth process, which involves the addition of monomers to growing free radicals or cationic or anionic chains. As the name implies, the chain of step-growth polymerization grows in a stepwise manner. Condensation polymerization and alkyne-based click polymerization usually exhibit the features of step-growth polymerization. Ring-opening polymerization (ROP) possesses many features of chain-growth polymerization, but it may also exhibit some features of step-growth polymerization. Free-radical polymerizations usually comprise four processes: initiation, propagation, chain transfer, and termination. If chain transfer and termination can be minimized during the process of polymerization, the resultant polymer may be controlled to a certain level. The corresponding polymerization is called controlled radical polymerization (CRP).

In this section, the research content includes the preparation of new functional monomers and the development of new polymerization reactions. It is essential to develop a polymerization reaction featuring advantages such as simplicity, efficiency, mild conditions, good selectivity, atomic economy, and easily available raw materials for the preparation of polymeric materials with newer structures and functions. Due to space limitations, we will discuss several representative polymerizations for the preparation of functional polymers: (1) CRP (chain-growth polymerization) for the preparation of olefin functional polymers; (2) alkyne-based click polymerization (step-growth polymerization); (3) multicomponent polymerization (MCP) based on triple-bond building blocks used for the synthesis of alkyne functional polymers (step-growth polymerization); and (4) ROP (possessing the features of both chain- and step-growth polymerizations) for the preparation of degradable and renewable polymer materials by the use of new catalysts and renewable biomonomers.

### 2.1 CRP

CRP or reversible-deactivation radical polymerization (RDRP) has reshaped the field of polymer chemistry and materials science by allowing nonexperts to access well-defined polymers that can be prepared under mild conditions for a myriad of applications. CRP relies on either reversible termination or degenerative chain transfer to control the equilibrium between dormant and active radicals; polymerization control is realized by biasing the equilibrium toward the dormant species. As such, the concentration of active radicals is minimized, and side reactions such as bimolecular termination and irreversible chain

transfer are diminished. Although CRP techniques are not living polymerizations, they do exhibit features analogous to living polymerization (*e.g.*, living anionic polymerization), such as pseudo-first-order polymerization kinetics, predictable molecular weight with a narrow distribution, and high end-group fidelity. Despite having been extensively studied for more than 20 years, some exciting new trends have recently emerged, such as externally regulated CRP and enzyme-participated CRP. These advances provide new opportunities to control polymer architectures and functions, as well as the development of sustainable pathways for synthesizing materials under greener conditions.

External stimuli, such as light, electric field, or ultrasound, can provide useful means to regulate the activation or alteration of the polymerization rate of CRP by switching the stimuli ON/OFF or changing the input intensity/magnitude. In 2011, Matyjaszewski *et al.* reported an electrochemically mediated atom transfer radical polymerization (eATRP) process, whereby applying suitable potential reduces the air-stable Cu(II)/ligand deactivator to the Cu(I)/ligand activator to induce polymerization, and the polymerization rate can be modulated by varying the applied potential to control the activator/deactivator ratio.<sup>27</sup> Inspired by the visible-light photoredox catalysis for organic transformations, Fors and Hawker *et al.* developed visible-light-mediated ATRP using *fac*-[Ir(ppy)<sub>3</sub>] as the photoredox catalyst, which allowed the photomodulation of polymerization kinetics by switching the light ON/OFF.<sup>28</sup> However, transition-metal photocatalysts are expensive and may potentially contaminate the obtained polymers: a similar problem was found in traditional ATRP when using a Cu/ligand catalyst. To address this challenge, the same team further developed a metal-free ATRP using a highly reducing organic photocatalyst, namely, PTH (Fig. 1).<sup>26</sup> In contrast to *fac*-[Ir(ppy)<sub>3</sub>], PTH was compatible with a broader range of functional monomers, such as 2-(dimethylamino)ethyl methacrylate. The field of using organic photocatalysts for conducting metal-free or organic-compound-catalyzed ATRP (O-ATRP) was further developed by Miyake *et al.*, who introduced diaryl dihydrophenazine as another class of strongly reducing photoredox catalysts.<sup>29</sup> On the basis of computational studies, it was suggested that the ability to access photoexcited intramolecular charge transfer states is important in minimizing fluorescence and facilitating electron transfer from the photoexcited photocatalyst to the substrate, thereby improving the efficiency and control of ATRP.<sup>30</sup> However, the light absorption

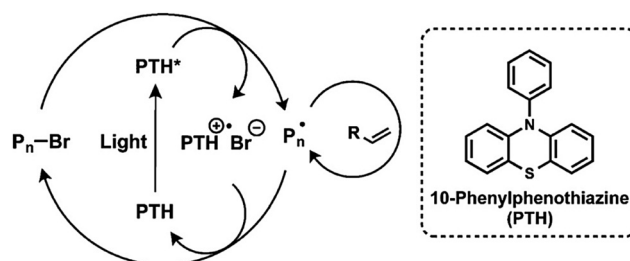


Fig. 1 Metal-free ATRP mediated by 10-phenylphenothiazine (PTH) as a photoredox catalyst. Adapted from ref. 26 with permission. Copyright (2014) American Chemical Society.

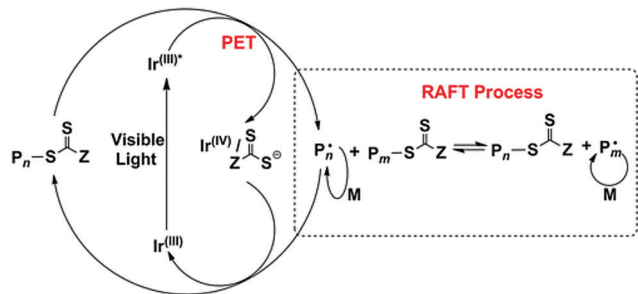


Fig. 2 Mechanism of PET-RAFT using *fac*-[Ir(ppy)<sub>3</sub>] as a photoredox catalyst. Adapted from ref. 33 with permission. Copyright (2014) American Chemical Society.

wavelength of existing organic photoredox catalysts for ATRP is limited to ultraviolet or blue light; therefore, significant efforts have been devoted toward the design and synthesis of long-wavelength-absorbing photoredox catalysts, *e.g.*, by extending the conjugation length.<sup>31,32</sup> Photoredox-catalyzed reactions have also been coupled with another type of widely used CRP, namely, reversible addition–fragmentation chain transfer (RAFT) polymerization, which was pioneered by Boyer *et al.* in 2014 and was called the photoinduced electron/energy transfer RAFT (PET-RAFT) (Fig. 2).<sup>33–40</sup> PET-RAFT is extremely versatile as evident by the variety of photocatalysts (metal complexes, organic molecules, biocatalysts, and nanoparticles (NPs)) and therefore a wide range of wavelengths covering the visible as well as near-infrared (NIR) spectra, a diverse families of monomers, and various polymerization media. In particular, some of these photocatalysts are capable of reducing triplet oxygen into its singlet state, thereby enabling polymerizations to be conducted with oxygen tolerance.<sup>33</sup> This trait has been harnessed to facilitate flow polymerization and high-throughput synthesis at lower volumes.<sup>41</sup>

Despite being highly versatile, only a few photocatalysts have been employed for aqueous polymerization. A limitation of aqueous polymerization when using photoredox catalysis is that the catalysts typically exhibit large conjugation and therefore tend to aggregate in water, which, in turn, reduces their performance. To address this problem, An *et al.* introduced a host–guest strategy to improve water solubility and reduce the aggregation of photoredox catalysts. Enhanced polymerization rates were demonstrated with respect to cucurbit[7]uril (CB[7]) binding with the photocatalyst, Zn(II) *meso*-tetra(4-naphthalylmethylpyridyl) porphyrin.<sup>42</sup> This host–guest strategy is simple and effective with minimal synthesis efforts and allows tuning the photochemical properties of supramolecular photocatalysts.

The rapid progress in visible-light-controlled CRP has offered exciting opportunities for the preparation of advanced materials under biologically friendly conditions. A notable example along this line was the work by Hawker and Soh *et al.*,<sup>43</sup> who reported a cytocompatible strategy using Eosin Y as the photocatalyst for directly grafting high-density structurally defined functional polymers from living cell surfaces with high cell viability. This advancement was enabled by fast polymerization kinetics using a biocompatible polyethylene glycol (PEG)–acrylamide monomer and equimolar chain transfer agent and cocatalyst,

which resulted in conversion of <30% within 5 min. In addition, ultrasound has been investigated as a mechanical stimulus to modulate the CRP, involving both ATRP and RAFT.<sup>44,45</sup>

Further, we have witnessed significant interest in biocatalyzed CRP in recent years, which promises greener pathways for precision polymer synthesis owing to the mildness and high efficiency of enzymatic catalysis.<sup>47,48</sup> The enzymes that have been explored to control radical polymerizations are mainly oxidoreductases such as metalloproteins or flavoproteins. Metalloproteins containing a transition-metal complex as the cofactor have been examined as ATRP catalysts given their similar properties to traditional ATRP catalysts (*e.g.*, Cu/ligand complex). However, metal complexes that are tightly bound to the protein matrix in metalloproteins can yield a catalyst with much lower toxicity and that is more robust toward functional monomers. For example, the polymerization of monomers with complexing capabilities toward transition metals is difficult to control by traditional ATRP. In a notable recent work, Bruns *et al.* reported the controlled polymerization of *N*-vinylimidazole under aqueous ATRP conditions catalyzed by laccase, a multi-copper-containing oxidoreductase.<sup>49</sup> Laccase could be quantitatively separated from the polymerization solution *via* ultracentrifugation fitted with filter membranes having a suitable molecular weight cutoff. The ability to access low-dispersity functional poly(*N*-vinylimidazole) without metal contamination permits its applications as biomaterials and energy materials. Some peroxidase metalloproteins such as horseradish peroxidase (HRP) catalyze the oxidation of electron-rich compounds and generate radical intermediates, which can be harnessed to initiate RAFT polymerization or activate ATRP by reducing a Cu(II)/ligand deactivator into a Cu(I)/ligand activator. This strategy was first introduced by An *et al.* in RAFT polymerization initiated by HRP/hydrogen peroxide/acetylacetone—a three-component system.<sup>50</sup> Predictable molecular weights ranging from  $5 \times 10^3$  to  $520 \times 10^3$  g mol<sup>-1</sup>, low narrow molecular weight distributions, and pseudo-first-order kinetics were observed for the polymerizations of various monomers in solution, dispersion, and biological media. Utilizing a biological Fenton reaction between hemoglobin and hydrogen peroxide to generate hydroxyl radicals, Qiao *et al.* reported a very interesting blood-catalyzed RAFT polymerization process, which may permit *in vivo* cell engineering using synthetic polymers.<sup>51</sup> One of the grand challenges of CRP is its oxygen sensitivity. Recently, this challenge has been successfully tackled by the use of flavin-containing oxidases, which can effectively remove dissolved oxygen from the polymerization solution, allowing CRP to be conducted without prior degassing *via* traditional inert gas purging or freeze–pump–thaw operations. Since the first demonstration of oxygen-tolerant RAFT enabled by glucose oxidase (GOx) deoxygenation by Stevens *et al.*,<sup>52</sup> applying oxidase to enable oxygen-tolerant RAFT and ATRP has attracted significant attention because this elegant strategy not only simplifies the synthesis operation but facilitates combinatorial synthesis methods to be developed, which is expected to play a key role in functional material discovery by high-throughput

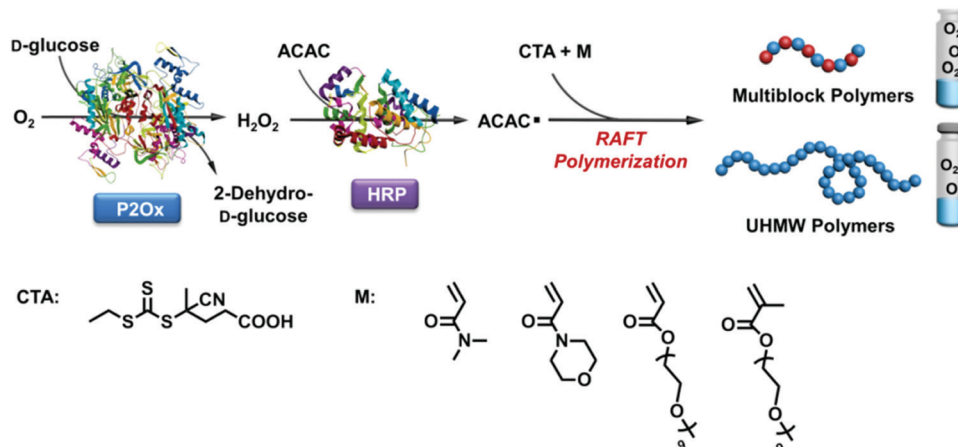


Fig. 3 Pyranose oxidase (P2Ox)–HRP enzymatic cascade catalysis for the synthesis of multiblock and ultrahigh-molecular-weight (co)polymers with oxygen tolerance. Adapted from ref. 46 with permission. Copyright (2017) Wiley-VCH.

structure–property relationship screening.<sup>53–56</sup> Remarkably, enzymatic cascade catalysis that involves oxidase deoxygenation and peroxidase radical generation has been demonstrated to be an elegant and powerful strategy for the efficient synthesis of well-defined polymers with oxygen tolerance. An *et al.* reported the use of GOx–HRP dual enzymes for controlled RAFT polymerization under ambient conditions, in which GOx deoxygenation provided hydrogen peroxide *in situ* for the HRP oxidation of acetylacetone to initiate well-controlled RAFT polymerization.<sup>50</sup> Later on, inspired by the oxidase-peroxidase cascade catalysis found in root fungi, they reported a new cascade involving P2Ox and HRP for the efficient synthesis of decablock copolymers in open vessels and ultrahigh-molecular-weight polymers (up to  $2.3 \times 10^6 \text{ g mol}^{-1}$ ) in close vessels without prior deoxygenation (Fig. 3).<sup>46</sup> The GOx–HRP cascade was also employed by Matyjaszewski *et al.* for oxygen-tolerant ATRP, in which the radicals generated by the cascade catalysis was used to reduce Cu(II)/ligand to activate the polymerization (Fig. 4).<sup>57</sup>

## 2.2 Alkyne-based click polymerization

Since it was coined by Sharpless *et al.* in 2001, click chemistry has inspired a lot of scientists in various fields.<sup>58</sup> Attracted by the outstanding merits of click reactions, such as high efficiency, atom economy, mild reaction conditions, good functional tolerance, simple isolation procedure, and regio-/stereoselectivity,

polymer chemists have made great effort to develop click reactions into versatile polymerization techniques, that is, click polymerizations.<sup>59,60</sup> Until now, besides Cu(I)-catalyzed azide–alkyne click polymerization (CuAAC) evolved from a typical click reaction of Cu(I)-catalyzed azide–alkyne cycloaddition (CuAAC), numerous click polymerizations such as Diels–Alder click polymerization, thiol-based click polymerization, and amino-based click polymerization have also been established, which inherit the fascinating features of click reactions. In particular, alkyne-based click polymerizations combining the advantages of click reaction and the rich chemistry of alkynes have drawn considerable attention of polymer chemists.<sup>61</sup> As shown in Fig. 5, diversified alkyne-based click polymerizations including azide–alkyne click polymerizations (AACPs), thiol–yne click polymerizations, and emerging amino–yne click polymerizations as well as hydroxyl–yne click polymerizations, have been exploited to fabricate functional polymers for specific applications.

Over the past few decades, CuAACs—the most famous click polymerizations—have been widely applied in the construction of linear and hyperbranched polytriazoles (PTAs) with varied properties, such as amphiphathy, self-assembly, self-healing, photoluminescence, aggregation-induced emission (AIE), chemosensing, ionic conductivity, light harvesting, optical nonlinearity, and catalytic activity (Fig. 6a).<sup>62</sup> In recent years, considerable progress in the methodology has also been made in the development of CuAACs. Generally, CuAACs is a step-growth process. By taking advantage of the complexation between triazole rings and Cu(I) catalyst, Gao *et al.* successfully established a novel chain-growth CuAAC of multifunctional  $AB_m$  ( $m \geq 2$ ) monomers and prepared a series of hyperbranched PTAs with high molecular weights (up to  $430 \times 10^3 \text{ g mol}^{-1}$ ), low dispersities, and high degrees of branching (DB).<sup>63–66</sup> Considering the fact that CuAACs could only afford 1,4-regioregular PTAs, Qin and Tang *et al.* developed the ligand-controlled regiodivergent Ru(II)-catalyzed azide–alkyne click polymerization (RuAACP), which can selectively offer 1,4- or 1,5-regioregular PTAs (Fig. 6b).<sup>67,68</sup> With this RuAACP in hand, they systematically investigated the structure–property

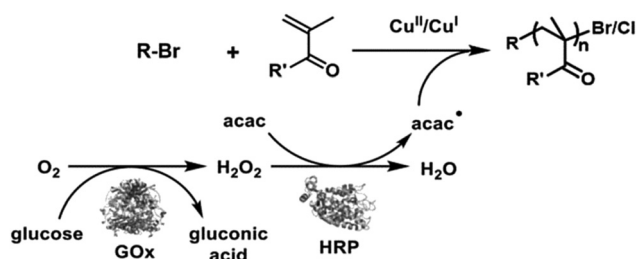


Fig. 4 “Oxygen-fueled” ATRP. Adapted from ref. 57 with permission. Copyright (2018) Wiley-VCH.

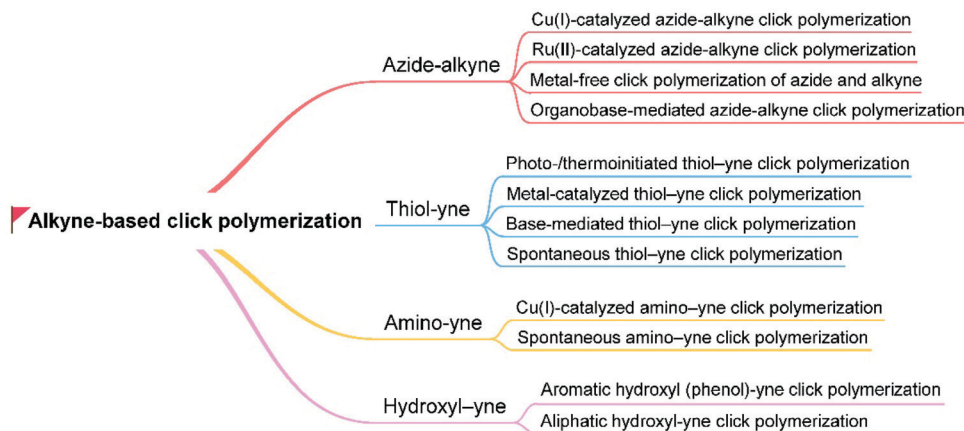


Fig. 5 Alkyne-based click polymerization.

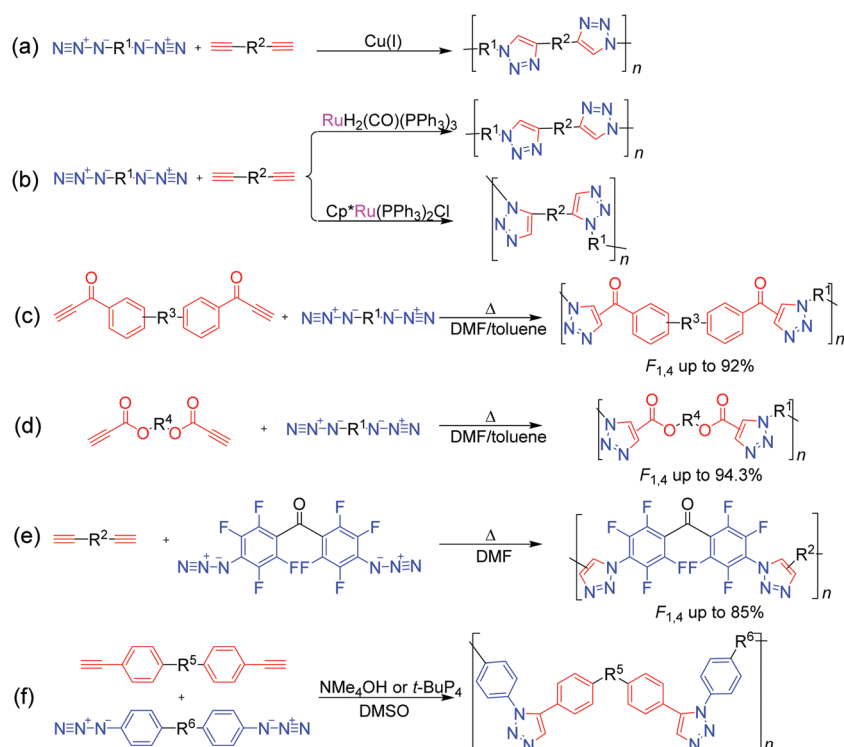


Fig. 6 AACPs: (a) CuAAC, (b) RuAAC, and (c) MFCP of arylacetylene and azide monomers; (d) MFCP of propiolate and azide monomers; (e) MFCP of 4,4'-diazidoperfluorobenzophenone and alkyne monomers; and (f) organobase-mediated AACPs.

relationship of regioregular PTAs. Although these transition-metal-catalyzed AACPs have become powerful tools for the preparation of functional PTAs with diverse structures, metal catalyst residues are difficult to be completely removed from PTAs owing to the strong coordination between the metallic species and nitrogen atoms in the triazole rings, which are detrimental to their applications in the areas of biomaterials and optoelectronic materials.<sup>69</sup> The usage of supported Cu(I) catalysts for AACPs can reduce the metal residue contents in the resultant PTAs, which stimulates the development of the metal-free click polymerization (MFCP) of azides and alkynes without the risk of metal catalyst residues.<sup>70,71</sup>

With arylacetylenes as activated alkyne monomers, Qin and Tang *et al.* successfully developed a thermal-initiated MFCP process for the preparation of 1,4-regioregular PTAs (Fig. 6c).<sup>72–75</sup> To avoid the tedious and difficult synthesis of arylacetylene monomers, readily available propiolate monomers with similar structures were then designed and applied in MFCP (Fig. 6d).<sup>76–78</sup> In addition to these MFCPs of azides and activated alkynes, the MFCPs of alkynes and activated azide of 4,4'-diazidoperfluorobenzophenone were also established by them (Fig. 6e).<sup>79,80</sup> They prepared several 1,4-regioregular PTAs with well-defined structures, which could be used in the areas of chemosensing, self-healing, photonic patterning, nonlinear optics, *etc.*

Recently, Qin and Tang *et al.* reported the AACPs mediated by the organobases of tetramethylammonium hydroxide (NMe<sub>4</sub>OH) and phosphazene base (*t*-BuP<sub>4</sub>); consequently, 1,5-regioregular PTAs were successfully produced (Fig. 6f).<sup>81,82</sup>

Another emerging alkyne-based click polymerization reaction is thiol- $\pi$  click polymerization, which can proceed in the bis-addition or mono-addition manner to afford polythioethers or poly(vinyl sulfide)s (PVSS).<sup>83</sup> In general, the mechanism of thiol- $\pi$  click polymerizations can be categorized into three types, namely, radical-initiated, transition-metal-catalyzed, and base-mediated processes. Inspired by the original work of Perrier *et al.* in 2009,<sup>84</sup> photo-/thermo-initiated thiol- $\pi$  click polymerizations, typical radical-initiated thiol- $\pi$  click polymerizations, and those occurring through a bis-addition process, were widely investigated and applied in the preparation of hyperbranched polythioethers with high DB and various other properties, such as high refractivity, biodegradability, and liquid crystallinity (Fig. 7a). By using disubstituted aromatic alkynes as monomers, Voit *et al.* successfully exploited a novel thermo-initiated thiol- $\pi$  click polymerization reaction, which proceeded *via* a mono-addition process to produce PVSS (Fig. 7b).<sup>85</sup> Diverse hyperbranched PVSSs with high refractive indices were obtained, which have been applied in the fabrication of planar all-polymer photonic crystals and organic light-emitting diodes.<sup>86,87</sup> Theoretically, the mono-addition of thiol to alkyne can occur in the form of Markovnikov or anti-Markovnikov addition, and the produced vinyl sulfides can be regio- and stereoisomers. To obtain PVSSs with well-defined structures, Tang *et al.* developed a transition-metal-catalyzed thiol- $\pi$  click polymerization reaction with regioselectivity and stereoselectivity.<sup>88</sup> By using the Rh(PPh<sub>3</sub>)<sub>3</sub>Cl catalyst, anti-Markovnikov additive PVSSs with high stereoregularities (*E*-isomer contents up to 100%) could be obtained in high yields (Fig. 7c). Meanwhile, with propiolates as activated monomers and diphenylamine as a promoter, organobase-mediated thiol- $\pi$  click polymerization was also established for the

construction of *Z*-stereoregular PVSSs (*Z*-isomer contents up to 81.4%) (Fig. 7d).<sup>89</sup> More recently, Qin and Tang *et al.* reported an efficient inorganic base-mediated thiol- $\pi$  click polymerization reaction in the presence of K<sub>3</sub>PO<sub>4</sub>, which afforded PVSSs with *Z*-stereoregularities up to 100% (Fig. 7e).<sup>90</sup> In particular, they also found that the click polymerization of thiols and aromatic alkynes could proceed in an anti-Markovnikov additive manner by simple mixing in tetrahydrofuran (THF) without the assistance of external stimuli and additional catalysts, indicating the spontaneity of this click polymerization reaction (Fig. 7f).<sup>91,92</sup> A series of PVSSs with linear and hyperbranched structures were prepared by this click polymerization reaction.

Besides the promising progress in AACPs and thiol- $\pi$  click polymerizations, new types of alkyne-based click polymerizations with great potential have also been developed in the past few years. In 2016, Qin and Tang *et al.* reported Cu(I)-catalyzed amino- $\pi$  click polymerization for the preparation of nitrogen-containing polymers (Fig. 8a).<sup>93</sup> The polymerizations of diamines and diynes were performed in bulk under the catalyst of CuI in a regioselective and stereoselective manner, producing *Z*-stereoregular polyenamines with high molecular weights (up to  $13.5 \times 10^3$  g mol<sup>-1</sup>) in high yields. In the subsequent year, a simple and powerful spontaneous amino- $\pi$  click polymerization reaction was successfully developed for the first time by them (Fig. 8b). Poly( $\beta$ -aminoacrylate)s with high molecular weights ( $M_w$  up to  $64.4 \times 10^3$  g mol<sup>-1</sup>) and excellent yields could be produced by this polymerization reaction under mild conditions, and the resultant polymers possessed good solubility and high thermal stability. Furthermore, the polymers with tetraphenylethylene (TPE) moieties in the main chains were AIE-active and could be used in the sensing of explosives and specific lysosome labeling, indicating that this click polymerization could be widely applied in diverse areas.<sup>94</sup>

Anti-Markovnikov additive poly( $\beta$ -aminoacrylate)s with *E*-isomer contents of 100% were afforded in excellent yields after stirring the mixtures of dipropiolates and diamines in DCM for 3 h at room temperature. Taking advantage of its high efficiency and fantastic spontaneity, this amino- $\pi$  click polymerization reaction was successfully developed into a fancy intracellular polymerization reaction by the use of acryloyl-tyrosine instead of propiolate as the monomer.<sup>95</sup> Interestingly, turn-on fluorescent imaging and *in situ* killing of cells could be realized by this *in vivo* amino- $\pi$  polymerization reaction. In spite of its several advantages, such propiolate-based spontaneous amino- $\pi$  click polymerization still has some shortcomings. For example, only aliphatic secondary diamines could polymerize with propiolate derivatives in a regio- and stereospecific manner. The polymerization lost its stereospecificity or even failed when aliphatic primary diamines or aromatic diamines were adopted to polymerize with propiolate derivatives, which limits its applications and promotes the development of new spontaneous amino- $\pi$  click polymerization. Considering the fact that the reactivity of ethynyl groups could be enhanced by increasing the electron-withdrawing ability of the adjacent groups, Qin and Tang *et al.* designed and synthesized highly active ethynylsulfone monomers in which

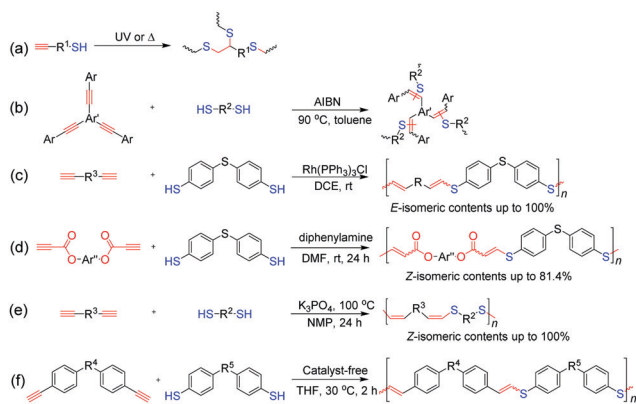


Fig. 7 Thiol- $\pi$  click polymerizations: (a) photo-/thermo-initiated thiol- $\pi$  click polymerization, (b) thermo-initiated thiol- $\pi$  click polymerization based on disubstituted aromatic alkynes, (c) transition-metal-catalyzed thiol- $\pi$  click polymerization, (d) organobase-mediated thiol- $\pi$  click polymerization, (e) K<sub>3</sub>PO<sub>4</sub>-mediated thiol- $\pi$  click polymerization, and (f) spontaneous thiol- $\pi$  click polymerization.

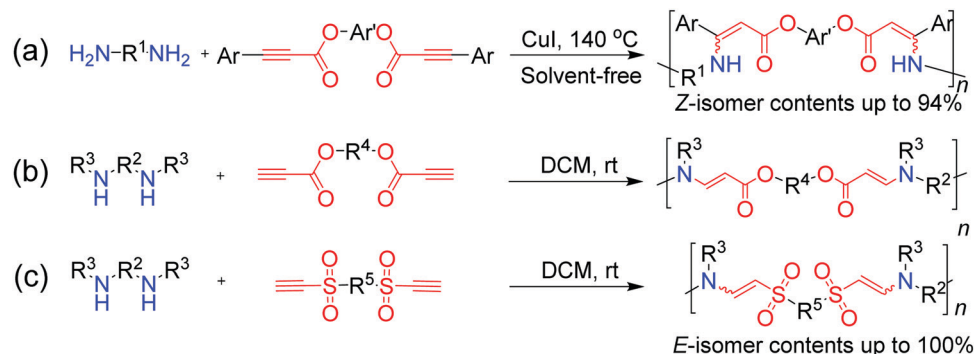


Fig. 8 Amino-yne click polymerizations: (a) Cu(I)-catalyzed amino-yne click polymerization, (b) propiolate-based spontaneous amino-yne click polymerization, and (c) ethynylsulfone-based spontaneous amino-yne click polymerization.

the ethynyl groups were connected with stronger sulfonyl groups instead of ester groups.<sup>96</sup> By the meticulous optimization of polymerization conditions, an ethynylsulfone-based spontaneous amino-yne click polymerization process was developed and regio-/stereoregular poly( $\beta$ -aminovinylsulfone)s (PAVSS) could be obtained by polymerizing ethynylsulfones with various amines including aliphatic/aromatic primary/secondary ones (Fig. 8c). Because of the strong electron-withdrawing ability of sulfonyl groups, a dynamic property of these PAVSS could be observed, which endows them with degradability *via* an amine-exchange process.

To further expand the structures of polymers based on triple-bond building blocks, hydroxyl-yne click polymerizations have also been developed by Qin and Tang *et al.* in recent years.<sup>97</sup> As shown in Fig. 9a, bis(aroylacetylene) and diphenol were polymerized in the presence of the Lewis base of 4-dimethylamino-pyridine (DMAP) in THF at room temperature, affording anti-Markovnikov additive poly(vinylene ether ketone)s (PVEKs) with high molecular weights (up to  $35.2 \times 10^3 \text{ g mol}^{-1}$ ) and *E*-isomer contents as high as 100% at higher yields. Subsequently, regiospecific hydroxyl-yne click polymerization was exploited by the use of superbases *t*-BuP<sub>4</sub> as the catalyst for the preparation of poly(vinyl ether)s (PVEs) (Fig. 9b).<sup>98</sup> Both PVEKs and PVEs containing abundant acid-labile vinyl ether sequences in the polymer backbones exhibit degradability under strong acid conditions, suggesting the promising potential of hydroxyl-yne click polymerizations in the biomedical field.

In short, the recent progress made in alkyne-based click polymerizations, including azide-alkyne, thiol-yne, amino-yne, and hydroxyl-yne click polymerizations and their subtypes, have been addressed. Moreover, the concept, mechanism, and design strategies of AIE-active polymers have also been introduced.

All the polymerization methods and reactions summarized by us may provide researchers and engineers with additional approaches to fabricate desirable advanced functional polymers.

### 2.3 MCP based on triple-bond building blocks

MCP, a new strategy in polymer science, provides a series of advantages, such as easy preparation of monomers, mild reaction conditions, controllable polymer structures, and atom economy. It is well known that both isocyanide and alkyne are active chemical compounds with triple bonds. Isocyanide-based MCPs have been used to synthesize functional polyamides, namely, poly(ester amide)s, polyoxazolines, polyesters, and hyper-branched polymers (HBPs). Meanwhile, alkyne-based MCPs have also been recently reported by polymerizing diynes-primary amines-aldehydes and diynes-disulfonylazide-amino esters. Dialkyl acetylenedicarboxylate (DAAD) is a commercial chemical and has been widely used in cycloaddition reactions. Isocyanides can attack DAAD to afford zwitterionic species, which are readily trapped by carbon electrophiles and then converted into a pyrrole, furan, ketenimine, dihydropyridine, *etc.*

Many limitations, such as narrow monomer scope, low conversion, and more possible side reactions, have hindered the development of MCP. The Cu(I)-catalyzed multicomponent reactions (MCRs) based on sulfonyl azide, alkyne, and third nucleophiles inherit the advantages of Cu(I)-catalyzed azide-alkyne cycloaddition and exhibit promising potential in polymer synthesis. Based on these reactions, Choi *et al.* were the first to develop the new click-reaction-based MCPs of diynes, sulfonyl azides, and diamines/diols.<sup>99,100</sup> These polymerizations perfectly adapted to the “graft through” strategy: a series of graft and dendronized polymers with huge bulky side chains were afforded with excellent results by the use of



Fig. 9 Hydroxyl-yne click polymerizations: (a) DMAP-catalyzed aromatic hydroxyl (phenol)-yne click polymerization and (b) superbases *t*-BuP<sub>4</sub>-catalyzed aliphatic hydroxyl-yne click polymerization.

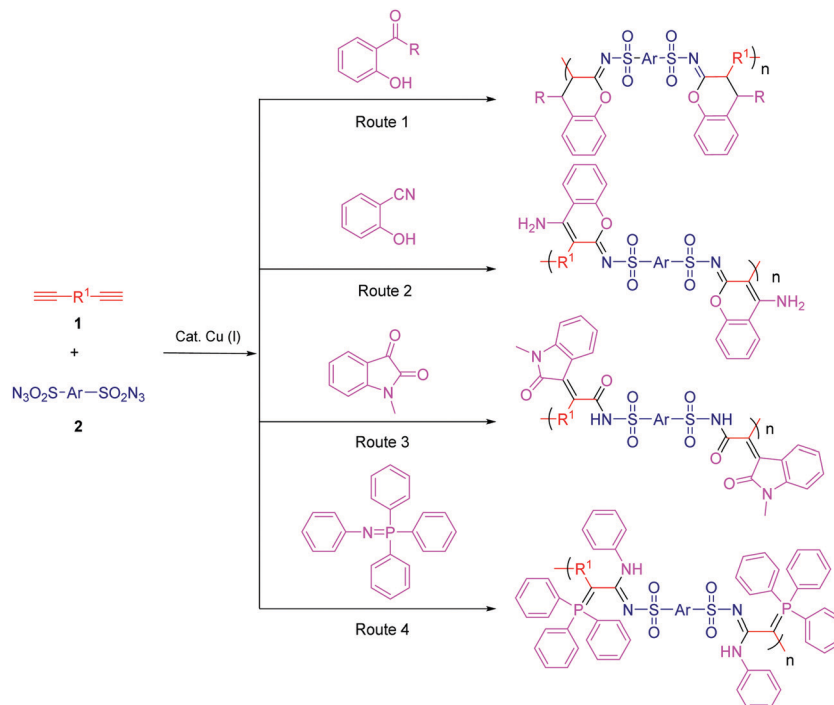


Fig. 10 Copper(I)-catalyzed click-polymerization-based MCPs of diynes, disulfonyl azides, and third components.

monofunctional alkyne macromolecules.<sup>101</sup> Tang *et al.* reported the facile and efficient click-polymerization-based MCP of diynes, disulfonyl azide, and salicylaldehyde, or *o*-hydroxyacetophenone catalyzed by Cu(I) and trimethylamine at room temperature (Fig. 10, Route 1). This polymerization approach inherits the remarkable merits of both MCRs and click reactions, such as simple operation, high reaction efficiency and isolation yields, mild reaction conditions, and common substrates. The resultant polymers with high molecular weights ( $M_w$  up to  $64.6 \times 10^3 \text{ g mol}^{-1}$ ) possessed outstanding film-forming ability, high thermal stability, and good morphological stability. The obtained polymers with bright film emission and high photosensitivity could be facilely fabricated into well-resolved 2D and 3D patterns by treating their films with UV light.<sup>102</sup>

Based on the strategy of changing the third components, various click-reaction-based MCPs have been developed to fabricate functional polymers with complex structures. Hydroxybenzotrile underwent a similar reaction mechanism to yield fused heterocycles-containing polymers (Fig. 10, Route 2).<sup>103</sup> Because of the presence of multiple heteroatoms and amino groups in the polymer chains, the resultant polymers enjoyed antibacterial properties and potential sensitivity and selectivity of  $\text{Ru}^{3+}$  detection. In addition, *N*-protected isatins can act as a nucleophile for attacking the highly reactive intermediate to yield oxindole-containing poly(*N*-acylsulfonamide)s (Fig. 10, Route 3). These polymers enjoyed reversibly tunable hydrophilicity by undergoing structural changes when using LiOH and HCl.<sup>104</sup> Heteroatoms can be conveniently introduced into the polymer chains *via* the third component. Iminophosphorane was utilized as the third monomer to develop a new MCP involving diynes and disulfonyl azides in order to fabricate N-, O-, S-, and P-containing

heteroatom-rich polymers (Fig. 10, Route 4).<sup>105</sup> The obtained polymers exhibited good solubility in polar solvents, good film-forming ability, and high refractivity. The possible interaction between the heteroatom and metal ion endowed the polymer with excellent selectivity in  $\text{Pd}^{2+}$  fluorescence detection. Moreover, this powerful MCP provides straightforward access to the synthesis of heteroatom-rich HBPs by the use of multifunctional alkynes and disulfonyl azides.<sup>106</sup>

When compared with five- or six-membered heterocycles, the *in situ* generation of small-ring heterocycles in polymer chains has encountered a formidable challenge due to difficulties during synthesis and inherent low stabilities. By replacing *o*-hydroxyacetophenone with carbodiimides, Tang *et al.* also developed a novel three-component polymerization route for fabricating functional polymers bearing heteroatom-rich multi-substituted azetidine rings (Fig. 11). This synthesis strategy exhibits advantages such as readily available monomers, simple operation, high efficiency, mild condition, environmental friendliness, and high atom economy. The strained rings in the polymer main chains enable them to efficiently undergo ring-opening post-modification under simple acidic conditions to transform into new polymers with amide and amidine units. It is noteworthy that the obtained azetidine-containing polymers showed obvious clusteroluminescence characteristic because the clustering of heteroatoms and conformational-rigidification-induced formation of efficient through-space conjugation played an important role in such an unconventional luminescence phenomenon.<sup>107</sup>

More recently, Tang *et al.* reported an efficient MCP strategy toward chalcogen-rich multifunctional poly(vinyl sulfone)s with regio-/stereoselectivity in high yields. This polymerization

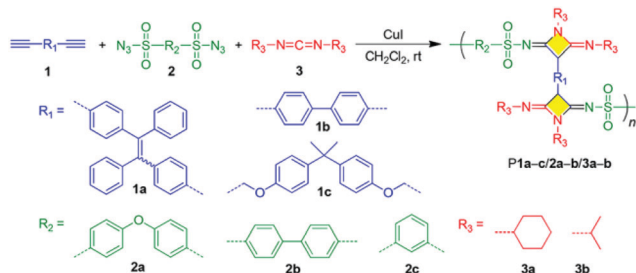


Fig. 11 Three-component polymerization using the small heterocycles route. Adapted from ref. 107 with permission. Copyright (2018) American Chemical Society.

proceeded smoothly at room temperature under the protection of nitrogen and could yield functional poly(vinyl sulfone)s that could serve as effective fluorescent bioprobes for use in cellular imaging.<sup>108</sup> Another significant work is the preparation of a series of multifunctional oxazine-containing polyheterocycles with AIE characteristics, which are generated *in situ* by metal-free one-pot A<sup>3</sup> polymerizations of terminal diynes, dialdehydes, and ureas. These obtained polymers exhibited outstanding fluorescence response to protonation and deprotonation. Based on this, a fast-response reversible fluorescent sensor for ammonia can be used to detect biogenic amines and seafood spoilage.<sup>109</sup>

In 2014, Tang *et al.* explored the first multicomponent tandem polymerizations (MCTPs) for the synthesis of conjugated polymers (CPs), which could yield functional CP materials by the three-component reaction of alkyne, acryloyl chloride, and ethyl 2-mercaptoacetate (Fig. 12, Route 1).<sup>110</sup> After that, by combining the Sonogashira coupling reaction and other reactions, a series of one-pot, two-step, three-component coupling–addition tandem reactions and polymerizations were developed by them. For example, by utilizing the Sonogashira coupling reaction and the hydrothiolation reaction of electron-deficient alkynone intermediates, the multicomponent tandem reactions (MCTRs) and the corresponding MCTPs of alkynes, carbonyl chlorides, and aliphatic/aromatic thiols were investigated,

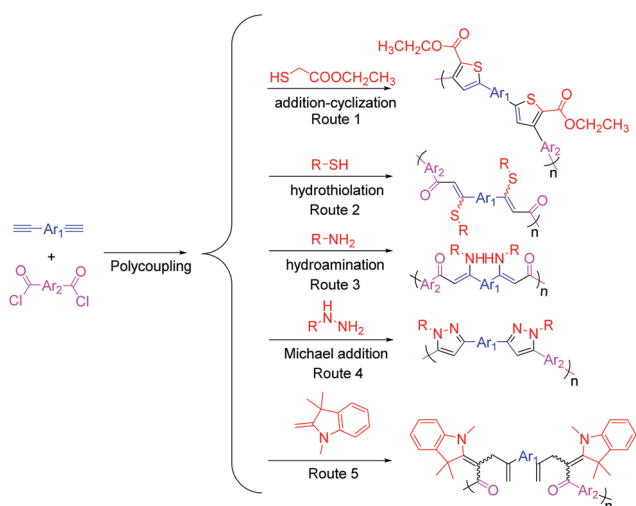


Fig. 12 MCTPs of alkynes, carbonyl chlorides, and third components.

which could afford sulfur-rich CPs with high molecular weights ( $M_w$  up to  $59.1 \times 10^3 \text{ g mol}^{-1}$ ), high regioselectivity, and good stereoregularity in high yields (Fig. 12, Route 2).<sup>111</sup> Through the one-pot sequential Sonogashira coupling of alkyne and carbonyl chloride, as well as the hydroamination of the internal alkyne, nitrogen-containing CPs with high regioselectivity and stereoselectivity could be obtained by the efficient MCTPs of alkynes, carbonyl chlorides, and primary amines (Fig. 12, Route 3).<sup>112</sup> Heterocycle-containing conjugated polypyrazoles were also synthesized by the efficient MCTPs of alkyne, carbonyl chloride, and hydrazines/aromatic diynes through the combination of Sonogashira coupling–Michael addition–cyclocondensation reactions in a one-pot procedure (Fig. 12, Route 4).<sup>113</sup> Furthermore, the one-pot MCTPs of alkynes, carbonyl chloride, and Fischer's base were developed to afford conjugated poly(diene-merocyanine)s with satisfactory molecular weights ( $M_w$  up to  $10.9 \times 10^3 \text{ g mol}^{-1}$ ) and higher yields (up to 81%) (Fig. 12, Route 5).<sup>114</sup> The MCTPs of *N*-(2-iodophenyl)-3-phenyl-*N*-tosylpropiolamide, aromatic terminal alkynes, and diamines were explored to afford poly(indolone)s with unique structures and high molecular weights ( $M_w$  up to  $30.4 \times 10^3 \text{ g mol}^{-1}$ ) in high yields.<sup>115</sup> In addition, by combining Glaser coupling–nucleophilic addition–heterocyclization–oxidation reactions, Hu and Tang *et al.* reported the four-step MCTPs of diyne, guanidine hydrochloride, DMSO, and  $O_2$ , affording conjugated poly(pyrimidine)s with well-defined and tunable structures.<sup>116</sup> These works have demonstrated the promising potential of MCTPs: various reactions with compatible conditions could be integrated into a single polymerization route and could afford polymers with diverse structures, which considerably extends the scope of such polymer structures.

Moreover, Hu and Tang *et al.* also developed the facile metal-free one-pot MCTPs of activated internal alkynes, aromatic diamines, and formaldehyde with high efficiency and enhanced convenience, which enabled the facile preparation of polyheterocycles with well-defined structures, regulated sequence, high molecular weights ( $M_w$  up to  $69.8 \times 10^3 \text{ g mol}^{-1}$ ), and high yields (up to 99%) (Fig. 13).<sup>117</sup> In these MCTPs, functional heterocycles were directly constructed from the polymerization of commercially available, inexpensive, simple monomers; further, the resultant polymer structures were inaccessible by other synthesis approaches. Moreover, disubstituted internal alkynes used as monomers could considerably enrich the monomer variety, product diversity, and functionalities.<sup>117</sup> Based on this, they constructed four HBPs with different topological structures having unique advantages and strong designability of these MCTPs; this provides a synthesis platform for the preparation of hyperbranched polyheterocycles with diverse functionalities.<sup>118</sup>

Inspired by the small molecule reaction reported by other groups and experiences in the triple-bond chemistry field, Dong *et al.* developed four new methods for preparing heterocyclic-containing functional polymers based on the multicomponent cyclopolymerization (MCC) of isocyanides, DAAD, and third component in the presence of a metal-free catalyst, which have been committed toward the development of new functional materials based on triple-bond reactions.<sup>119,120</sup> Indeed, a series of heterocyclic-containing functional polymers

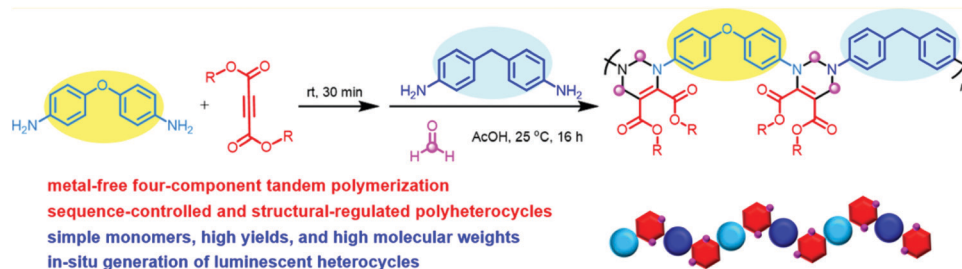


Fig. 13 Metal-free MCTPs of alkynes, aromatic diamines, and formaldehyde. Adapted from ref. 117 with permission. Copyright (2017) American Chemical Society.

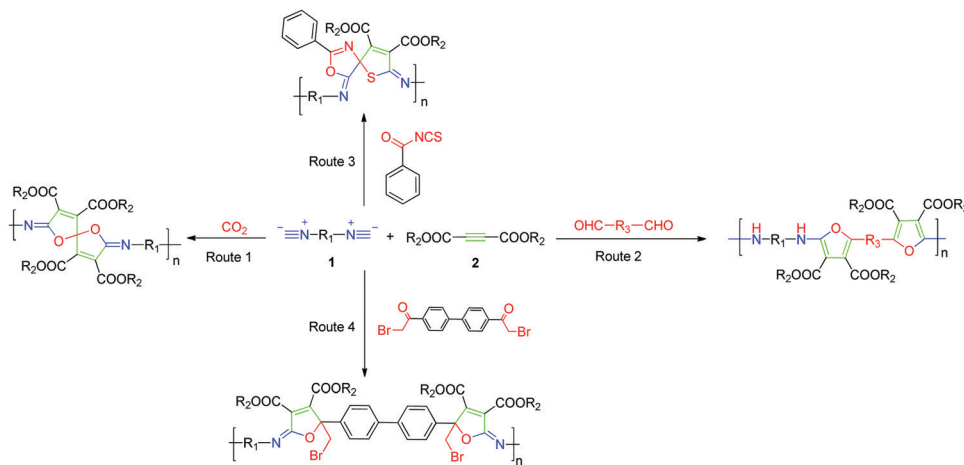


Fig. 14 MCCs based on isocyanides, DAAD, and third component.

with good solubility and thermal stability were successfully prepared in high yields (Fig. 14). Therefore, MCC may be a powerful tool for fabricating new types of multifunctional polymers.

First, catalyst-free one-pot multicomponent spiropolymerization (MCS) was developed for spiropolymer synthesis by using diisocyanides and alkynes in the presence of  $\text{CO}_2$  as the monomer (Fig. 14, Route 1). The target spiropolymers contained 1,6-dioxospiro[4,4]nonane-3,8-diene as the spirostructural unit in the main chain with different types of spacer groups.<sup>121</sup> As a kind of atom-economical polymerization process, this new MCS reaction can serve as the beginning of spiropolymer preparation and can have wide potential applications in materials science. This work inspired the use of dialdehydes instead of carbon dioxide to investigate the possibility of multicomponent cyclopolymerization (Fig. 14, Route 2). This approach afforded poly(amine-furan-arylene)s (PAFAs) with high molecular weights ( $M_w$  up to  $76.4 \times 10^3 \text{ g mol}^{-1}$ ), comprising 3,4-substituted furan that was formed *in situ* during the polymerization process, with good thermal stability and film-processing properties. The single crystal of the model compound (MC) indicated that the carbonyl group on the furan ring and the adjacent amino group had an intramolecular hydrogen bond to enhance the conjugation of the main chain, which exhibited the characteristic of black materials that have high absorption property and high thermal stability.<sup>122</sup> Recently, they extended this system to additional aromatic dialdehyde monomers,

including those with other heterocyclic spacers (e.g., pyrimidine, quinoxaline, pyridine, and thiophene). A series of poly(furan-amine)s (PFAs) based on these aromatic heterocyclic dialdehydes were successfully prepared under catalyst-free polymerization conditions. Interestingly, these PFAs can be degraded into volatile products with low molecular weights under UV irradiation. Therefore, this MCC process can yield useful photodegradable materials, which may offer a new platform for the design of multifunctional PAFs.<sup>123</sup>

Second, a catalyst-free MCS route for the facile construction of heteroatom-containing spiropolymers has been developed by Dong *et al.* (Fig. 14, Route 3). Here, 4,7-bis[alkyl(aryl)imino]-2-phenyl-3-oxa-6-thia-1-azaspiro[4.4]nona-1,8-dienes as a structural unit was instantly formed by the conversion of benzoyl isothiocyanate into spirosegments. With respect to monomer versatility, procedure simplicity, and high atom economy of the spiropolymerization process, this catalyst-free catalyzed MCS process opens up newer opportunities for the preparation of advanced spiropolymer materials. They further replaced benzoyl isothiocyanate with 4,4'-bis(2-bromoacetyl)biphenyl and constructed polymers containing iminolactone arylene (PIAs) by using catalyst-free MCP (Fig. 14, Route 4). The results of this analysis indicated that PIAs exhibited good solubility, aggregation-enhanced emission (AEE) properties, and high refractive indices under visible light (400–800 nm). Moreover, in the presence of bromomethyl groups in the side chain, PIAs

were endowed with better performance because of post-functionalization.<sup>124</sup>

The utilization of sulfur remains a challenge for the efficient and environmentally friendly transformation of byproducts into commercial offerings based on their abundant storage but limited consumption in the petroleum industry. Polysulfide anions with high nucleophile content generated from elemental sulfur and aliphatic amines can become involved in attacks on both alkynes and isocyanides to obtain sulfur-based functional groups. Hu and Tang *et al.* reported the efficient and metal-free MCPs of elemental sulfur, diynes, and aliphatic diamines in pyridine at 100 °C to obtain well-defined polythioamides with high molecular weights ( $M_w$  up to  $127.9 \times 10^3 \text{ g mol}^{-1}$ ) (Fig. 15, Route 1).<sup>125</sup> The resulting polythioamides exhibited unconventional luminescence behavior without any typical fluorophores, which can be attributed to the presence of “heterodox clusters” formed with numerous lone-pair-containing electron-rich heteroatoms. With the presence of sulfur in polymer chains, they possessed high refractivity index of 1.80–1.66 in a wide wavelength ranging from 400 to 1700 nm.

Polythioureas have attracted considerable attention in many fields due to their special properties such as self-healing characteristic, excellent electrical performance, and metal removal ability. Inspired by the polymerization reaction discussed earlier, isocyanides were utilized to replace alkynes as monomers to develop another new element, namely, sulfur-based auto-catalyzed MCP comprising diisocyanides and aliphatic diamines for the synthesis of polythioureas (Fig. 15, Route 2).<sup>126</sup> The analysis based on *in situ* IR technique indicated that the high reactivity of isocyanide endowed the polymerization with high efficiency and quick reaction at both ambient temperature and 100 °C, affording a series of high-molecular-weight polythioureas ( $M_w$  up to  $242.5 \times 10^3 \text{ g mol}^{-1}$ ) with well-defined structures. Due to the high affinity and selectivity of thiourea to mercury(II), the obtained polythioureas exhibited high sensitivity and selectivity in mercury fluorescence detection and excellent removal efficiency (>99.99%) toward  $\text{Hg}^{2+}$  pollution, which implied that these sulfur-based materials have feasible potential applications in the recovery of heavy metals.

In summary, several types of MCPs based on triple-bond building blocks by Tang and Dong *et al.* are briefly introduced, including Cu(I)-catalyzed MCPs, MCTPs, MCCs, MCPs based on elemental sulfur, *etc.* All these MCPs may supply many useful tools for designing and fabricating advanced functional polymers.

#### 2.4 ROP of biorenewable poly( $\gamma$ -butyrolactone) (P $\gamma$ BL)

Preparation methods for olefin and alkyne polymers, which are nonrenewable and nondegradable, have been addressed

earlier. In this subsection, the ROP of renewable monomers is addressed. Aliphatic polyesters have become a class of important biomaterials that have been widely used as drug delivery carriers, tissue engineering scaffolds, and food packaging materials because of their promising biocompatibility and biodegradability. In particular, the ROP of cyclic esters or lactones has been applied in the preparation of polyesters with well-defined structures on a large scale.<sup>127</sup> Recently, P $\gamma$ BL has attracted considerable interest since  $\gamma$ -butyrolactone ( $\gamma$ -BL) can be obtained from succinic acid, which is the most promising biomass-derived compound to replace petroleum-derived chemicals.<sup>128,129</sup> On the other hand, poly(4-hydroxybutyrate) (P4HB), chemically identical to P $\gamma$ BL, exhibits a suitable degradation rate *in vivo* and can avoid the undesirable accumulation of acidic degradation products as compared to other aliphatic polyesters. Remarkably, P $\gamma$ BL can be completely depolymerized back to its monomeric form of  $\gamma$ -BL by simple heating. Therefore, P $\gamma$ BL is a truly sustainable polymer, which is fully biorenewable and completely recyclable.<sup>130</sup>

However,  $\gamma$ -BL has been ignored as a monomer in polymer science for a long time due to the presence of a five-membered ring with low strain energy and related unfavorable thermodynamics involved in the ROP process. In 2016, Hong and Chen *et al.* demonstrated a milestone work and successfully prepared high-molecular-weight P $\gamma$ BL *via* ROP at lower temperatures.<sup>130</sup> P $\gamma$ BL with high molecular weight (up to  $30.0 \times 10^3 \text{ g mol}^{-1}$ ) and high monomer conversion (up to 90%) could be fabricated for the first time by utilizing  $\text{Ln}[\text{N}(\text{SiMe}_3)_2]_3$  or yttrium-amide-catalyzed coordination ROP at  $-40$  °C (Fig. 16a). The resulting P $\gamma$ BL possessed either linear or cyclic topology (Fig. 16b) depending on the catalyst/initiator nature and ratio. Subsequently, they demonstrated that the organic superbase of 1-*tert*-butyl-4,4,4-tris(dimethylamino)-2,2-bis[tris(dimethylamino)-phosphoranylideneamino]-2 $\lambda^5$ ,4 $\lambda^5$ -catenadi (*t*-BuP $_4$ , Fig. 16a) can also catalyze the ROP of  $\gamma$ -BL at lower temperatures.<sup>131</sup> In this case, the superbase *t*-BuP $_4$  can abstract the  $\alpha$ -proton of the monomer to generate the reactive enolate species and therefore directly initiate the ROP of  $\gamma$ -BL. For example, conversion of 30.4% after 12 h could be achieved, and a P $\gamma$ BL sample with  $M_n$  of  $26.4 \text{ kg mol}^{-1}$  and  $D$  ( $M_w/M_n$ ) of 1.79 was obtained when using *t*-BuP $_4$  alone (1.0 mol% loading) in toluene at  $-40$  °C. Moreover, a combination of alcohol with the initiator formed the ion pair [*t*-BuP $_4\text{H}^+ \cdots \text{OBn}^-$ ] and led to an even more effective ROP system, as evidenced by the high monomer conversion (up to 90%) and high-molecular-weight P $\gamma$ BL ( $M_n$  up to  $26.7 \text{ kg mol}^{-1}$ ). Recently, Zhang *et al.* reported the dual organocatalyst *t*-BuP $_4$ / (thio)urea for the ROP of  $\gamma$ -BL with increased activity and enhanced selectivity in comparison to those obtained when using *t*-BuP $_4$  alone.<sup>132</sup> The turnover frequencies (TOFs) of these

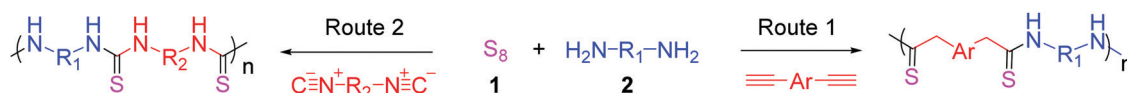


Fig. 15 MCPs based on elemental sulfur, aliphatic diamines, and diynes/isocyanide. Reproduced from ref. 125 and 126 with permission. Copyright (2015 and 2018) from American Chemical Society.

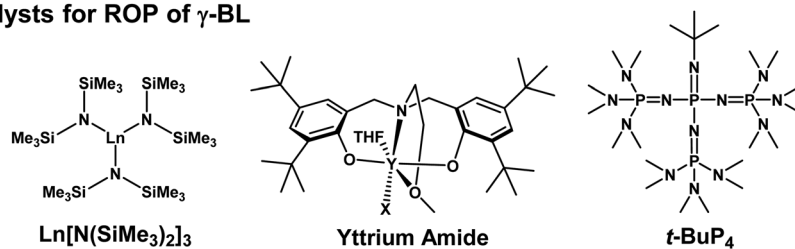
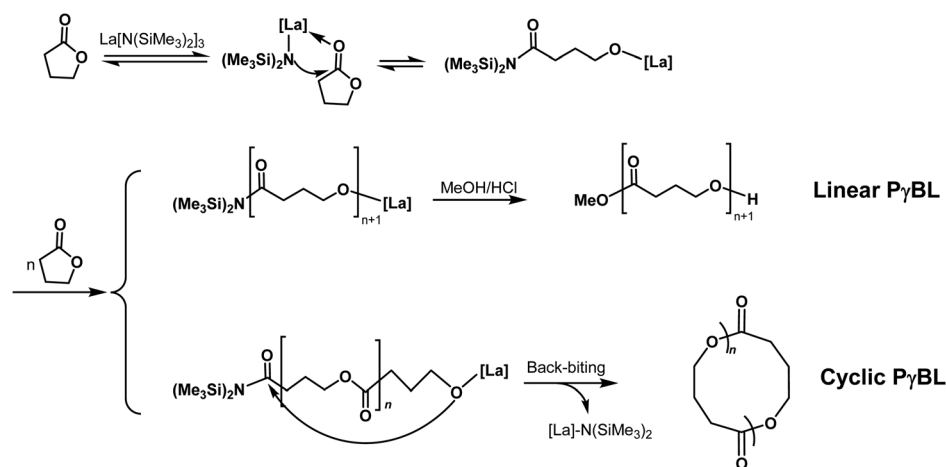
(a) Catalysts for ROP of  $\gamma$ -BL(b) Proposed mechanism for ROP of  $\gamma$ -BL by  $\text{La}[\text{N}(\text{SiMe}_3)_2]_3$ 

Fig. 16 (a) Metal catalysts and organocatalysts for the ROP of  $\gamma$ -BL; (b) proposed mechanism for the ROP of  $\gamma$ -BL by  $\text{La}[\text{N}(\text{SiMe}_3)_2]_3$ . Reproduced from ref. 130 and 131 with permission. Copyright (2016) from Nature Publishing Group.

$t\text{-BuP}_4$ /(thio)urea pair-catalyzed processes were as high as  $125 \text{ h}^{-1}$ , while the resultant P $\gamma$ BLs had molecular weights as high as  $64.3 \text{ kg mol}^{-1}$ . Besides the homopolymerization of  $\gamma$ -BL, Hong and Chen *et al.* also reported the effective copolymerization of  $\gamma$ -BL with  $\epsilon$ -CL or  $\delta$ -VL using  $t\text{-BuP}_4$  as the catalyst.<sup>133</sup> The copolymerization produced random copolyesters with high molecular weights (up to  $135 \text{ kg mol}^{-1}$ ) as well as an unprecedented level of  $\gamma$ -BL incorporation (up to 84.0 mol%). The successful synthesis of these copolyesters with high  $\gamma$ -BL incorporations of > 50 mol% led to the discovery of the eutectic phase of the  $\gamma$ -BL/ $\epsilon$ -CL copolymer with a eutectic temperature,  $T_{\text{eu}}$ , of  $11.0 \text{ }^\circ\text{C}$  and eutectic composition,  $X_{\text{eu}}$ , of 66.0 mol%  $\gamma$ -BL; therefore, the copolyester with this composition was a viscous liquid at room temperature, although the two constituent homopolyesters were semicrystalline solids.

On the other hand, Li *et al.* reported that the cyclic trimeric phosphazene base (CTPB, Fig. 17a) was a highly efficient organocatalyst for the ROP of  $\gamma$ -BL and can be used to prepare well-defined PBL with high conversions (up to 98%) below  $-50 \text{ }^\circ\text{C}$  within 4 h.<sup>134</sup> The resulting P $\gamma$ BLs have high molecular weights (up to  $22.9 \text{ kg mol}^{-1}$ ) and relatively narrow mass distributions ( $D = 1.27\text{--}1.50$ ). In contrast to  $t\text{-BuP}_4$ , CTPB alone could not activate the monomer and therefore showed no activity in the absence of alcohol, as indicated by the *in situ* NMR analyses. The structural analyses of a low-molecular-weight polymer obtained in the presence of BnOH by MALDI-TOF

suggested that the linear polymers with BnO/H chain ends were the primary products. More recently, Li *et al.* successfully synthesized a series of PBL-*b*-poly(*L*-lactide) (PLLA) diblock copolyesters with different compositions *via* a one-pot method by the sequential ROP of  $\gamma$ -BL and *L*-LA in the presence of CTPB as the catalyst (Fig. 17c).<sup>135</sup> Although the first ROP of  $\gamma$ -BL could not consume all the monomers, the unreacted  $\gamma$ -BL could not homopolymerize or copolymerize with *L*-LA after the addition of the *L*-LA/THF solution. The PBL-*b*-PLLA copolyesters showed improved thermal stability as compared to the PBL homopolymer according to the thermogravimetric analysis (TGA) results. For example, the TGA curve of P $\gamma$ BL<sub>53</sub>-*b*-PLLA<sub>97</sub> displayed a one-step decomposition profile with  $T_{\text{d},5\%}$  value of  $326 \text{ }^\circ\text{C}$  and  $T_{\text{d,max}}$  value of  $368 \text{ }^\circ\text{C}$ , both of which are higher as compared to those of the P $\gamma$ BL homopolymer ( $T_{\text{d},5\%}$  of  $273 \text{ }^\circ\text{C}$  and  $T_{\text{d,max}}$  of  $334 \text{ }^\circ\text{C}$ ). Differential scanning calorimetry (DSC) results revealed the microphase separation in the P $\gamma$ BL-*b*-PLLA block copolymer (BCP), in which two separate melting transitions were present. Amphiphilic copolymers with P $\gamma$ BL as the hydrophobic segment were also successfully prepared by the use of PEG as the macroinitiator and CTPB or sodium hydride as the catalyst (Fig. 17d).<sup>136</sup> Both catalytic/initiating systems showed moderate control over the ROP of  $\gamma$ -BL and successfully produced PEG-*b*-P $\gamma$ BL diblock copolymers with varied molecular weights. A suitable activation temperature ( $-40 \text{ }^\circ\text{C}$ ) was crucial to obtain moderate to high  $\gamma$ -BL conversions and satisfactory molecular

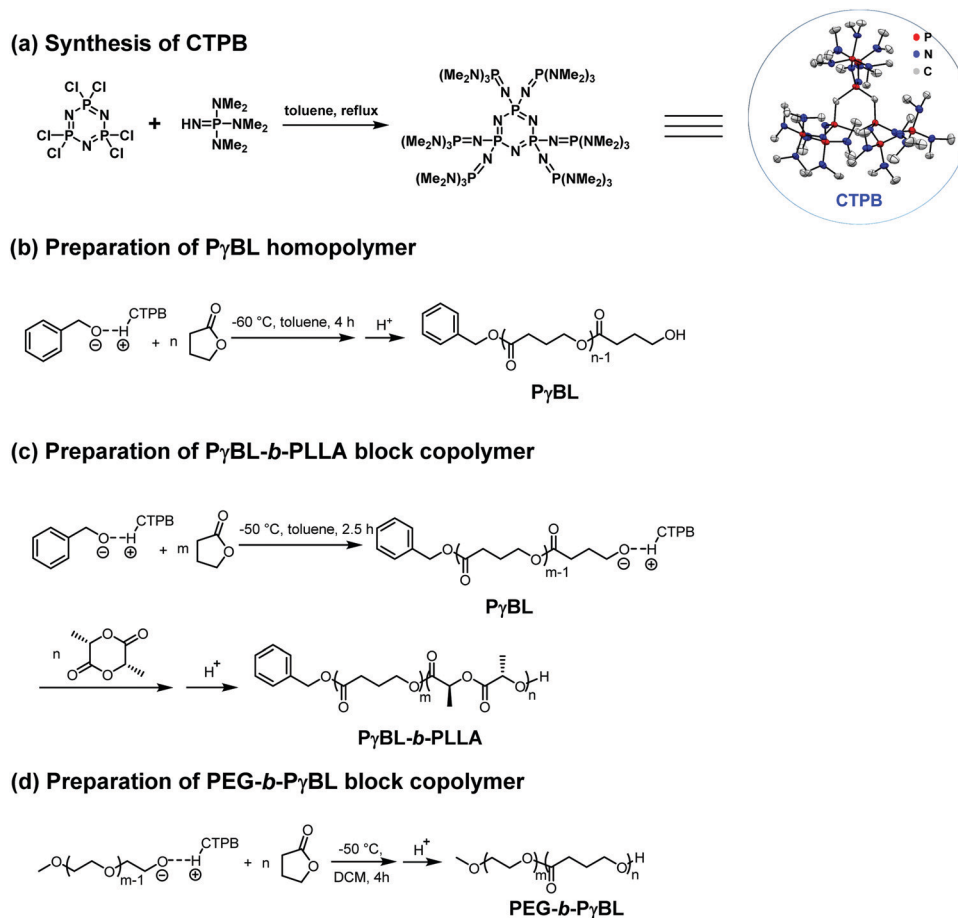


Fig. 17 (a) Synthesis of CTPB and using it for the preparation of P $\gamma$ BL-containing (co)polymers (b–d). Reproduced from ref. 134–136 with permission. Copyright (2016, 2017, and 2018) from Wiley-VCH, Royal Society of Chemistry, and American Chemical Society.

weight of PEG-*b*-P $\gamma$ BL with CTPB as the catalyst. The self-assembly behavior of the obtained amphiphilic PEG-*b*-P $\gamma$ BL diblock copolymer was preliminarily investigated with dynamic light scattering (DLS) and transmission electron microscopy (TEM) modalities. The TEM results revealed that the amphiphilic PEG-*b*-P $\gamma$ BL copolymer could self-assemble into micelles and vesicles in water, making it to be a promising candidate as a drug delivery carrier. Given the advantages of P $\gamma$ BL mentioned above, amphiphilic PEG-*b*-P $\gamma$ BL is expected to exhibit superior performances as compared to other existing amphiphilic aliphatic polyesters in biomedical applications.

Although comprehensive studies have been conducted in the past few years, the development of new catalysts having both high activity and selectivity for the ROP of  $\gamma$ -BL remains a challenging subject. Dual catalytic systems would be a valuable alternative. The construction of P $\gamma$ BL-containing copolymers with complex but well-defined monomer sequences and varying topologies, *e.g.*, graft or comb-like copolymers, should be paid more attention.

In this section, the developed preparation methods and polymerization reactions of functional polymer materials in the past few years are addressed, including CRP, alkyne-based click polymerization, MCP based on triple-bond building blocks,

and ROP. Although new progresses in the preparation methods have been achieved in the past few years, particularly with regard to sequence-controlled polymerization and MCP, there are still a number of challenges in this field, such as building relationships between the polymer structures and functions, realizing controllable synthesis and function integration, and tailoring and modifying functional polymers. Therefore, more efforts should be directed toward the development of synthesis methodologies, allowing us to prepare more varied functional polymer materials, improve their performances, and reduce their costs of preparation.

### 3. Luminescent polymer materials

Luminescent materials are capable of converting other energy forms into light energy and subsequent light emission. They can be classified into three types: inorganics, organic molecules, and polymeric materials. Luminescent polymers can be classified into two subtypes based on the luminescent principle: fluorescent and phosphorescent polymers. According to the spin statistical distribution, only 25% of singlet (S) excitons and 75% of triplet (T) excitons are generated when a molecule in the ground

state ( $S_0$ ) is excited by external light or electricity. For fluorescent polymers, only S excitons can be utilized, while T excitons can only be dissipated as thermal energy due to their spin-forbidden characteristic. Phosphorescent polymers can use the spin-forbidden triplet state under the action of the rotating orbital coupling of heavy metal atoms. Therefore, 100% internal quantum efficiency (IQE) can be achieved. Although many methods have been used to improve the device efficiency of fluorescent polymers, all of them are based on the use of T excitons. The most popular one is that the T excitons are converted into S excitons by an upconversion process, generating fluorescence ( $S_1 \rightarrow S_0$ ). Many theories have been developed in recent years, including thermally activated delayed fluorescence (TADF), hybridized local charge transfer (HLCT), and triplet-triplet annihilation (TTA). Theoretically, 100% IQE can be achieved when using the above methods. The actual device efficiency of fluorescent polymers obtained by the use of the above methods is usually lower than that of phosphorescent polymers. In this section, phosphorescent polymers such as carbonized polymer dots (CPDs) and electrophosphorescent polymers (PhPs) have been discussed. For fluorescent polymers, only AIE polymers are addressed; interested readers can refer to other reviews.<sup>137,138</sup>

### 3.1 AIE polymers

AIE, a groundbreaking concept coined by Tang *et al.* in 2001, has attracted considerable attention because of its attractive applications in diverse areas.<sup>139</sup> Researchers from all over the world have been working in this promising area, and significant progresses have been made. A large variety of AIE molecules with promising structural diversity have been developed, such as TPE, hexaphenylsilole (HPS), distyreneanthracene (DSA), and tetraphenylpyrazine (TPP) (Fig. 18).<sup>140–143</sup>

When compared with AIE-active low-mass molecules, AIE polymers have been less explored.<sup>144</sup> However, with the research of AIE polymers being constantly performed, an increased number of superior performances have been found, which, in turn, promote their development. According to the principles involved in the restriction of intramolecular motion as well as structural rigidification, the intramolecular motion of AIE units in polymer chains can be further restricted in comparison to their solid-state counterparts, leading to stronger emissions. Therefore, when luminogens with AIE characteristics (referred to as “AIEgens”) are involved in polymer chains, steric

hindrance can be enhanced because of the long-chain segment of AIE polymers, hindering the free rotation of the luminogenic units. Evidently, intramolecular motion can be more easily restricted in polymers than that in low-mass molecules, resulting in a relatively more emissive solution state for polymers.<sup>145</sup> Furthermore, AIE polymers possess many advantages over low-mass molecules, such as processability, ease of functionalization, structural diversity, and good thermal stability, facilitating their potential practical applications. Until now, a large number of AIE polymers have been reported, such as nonconjugated linear AIE polymers, conjugated linear AIE polymers, and nonlinear AIE polymers. In addition, the applications of AIE polymers in optoelectronic, fluorescent chemosensor, biological, and other fields have been summarized in some excellent reviews.<sup>146,147</sup>

Although various synthesis strategies have been used to fabricate AIE polymers, all of them have the same core principle, *i.e.*, incorporating AIEgens into polymer structures, such as TPE, HPS, DSA, and TPP. Generally, two methods have been employed to incorporate AIEgens into polymers, *i.e.*, post-functionalization and direct polymerization.<sup>147</sup> The post-functionalization method can transform AIE-inactive polymers into AIE polymers *via* different polymer reactions. For example, AIEgens are usually linked to polymers as pendants or connectors *via* chemical reactions. Jin *et al.* converted the synthesized zwitterionic copolymer into an AIE polymer by grafting TPE molecules to the polymer side chains *via* pH-cleavable hydrazine bonds. The resultant AIE copolymer could self-assemble into spherical micelles and it could swell under acidic conditions after the cleavage of TPE, which may find applications in bioimaging and drug delivery.<sup>148</sup> In addition, the direct polymerization method can also be used to fabricate AIE polymers *via* the polymerization of low-mass molecules. Tang *et al.* designed and prepared a series of AIE-active polymers by polymerizing aliphatic hydroxyl groups and AIEgen-containing diynes. The obtained PVEs exhibit typical AIE characteristics, *i.e.*, stronger emissions in the aggregate state than that in the solution state, which can be used in the detection of explosives.<sup>98,149,150</sup>

Besides fluorescence, room-temperature phosphorescence (RTP) has also been reported in natural compounds and polymers, such as starch, cellulose, and chitosan. These natural macromolecules cannot emit in solutions, but they are capable of emitting bright phosphorescence in the solid states. Therefore, AIE polymers can emit not just fluorescence but also RTP, which is an important parameter for exploring the unknown territory of molecular luminescence.

### 3.2 CPDs

RTP has attracted increasing attention owing to its long-lived luminescence, large Stokes shift, and high signal-to-noise ratio, facilitating its applications in the fields of optoelectronics, photocatalytic reactions, molecular imaging, security aspects, chemical sensors, and so forth.<sup>151,152</sup> Currently available RTP materials suffer from problems such as scarce resources, high cost, toxicity, limited species, and complex synthesis. Therefore, there is an urgent need to develop new classes of metal-free RTP materials.

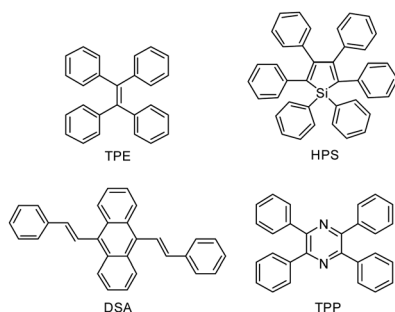


Fig. 18 Chemical structures of typical AIE luminogens.

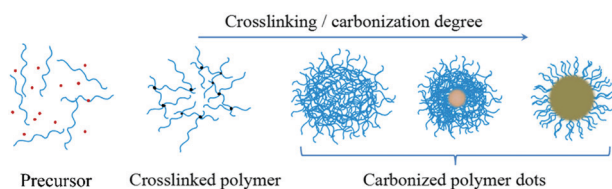


Fig. 19 Schematic illustrations of the formation process and structure of CPDs.

Carbon dots (CDs) have been recognized and proposed as luminescent nanomaterials since 2004. Until now, types of CDs have been continuously increasing, and they can be mainly divided as follows: graphene quantum dots, carbon quantum dots, and CPDs. Emerging CPDs can most likely resolve the abovementioned problems. In fact, CPDs are carbon-based nanodots with obviously polymeric, slightly graphitized, and inaccurate chemical structures. Their interior is a highly cross-linked network structure, and the outer part retains a large number of hydrophilic flexible polymer chains and functional groups (Fig. 19).

The formation of CPDs usually necessitates four processes, namely, dehydration, condensation, crosslinking, and carbonization, which enable them to inherit the molecular properties of their raw materials, as well as the properties of both polymers and quantum dots induced by the polymerization and carbonization reactions. More importantly, the abundant energy levels of CPDs increase the probability of intersystem crossing (ISC), and their covalently crosslinked framework structures considerably suppress nonradiative transitions, exactly following the generation principle of phosphorescence.

Therefore, promising RTP can be probably achieved from CPDs by means of smarter design.

Yang *et al.* were the first to describe a feasible and facile route to prepare metal-free RTP materials by directly constructing CPDs *via* hydrothermal synthesis without the compositing of additional matrices.<sup>153</sup> As a model system, CPDs synthesized from polyacrylic acid and ethylenediamine (EDA) exhibited unique RTP properties in air, as expected, with lifetime up to 658.11 ms. These CPDs synthesized with amide or imide structures, alongside the crosslink enhanced emission (CEE) effect, were confirmed to contribute toward the generation of RTP in CPDs by both contrast experiments and computational simulations (Fig. 20). First, chemical reactions easily occurred among the abundant functional groups. EDA serving as a crosslinker reacted with extensive dangling carboxyl groups on the polymer chains, forming luminescent centers and covalently crosslinked self-matrix frameworks. Second, effective crosslinking could shorten the distance between the functional groups. New distributions of energy levels can be formed by electron overlap from coupled luminescence units in nano-confined space that can effectively reduce the energy gap ( $\Delta E_{ST}$ ) and facilitate the generation of intrinsic triplets. Third, the widespread existence of supramolecular interactions in the interior of CPDs can also decrease vibrations and rotations, thereby inhibiting any nonradiative relaxation. The reported guiding results demonstrate the potential of CPDs as a universal route for achieving effective metal-free RTP. Recently, they expanded the method from polycondensation to poly-addition, confirming the universality and feasibility of the hydrothermal crosslinking and carbonization processes for CPDs to achieve the RTP property.<sup>154</sup>

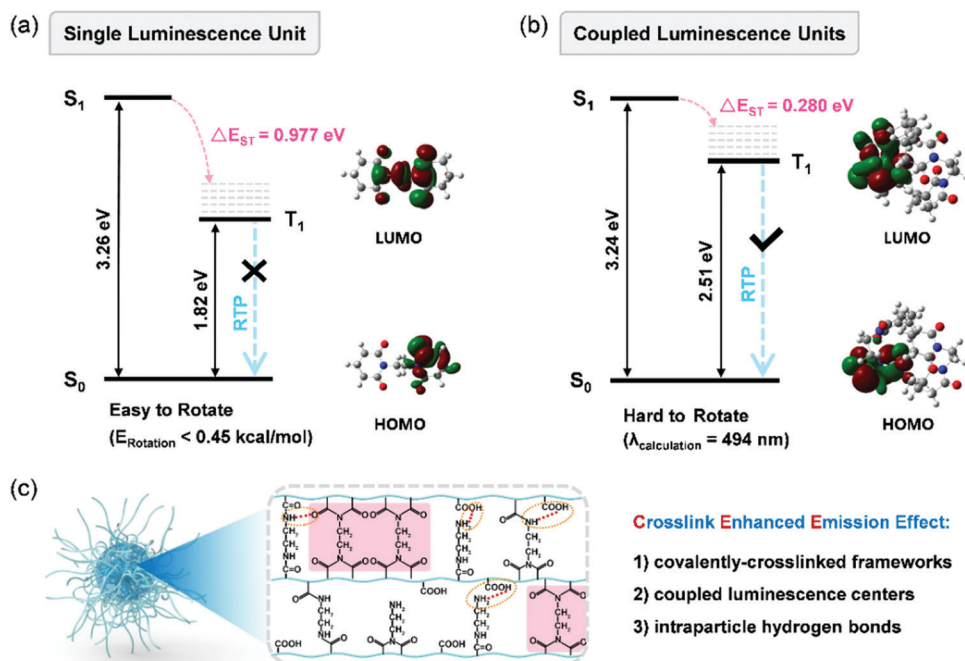


Fig. 20 Theoretical analysis of the CEE effect: (a) energy level diagrams of single luminescence units and (b) coupled luminescence units. (c) Schematic illustration of the CEE effect on RTP properties in CPDs. Reprinted from ref. 153 with permission. Copyright (2018) Wiley-VCH.

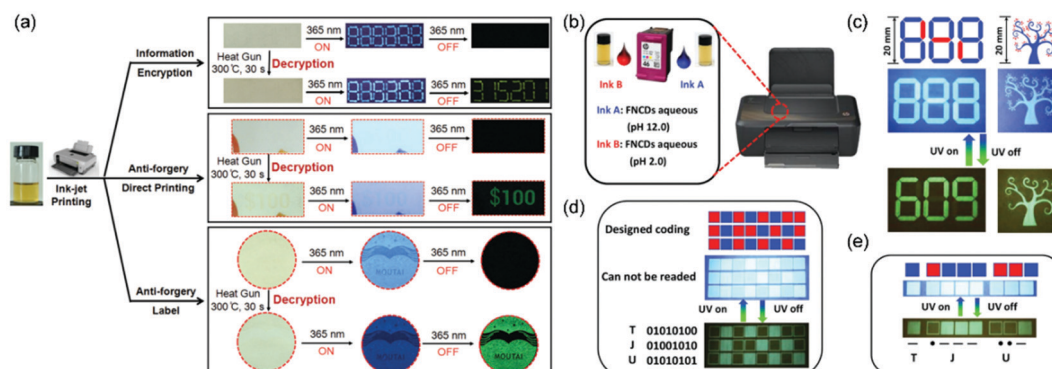
Inspired by related reports, increasing number of researchers have noticed the advantages of CPDs on RTP performance and a range of significant studies have been rapidly performed. For example, Lin *et al.* successively achieved RTP from two similar CPD systems based on the aforementioned CEE effect. One involved the preparation of CPDs with RTP performance by the microwave irradiation of ethanolamine and phosphoric acid. Gram-scale CPDs and the longest RTP lifetime (1.46 s) could be achieved.<sup>155</sup> Another study described the hydrothermal treatment of EDA and phosphoric acid for obtaining CPDs. The as-synthesized CPDs could realize the conversion from fluorescence (180 °C) to ultralong RTP (280 °C) by an external heating stimulus. They supposed that the relatively low temperature afforded intertwined CEE-type CPDs with abundant sub-fluorophores. After heating to a higher temperature, parts of the intertwined polymer chains in the CPDs could be further reacted *via* crosslinking and dehydration/carbonization to form compact cores, which behaved as a matrix to immobilize the sub-fluorophores, eventually inducing RTP performance. The system demonstrated the potential of CPDs as stimuli-responsive optical materials (Fig. 21a).<sup>156</sup> Subsequently, Qu *et al.* utilized the difference in luminescence responses between several CPDs and polyvinyl alcohol composites annealed at different temperatures. They proposed the concept of thermal-treatment-controlled multilevel fluorescence/phosphorescence data encryption and further developed the application of CPDs in the stimuli-responsive field.<sup>157</sup> Moreover, Feng *et al.* designed and synthesized novel fluorine and nitrogen codoped CPDs *via* one-step solvothermal treatment. The CPDs possessed good aqueous solubility, blue pH-stabilized fluorescence, and green pH-responsive RTP with outstanding reversibility. Based on these interesting properties, they demonstrated the data encoding/reading strategy using basic or acidic aqueous CPD solutions for advanced anticounterfeiting *via* time-resolved luminescence imaging techniques, as well as the steganography of complex patterns (Fig. 21b–e).<sup>158</sup>

Given the wide range of chemical precursors and possible synthesis conditions, these approaches can lead to a wide

variety of CPD materials with diverse engineered RTP properties. A careful selection of precursors (such as combinations of poly-acid, polyamine, polyol, diacid, diamine, and other potential multifunctional compounds) and different synthesis routes (such as solvothermal and ultrasonic-/microwave-assisted hydrothermal/solvothermal treatment), as well as even some specific addition polymerization reactions, can provide a wide range of potential routes for the formation of CPD materials for use in particular applications. Further studies should focus on making RTP materials brighter, with different emission colors and improved quenching resistance, as well as simultaneously being observable in aqueous solutions. Moreover, the emergence of novel biosystems is also anticipated, which can further expand the RTP properties for specific biomaterials.

### 3.3 PhPs

PhPs, another important class of phosphorescent polymers, refer to instances in which phosphorescent emitters (*e.g.*, Ir(III) and Pt(II) complexes) are incorporated into the polymer main chains or side chains *via* covalent bonds.<sup>159</sup> Owing to the strong spin–orbital coupling of the phosphorescent emitters, PhPs can harvest both S and T excitons to achieve 100% IQE, which is four-fold higher than that of fluorescent polymers. To develop efficient PhPs, several requirements need to be addressed. First, the triplet energy level ( $E_T$ ) of polymer hosts should be higher than that of the phosphorescent emitters; otherwise, triplet back energy transfer from the phosphorescent emitters to the polymer hosts can occur, and the emission of phosphorescent emitters can get quenched. Second, the polymers should have suitable highest occupied molecular orbital (HOMO)/lowest unoccupied molecular orbital (LUMO) energy levels to facilitate charge injection from the electrode to the emissive layer and then create charge transport to the recombination zone. Third, phosphorescent emitters should exhibit high photoluminescent quantum efficiency (PLQY) to convert all the generated excitons into photons in order to achieve efficient electroluminescence. Finally, PhPs should exhibit excellent solubility in common solvents to form high-quality films and



**Fig. 21** (a) Potential applications of F-CD-based ink in information encryption, direct printing on a coupon, and Moutai Liquor anticounterfeiting tags. (b) Schematic illustrations of a representative customized tricolor inkjet cartridge. (c) Demonstration of lifetime encoding for security applications. Digital photographs of (d) binary-coded and (e) Morse-coded two-color microarray data storage chips. Reprinted from ref. 156 and 158 with permission. Copyright (2018) Wiley-VCH.

good thermal and morphological stabilities for the fabrication of long-durable light-emitting devices. In the past few years, several efficient PhPs have been synthesized by incorporating phosphorescent emitters into conjugated or nonconjugated polymer hosts to form a host/dopant system. In this subsection, recent progresses made in the two abovementioned kinds of PhPs, *i.e.*, full-color emission and white-light emission PhPs, is briefly addressed.

**Full-color-emitting PhPs.** To develop efficient PhPs with full-color emission, polymer hosts with different  $E_T$  levels are desirable in order to confine the T excitons on phosphorescent emitters. Generally, there are two main kinds of polymer hosts: conjugated and nonconjugated polymer hosts. To realize blue triplet emission, the  $E_T$  value of the polymer hosts should be higher than those of blue phosphorescent emitters (2.60–2.70 eV). In fact, nonconjugated polymers having higher  $E_T$  levels have been used as polymer hosts for the design of blue-emitting PhPs (PhP-Bs). For example, poly(*N*-vinylcarbazole) (PVK) with a high  $E_T$  value of 3.0 eV was used as the backbone for PhP-Bs.<sup>159</sup> However, PVK has a deep HOMO level of  $-5.90$  eV and a shallow LUMO level of  $-2.10$  eV, which leads to large charge injection barriers for air-stable electrodes. To resolve this problem, a nonconjugated, bipolar poly(arylene ether phosphine oxide) host (PCzPO) was reported by incorporating a carbazole/triaryl phosphine oxide hybrid into a poly(arylene ether) scaffold. For this polymer host, a tradeoff between high triplet energy (2.96 eV) and suitable HOMO/LUMO levels ( $-5.70/-2.30$  eV) could be achieved. By physically doping iridium(III)[bis(4,6-difluorophenyl)-pyridinato-*N,C*<sup>2</sup>]-picolinate (FIrpic, 5 wt%) into PCzPO, a single-layer blue-emitting device could be fabricated with the maximum luminous efficiency (LE) of  $1.27$  cd A<sup>-1</sup>, which is 9-fold higher than

that of the PVK-based device. When a hole-/exciton-blocking layer was utilized for the PCzPO device, the LE was further improved to  $23.3$  cd A<sup>-1</sup>. This result showed that PCzPO was a promising polymer host for blue phosphorescent emitters.<sup>160</sup> By covalently tethering FIrpic as the blue phosphorescent emitter into the side chain of a poly(arylene ether) host, a series of PhP-Bs (Fig. 22a) were developed. By tuning the content of the blue phosphorescent emitter, complete energy transfer from the polymer host to the dopant was observed. When the phosphor content was 5 mol%, the PhPs afforded the best device performance with the maximum LE of  $19.4$  cd A<sup>-1</sup>, which was  $\sim 3.5$  times higher than that of PhP-Bs with PVK as the host. It should be noted that a low-temperature condensation polymerization reaction enabled by a fluorinated poly(arylene ether phosphine oxide) backbone (FPCzPO) was the key to the successful synthesis of PhP-Bs. At a high temperature ( $165$  °C), the phosphorescent emitter, FIrpic, may undergo structural decomposition, which might lead to an unwanted bathochromic shift in the resulting PhPs. In contrast, under a lower temperature of  $120$  °C, the structure of FIrpic remained unchanged, and the emission from the resulting PhPs matched well with the characteristic emissions from a phosphorescent emitter.<sup>161</sup> This strategy has also been used for the design of yellow-emitting PhPs (PhP-Ys) by introducing the 2-(fluorene-2-yl)-1*H*-benzimidazole-based iridium complex, (fbi)<sub>2</sub>Ir(acac), into the FPCzPO backbone (Fig. 22c). Due to the efficient intermolecular Förster energy transfer from FPCzPO to (fbi)<sub>2</sub>Ir(acac) and charge trapping on (fbi)<sub>2</sub>Ir(acac), the electroluminescence from FPCzPO could be completely quenched even at low Ir-complex contents. With Ir loading of 3 mol%, the polymer exhibited the best device performance with peak LE of  $10.4$  cd A<sup>-1</sup> and Commission internationale de l'éclairage (CIE) coordinates of (0.53, 0.46).<sup>162</sup>

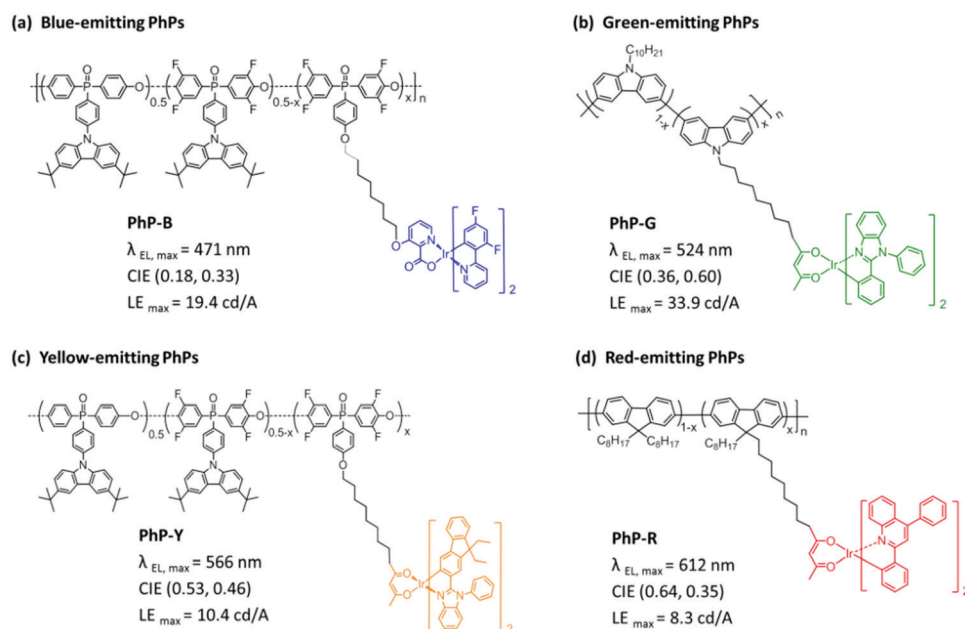


Fig. 22 Full-color-emitting PhPs. Reproduced from ref. 161–163 and 165 with permission. Copyright (2012, 2012, 2010, and 2011) from American Chemical Society, Royal Society of Chemistry, and Wiley-VCH.

To design red-emitting PhPs (PhP-Rs) and green-emitting PhPs (PhP-Gs), CPs with lower  $E_T$  values (2.10–2.60 eV) can be selected as the hosts. For instance, PhP-Rs (Fig. 22d) were reported by the use of polyfluorene (PF) with  $E_T$  value of 2.10 eV as the host and Ir(ppq)<sub>2</sub>(acac) (ppq: 2,4-diphenylquinolyl-N-C<sup>2'</sup>) as the phosphorescent emitter. A single-layer device based on this polymer exhibited superior EL performance with the maximum LE of 5.0 cd A<sup>-1</sup> and CIE coordinates of (0.63, 0.35). A further optimization of the device structure by inserting an alcohol-soluble electron injection layer could improve the LE to 8.3 cd A<sup>-1</sup>.<sup>163</sup> In addition, it was found that the charge balance of the polymer host played a crucial role in determining the device efficiency of the PhP-Rs. By alteration of the polymer host from PF to poly(fluorene-*alt*-carbazole) (PFCz) and polycarbazole (PCz), the LE decreased from 5.0 to 0.3 cd A<sup>-1</sup> and 0.04 cd A<sup>-1</sup>, respectively, which could be attributed to the imbalanced hole and electron flux in PFCz and PCz caused by their higher hole mobilities than electron mobilities.<sup>164</sup> For the development of PhP-Gs, polycarbazoles are selected as the hosts owing to their higher  $E_T$  values (2.40 eV). For example, three kinds of PhP-Gs have been developed by using PF, PFCz, and PCz with different  $E_T$  values as the polymer hosts and 1,2-diphenyl-benzimidazole-based Ir complexes as the green phosphorescent emitter (Fig. 22b, PhP-G). Since the  $E_T$  level gradually increased from 2.20 to 2.40 and 2.60 eV from PF to PFCz and PCz, respectively, the triplet back energy transfer from the phosphorescent emitter to the polymer host could be effectively inhibited. As a result, the maximum LE of the PhPs with a double-layer device configuration was significantly improved from 0.3 (PF) to 33.9 cd A<sup>-1</sup> (PCz).<sup>165</sup>

**White-emitting PhPs.** White polymer light-emitting diodes (WPLEDs) have attracted considerable attention owing to their applications in fabricating low-cost light sources and full-color displays by means of solution processes such as inkjet printing and roll-to-roll printing technologies. In general, there are two main approaches to obtain white emission from polymers. One is polymer blend systems, *e.g.*, polymer/polymer or polymer/low-mass molecules with complementary colors (blue and yellow for two-color white emission; blue, green, and red for three-color white emission) to cover the entire visible region. These multi-component systems may suffer from the phase separation of different emitters, which can lead to the deterioration of device performance. An alternative approach is to develop single white-emitting polymers (SWPs) by covalently incorporating complementary emitters into a single polymer, which produces broadband emissions from all the emitters *via* partial energy transfer from the high-energy emitter (blue emitter) to the lower-energy ones (yellow, green, or red emitters). Because all the emitters are covalently bonded to the polymer backbone, the risk of phase separation can be avoided. Since the first report on SWPs with individual blue and yellow fluorescent emissions was published by Wang *et al.* in 2004,<sup>166</sup> a number of SWPs, namely, all-fluorescent SWPs and fluorescent/phosphorescent hybrid ones, have been reported. However, their device performances, such as driving voltage, external quantum efficiency (EQE), and power efficiency (PE), have not been comparable to those of low-mass-molecule organic light-emitting diodes (OLEDs).

To improve device efficiency, all-phosphorescent SWPs were reported by simultaneously grafting blue- and yellow-emitting phosphors into a bipolar, high- $E_T$  poly(arylene ether) host. Different from all-fluorescent and fluorescent/phosphorescent hybrid ones, all-phosphorescent SWPs were supposed to fully utilize the S and T excitons by phosphorescent dopants. Based on this consideration, FIrpic and (fbi)<sub>2</sub>Ir(acac) were selected as blue- and yellow-emitting phosphors, respectively, for the design of all-phosphorescent SWPs. A series of all-phosphorescent SWPs (Fig. 23, WPB75Y7) were successfully prepared by tailoring the feed ratios of monomers. The obtained polymers showed balanced blue emissions from FIrpic and yellow emissions from (fbi)<sub>2</sub>Ir(acac) to yield white emission. Among the SWPs, polymers containing 7.5 mol% FIrpic and 0.7 mol% (fbi)<sub>2</sub>Ir(acac) exhibited the best device performance with the maximum LE of 18.4 cd A<sup>-1</sup>, maximum PE of 8.5 lm W<sup>-1</sup>, peak EQE of 7.1%, and CIE coordinates of (0.31, 0.43). Even at a brightness level of 1000 cd m<sup>-2</sup>, the LE was as high as 14.2 cd A<sup>-1</sup>, indicating slow efficiency roll-off. These LE and PE values were much higher than those of all-fluorescent and fluorescent/phosphorescent hybrid SWPs, shedding light on the significance of all-phosphorescent SWPs.<sup>167</sup>

To realize high PE, new all-phosphorescent SWPs (Fig. 23, SWP1–FIrpic, SWP1–FIrOBu, SWP2–FIrOBu, and SWP3–FIrOBu) were synthesized by the use of a combination of high-HOMO-level blue-emitting phosphor and high-HOMO-level nonconjugated poly(arylene phosphine oxide) host. On one hand, by changing the common blue-emitting phosphor (FIrpic) to a homemade blue-emitting phosphor (FIrOBu) containing electron-donating alkoxy groups on ligands, the HOMO level could be elevated from -5.64 to -5.28 eV, which matches well with the HOMO level of the polymer host. Therefore, hole scattering between the blue-emitting phosphors and polymer hosts can be eliminated, resulting in a reduced turn-on voltage from 4.2 to 3.4 V. On the other hand, the HOMO level of the polymer host can also be tuned from -5.64 to -5.20 eV by the use of alkoxy-substituted carbazole dendrimers instead of using the *tert*-butyl-capped carbazole as the side chain, coming close to the work function of the anode. Consequently, the hole injection barrier at the polymer/anode interface was lowered, and the turn-on voltage was reduced to 2.8 V. As a result, a low turn-on voltage of 2.8 V, record EQE of 18.0%, and forward PE of 52.1 lm W<sup>-1</sup> could be achieved for SWPs. Such remarkable progress makes SWPs outperform currently used polymer blends for solution-processed white organic light-emitting diodes (WOLEDs) and comparable with the low-mass molecules used for vacuum-deposited WOLEDs.<sup>168</sup>

Recent progresses made in AIE polymers, CPDs, and PhPs are introduced in this section. The preparation methods, design strategies, material systems, and device performances of these luminescent polymers are emphasized. Although significant advances have been made, many challenges require redressal for the development of phosphorescent polymers. For example, their material design, photophysical properties, and device processes need to be improved. The future directions for phosphorescent polymers can focus on the following

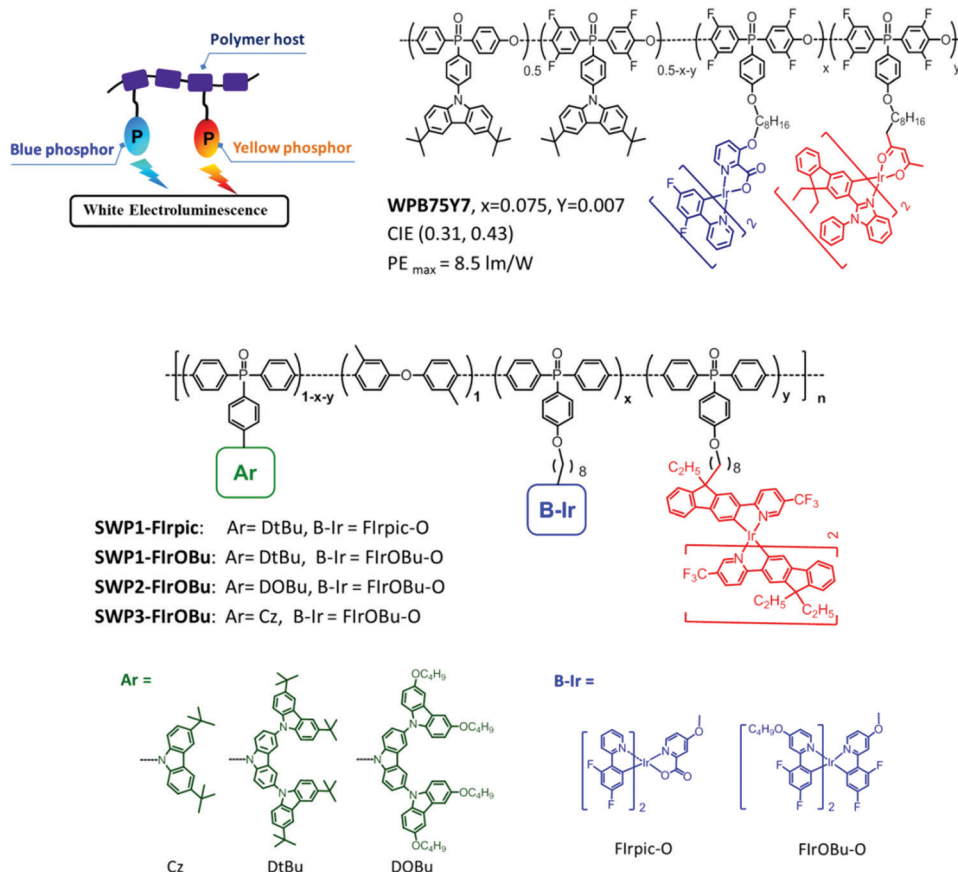


Fig. 23 White-emitting PhPs. Reproduced from ref. 167 and 168 with permission. Copyright (2012 and 2018) from American Chemical Society and Royal Society of Chemistry.

aspects: long-life blue emissions, high-efficiency white lighting, and low-cost nonprecious metal-containing ones.

## 4. Photovoltaic polymer materials

One of the strategies to resolve energy shortage and environmental pollution is to develop solar energy conversion. Solar cells are classified into two types: inorganic solar cells and OSCs. Inorganic solar cells that are based on inorganic semiconducting materials have achieved great advances. For example, crystalline silicon solar cells have been commercialized for many years. When compared with inorganic solar cells, OSCs based on polymer materials have many advantages, such as low cost, lightweight, low toxicity (or nontoxic), flexibility, high absorption coefficient, semitransparency, and easy and large-area fabrication.<sup>169</sup> Moreover, photovoltaic functional polymers have a wide range of properties, and their optical properties (absorption range and intensity), electronic properties (energy level and carrier mobility), and film morphology can be effectively controlled by chemical methods.

Conventional OSC devices generally comprise the active layer, interfacial layers, and electrodes (Fig. 24). The active layer of bulk-heterojunction (BHJ) OSCs is generally composed

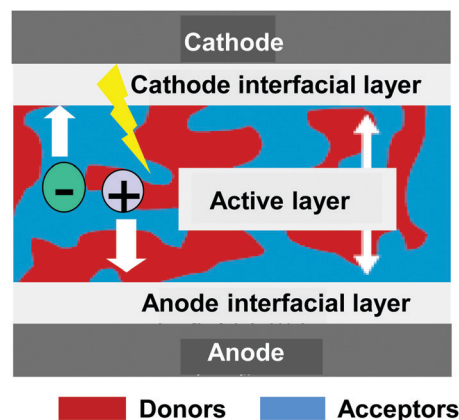


Fig. 24 Structure of a BHJ solar cell.

of two kinds of semiconductor materials, namely, electron donor (D) and electron acceptor (A), which exhibit microphase separation between the donors and acceptors at the scale of 10–20 nm. Many strategies have been used to obtain OSCs with excellent power conversion efficiencies (PCEs), such as designing active layer materials (including donors and acceptors), controlling the morphology of the active layer, and optimizing the fabrication methods as well as structures of the OSC devices.

Exciton diffusion, charge separation, and charge transport, which occur in the active layer, require appropriate nanoscale phase-separated interpenetrating network (IPN) structures. Therefore, controlling the morphology of the active layer is crucial. In addition, water/alcohol-soluble conjugated polymers (WSCPs) can be used as interfacial layer materials to optimize the processing methods of OSC devices. The development of electron donors has been more comprehensively investigated than electron acceptors for the past decade; interested readers should refer to other reviews in relation to electron donors.<sup>170,171</sup>

Among acceptors, fullerene derivatives have dominated this field for two decades, and single-junction fullerene-based devices have exhibited PCEs up to 11%.<sup>172</sup> However, shortcomings of fullerene derivatives, such as weak absorption in the visible region, difficult synthesis and purification, and low tunability of energy levels, hinder further improvements in device performance. In recent years, non-fullerene acceptors (NFAs) have attracted increasing attention, among which rylene diimides and fused-ring electron acceptors (FREAs) are two promising material categories.<sup>9,173</sup> When compared with fullerenes and rylene diimides, FREAs exhibit several superiorities, such as stronger and red-shifted absorption in the visible and NIR regions and more tunable energy levels. Currently, OSCs based on FREAs have achieved PCEs of over 17%.<sup>174</sup> In this section, NFAs, control morphology of the active layer, and solution processing using WSCPs have been addressed.

#### 4.1 ITIC-based fused-ring electron acceptors

In 2015, Zhan *et al.* reported the star acceptor of ITIC based on a fused-ring core, indacenodithieno[3,2-*b*]thiophene (IDTT),

with four 4-hexylphenyl side chains and two 2-(3-oxo-2,3-dihydro-1*H*-inden-1-ylidene)malononitrile (IC) end groups (Fig. 25).<sup>175</sup> As-cast devices based on the polymer donor PTB7-Th and ITIC exhibited PCEs up to 6.8%, which was the new record for fullerene-free OSCs at that time. Since then, the ITIC family has begun to replace the dominant position of fullerene acceptors, inaugurating a new era—the era of NFAs. In general, the FREA molecule consists of three parts: electron-donating aromatic fused-ring core, electron-withdrawing terminal groups, and side chains. FREAs exhibit good solubility in chloroform, chlorobenzene, and *o*-dichlorobenzene, good thermal stability with a high decomposition temperature of over 300 °C, and tunable electronic properties with different energy levels and variable absorption edges from 700 to 1100 nm. Meanwhile, their solubility, crystallinity, energy level, absorption, and charge mobility can be modulated by fused-ring core engineering, end-group engineering, and side-chain engineering.

**Fused-ring core engineering.** First, a series of FREAs based on indacenodithiophene (IDT) oligomers were designed to investigate the effects of fused-ring numbers (Fig. 25).<sup>176</sup> By increasing the number of IDTs, the absorption spectrum experiences red-shifts, the HOMO experiences upshifts, and electron mobility decreases due to the increased twisted main chain and weaker  $\pi$ - $\pi$  stacking. The thieno[3,2-*b*]thiophene (TT) unit possesses a symmetrical and coplanar structure, large electron delocalization, and strong electron-donating ability. IHIC and FOIC were designed, where the benzene ring in IDT and IDTT were replaced by TT, respectively (Fig. 25).<sup>177,178</sup> Both IHIC and FOIC showed stronger NIR absorption, upshifted HOMO level, and higher electron mobility than those of their

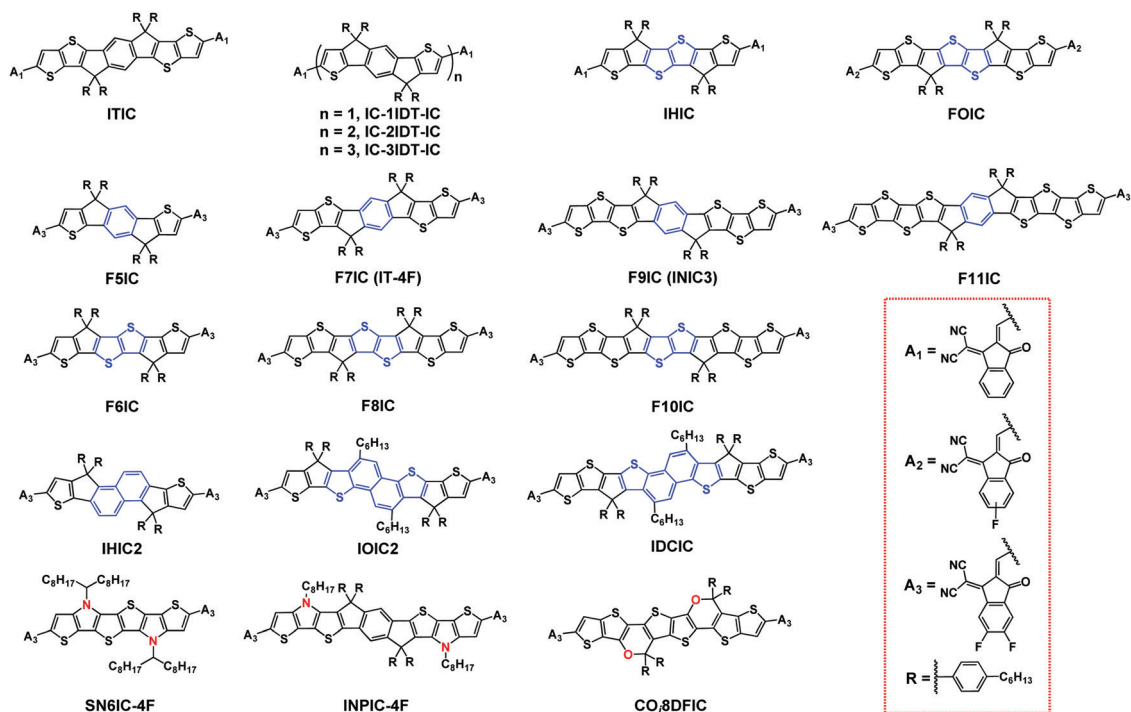


Fig. 25 Fused-ring core engineering of FREAs.

benzene-based counterparts, leading to a higher short-circuit current density ( $J_{SC}$ ).<sup>177,178</sup> Meanwhile, IHIC2 and IOIC2 were reported, with the substitution of the benzene ring in the IDT unit with naphthalene and naphtho[1,2-*b*:5,6-*b'*]dithiophene (NTT), respectively.<sup>179</sup> When compared with IHIC2, IOIC2 showed higher energy levels, broader absorption spectrum, and higher electron mobility, leading to higher values of open-circuit voltage ( $V_{OC}$ ),  $J_{SC}$ , fill factor (FF), and ultimately much higher PCE. Subsequently, IDCIC with a fused-10-ring core was developed, where the outermost thiophene rings in IOIC2 were replaced with TT units.<sup>180</sup> The FTAZ:IDCIC-based OSCs yielded PCE of 13.58% due to the higher energy levels and red-shifted absorption spectrum than those for IOIC2. In order to systematically study the effects of core size, seven FREAs (F5-11IC)<sup>181,182</sup> with the same end groups and side chains but different cores were synthesized, whose sizes ranged from 5 to 11 fused rings. Increasing the number of outermost fused-thiophene rings and replacing the benzene ring with a TT unit led to a red-shift in the absorption spectra, upshift in the energy levels, and synergistic enhancement in molecular packing and electron mobility. Heteroatoms (*e.g.*, O and N) also played an important role in the fused-ring core engineering due to the  $\pi$ -conjugative effect and  $\sigma$ -inductive effect. Pyrrole rings were introduced in the backbone to enhance the electron-donating ability of the cores in SN6IC-4F and INPIC-4F (Fig. 25).<sup>183,184</sup> Both SN6IC-4F and INPIC-4F exhibited strong absorption characteristics in the NIR region and the PCEs of devices blended with the polymer PBDB-T donor were higher than 13%. Oxygen atoms were added into the fused-ring core of CO<sub>8</sub>DFIC by using the “carbon–oxygen bridge” to replace the C bridge in F8IC, thereby enhancing the electron-donating capability of the core; this led to a very small optical bandgap (1.26 eV).<sup>185</sup> A  $J_{SC}$  value of 26.12 mA cm<sup>-2</sup> and PCE of 12.16% were achieved from a PTB7-Th:CO<sub>8</sub>DFIC-based device, and PCE of 17.36% was achieved from a tandem device (Fig. 25).<sup>174</sup>

**End-group engineering.** IDT-2BR, using an IDT unit as the core and 5-(benzo[*c*][1,2,5]thiadiazol-4-ylmethylene)-3-ethyl-2-thioxothiazolidin-4-one (BR) units as the end-capped

electron-withdrawing groups, achieved PCE of 5.12% when combined with poly(3-hexylthiophene) (P3HT) (Fig. 26).<sup>186</sup> Furthermore, some similar FREAs such as ATT-3 and ORCN were developed by changing the end groups, which afforded PCEs >6%.<sup>187,188</sup> IEIC with an alkyl-substituted thiophene spacer between the IDT and IC groups afforded PCE of 6.31% when blended with PTB7-Th.<sup>189</sup> Later, some modifications were made on the structure of IEIC, such as substituting the alkyl on the thiophene with alkoxy and adopting a fluorinated IC group. The PCEs of devices based on PTB7-Th:IEICO-4F and PBDB-T:ITOIC-2F blends are ~12% due to the stronger NIR absorption and  $\pi$ - $\pi$  stacking.<sup>190,191</sup> Considerable efforts have been devoted toward the modulation of the end groups of ITIC. One and two methyl groups were introduced onto the IC groups to afford IT-M and IT-DM, respectively.<sup>192</sup> OSCs based on PBDB-T:IT-M exhibited high PCE of 12.05% with higher  $V_{OC}$  and  $J_{SC}$  as compared to those for the control device based on PBDB-T:ITIC. Meanwhile, the benzene unit of the IC group was replaced with a thiophene ring, *e.g.*, ITCC, ITCPTC, and MeIC.<sup>193–195</sup> The substitution with a thiophene ring affected the electronic properties and enhanced the intermolecular interactions, leading to higher FF and PCE. INIC and three fluorinated analogs (INIC1–3) were obtained with IC and fluorinated ICs, respectively.<sup>196</sup> Fluorine substitution downshifted the LUMO energy level, red-shifted the absorption spectrum, and enhanced the electron mobility. OSCs based on fluorinated FREAs showed PCEs as high as 11.5%, which is much higher than that of its nonfluorinated counterpart (7.7%).

**Side-chain engineering.** Side chains on the core unit have important effects on the solubility, crystallinity, intermolecular interactions, and electron mobility of the acceptors. IDIC was synthesized by the use of an alkyl instead of alkylphenyl groups of IC-1IDT-IC, which afforded a red-shifted absorption spectrum, more ordered molecular packing, and improved electron mobility than those from IC-1IDT-IC, leading to better performance (Fig. 27).<sup>197</sup> O-IDTBR was synthesized with *n*-octyl instead of *n*-hexylphenyl in IDT-2BR, which showed red-shifted absorption and enhanced intermolecular packing than those in

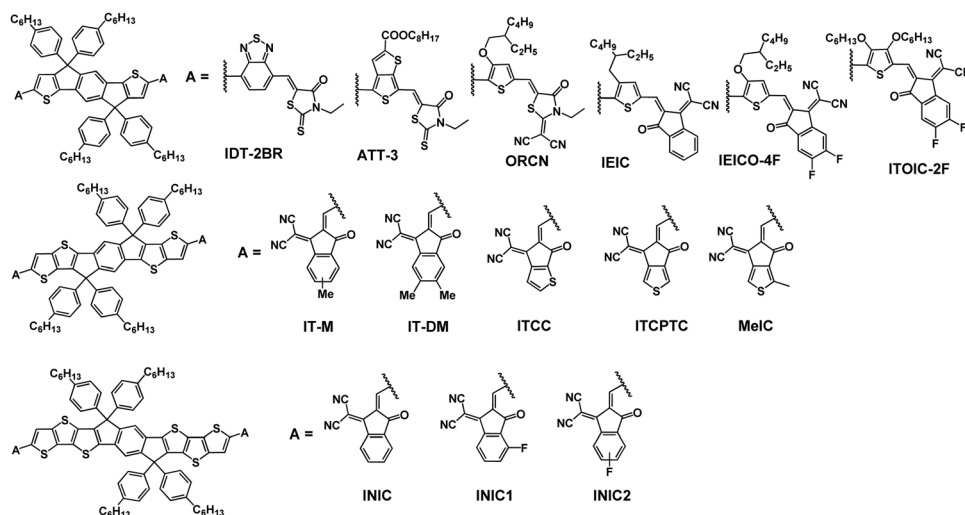


Fig. 26 End-group engineering of FREAs.



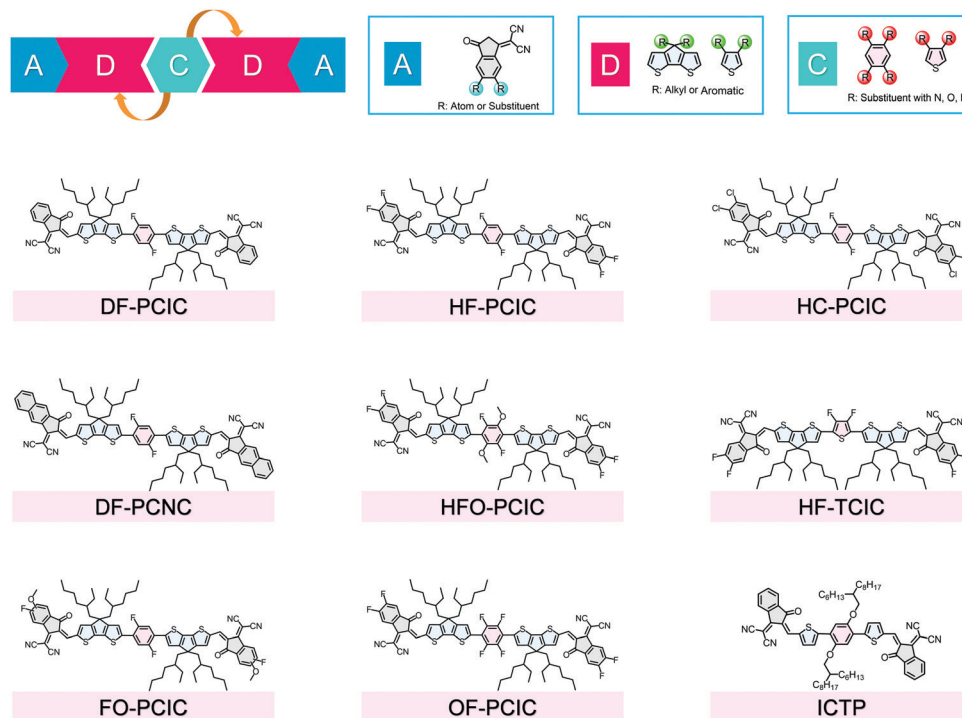


Fig. 28 Diagram of UCEAs and chemical structures of representative molecules.

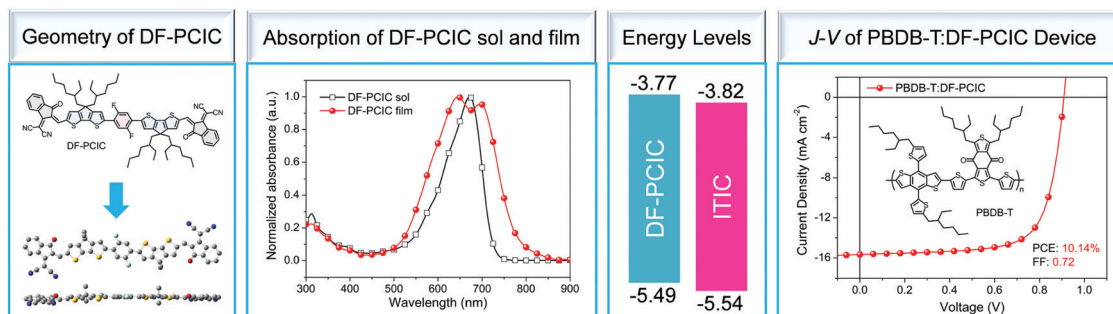


Fig. 29 Geometry, absorption, and energy levels of DF-PCIC and device performance of PBDB-T:DF-PCIC-based OSCs.

group on the benzene-based core (HFO-PCIC), extension of the conjugated length of IC terminals (DF-PCNC), replacement of benzene-based core with thiophene-based core (HF-TCIC), and so on.<sup>211–215</sup> After selecting suitable polymer donors, such as PBDB-T or its fluorinated analog PBDB-TF, obvious enhancement in  $J_{SC}$  can be observed (over 17 mA cm<sup>-2</sup> for HF-PCIC-based OSCs; over 18 mA cm<sup>-2</sup> for HC-PCIC- and DF-PCNC-based OSCs; over 20 mA cm<sup>-2</sup> for HF-TCIC-based OSCs) for the relevant devices. Therefore, HF-PCIC-, HC-PCIC-, and DF-PCNC-based OSCs could yield PCEs over 11%. For PBDB-TF:HC-PCIC-based OSCs, the subsequent use of PC<sub>71</sub>BM as the third component to optimize the morphology and enhance the charge transfer can yield solar cells that exhibit large improvements in quantum efficiencies (from ~70% to over 80%) with outstanding PCE of 12.36%.<sup>212</sup> The above efforts from Chen *et al.* demonstrate that UCEAs possess good structural variability for tuning the material properties for obtaining high-performance OSCs.

New acceptors with fully unfused structures have also been recently developed by Chen *et al.*, such as ICTP that is constructed from single aromatic units within 2 or 3 synthesis steps.<sup>216</sup> With the assistance of noncovalent interactions, they identified that these potentially rotatable acceptor structures allow adapting stackable yet thermally stable conformation in a condensed solid, facilitating the new molecules to not only exhibit feasible solution processability but also excellent film characteristics. PCE of 4.43% was achieved for ICTP with the PBDB-T polymer. Further, PCE of over 10% can also be obtained for such a type of simple electron acceptor, revealing that there is still considerable space for achieving better device performance. Notably, such unfused acceptors can be synthesized with C–H activation coupling reaction and purified with recrystallization, largely reducing the synthesis complexity of electron acceptors.

To enable the efficient working of fullerene-free OSCs, allowing small driving forces is essential to achieve higher

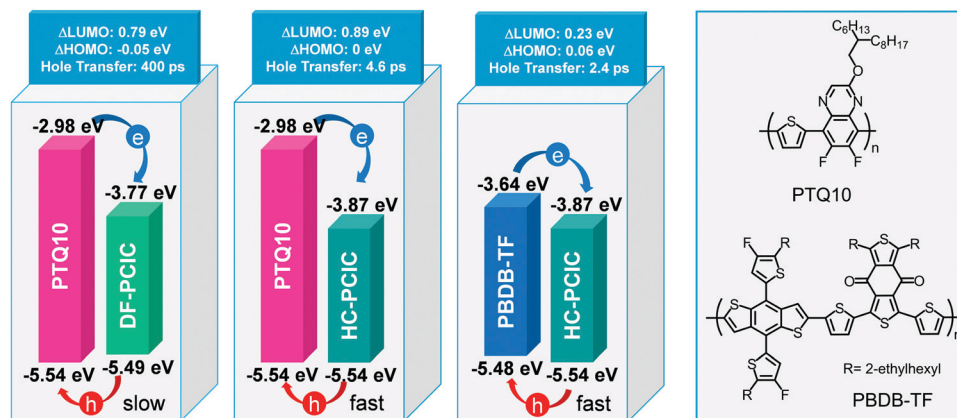


Fig. 30 Diagram of hole transfer for different blends with various HOMO offsets and hole transfer rates. Adapted from ref. 217 with permission. Copyright (2019) American Chemical Society.

voltages and better performance for the OSCs. Very recently, Chen *et al.* performed a systematic study on the above issue with UCEAs and revealed profound understandings on the charge transfer process in non-fullerene blends. Due to the adjustable energy levels of UCEAs, they chose three UCEAs, namely, DF-PCIC, HC-PCIC, and FO-PCIC, developed by their group, as well as three polymer donors, namely, PTQ10, PBDB-TF, and PBDB-T; therefore, six blends with various HOMO offsets ranging from  $-0.05$  to  $0.21$  eV were formulated (Fig. 30).<sup>217</sup> Through the aid of bilayer hole-only devices and transient absorption spectra, they found that the occurrence of hole transfer is independent of the HOMO offsets, but the hole transfer rate was influenced by the HOMO offsets. The blend of PTQ10/DF-PCIC with slightly negative HOMO offset of  $-0.05$  eV yielded a really slow hole transfer rate of  $\sim 400$  ps, while blends such as PTQ10/HC-PCIC and PBDB-TF/HC-PCIC with HOMO offsets of  $\geq 0$  eV could yield faster hole transfer rates of  $\leq 4.6$  ps. As a result, small HOMO offsets of 0 and 0.06 eV can enable achieving higher PCEs of 10.42% and 11.75% for the PTQ10:HC-PCIC blend and PBDB-TF:HC-PCIC blend, respectively. This work illustrates the fact that the modulation of HOMO offsets ranging between 0 and 0.1 eV has promising potential to realize breakthroughs in OSC efficiencies and energy loss.

In summary, from the abovementioned discussion, we can understand that UCEAs possess advantages with respect to the

following aspects: (1) flexibility and diversity in molecular design, (2) reduction in synthesis complexity of new materials, and (3) efficient performance and good stability. UCEAs would also be good choices for the design of high-performance and stable NFAs.

### 4.3 Control morphology of active layer during solution processing

CPs and their corresponding blends have promising potential applications in organic optoelectronic devices, such as organic thin-film field-effect transistors (OTFTs), OSCs, and OLEDs, due to their advantages such as a combination of optoelectronic properties and solution processing, *e.g.*, inkjet printing, roll-to-roll printing, *etc.* Part of the challenge of the application of CP semiconductors is the deeper understanding of the effect of the microstructure on the macroscopic electrical properties. Different device applications have different requirements for the polymer microstructures (Fig. 31). In this subsection, we focus on how to control the morphology of the polymers from the viewpoint of thermodynamics or kinetics during solution processing, and the relationship between morphology and device performance is also established.

Charge transport in semiconducting polymers occurs within the polymer backbones and between different chains due to the electronic coupling between the rings, which is related to their

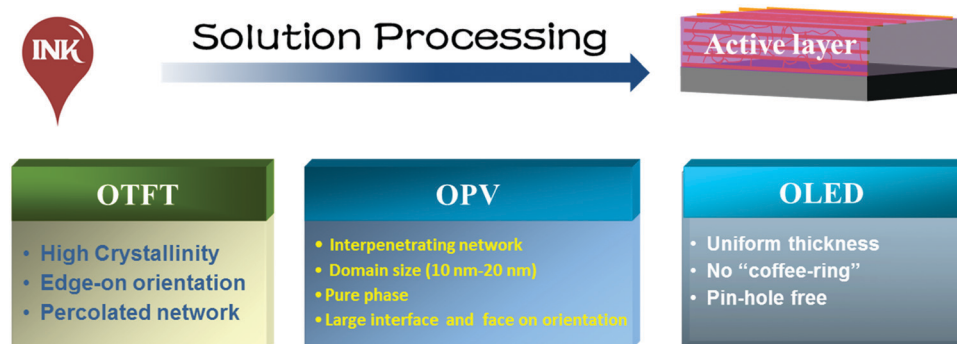


Fig. 31 Requirements of the active layer for OTFTs, oligo(*p*-phenylene vinylene) (OPVs), and OLEDs.

microstructure. In order to obtain higher carrier mobility, the CP films should possess a percolated network with long-range order, high crystallinity, and appropriate crystal orientation. However, the entanglement of the CP caused by the flexible side chain and rigid backbone suppresses the intrachain mechanism. Furthermore, the fast crystallization kinetics during the film-forming process yields several grain boundaries among the crystals. Hence, the high-density nanowires of the CP are desirable to improve the charge transport. However, the entanglement of the CP chain severely impedes the nucleation to grow nanowires at a higher solution concentration. Han *et al.* controlled the equilibrium of entanglement, disentanglement, nucleation, and growth by tuning the interaction between the solvent and CP, temperature, ultrasonic oscillation, adding additives, blending with flexible polymer, *etc.*<sup>218–221</sup> Once the disentangled polymer chains could nucleate to form nanowires, the equilibrium between entanglement and disentanglement is broken and the entangled polymer chains get further disentangled to self-assemble into nanowires in the solution or during the film-forming process. Furthermore, directional solvent evaporation and zone casting were employed to fabricate aligned fibers. Evidently, when the molecular crystallization rate matched the solvent evaporation rate, the polymers tended to form a highly oriented film and the order parameter reached as high as 0.97.

The morphology of the active layer of OSCs plays a crucial role in determining the PCE of the devices since it is closely related to the photoelectric conversion process. The ideal morphology of the active layer should comprise a pure donor and acceptor phase with a domain size of 10–20 nm and a large interfacial mixed phase, which guarantees efficient exciton separation, carrier transport, and carrier collection. However, due to the competition and coupling between crystallization and phase separation, the active layer usually possessed poor morphology, including the “sea-island” phase separation structure, low crystallinity, and inconsistent molecular orientation. Han *et al.* proposed that the application of the liquid–solid solution phase transformation resulted in phase separation, fractional crystallization, and confined crystallization; further, controlled

film-forming kinetics could optimize the film morphology.<sup>221–223</sup> They selected the P3HT:O-IDTBR blend as a model system, and they proposed to control the film-forming kinetics, *i.e.*, the crystallization of P3HT occurred ahead of O-IDTBR, to get an IPN with higher crystallinity and optimized lateral and vertical phase separations. The optimized film morphology boosted the PCE from 4.45 to 7.18%, which is the highest performance in P3HT binary non-fullerene solar cells. In addition, the molecular orientation in polymer blends is decisive to the charge transfer efficiency of these  $\pi$ -conjugated photovoltaic devices. They used epitaxial crystallization, confined crystallization, and tuning the solution state or film-drying time to control the D/A molecular orientation.<sup>224,225</sup> In P3HT/N2200 blends, molecular orientation and phase separation were controlled by chain segment and molecular movement. P3HT crystallization with face-on or edge-on orientation for different post-annealing processes is dependent on the P3HT chain segment and molecular movement confined by crystalline N2200 or broken out of N2200 crystalline confinement. For the conjugated PTB7-Th and N2200 polymers, the transformation of preferential molecular orientation from the face-on state to the edge-on state occurs when extending the film-drying time and changing the processing solvents. Therefore, the PCE of the devices was improved from 0.53 to 3.52%.

#### 4.4 WSCPs

WSCPs are a type of CP that possess solubility and processability in water-/alcohol-like polar solvents. When compared with most traditional CPs with hydrophobic alkyl groups that impart solubility in nonpolar/low-polar solvents, WSCPs possess highly polar or ionic side chains that facilitate excellent water/alcohol solubility and processability.<sup>226–228</sup> The water/alcohol solubility of WSCPs provides a promising opportunity to fabricate multilayer films using orthogonal solvents, which can be compatible with the multilayered fabrication of optoelectronic devices (Fig. 32a).<sup>229–231</sup> Moreover, the conjugated backbones of WSCPs endow them with semiconductivity that can facilitate the charge transport characteristics for various applications.<sup>232–234</sup> WSCPs have been widely used as electron/hole transport materials (ETMs/HTMs) in optoelectronic devices,<sup>226,227</sup>

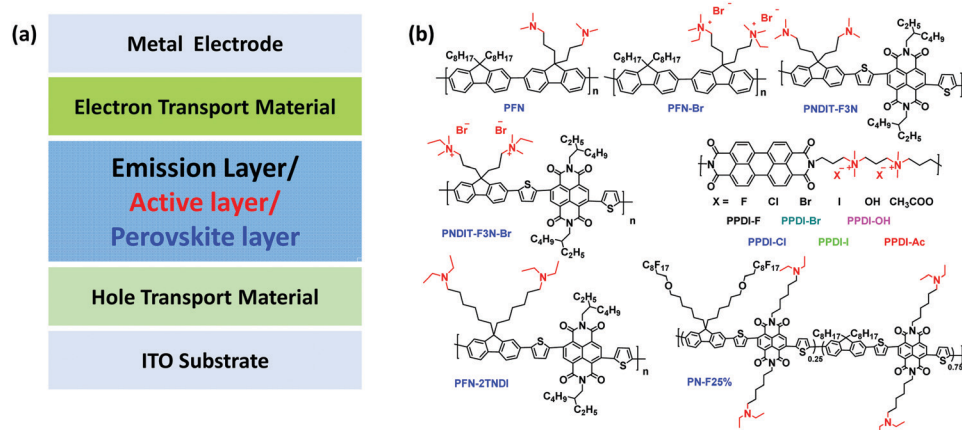


Fig. 32 (a) Structure of optoelectronic devices; (b) representative WSCPs.

including OLEDs, OSCs, perovskite solar cells (PVSCs), and organic field-effect transistors (OFETs). For example, in OSC devices, a thin layer of PFN or PFN-Br acting as the ETM can improve electron collection at the cathode.<sup>235</sup> Evidently, WSCPs in OSCs can reduce the work function of the metal electrode and optimize the active layer morphology, block holes, and dope PCBM, all of which contribute toward the enhancement of device performance.<sup>236,237</sup> However, the low electron mobilities of PFN and PFN-Br hinder their application as thick ETMs in large-area OSCs, while a thicker ETM is much more compatible with large-area OSCs processed by the roll-to-roll technique.<sup>238</sup>

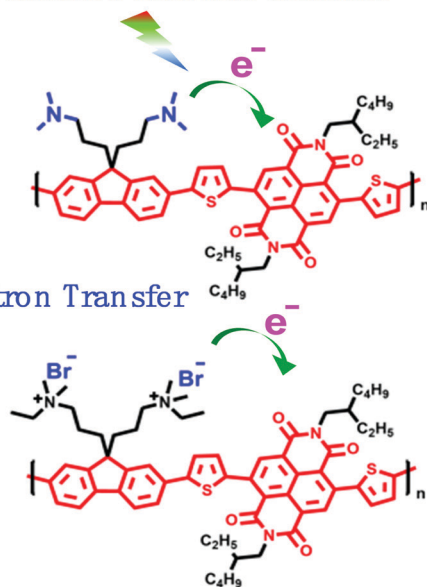
To overcome this problem, a series of novel WSCPs with self-doping behaviors have been recently developed by rational design on the polar side chains and n-type conjugated backbones.<sup>239</sup> Self-doping behaviors were observed in these n-type WSCPs due to the electron transfer from polar groups to n-type conjugated backbones. A special finding in these WSCPs was that their self-doping behaviors could be regulated by changing the polar group species. Wu *et al.* reported that neutral amino-group-functionalized WSCPs (PNDIT-F3N, Fig. 32b) employed a light-induced doping mechanism, while the ammonium-bromide-group-functionalized WSCPs (PNDIT-F3N-Br) could be self-doped without light (Fig. 33).<sup>239</sup> Consequently, PNDIT-F3N exhibited a photoinduced conductivity-enhancing property, while the conductivity of PNDIT-F3N-Br showed no difference in dark and light (Fig. 33). Moreover, the self-doping behaviors and charge transport properties of WSCPs could be readily regulated by changing the counterion type and energy levels of the conjugated backbones in WSCPs.<sup>240–242</sup> For example, Hu *et al.* prepared a series of perylene-diimide-based polyelectrolytes (PPDI-X, Fig. 32b)

with different counterions ( $F^-$ ,  $Cl^-$ ,  $Br^-$ ,  $I^-$ ,  $CH_3COO^-$ , and  $OH^-$ ) and investigated their self-doping behaviors. Evidently, polyelectrolytes with  $F^-$ ,  $OH^-$ , and  $CH_3COO^-$  as anions are strongly self-doped, and polyelectrolytes with  $Cl^-$ ,  $Br^-$ , and  $I^-$  possess weak self-doping properties.<sup>240</sup>

These self-doped WSCPs showed great promise as thickness-insensitive ETMs to enable highly efficient OSCs.<sup>239,240</sup> With PffBT4T-2OD/PC<sub>71</sub>BM as an active layer, high PCE of 10.11% for devices with 5 nm PNDIT-F3N-Br and prominent PCE of 8.04% for devices with 100 nm PNDIT-F3N-Br can be achieved.<sup>239</sup> Moreover, these self-doped WSCPs also worked well in non-fullerene solar cells.<sup>243,244</sup> Sun *et al.* reported that PNDIT-F3N largely improved electron collection and reduced charge recombination at the cathode of non-fullerene solar cells. One particular finding was that PNDIT-F3N in the interlayer also worked as the third component and induced charge transfer from the donor in the active layer to PNDIT-F3N, contributing to the photocurrent output of non-fullerene solar cells.<sup>243</sup> Moreover, it was found that a thicker ETM could also slow down and block the diffusion of metal atoms (from the metal electrode), resulting in much-improved device stability.<sup>245</sup>

In addition to the successful application of self-doped WSCPs in single-junction OSCs, WSCPs also showed promising applicability as interconnection materials for tandem OSCs.<sup>246,247</sup> Zhang *et al.* reported that an interconnection layer constructed by PNDIT-F3N-Br/Ag (2 nm)/poly(3,4-ethylenedioxythiophene) polystyrene sulfonate (PEDOT:PSS) could efficiently protect the front cell from the erosion caused by the solutions used for processing the back cell, enabling the successful fabrication of tandem OSCs.<sup>246</sup> More recent works have shown that all the

### Photo induced Electron Transfer



### Electron Transfer

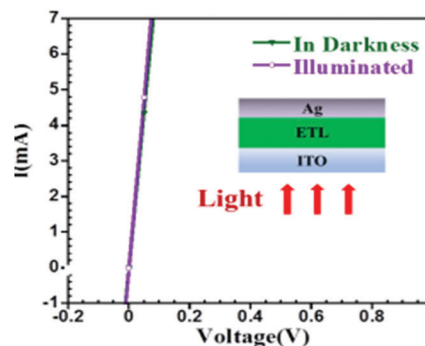
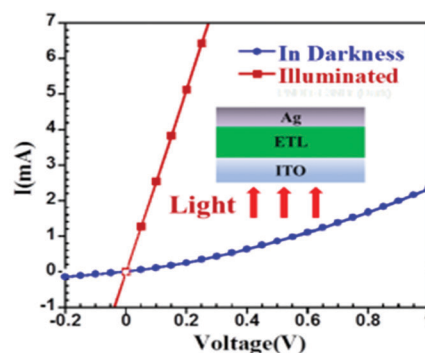


Fig. 33 Different doping processes of PNDIT-F3N and PNDIT-F3N-Br. Adapted from ref. 239 with permission. Copyright (2016) American Chemical Society.

solution-processable interconnection layers for tandem OSCs could be realized by combining a WSCP-based HTM and ETM, affording highly efficient electron–hole recombination as well as much improved photovoltaic performance.<sup>247</sup>

WSCPs were also discovered as efficient ETMs for PVSCs due to their excellent interface modification capability. Amino-functionalized WSCPs can improve interfacial contact between the perovskite layer and metal cathode, resulting in more efficient electron collection at the cathode.<sup>248,249</sup> Sun *et al.* found that the amino groups in PFN-2TNDI (Fig. 32b) could passivate the interface traps of perovskite layer, reducing interface recombinations and improving the device performance of PVSCs.<sup>248</sup> Tian *et al.* further introduced perfluorooctane side chains into a self-doped CP and obtained the hydrophobic polymer PN-F25% (Fig. 32b) with excellent water resistance capability. PN-F25% could act as a high-performance bifunctional ETM for PVSCs due to its amino and fluoro side chains, which not only enhanced electron collection but also significantly improved the device stability of PVSCs.<sup>249</sup>

In general, the design of electron acceptors in the active layer, including fused-ring and UCEAs, control morphology of the active layer using solution processing, and WSCPs are summarized. Recent progresses, molecular design strategies, and structure–property relationships of fused-ring and UCEAs are initially addressed briefly. The factors affecting the morphology of the active layer and the corresponding strategies are subsequently introduced. After that, the progress of WSCPs that can act as interfacial layers is presented. In the future, the development of high-performance electron donors and acceptors will still remain an important research direction. For instance, the high carrier mobility of electron acceptors can meet the needs of high-efficiency thick-film OPV devices. Acceptor materials that can be processed by environmentally friendly solvents can meet the requirements for the preparation of low or nontoxic OPV devices. Moreover, improvements in device stability and lifetime of OPV devices can be another important research direction.

## 5. Other electrical and optical polymer materials

Besides the aforementioned polymer materials, there are still other electrical and optical polymers that are very important for basic research and practical applications, such as conducting polymers and photosensitive polymers. These polymers can also be divided into numerous subtypes. For example, conducting polymers include two different subtypes of polymer materials. One is a polymer composite filled with conductive fillers such as carbon nanotubes (CNTs), graphene oxide, metal particles, *etc.* The other subtype is a polymer that can generate and propagate charge carriers along its backbones. In this section, we will introduce some other functional polymers that are also very important for applications in our daily life. Polyimide can be used to fabricate the flexible electrode of lithium-ion batteries (LIBs) with carbon materials,<sup>250</sup> it can also be used in the

wireless communications industry due to its low dielectric constant ( $k$ ). Second-order nonlinear optical (NLO) polymers have been drawing considerable attention due to their potential applications in photonic devices. Therefore, we address these functional polymers with respect to their preparation methods, design strategies, and potential applications in this section.

### 5.1 Intrinsic low- $k$ polyimide materials

With the development of the wireless communication industry, the high speed and high-frequency transmission, particularly the fifth generation of wireless communications technology (5G technology), has been a hot research subject.<sup>251</sup> Both signal transmission speed ( $V$ ) and signal propagation loss rate ( $\alpha$ ) are closely related to  $k$  and dielectric loss of the substrate materials, which can effectively reduce the resistance–capacitance delay, line-to-line crosstalk noise, and power dissipation. As a result, the accuracy and integrity of signal transmission can be ensured, and the service life of electronic components can also be improved.<sup>252</sup> Therefore, low- $k$  and low-loss dielectric materials are highly desirable in the wireless communications industry.

There are two ways to design low- $k$  dielectric polymers. One is the introduction of nanosized air voids ( $k \sim 1$ ) into the bulk polymers. The dielectric constant of these porous polymers can be reduced to a certain level, but the overall properties of the materials can be considerably damaged, limiting their practical applications.<sup>253</sup> The other is to design the dielectric properties of materials at the molecular level. For example, some chemical bonds with lower molar polarizability (such as the  $-C-F$  bond) or some rigid and shape-persistent components (such as triptycene moiety) can be introduced into the molecular structures of the polymer.<sup>254</sup> When compared with porous polyimide materials, the intrinsic low- $k$  polyimide materials have practical application value with regard to both preparation process and comprehensive performance. However, the  $k$  values of most of the developed low- $k$  polyimide materials can only reach around 2.3. Therefore, the development of flexible high-performance intrinsic low- $k$  or ultralow- $k$  polymer materials remains a formidable challenge in the microelectronics industry.

Recently, a highly efficient design strategy to fabricate high-performance intrinsic low- $k$  polyimides has been developed by Zhang and Xu *et al.* by the use of propeller-like structures (such as triphenylamine, triphenylmethane, and tristyrene units) in order to form a series of novel functional diamine monomers with rigid nonplanar large conjugated structures.<sup>255–261</sup> By regulating the chemical structures and aggregation structures of the polymers at the molecular level, low- and even ultralow- $k$  polyimide films with excellent comprehensive properties can be obtained (Fig. 34).<sup>260</sup>

Liu *et al.* designed and synthesized novel, simple, and efficient diamine (TriPEDA and TPEDA)-containing rigid nonplanar conjugated triphenylethylene and TPE moieties through the Wittig–Horner and Suzuki coupling reactions.<sup>255,256</sup> A series of high-performance functional polyimides were consequently prepared by the dipolymerization of TriPEDA/TPEDA and four dianhydrides (Fig. 35). Because of the introduction of the aromatic rigid nonplanar triphenylethylene/TPE structure,

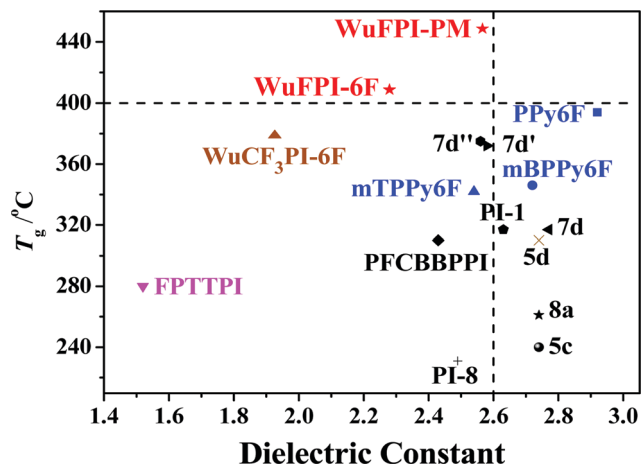


Fig. 34 Relationship between  $k$  and  $T_g$  of the polyimides developed by Zhang's group and the reported intrinsic low- $k$  polyimides. Reprinted from ref. 260 with permission. Copyright (2019) from Springer.

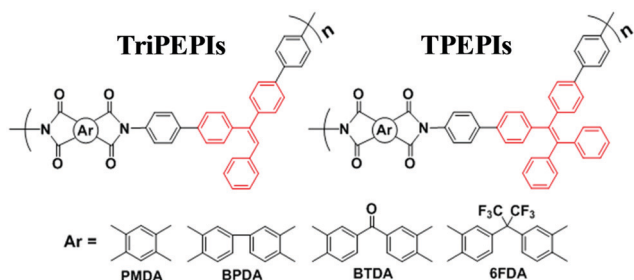


Fig. 35 Low- $k$  polyimides containing triphenylethylene and TPE moieties. Adapted from ref. 255 and 256 with permission. Copyright (2012 and 2013) American Chemical Society and Wiley-VCH.

the polyimides exhibited the following: low  $k$  (2.63–2.76); excellent mechanical properties with high tensile strength and high tensile modulus in the range of 115–130 MPa and 2.2–2.7 GPa, respectively; and high glass transition temperatures ranging between 359 and 443 °C. Moreover, these organo-soluble polyimides exhibited special fluorescent and resistive switching (ON/OFF) characteristics, as the maximum fluorescence emission of the four polyimides was observed at 425–505 nm in the NMP solution and 470–541 nm in the film states.

Chen *et al.* prepared novel fluorinated aromatic polyimides by the conventional two-step imidization of 4,4'-(hexafluoroisopropylidene)diphthalic anhydride (6FDA) and diamines (TriPMPDA and TriPMMDA)-bearing triphenyl methane moiety (Fig. 36).<sup>257</sup> Both flexible and tough polyimide films exhibited intrinsic low  $k$  values of 2.56 and 2.33 at a frequency of 10 kHz, respectively, due to the introduction of bulky triphenyl methane side groups as well as tortuous backbone structures. In addition, they showed light color, high thermal stability, moderate mechanical property, and more importantly, excellent solubility in common organic solvents. Therefore, both these functional polyimides possessed attractive potential applications in the field of high-performance flexible polymer interlayer materials.

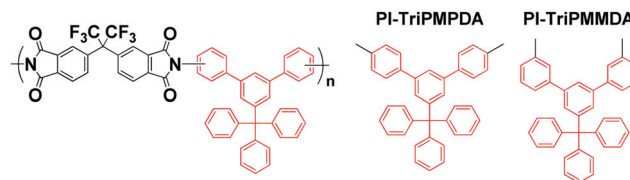


Fig. 36 Low- $k$  polyimides containing triphenylmethane moieties. Adapted from ref. 257 with permission. Copyright (2016) Elsevier Ltd.

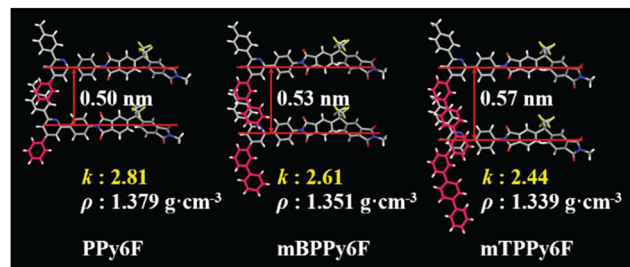


Fig. 37 Intrinsic low- $k$  polyimide containing the diphenylpyridine structure. Adapted from ref. 258 with permission. Copyright (2017) Royal Society of Chemistry.

Bei *et al.* successfully synthesized three polyimides (PPy6F, mBPPy6F, and mTPPy6F) containing the same rigid nonplanar conjugated diphenylpyridine structure in the main chains and a different number of benzene rings in the side chains (Fig. 37).<sup>258</sup> The polyimides having one, two, and three benzene rings in the pendant group showed intrinsic  $k$  values of 2.81, 2.61, and 2.44, respectively. Owing to the rigid nonplanar conjugated diphenylpyridine structure in the main chain, PPy6F had a lower  $k$  value of 2.81 as compared to that of the traditional polyimide 6FDA-ODA (3.0). The  $k$  values of PPy6F, mBPPy6F, and mTPPy6F presented regular decreasing tendency. Based on the results of WAXD, morphology, and density studies, it was revealed that the free volume of the polyimides increased with an increase in the number of benzene rings in the pendant group. Meanwhile, mTPPy6F showed excellent comprehensive properties, such as  $T_g$  of 342 °C, decomposition temperature ( $T_{d,5\%}$ ) of 551 °C, tensile strength of 105 MPa, low moisture absorption of 0.61%, and good solubility, making it a competitive material for potential applications in microelectronics industries.

Furthermore, a series of high-performance multifunctional polyimides with intrinsic ultralow- $k$  values ( $\sim 1.93$ ) as well as exceptional thermostability and solubility ( $T_g$  as high as 494 °C, and, at the same time, worthwhile solubility in common organic solvents) were successfully designed and synthesized by introducing a typical aromatic rigid trifluoromethyl-containing moiety with special nonplanar and conjugated characteristics into the polymer backbone (Fig. 38).<sup>259,260</sup> In addition, these polyimides showed light color (even colorlessness in one case) and transparency as well as electrical bistability characteristics (ON/OFF ratio as high as  $10^7$ ; working voltage as low as 1.5 V). The excellent thermal stability and solubility allowed them to undergo the high-temperature process (over 400 °C) in the

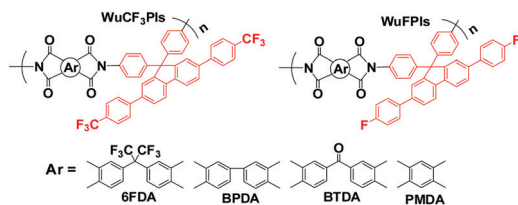


Fig. 38 Multifunctional polyimides containing trifluoromethyl moieties. Adapted from ref. 259 and 260 with permission. Copyright (2017 and 2019) Royal Society of Chemistry and Springer.

preparation of photoelectric devices (such as PVD and PECVD) or the highly efficient, continuous roll-to-roll process. Ideally, the as-synthesized polymers are potential candidates for practical applications in the fields of ultra-large-scale integration (ULSI), high-performance polymer memory devices, flexible displays, thin-film photovoltaic industries, and wearable electronics.

A bulk dielectric polymer film with an intrinsic ultralow- $k$  value of 1.52 at 10 kHz was successfully synthesized based on the novel polyimide, FPTTPI (Fig. 39).<sup>261</sup> It is well known that this  $k$  value is the highest among the intrinsic ultralow- $k$  polymers in the literature. More importantly, such outstanding dielectric properties remained stable until up to 280 °C. The excellent ultralow dielectric properties could be mainly attributed to the large free volume (sub-nanoscale), which intrinsically existed in the amorphous region of polymeric materials. Meanwhile, FPTTPI also showed excellent thermal stability and mechanical properties,  $T_g$  of 280 °C, decomposition temperature of 530 °C, and residual of 63% at 800 °C under N<sub>2</sub>. It was soluble in common solvents, which makes it possible to be used in simple spin-on or efficient, low-cost, and continuous roll-to-roll processes.

In general, polyimides have shown potential applications in the wireless communications industry due to their low- $k$  values and low dielectric losses. Therefore, the design strategies and preparation methods for intrinsic low- $k$  polyimides with good properties are discussed. Although a polyimide with an

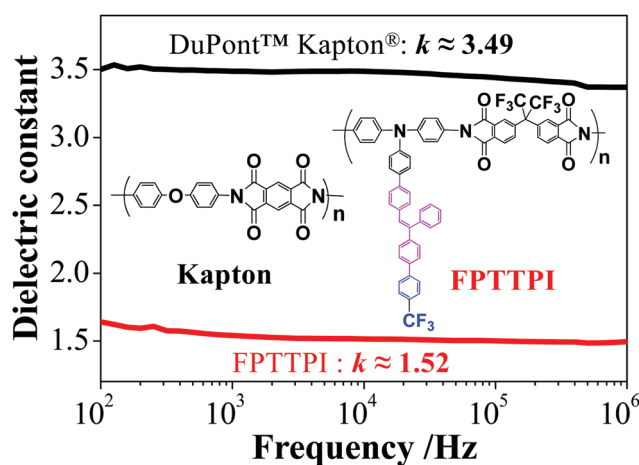


Fig. 39 Dielectric properties of the FPTTPI film and a DuPont Kapton polyimide film. Reprinted from ref. 261 with permission. Copyright (2015) American Chemical Society.

intrinsic ultralow- $k$  value (as low as 1.52) has been obtained, it still remains a challenge to obtain a lower  $k$  value while retaining other worthwhile properties.

## 5.2 Second-order NLO polymers and dendrimers

Organic second-order NLO materials have attracted considerable attention owing to their potential applications in photonic devices, such as high-speed electrooptic modulators, optical switches, and frequency converters. Until now, many strategies and approaches have been proposed to meet the basic requirements of practical applications, including large optical non-linearity, low optical loss, and excellent temporal stability of dipole orientation. A major problem encountered in organic NLO materials is to overcome the strong dipole-dipole interactions among the chromophores with donor- $\pi$ -acceptor (D- $\pi$ -A) structures. Therefore, it is crucial to efficiently translate the large  $\beta$  values of organic chromophores into high macroscopic NLO activities of polymers, which is mainly based on the orderly arrangement of the chromophore moieties with noncentrosymmetric forms. However, due to the strong electrostatic interaction of the chromophore moieties with electronic push-pull structures, the centrosymmetric arrangement is the natural state with nearly no macroscopic second-order NLO coefficient. Therefore, many approaches have been explored to reverse the orientation of such chromophores, with the aim to achieve favorable molecular arrangement, for instance, a poling process. Further, the optimized molecular shapes and electronic properties of chromophores play essential roles in molecular packing of thin films, which are mainly based on the site isolation principle, that is, the more spherical the molecular structure, the larger is the reduction in dipole-dipole interactions.

Hence, systematical studies were conducted by Li *et al.* to optimize the molecular alignment of organic chromophores in both polymers and dendrimers. First, according to the site isolation principle, the larger the sizes of the side groups, the better was the isolation effect; this led to the noncentrosymmetric arrangement of the chromophores. On the other hand, this decreased the loading density of the chromophore moieties, which could weaken the NLO coefficient ( $d_{33}$ ). Therefore, there could exist a most suitable state in which the minimum size of the isolation group could prevent interactions between the adjacent chromophore moieties, and the maximum loading density could be guaranteed at the same time. To reach this state, Li *et al.* conducted systematic work and proposed the concept of a “suitable isolation group (SIG).” That is, for a given chromophore, there could be a SIG present to ensure the noncentrosymmetric arrangement of the chromophores, thereby achieving possibly improved macroscopic NLO properties in the organic materials. For example, various NLO effects could be obtained in linear polymers bearing the same nitroazobenzene chromophore modified by different isolation groups and different linkage modes. In most of the cases, it was clear that the NLO effect initially increased and then decreased with greater sizes of isolation groups: P2 exhibited the largest  $d_{33}$  value with the phenyl moiety as the SIG. However, for different

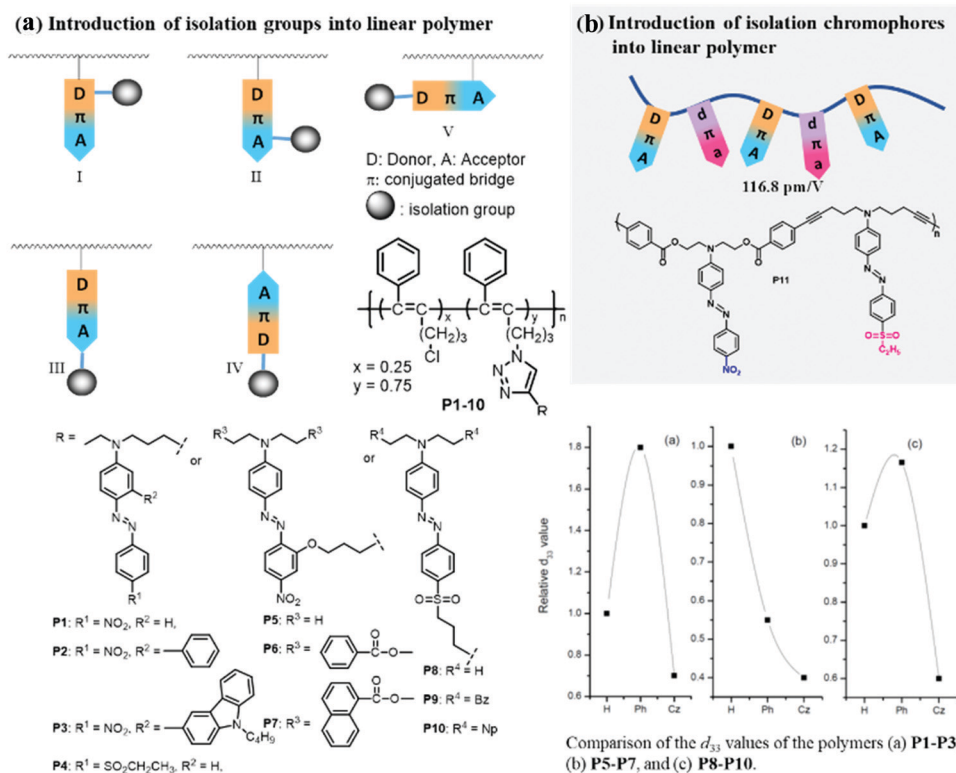


Fig. 40 Structures of linear polymers P1–11 and schematic illustrations of isolation groups and isolation chromophores. Adapted from ref. 262 and 263 with permission. Copyright (2007 and 2013) American Chemical Society and Royal Society of Chemistry.

linkage modes, SIG is considerably different for the different arrangements of chromophores as the natural state (Fig. 40).<sup>262</sup> Since the isolation group was not the valid unit for the NLO effect, they subsequently proposed a more efficient strategy by the introduction of isolated chromophores bearing different polarities with the main ones. As shown in Fig. 40, P11 with alternative chromophore moieties (nitroazobenzene and sulfonylazobenzene ones) afforded a larger  $d_{33}$  value ( $116.8 \text{ pm V}^{-1}$ ) as compared to those obtained from polymers with single chromophores ( $78.1$  and  $34.3 \text{ pm V}^{-1}$ ), confirming that favorable arrangement can be achieved with tunable dipole–dipole interactions.<sup>263</sup>

As the most promising candidates for second-order NLO materials, dendrimers with a special topological structure were also employed with the optimized chromophores as repeat units (Fig. 41). Excitingly, because the SIG concept was utilized, the NLO effect could be dramatically enhanced with increasing dendritic generations because of the dendrimer (dendritic) effect.<sup>264</sup> Moreover, this could be further enhanced by modifying the topological structures. As shown in Fig. 41, X-type dendrimers exhibited almost parallel alignment of the chromophore units with fixed orientation as the initial state, which could help in maintaining the noncentrosymmetric arrangement during the electronic poling process.<sup>265,266</sup> Accordingly, the Janus dendrimer X-1 possessed a  $d_{33}$  value of  $299 \text{ pm V}^{-1}$ , which is the highest value reported so far for polymers containing a simple azo-chromophore moiety, further confirming the advantage of Janus dendrimers and the key role of the molecular arrangement for the NLO property. With structures similar to

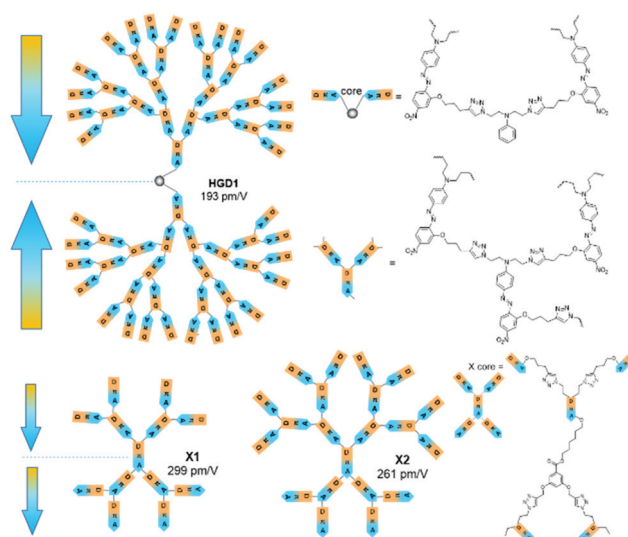


Fig. 41 Chemical structures of the high-generation dendrimer HGD1 and X1 and X2 with a Janus structure.

those of dendrimers, HBPs can be prepared very easily, typically by one-pot syntheses, which are likely to have more industrial and commercial values.<sup>267</sup> By introducing isolation chromophores derived from  $AB_4$ -type monomers by means of click chemistry, the resultant HBPs demonstrated good NLO properties and optical transparency, making them promising candidates for practical applications.<sup>268</sup>

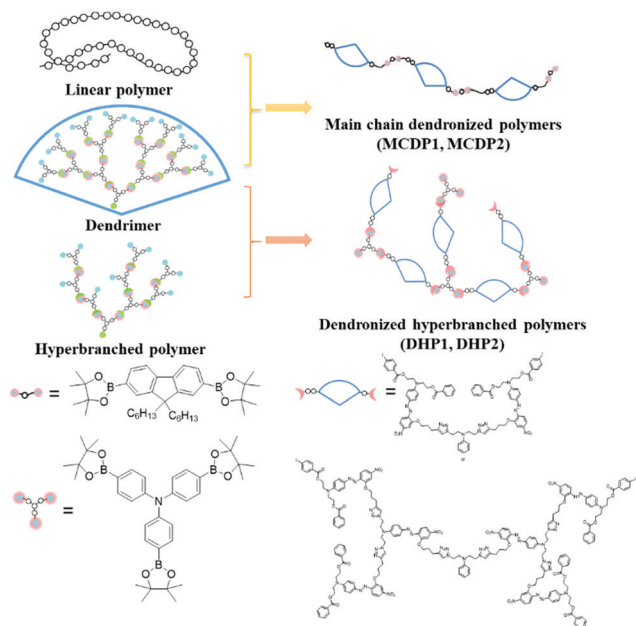


Fig. 42 Chemical structures of the main-chain dendronized polymers and DHPs.

Furthermore, when considering dendrimers with perfect conformations and linear and HBPs with easy synthesis strategies, their combined advantages can be exploited by the introduction of dendrimers into linear or HBPs with rational molecular designs. Main-chain dendronized polymers, namely, MDPG1 and MDPG2 (Fig. 42), which were prepared by the macromonomer approach, simultaneously exhibited the advantages of both main-chain polymers (which usually demonstrate good stability of the NLO effect) and dendronized polymers (which usually possess a large NLO coefficient).<sup>269</sup> With the incorporation of low-generation dendrimers as the monomer in the HBP backbone, a novel kind of polymer, namely, dendronized hyperbranched polymers (DHPs) (Fig. 42), was proposed by Li *et al.* in 2013.<sup>270,271</sup> This special system could maintain both high NLO efficiency and improved NLO thermal stability, partially combining the advantages of HBPs and dendrimers. For an A<sub>2</sub>-type monomer with a dendrimer as the side chain, after being polymerized with a B<sub>3</sub>-type monomer of the TPA–borate ester, polymer DHP1 exhibited a high  $d_{33}$  value of 133 pm V<sup>-1</sup>, and the decay temperature was as high as 110 °C, which were much better than those of the corresponding dendrimer (100 pm V<sup>-1</sup>, 30 °C).

In summary, based on some typical examples, recent progresses made in NLO dendrimers and polymers as developed by Li *et al.* are demonstrated, mainly focusing on molecular design ideas at the molecular level, as well as an attempt to determine the structure–property relationships to aid the rational design of better NLO materials for further investigations and possibly render designs that can be emulated by other functional polymers.

## 6. Biorelated polymer materials

Due to the aging of the global population, environmental problems caused by economic development have resulted in

higher incidence of major diseases such as cancer, cardiovascular diseases, and diabetes; further, these patients tend to be younger. It is highly desirable to develop new technologies for diagnosing and treating diseases. Polymer materials have been playing an increasing important role in the diagnosis and treatment of major diseases such as cancer, cardiovascular and cerebrovascular diseases, and infectious diseases. A large number of functional polymers, such as micelles, gels, composites, fibers, particles, brush polymers, SMPs, *etc.*, have been investigated for biomedical applications for many years. Their characteristics such as thermal and chemical stabilities, shape, dimensions, surface charge, surface chemistry, mechanical properties, and porosity may be tuned by using certain strategies to achieve the targeted functionalities that can meet the demands of specific biomedical applications. Therefore, recent progress in functional polymers and their corresponding strategies have become highly desirable. In this section, we only introduce polymers that are still in the state of basic research, including polymerization for protein modification, medical imaging materials, drug and gene delivery materials, biomedical surface–interface materials, integrated disease diagnosis and treatment materials, *etc.* Marine biofouling can accelerate bio-corrosion and enhance hydrodynamic drag; therefore, it is highly desirable to develop new coatings for antibiofouling. Hence, we also address this type of functional polymers on the basis of preparation methods, design strategies, and applications in this section. Polymer gels and SMPs for biomedical applications will be addressed in the subsequent sections.

### 6.1 *In situ* polymerization for protein modification

After thousands of years of evolution, the natural structures of proteins are considered to be in their most optimized form. When used in therapeutics, biosensors, or industrial catalysts, catalytical proteins play key roles in enhancing the quality of life. Most of these proteins, however, have currently been utilized in brand new areas, far away from their original ecologies, resulting in an inevitable need for structural modification. For example, cells make superoxide dismutase (SOD) to control intracellular superoxide from DNA mutation, while SOD has been used in sunblock to protect our skin from sunburn. A new application environment breeds additional modifications to the protein structure. Until now, various chemical strategies have been explored to facilitate protein modification for better functions, including liposome encapsulation, polymeric conjugation, supramolecular encapsulation, inorganic adsorption, and so on.<sup>272,273</sup>

Polymeric materials are the most popular choices for protein modification for obtaining diverse functions: *in situ* polymerization of the protein structure is a feasible and efficient method. A generic *in situ* polymerization reaction for proteins comprises two steps: (1) amine-reactive acrylates are conjugated to the protein surface in order to introduce polymerizable groups; (2) initiating the *in situ* polymerization of hydrophilic monomers in aqueous media at room temperature yields a thin polymeric shell around each protein molecule. Such a polymeric shell could offer not only enhanced thermodynamic

stability to the protein but also additional functions depending on the type of monomer.<sup>274</sup>

For example, protein therapeutics usually suffer from high immunogenicity, which might result in shorter circulation times and allergies; moreover, the *in situ* polymerization of a nonfouling polymeric shell, such as poly(2-methylacryloyloxyethyl phosphorylcholine) (pMPC) or poly(*N*-vinylpyrrolidone) (PVP) on the surface of protein therapeutics, can shield immunological recognition and clearance, yielding high biological stability for protein therapeutics (Fig. 43).<sup>275–278</sup>

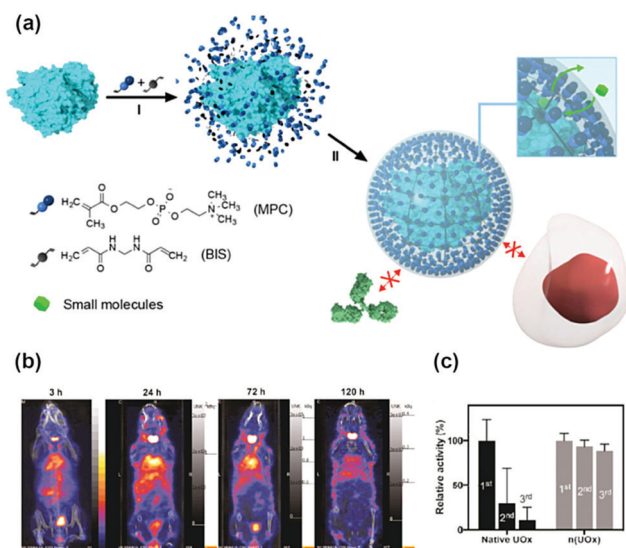


Fig. 43 (a) *In situ* polymerization route of the MPC monomer for building nonfouling surfaces on proteins; (b) SPECT/CT images of a mouse after the intravenous administration of radioactive <sup>125</sup>I-labeled pMPC-modified protein; a significant concentration of protein persists even after 120 h; (c) serum urate oxidase (UOx) activities after the repetitive administration of native UOx or UOx nanocapsules (n(UOx)), confirming the ability of the nanocapsules to evade an adaptive immune system. Reprinted from ref. 276 with permission. Copyright (2016) from Springer.

A prolonged plasma circulation time can be observed (as high as 7 days) with low level of immunogenicity. This *in situ* polymerization technique can be universally applied to a library of proteins, with different sizes, surface charges, and structures.

Furthermore, the *in situ* copolymerization of two or three monomers can also ensure that the protein structure exhibits multiple functions. For example, the copolymerization of MPC and specific monomers can induce additional functions to protein therapeutics, such as targeted delivery (with folic-acid-modified monomers) as cancer therapy or blood–brain barrier (BBB) penetration (with TAT-modified monomer or mustard-modified monomers) as a part of Parkinson's therapy (Fig. 44).<sup>279–282</sup> Based on a prolonged circulation time as well as low immunogenicity, all these therapies exhibited significant delivery efficiency. A proper mechanism might involve a longer circulation time that can statistically provide more possibilities; based on this, the specific interaction between the protein and targets becomes significant.

As most functional proteins favor aqueous media, the hydrophobic modification of proteins is always cumbersome, which might afford an unstable dispersed system with protein denaturation. However, hydrophobic modification is desirable for certain applications, such as heterogeneous catalysis and electrochemistry applications. The *in situ* copolymerization of one hydrophobic monomer with one hydrophilic monomer can offer the opportunity to introduce hydrophobic segments into the protein structure, which might induce additional photo-electronic properties to the protein functions.

For example, in the electrochemistry applications of proteins, such as biosensors and biofuels, the protein converts matter signals (substrate concentrations) into electrical signals *via* substrate catalysis. Most proteins yield a fast electron turnover rate, resulting in high sensitivity or power output. However, the protein structures have super low conductivity, and the distance between the active center and protein surface is considerably higher than the electron tunneling distance

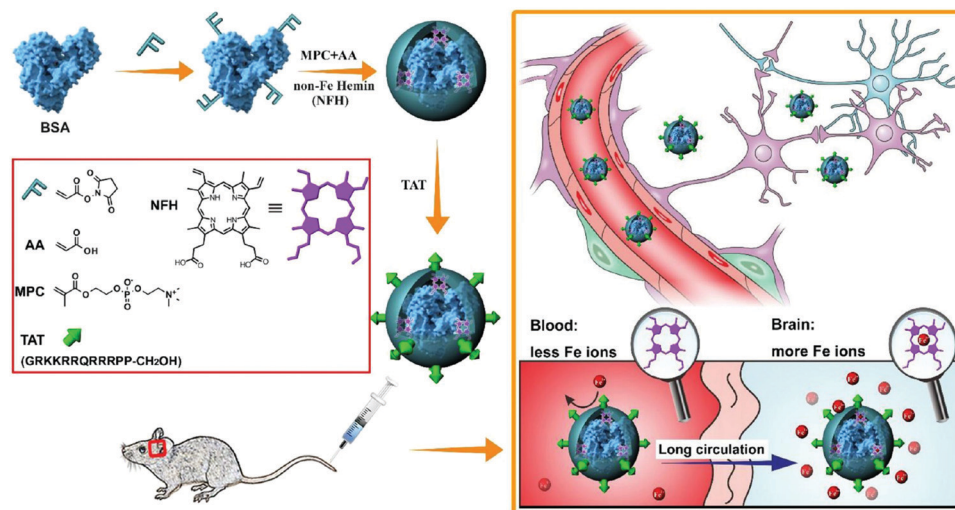


Fig. 44 TAT-modified protein obtained by *in situ* copolymerization and its mechanism of BBB penetration to treat Parkinson's disease. Reprinted from ref. 280 with permission. Copyright (2017) American Chemical Society.

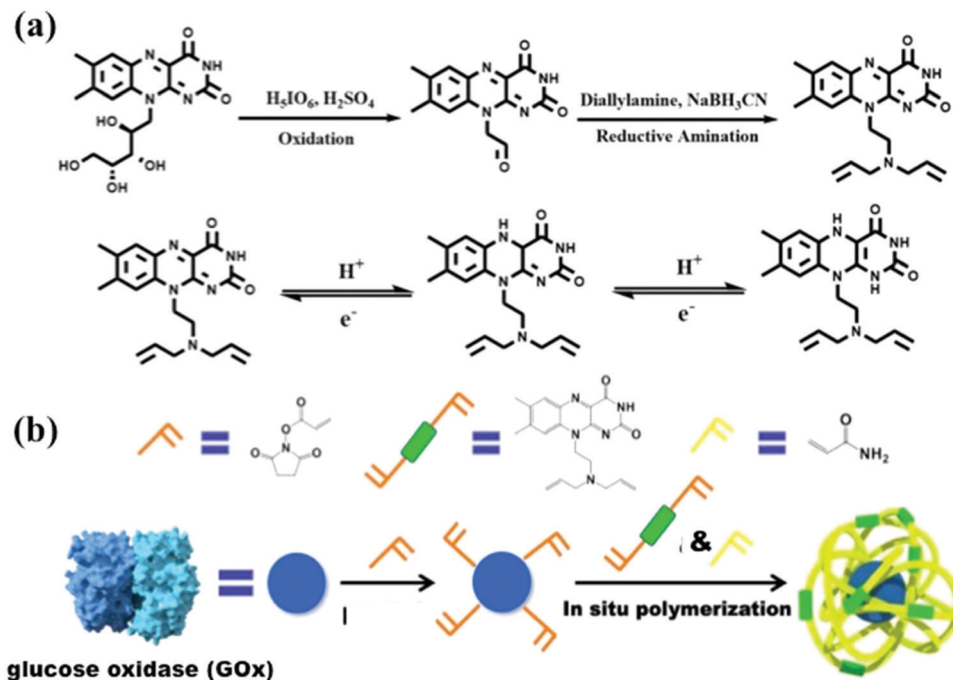


Fig. 45 (a) Synthesis routes of DAA-Flavin and its redox mechanism; (b) *in situ* copolymerization of a hydrophobic monomer and a hydrophilic monomer. Reprinted from ref. 283 with permission. Copyright (2018) American Chemical Society.

obtained from the Marcus theory; therefore, the electron transfer rate is much lower in applications. Jin *et al.* designed a cofactor-like molecule, 2'-diallylamino-ethyl flavin (DAA-Flavin), derived from riboflavin and incorporated as a crosslinker into the polyacrylamide (PAAm) network around the GOx surface by *in situ* polymerization (Fig. 45).<sup>283</sup> This incorporated DAA-Flavin could act as an alternative electron acceptor to interact with the active centers of GOx in the presence of the substrate, as well as initiate direct electron transfer to the protein surface. Based on these functions, DAA-Flavin-modified GOx can yield enhanced catalytic activity toward glucose as well as the direct sensing of glucose concentration *via* fluorescence signal changes.

A variety of polymerization techniques, such as free radical polymerization, ATRP, and RAFT, can be applied *in situ* for protein modification under mild reaction conditions. Based on these mature polymerization processes, the modification of proteins has become controllable and variable. Because of the diverse choice of polymeric materials, *in situ* polymerization offers promising diversities in protein structures and functions. Last but not least, the scale-up manufacturing of such modified proteins has become bright and worthwhile.

## 6.2 CPs for biosensing, imaging, and therapy

CPs are organic macromolecules that are characterized by large  $\pi$ -electronic delocalized backbones, which are capable of effectively coordinating electronic coupling to enhance the light-harvesting and light-amplifying abilities, as discussed in Section 4. It is worth mentioning that the tuning of the compositions and structures of the polymer backbones and pendants allows control over their optical properties. As a result, CPs provide a versatile platform in resolving biomedical

issues, such as designing optical biosensors as well as fluorescence imaging and therapeutics.<sup>284–287</sup>

**DNA sensing.** DNA methylation is an important component of epigenetic regulation, and hypermethylation ultimately leads to aberrant protein expression as well as cancer formation and progression. A protocol for site-specific CpG methylation detection has been established by Wang *et al.*, as shown in Fig. 46.<sup>288</sup> In this protocol, a methylation-sensitive restriction endonuclease (HpaII) was introduced to specifically digest unmethylated recognition sites that were directly derived from a small amount of genomic DNA, leaving the methylated DNA intact. The subsequent nested PCR amplification can exclusively incorporate Fl-dNTPs into the PCR products for methylated DNA, but not for the unmethylated one. Upon the addition of a cationic conjugated polymer (CCP), distinct fluorescence resonance energy transfer (FRET) from the CCP to Fl could be observed for situation B. The DNA methylation levels of seven colon-cancer-related genes in the Chinese population have been quantitatively analyzed using dyes exploiting FRET from the CCP to Fl. By means of a stepwise discriminant analysis and the cumulative detection of methylation alterations, high accuracy and sensitivity for colon cancer detection (86.3 and 86.7%) and for differential diagnosis (97.5 and 94%) could be obtained. Moreover, a correlation between the CpG island methylator phenotype and clinically important parameters in patients with colon cancer was determined. The cumulative analysis of promoter methylation alterations by the CCP-based FRET may be useful for the screening and differential diagnosis of patients with colon cancer, as well as for performing clinical correlation analyses.

**Bioimaging.** Imaging techniques have attracted considerable attention as a visualization tool for their potential

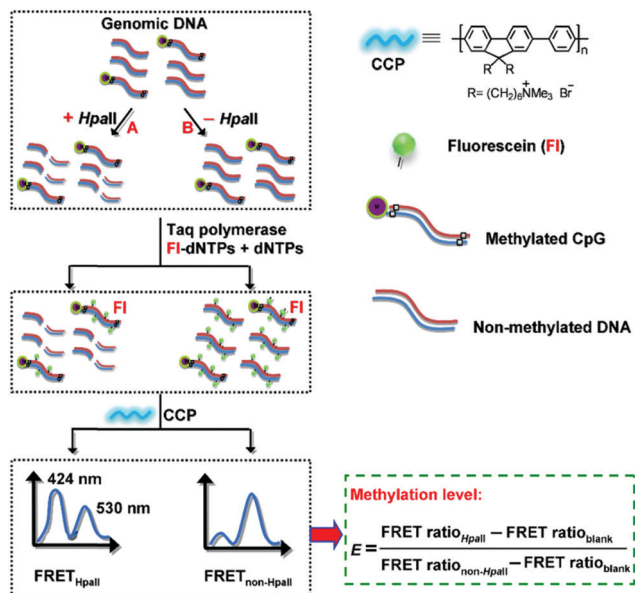


Fig. 46 Schematic illustration of the CCP-based FRET technique for DNA methylation detection. The methylation level ( $E$ ) of DNA is calculated according to FRET ratios ( $I_{530\text{ nm}}/I_{424\text{ nm}}$ ) of HpaII-treated, untreated, and blank control samples. Reprinted from ref. 288 with permission. Copyright (2012) from Nature Publishing Group.

applications in the biomedical field due to their distinguished advantages in the development of imaging agents. CPs with excellent optical properties have been regarded as imaging celebrities and elitists in multiple domains of bioimaging, such as multicolor and NIR imaging, two-photon imaging, super-resolution imaging, persistent luminescence imaging, photoacoustic imaging, and Raman imaging.<sup>285–287</sup> Considering the fact that each commercial optical instrument has a different excitation source, a targeted fluorescent material that could match such diverse excitation sources to yield multicolor emissions has become imperative. Conjugated polymer nanoparticles (CPNs) with obvious advantages have been developed as a class of promising fluorescence materials. Simple synthesis and versatile surface modification endow CPNs with the potential for multiplexed and targeted imaging. Wang *et al.* reported a method for the targeted imaging of tumor cells based on multicolor CPNs.<sup>289</sup> Four CPs with blue, green, yellow, and red emissions were synthesized to coprecipitate with poly(styrene-*co*-maleic anhydride) (PSMA), forming carboxyl-functionalized CPNs (Fig. 47a). Through multistep FRET, CPs with shorter wavelength emissions could serve as the donor to transfer the energy to longer-wavelength-emissive ones, resulting in multicolor CPNs that can absorb and emit in the entire visible-light region under a single excitation wavelength of 360 nm (Fig. 47b and c). To improve the specificity of the targeted imaging of tumor cells, the obtained CPNs were further modified with the antibody, such as anti-EpCAM. The targeted imaging of MCF-7 cells showed that CPNs could be excited at 405, 488, and 559 nm, which emitted blue, green, and red fluorescence, respectively (Fig. 47d). Raman imaging is considered to be a revolutionary imaging technique for providing

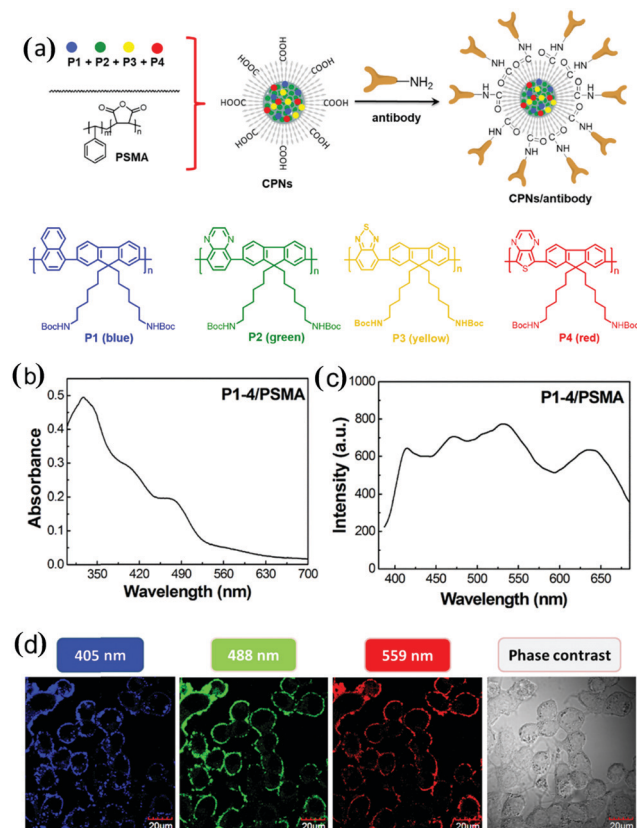


Fig. 47 (a) Schematic illustration of the preparation of P5–8/PSMA CPNs and the conjugation of antibody to the CNPs for target imaging. (b) UV-vis absorption and (c) fluorescence emission spectra of CPNs in aqueous solutions ( $\lambda_{\text{ex}} = 360\text{ nm}$ ). (d) Multichannel fluorescence images of MCF-7 cells incubated with CPNs/anti-EpCAM. The excitation wavelengths are 405, 488, and 559 nm, respectively. Adapted from ref. 289 with permission. Copyright (2014) Wiley-VCH.

direct vibrational information in the molecule fingerprint. Because of its inherent advantage, Raman imaging has a narrow-resolved band (0.1 nm) when compared with its fluorescence (10–50 nm), which can facilitate the simultaneous visualization of multiple molecules or bands. Recently, CPs with enhanced Raman activity were explored by Wang *et al.*, which opens up a new avenue into the high-speed development of Raman-responsive probes based on CP materials.<sup>290</sup>

**Therapeutics.** Significant advances of CPs for use in disease therapeutics have also been achieved in recent years, providing new tools for resolving the intractable puzzles involving pathogens and cancer treatments. Recent efforts have been focused on exploiting new photoactivable CPs and developing new phototherapy for pathogen and cancer, where photon transformation is employed to kill the pathogen and tumor cells in a remote-controlled manner. Photodynamic therapy (PDT) is an advanced treatment modality that normally employs cytotoxic reactive oxygen species (ROS) to eliminate pathogens and cancer cells, where CPs function as photosensitizers and are activated by light. The requirement of an outer light source limits the effective application of PDT to lesions in deeper tissues. To eliminate this limitation, Wang *et al.* developed a

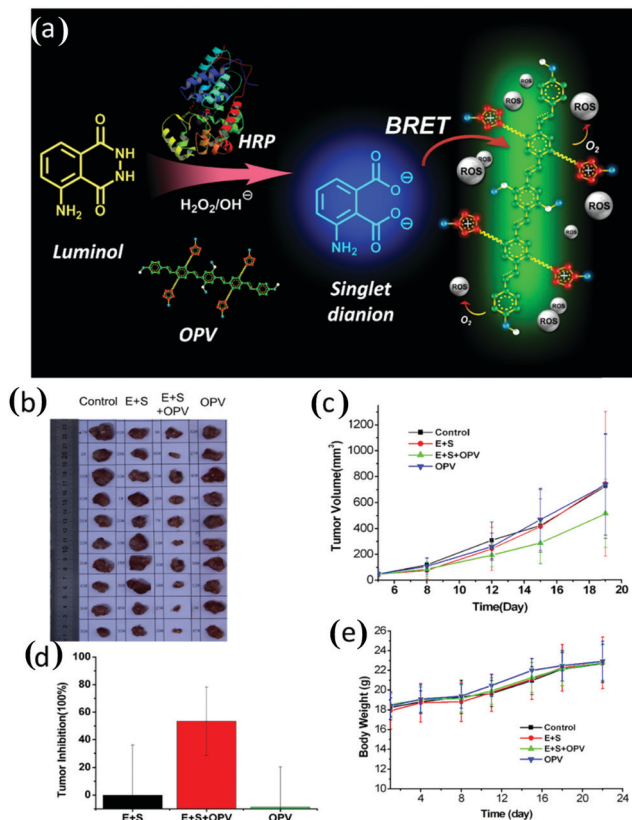


Fig. 48 (a) Schematic illustration of the BRET system for PDT and the synthesis of cationic OPV as a photosensitizer. (b) Representative photographs of mice and tumors collected from the tumor-bearing mice before and after 18 days of treatments with the BRET system. (c) Tumor volume as a function of treatment time. (d) Tumor inhibition of mice after 18 days of treatment. (e) Body weight curves after various treatments. Adapted from ref. 291 with permission. Copyright (2012) American Chemical Society.

novel PDT system for killing cancer cells in which the photosensitizer was activated by chemical molecules instead of a light source (Fig. 48a).<sup>291</sup> To demonstrate this concept, luminol, hydrogen peroxide, and HRP were used as the bioluminescent molecules and a cationic OPV was used as the photosensitizer. Bioluminescence resonance energy transfer (BRET) occurs between luminol and OPV since they meet the spectral overlap requirement. The excited OPV sensitizes the surrounding oxygen molecules to generate ROS that kill the adjacent cancer cells both *in vitro* and *in vivo*. The antitumor therapeutic efficacy of such a novel BRET system was evaluated in HeLa cell tumor-bearing nude mice *via* intratumoral injection; a tumor inhibition ratio of 55% could be obtained (Fig. 48b–d). By avoiding the use of light irradiation, this work opens up a new therapy modality to treat tumors and pathogen infections.

Recently, Wang *et al.* constructed, for the first time, an electricity-driven luminous system based on electrochemiluminescence (ECL) for killing pathogenic bacteria, where ECL was used as excitation of the photosensitizer instead of a physical light source in order to produce ROS (Fig. 49).<sup>292</sup> The mechanism for ECL therapeutics depends on the perfect spectral overlap and energy transfer from the ECL generated by luminol

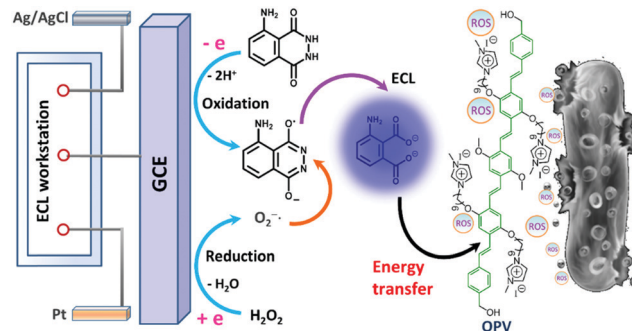


Fig. 49 Schematic illustration of electricity-driven mechanism for ECL therapeutics to produce ROS and kill pathogenic bacteria. Ag/AgCl electrode: reference electrode; Pt electrode: counter electrode; glassy carbon electrode (GCE): working electrode. Reprinted from ref. 292 with permission. Copyright (2018) American Chemical Society.

to the photosensitizer (cationic OPV), thereby sensitizing the surrounding oxygen molecule into ROS. Furthermore, stretchable and persistent hydrogel devices could be designed by integrating stretchable hydrogel, persistent ECL, and antibacterial function into hydrogel matrices. This novel strategy avoids the use of an external light source, making the process simple, convenient, and controllable. This affords new fields for ECL beyond sensors and also opens up another novel model for PDT.

CPs are rapidly emerging as a new photonic platform that can not only be used to monitor and image biophysical markers and tumor metastasis both *in vitro* and *in vivo* but also deliver drugs and intelligently trigger photo-attacks. It is believed that the considerable progress made in CPs and their hybrid materials can result in tremendous advances in noninvasive diagnosis, synergistic therapy, and theranostics. There is still plenty of space for further improvement, and further development in this field can promise fascinating applications that can overthrow the currently used operation modes in the field of disease imaging and therapeutics.

### 6.3 Stimulus-responsive polymers for anticancer drug and gene delivery

It is difficult for cancer nanomedicine to deliver therapeutic agents from the injection site to the action site inside tumor cells due to the presence of multiple physiological barriers in the delivery steps. Earlier studies have shown that the optimal nanoproperties to achieve high delivery efficacy vary for the different delivery steps. Therefore, stimulus-responsive polymers have been developed to endow cancer nanomedicines with self-adaptive nanoproperties to maximize their efficiency at each step. In this subsection, the current designs of stimulus-responsive polymers for cancer nanomedicine to realize 3S transitions have been summarized.

**Cancer drug delivery and nanomedicine design.** Nanomedicines have shown promise in cancer therapy, some of which have already been in clinical use.<sup>293</sup> Current FDA-approved nanomedicines, such as Doxil, have succeeded in alleviating drug side-effects, but they have failed to significantly

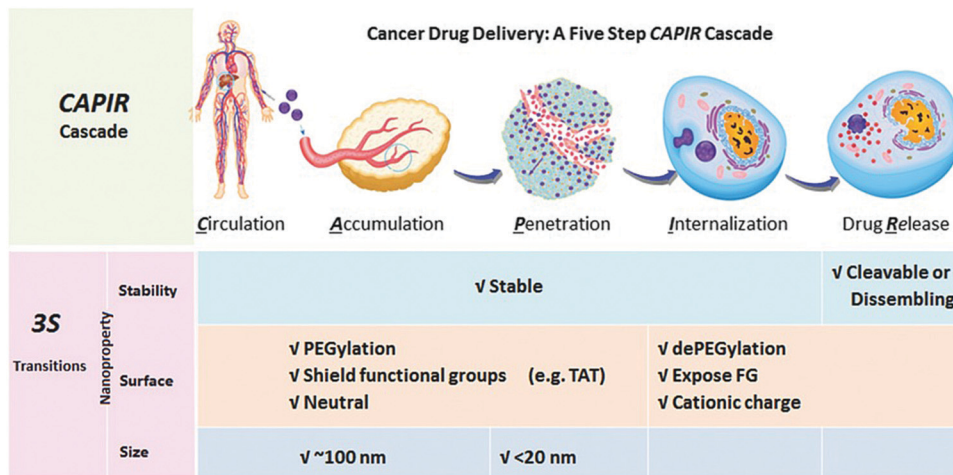


Fig. 50 Cancer drug delivery: the *CAPIR* cascade and 3S nanoproperty transitions for nanomedicines to obtain high overall efficiency. Reprinted from ref. 295 with permission. Copyright (2017) from Wiley-VCH.

improve the therapeutic efficacy.<sup>294</sup> A large body of evidence has indicated that failure in achieving the expected efficiency is associated with the incomplete delivery of nanomedicines. The journey of an intravenously administered nanomedicine to the cytosol of cells in a solid tumor includes circulation in the vasculature, accumulation in tumors by the enhanced permeability and retention (EPR) effect, penetration away from the perivascular to avascular regions to reach the cancer cells for cellular internalization, and intracellular release of the cargos. This arduous five-step journey was summarized as the *CAPIR* cascade (Fig. 50), where any inefficient step may affect the overall payload delivery efficiency and consequently the therapeutic efficacy.<sup>295</sup>

### 3S nanoproperty transitions for effective cancer nanomedicine.

The *CAPIR* cascade consists of complicated physiological barriers at each step for nanomedicines.<sup>296,297</sup> The nanoproperties needed to overcome these barriers at each step are distinct and even dilemmatic with the other options. For example, in the *CAPI* steps, nanomedicines are required to tightly hold the drug molecules to prevent premature leakage into blood; once accumulated in the cancer cells, the payload must be immediately unloaded (*R* step). In the *CA* steps, nanomedicines should be stealthy to immune systems (particularly the liver and spleen) to have long blood circulation for increased accumulation in the tumor by the EPR effect; once close to the tumor cells, they should be able to interact with the cell membrane (*i.e.*, sticky) for internalization. Furthermore, a nanomedicine with particle sizes of around 100 nm is advantageous to achieve longer circulation times and higher tumor accumulation in the *CA* steps, but a nanomedicine with a sufficiently small size can facilitate the tumor penetration ability. That is, the nanoproperties of a nanomedicine should be self-adaptive to each *CAPIR* step to overcome its barriers. Shen *et al.* proposed that the stability, surface, and size nanoproperty transitions (3S transitions) are necessary for the nanomedicine to be self-adaptive to overcome the barriers in the different steps and efficiently accomplish the *CAPIR* cascade delivery steps.<sup>295,298</sup> In brief, the 3S transitions involve stability transition from being

stable in the *CAPI* steps to being unstable in the *R* step, surface transition from being stealthy in the *CA* steps to being sticky in the *I* step, and size transition from large sizes (~100 nm) in the *CA* steps to small sizes in the *P* step. Therefore, various stimuli-responsive polymers have been designed and used to realize 3S transitions.<sup>293,299,300</sup>

**Stimulus-responsive polymers for surface nanoproperty transition.** To address the contrary demand of polymeric nanomedicines in the *CA* steps and *I* step and to realize the stealthy-to-sticky transition, responsive polymers have been designed to realize three strategies, namely, PEGylation/dePEGylation, hiding/exposing functional ligands, and surface charge reversal to generate cationic charges.

**PEGylation/dePEGylation.** Coating nanomedicine with PEG or “PEGylation” is a generally used approach to obtain a stealthy surface for long circulation, but it also decreases the cancer cell uptake. Therefore, PEGylation/dePEGylation transition is needed to resolve this PEG dilemma.<sup>301</sup> This can be realized by the use of labile linkages to connect the PEG chains to a nanomedicine, which are cleavable in response to different stimuli in the tumor microenvironment (such as acidic pH, altered enzymes, and ROS concentrations). Various acidic pH-labile bonds such as  $\beta$ -carboxylic acid amides, benzoic-imine-based materials, and borate-based coordination bonds have been used to develop PEG-detachable nanomedicines.<sup>302–307</sup> For instance, a layer of catechol-containing PEG was coated on a phenylboronic acid (PBA)-functionalized polydopamine (PDA) nanoparticle *via* a reversible covalent interaction. Due to the acid-labile PBA/catechol complex, the PEG corona of the NPs can be removed in response to the acidic milieu of the tumor (Fig. 51a).<sup>307</sup> Likewise, enzyme-triggered dePEGylation can be easily obtained by conjugating PEG *via* short peptides that are cleavable by enzymes such as matrix metalloproteinases (MMPs) overexpressed in many tumors.<sup>308</sup> ROS-triggered dePEGylation can be realized by conjugating PEG with ROS-sensitive thioketal bond. Normally, intercellular ROS concentration is not sufficiently high to detach the PEG. Wang *et al.* successfully

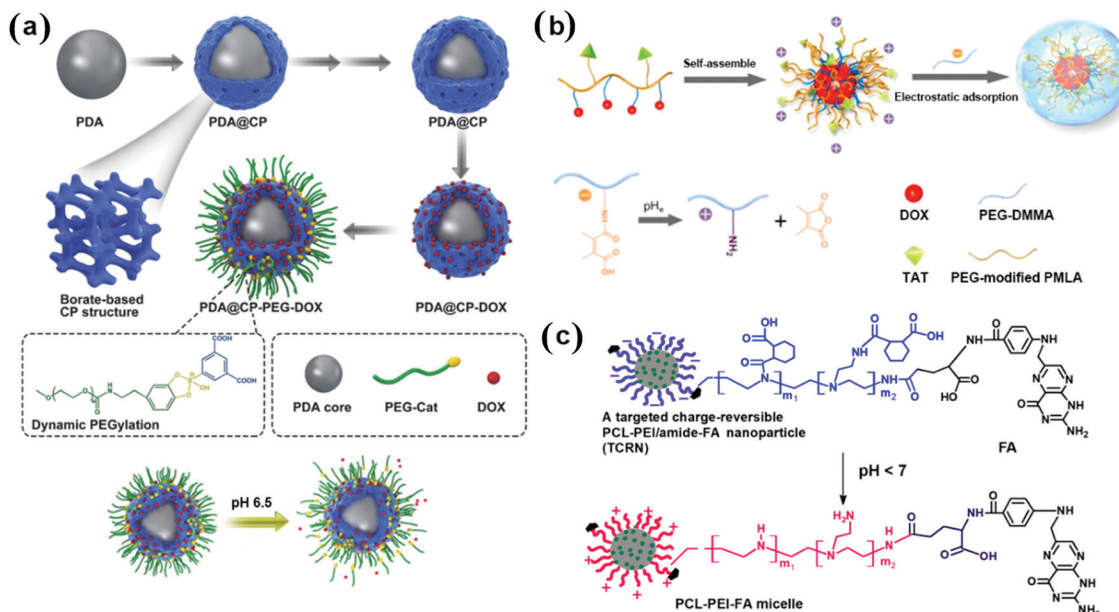


Fig. 51 Representative stimulus-responsive polymers for surface nanoproperty transition. (a) Fabrication of dynamically PEGylated, borate-coordination-polymer-coated PDA nanoparticle. (b) Preparation of PEG-DMMA-coated nanocomplex. (c) Structure of the targeted charge-reversal nanoparticle. Reprinted from ref. 307, 310, and 322 with permission. Copyright (2018, 2017, and 2007) from Wiley-VCH and Ivyspring International.

cleaved the thioketal-bond-linked PEG by encapsulating a ROS generator (chlorin e6, a photosensitizer) in a micelle with the help of NIR radiation.<sup>309</sup>

**Shielding/exposing targeting ligands.** It is an effective strategy to enhance tumor accumulation and cellular uptake of nanomedicines by tethering targets and other functional ligands on the surfaces. However, the hydrophobic and/or cationic ligands attached on the surfaces can deteriorate the nanomedicine's blood circulation time due to the increased nonspecific interaction with the cells and proteins in the *C* step. For instance, cell-membrane transduction peptides (CPPs), a class of small peptides capable of translocating across the cell membrane, are often rich in arginine and lysine residues and therefore carry positive charges. To resolve this problem, such ligands need to be hidden in the *C* step and exposed in the *I* step.

Two general approaches have been used to realize this transition. The first approach is to mask the cationic CPPs *via* electrostatic interactions by polyanions or polyanions-conjugated PEG. The surface polyanions were designed to shed off in response to pH or enzymes. For instance, 2,3-dimethyl-maleic anhydride (DMMA)-amide functionalized PEG and poly(methacryloyl sulfadimethoxine) (PSD)-linked PEG were used to mask the CPPs on NPs.<sup>310,311</sup> The DMMA amides underwent a negative-to-positive charge reversal in the acidic tumor microenvironment and therefore abolished their electrostatic interactions with CPPs (Fig. 51b). Hyaluronic acid (HA) was also used to complex with the CPP-functionalized NPs. In tumors, the surface HA degraded by the overexpressed hyaluronidases (HAase) and the exposed CPP for fast tumor cell internalization.<sup>312</sup> Another approach is to directly connect the cationic CPPs with short zwitterions through cleavable linkers

so that the polyanions can stick on the CPPs and mask the cationic charges. The linkers are chosen to be cleavable by pH, enzymes, or light.<sup>313–318</sup> The breakage of linkers weakens the interaction and makes the polyanions fall off. The other strategy is dePEGylation. Apparently, removing the surface PEG chains would expose the target motifs such as folic acid and Arg-Gly-Asp (RGD) anchored with stable linkers.<sup>305,319,320</sup>

**Surface charge transition.** Surface charge of a nanomedicine has a paramount influence on its *in vivo* fate. Slightly negative or neutral surface of a nanomedicine favors the *CA* steps, while positive charges facilitate the endocytosis by cancer cells in the *I* step. Therefore, a surface charge transition from negative/neutral to positive is needed to resolve this dilemma.

Acidity-triggered charge reversal is commonly used to realize this transition. Amine-containing polymers, such as polyethyl-imine (PEI), poly(L-lysine) (PLL), poly(2-aminoethyl methacrylate hydrochloride) (PAMA), amino-functionalized polyphosphoester, and chitosan, were reacted with anhydrides to produce acid-labile  $\beta$ -carboxylic amides, which decomposed in weak acidity to regenerate the amines carrying cationic charges.<sup>321–330</sup> As an example, Shen *et al.* were the first to demonstrate the charge-reversal concept using acid-labile  $\beta$ -carboxylic-amides-functionalized poly( $\epsilon$ -caprolactone) (PCL)-PEI nanoparticle (Fig. 51c). The resulting PCL-PEI/amide micelles underwent a surface charge transition as the pH decreased from 7.4 to 6.0. Subsequently, it was demonstrated that the pH sensitivity of the amides could be tuned by introducing substituents or unsaturated double bonds.<sup>322,331</sup> Subsequently, Wang *et al.* designed a series of charge-reversal systems using more labile amides.<sup>303,328,329,332</sup>

As alternatives, the protonation/deprotonation of carboxyl or amino groups has been applied to charge-reversal nanomedicine design.<sup>333</sup> Tuning the ratios of carboxyl to amino

groups or using zwitterionic groups afforded materials that were capable of charge transition.<sup>334–340</sup> The advantage of such charge transition is its instant response to pH variations. In a recent study,<sup>340</sup> a gene delivery system was fabricated *via* complexing PEI, PEG–histidine (His)–glutamic acid (Glu) copolymer, and p53 plasmid. By adjusting the ratio of PEI to copolymer, the resulting DNA complex could undergo a negative-to-positive charge reversal as the pH decreased from 7.4 to 6.8. In addition, charge-reversal nanomedicines triggered by other stimuli, such as enzymes or gas molecules, were also reported.<sup>308,341</sup>

**Responsive polymers for size transition.** Particle size is another key parameter affecting the *in vivo* fate of a nanomedicine.<sup>296,342,343</sup> It affects the nanomedicine's circulation, tumor accumulation, and penetration. Generally, nanomedicines with the size of ~100 nm are favorable for longer circulation and tumor accumulation, while the small-sized ones (<30 nm) are more efficient with respect to tumor penetration.<sup>344,345</sup> Therefore, it is desirable to develop nanomedicines with the size transition property from large to small to resolve this dilemma. Multistage delivery systems with the size transition ability, which maintain the large particle sizes for circulation and change to small particles in response to a variety of stimuli, have therefore been developed.

Dendrimers have mostly been used to develop size-transitional multistage delivery systems.<sup>346–351</sup> As an example, Shen *et al.* engineered a lipid-coated dendrimer nanoassembly system to address the size dilemma.<sup>346</sup> The nanoassembly was

a 45 nm-diameter PEGylated liposome encapsulated with almost 27 dendrimers (5 nm in diameter). Once accumulated at the tumor site, the lipid membrane fused with the cancer cell membrane to induce the rupture of the nanoassembly. The released small dendrimers then penetrated into deeper areas of the tumor. Meanwhile, the 2-(*N,N*-diethylamino)ethyl termini on the dendrimers protonated in the acidic tumor microenvironment, triggering cellular uptake and release of the hydrophobic payloads (Fig. 52). Likewise, the dendrimer complex was also designed to disassemble in response to a photoacoustic shockwave.<sup>352</sup> Recently, a micelle-in-liposome hybrid nanomedicine was designed to release micelles (~42 nm) from the liposomes (~94 nm) at the tumor microenvironment by responding to both MMP-9 and low pH.<sup>353</sup> In addition, size-shrinkable nanomedicines were designed by the use of enzyme-degradable substrates such as MMP-2-cleavable peptides and gelatin.<sup>354–356</sup> The degradation of polymer particles is another approach for size transition. For instance, a nanomedicine composed of ROS-responsive poly(thioacetal phosphoester), chlorin e6, and doxorubicin (DOX) could shrink from 154 to 72 nm after the degradation of thioacetal backbones by ROS produced under light irradiation.<sup>357</sup>

**Responsive polymers for stability transition.** Generally, therapeutic payloads are either covalently conjugated to or physically encapsulated in polymeric nanomedicines. For the efficient delivery of drugs to the target, a nanomedicine must tightly retain the drug molecules during the *CAP1* steps and quickly release them inside the cancer cells (*R* step). Polymer–drug

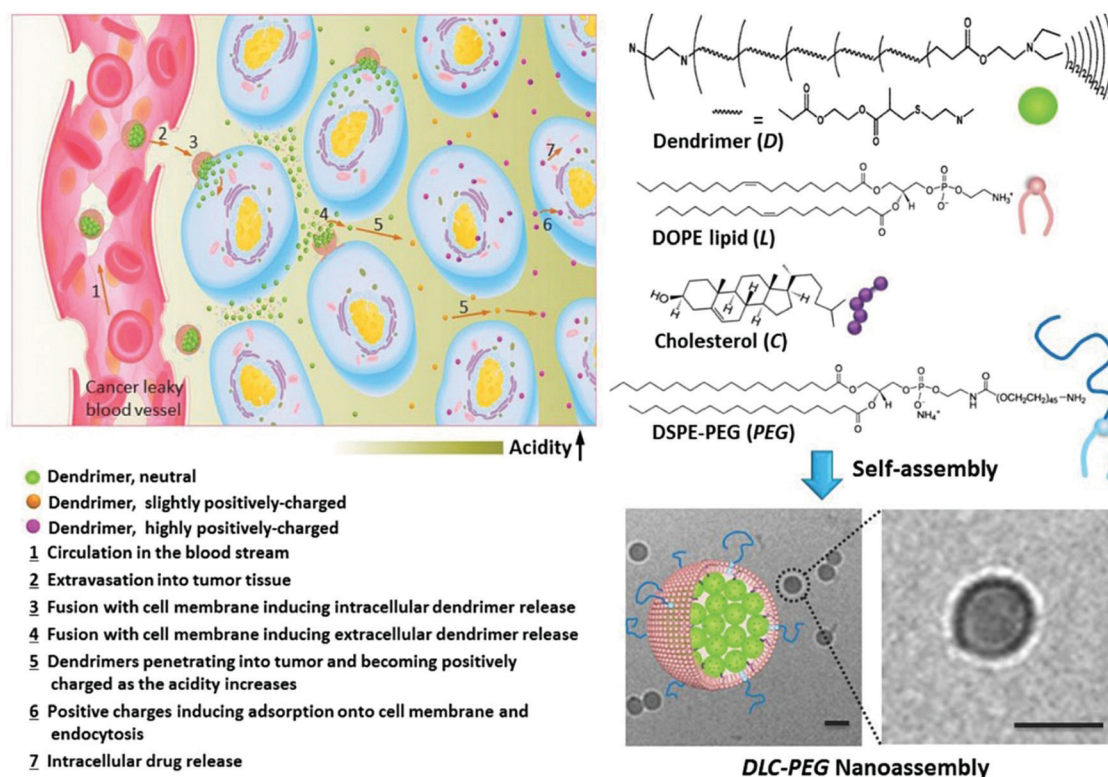


Fig. 52 Schematic of the cluster-bomb-like nanoassembly. Reprinted from ref. 346 with permission from Wiley-VCH.

conjugates can induce this stability transition using intracellularly cleavable linkages, including acid-labile bonds (*e.g.*, hydrazone and  $\beta$ -carboxylic amides), redox-sensitive bonds, and enzyme-sensitive linkers.<sup>358–371</sup>

For noncovalently encapsulated drugs, a transition can be realized by the swelling or dissociation of the hydrophobic cores in response to low pH, ROS, enzymes, or adenosine-5'-triphosphate (ATP).<sup>372–378</sup> For instance, in the ATP-triggered stability transition, ATP-binding moieties (such as aptamers, chaperonin, phenylboronic acid, or metal ions) were integrated into the cores of the nanomedicines.<sup>379–383</sup> After being internalized into the cancer cells, the concentrated ATP molecules were competitively bound to the motifs, causing the disassociation or conformational transition of nanomedicines followed by drug release. Recently, prodrugs, which are converted into active forms by responding to specific signals, loaded into polymeric NPs form a special type of nanomedicine. For instance, Shen *et al.* developed an enzyme and ROS-dependent cascade amplification strategy to specifically release the drug to cancer cells.<sup>384</sup>  $\beta$ -Lapachone, a ROS generator, was coencapsulated with a ROS-sensitive DOX prodrug into the micelles. In cancer cells, the overexpressed NAD(P)H:quinone oxidoreductase-1 (NQO1) enzyme enabled  $\beta$ -lapachone to produce an elevated level of ROS, which decomposed the prodrug and released free DOX in the cascade amplification strategy. This was also demonstrated in other systems.<sup>385–388</sup>

Polymer gene delivery systems generally have low gene expression efficiency mainly because the tight packing *via* strong interactions between the cationic polymers and genes makes it difficult to release DNA in the cells for expression.<sup>293,298,389</sup> Degradable cationic polymers, which can degrade into low-molecular-weight chains and therefore loose binding to the DNA, have been widely explored to enhance gene release.<sup>340,390–394</sup> However, the resulting positively charged short chains may still interfere with gene transfection. To overcome this limitation, Shen *et al.* developed intracellular positive-to-negative charge-reversal polymers to abolish the binding to complete DNA release, thereby yielding enhanced gene expression.<sup>395–398</sup> In one design, the cationic polymers were made to respond to intracellular ROS, which converted them into negatively or neutrally charged (Fig. 53).<sup>395,396,399</sup> The polymers degraded quickly in half an hour in the presence of  $H_2O_2$ . In another design, the cationic polymers were made susceptible to esterase-catalyzed hydrolysis (Fig. 54). The hydrolysis of the phenolic acetate triggered charge reversal.<sup>397</sup>

Complicated physiological environments in the human body yield a diverse array of barriers that need to be efficiently overcome by cancer nanomedicines with multiple functions. The 3S nanoproperty transition concept provides a design principle for the integration of necessary functions into a single nanomedicine. Stimuli-responsive polymers are excellent materials to realize such 3S nanoproperty transitions. Further design of biocompatible, highly selective and sensitive, and simply structured polymers is needed for developing highly potent and translational nanomedicines.

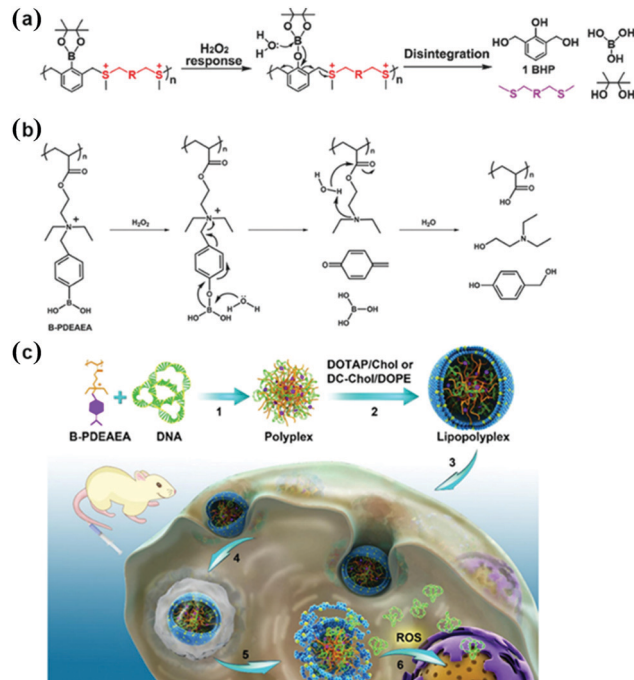


Fig. 53 (a and b) A positive-to-negative convention mechanism of surface charge for gene carriers. (c) Schematic representation of cationic-lipid-coated ROS-responsive polyplexes for gene delivery. Reprinted from ref. 395, 396 and 399 with permission. Copyright (2017, 2016, and 2018) from Wiley-VCH and American Chemical Society.

#### 6.4 Polymeric vectors for drug and gene delivery

Therapeutic agents, such as chemical compounds, genes, photosensitizers, and proteins, are fragile. They can undergo biodegradation in the organism's physiological environment before they can reach their targets. Therefore, powerful "body-guard" vectors have seen a rising demand, which can protect their clients from both *in vitro* and *in vivo* dangers and facilitate their delivery to the target sites. In the past few decades, the field of drug and gene delivery has grown tremendously by the development of a wide range of advanced delivery systems. Such delivery systems offer novel technologies and promising strategies for precise diagnosis and therapy, including chemotherapy, radiotherapy, immunotherapy, PDT, photothermal therapy (PTT), and ultrasound therapy. Polymer-based delivery systems have been continuously receiving significant attention due to their biocompatibility, higher carrying capacity, structure controllability, low cost, and production ability in large amounts. A number of polymeric delivery systems have been designed for controlled drug and gene release applications, which are capable of responding to a range of external stimuli (*e.g.*, light, heat, magnetic fields, and ultrasound) as well as internal stimuli (*e.g.*, pH, ionic strength, metabolic enzymes, glutathione (GSH), ROS, and ATP).

Each disease has its own characteristics as well as micro-environments, which play key roles in disease development and progression. For instance, the tumor microenvironment is a unique cellular environment in which the tumor exists, and it comprises the surrounding blood vessels, immune cells,

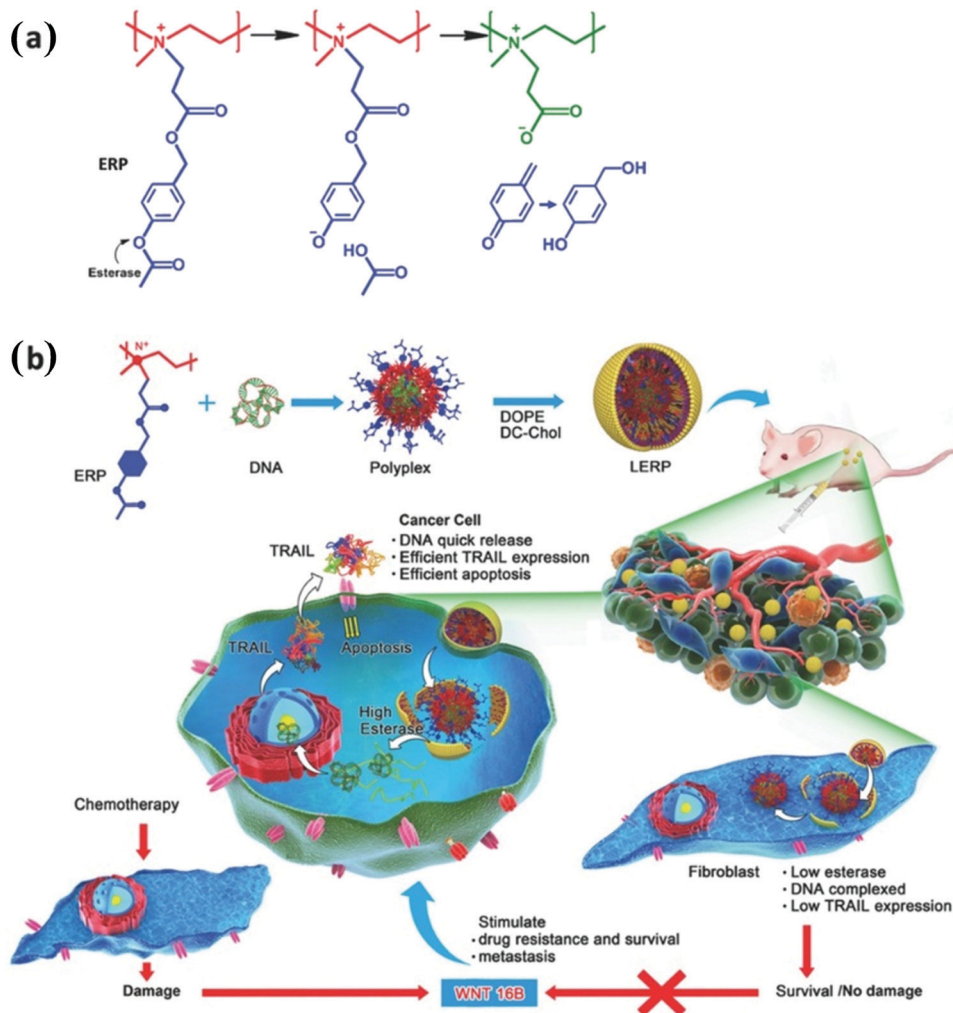


Fig. 54 Esterase-responsive charge-reversal polymer. (a) Mechanism of esterase-induced charge reversal of the gene carrier. (b) Illustration of selective gene delivery to cancer cells. Reprinted from ref. 397 with permission. Copyright (2016) Wiley-VCH.

fibroblasts, bone-marrow-derived inflammatory cells, lymphocytes, signaling molecules, and extracellular matrix (ECM). A tumor can change its microenvironment, which, in turn, can affect the tumor progression and metastasis as a result of its interactions with the host. Accordingly, the microenvironment of the disease provides various internal stimuli, which can be considered as appropriate triggers for smart delivery systems that may cause changes in the phase, size/shape, reactivity, permeability, and surface wettability. Moreover, after conjugating with the targeting ligands (*e.g.*, monoclonal antibodies, peptides, or small molecules), such vectors can be used to target the microenvironment with high specificity and greater affinity. This subsection discusses the use of polymeric vectors to deliver therapeutic drug molecules and bioactive genes inside the organism. The recent contribution by Zhang *et al.* to this field is summarized, and promising applications in cancer immunotherapy, targeted drug delivery, and highly efficient gene transfection are also demonstrated.

Light-sensitive systems have been used in diverse applications for the development of novel intelligent materials and systems.

Light or thermal stimuli can be used to tailor the physical and chemical properties of vectors, such as viscosity, refractive index, conductivity, pH, solubility, wettability, mechanical properties, and polymer morphology. Following this idea, Zhang *et al.* developed photoresponsive polymers possessing very attractive characteristics. Taking inspiration from nature, the biomimetic concept has been integrated into drug delivery systems. In a recent study by Zhang *et al.*, spinach thylakoid fragments were recombined with lipid molecules to synthesize a hybrid proteoliposome, also called a highly efficient life-support intracellular opto-driven system (HELIOS), for the generation of ATP.<sup>400</sup> With red-light irradiation, HELIOS could improve the intracellular ATP concentration to 1.38–2.45 times in various cell lines. Moreover, it was noticed that HELIOS-mediated ATP generation could comprehensively promote the cell functions such as protein synthesis and insulin secretion. At the organ and individual levels, it was proven that HELIOS rescued a mouse heart from myocardial infarction and sustained the life of fasting zebrafish models.

In addition, tumor hypoxia has attained the status of a core hallmark of cancer that globally affects the entire tumor phenotype.

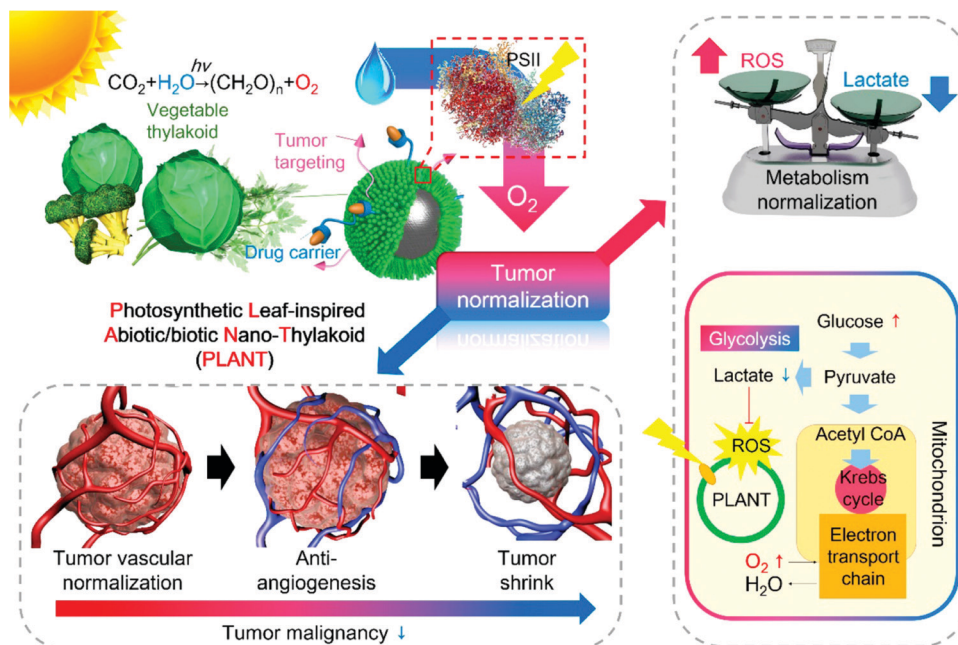


Fig. 55 Schematic diagram of tumor normalization induced by the PLANT system. Reprinted from ref. 401 with permission. Copyright (2018) American Chemical Society.

Reversing tumor hypoxia might offer alternative therapeutic opportunities for current anticancer therapies. To address this challenge, Zhang *et al.* further designed a photosynthetic leaf-inspired abiotic/biotic nanothylakoid (PLANT) system by fusing the thylakoid membrane with synthetic NPs for efficient  $O_2$  generation *in vivo* (Fig. 55).<sup>401</sup> Under laser irradiation at 660 nm, the PLANT system exhibited intracellular  $O_2$  generation. *In vivo*, it was found that this PLANT system could not only normalize the entire metabolic network but also adjust the abnormal structure and function of the tumor vasculature. This PLANT system could significantly enhance the efficacy of phototherapy or antiangiogenesis therapy. This facile approach for normalizing the tumor microenvironment can have promising potential in tumor therapies.

Tumor hypoxia can also considerably suppress the therapeutic efficacy of PDT, mainly because the generation of toxic ROS in PDT is highly oxygen-dependent. Different from ROS, the generation of oxygen-irrelevant free radicals is oxygen-independent. Zhang *et al.* synthesized a therapeutic strategy based on the light-induced generation of free radicals for cancer therapy.<sup>402</sup> An initiator of 2,2-azobis[2-(2-imidazolin-2-yl)propane]dihydrochloride (AIBI) was chosen as the radical source, which was loaded in the hollow interiors of gold nanoclusters

(AuNCs), followed by the coating of thermal-responsive poly(*N*-isopropylacrylamide-*co*-acrylamide) as the gatekeeper to form the AIBI@AuNC copolymer. Under NIR irradiation, the plasmonic heating effect of AuNCs could induce the decomposition of the initiator to generate alkyl radicals ( $R^\bullet$ ), which could increase the oxidative stress (OS) and cause DNA damage in the cancer cells, finally leading to apoptotic cell death under different oxygen tensions. As a proof of concept, this research opens up a new field to use various free radicals for cancer therapy.

Recently, bacteria-mediated drug delivery has attracted considerable attention, particularly for tumor therapy. Beyond utilizing their tumor-targeting ability, chemical explorations have been made to directly conjugate drug-loaded nanomaterials and tumor-targeting bacteria. As a biological strategy, genetic engineering has also been used to program bacteria to become *in situ* anticancer agents. As a further extension, the same group designed a hybrid system of CCN@*E. coli* to achieve photocontrolled bacterial metabolite therapy (Fig. 56).<sup>403</sup> By combining both selectivity of biosynthetic enzymes and robustness of photocatalytic nanomaterials, the bacterial NO generation could be controlled with light irradiation. In addition, self-powered nonpathogenic microorganisms were used to

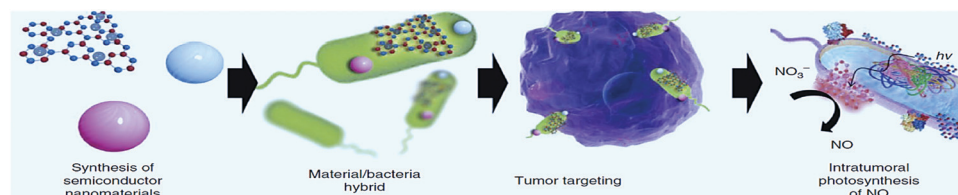


Fig. 56 Schematic diagram of the preparation of a PMT system. Reprinted from ref. 403 with permission. Copyright (2018) Nature Publishing Group.

overcome the shortcomings of classical nanomaterials in tumor targeting and organ penetration. The results obtained from proteomics also suggested that the immune response may get involved in this therapy, and the PMT strategy may lead to useful actuation in cancer immunotherapy.

Oral drug administration is widely adopted for diverse drugs and is convenient to use due to its capability to reach different parts of the body *via* the bloodstream. However, it is generally not feasible for biomacromolecular antitumor drugs, such as proteins and nucleic acids, due to the limited absorption through the gastrointestinal tract and poor tumor targeting. Zhang *et al.* recently engineered a noninvasive thermally sensitive programmable therapeutic system using *E. coli* MG1655 as a vehicle for tumor treatments *via* oral administration.<sup>404</sup> Thermally sensitive programmable bacteria (TPB) are transformed with plasmids expressing the therapeutic protein TNF- $\alpha$  and then decorated with biomineralized gold nanoparticles (AuNPs) to afford TPB@Au. AuNPs and TNF- $\alpha$  plasmids efficaciously protected by TPB in the gut can be transported into internal microcirculation *via* the transcytosis of microfold cells (M cells). The antitumor vehicles accumulate at the tumor sites due to the anaerobic bacterial feature of homing onto the tumor microenvironments. *In vitro* and *in vivo* experiments verify the successful delivery of AuNPs and TNF- $\alpha$  plasmids by TPB. Importantly, under remote activation, the expression of TNF- $\alpha$  at the tumor sites can be precisely controlled by the heat generated from the photothermal AuNPs to induce therapeutic actions.

The ability of the engineered cells to communicate with other cells *in vivo* and interact with the microenvironment of the disease allows them to perform many complex tasks. Inspired by the biological nature of red blood cells (RBCs) as the primary oxygen supplier in mammals, they engineered an aggressive manmade RBC (AmmRBC) to combat the hypoxia-mediated resistance of the tumors toward PDT.<sup>405</sup> In particular, the complex formed between hemoglobin and enzyme-mimicking PDA and the PDA-carrying photosensitizer were encapsulated inside the biovesicle that was engineered from recombined RBC membranes. Owing to the same origin of the outer membranes, AmmRBCs shared excellent biocompatibility with the parent RBCs. The introduced PDA played the role of antioxidative enzymes existing inside the RBCs to effectively prevent the oxygen-carrying hemoglobin from oxidation damage during circulation. This biomimetic engineering could facilitate accumulation in tumors, permit efficient *in situ* oxygen supply, and impose strong PDT efficacy toward the extremely hypoxic tumor with complete tumor elimination.

The development of multifunctional delivery systems that combine targeting, diagnostic, and therapeutic functions within single nanoscale agents is another design trend. Zhang *et al.* synthesized telluride molybdenum (MoTe<sub>2</sub>) nanosheets with wide NIR absorbance functionalized with PEG-cyclic arginine-glycineaspartic acid tripeptide (PEG-cRGD).<sup>406</sup> After loading a chemotherapeutic drug (DOX), MoTe<sub>2</sub>-PEG-cRGD/DOX was used for combined PTT and chemotherapy. With high photothermal conversion efficiency, this system exhibited favorable cell-killing ability under NIR irradiation. Owing to

the cRGD-mediated specific tumor targeting, it showed efficient accumulation in tumors to induce a strong tumor ablation effect. The *in vitro* and *in vivo* experimental results demonstrated that this theranostic nanoagent could be readily degradable in normal organs to enable rapid excretion and avoid long-term retention/toxicity, holding promising potential to effectively treat tumors.

Metastasis and recurrence are two unavoidable and intractable problems in cancer therapy. Immunotherapy might be an effective approach to resolve these problems, but the high heterogeneity of the tumor tissue, inefficient presentation of tumor antigen, and deficient targeting ability of therapy can usually blunt the efficacy of immunotherapy and hinder its clinical applications. In such a circumstance, Zhang *et al.* developed an approach based on combining photodynamic and immunological therapy.<sup>407</sup> The chimeric peptide PpIX-1MT was synthesized, which integrates the photosensitizer PpIX with the immune checkpoint inhibitor 1MT *via* a caspase-responsive peptide sequence, Asp-Glu-Val-Asp (DEVD), to realize a cascaded synergistic effect. The functional peptide could form NPs in PBS and accumulate in the tumor areas *via* the EPR effect. Upon light irradiation at 630 nm, the PpIX-1MT NPs produced ROS, induced the apoptosis of cancer cells, and facilitated the expression of caspase-3 and production of tumor antigens; these triggered an intense immune response. The subsequently released 1MT upon caspase-3 cleavage could further strengthen the immune system and effectively facilitate the activation of CD8<sup>+</sup> T cells. This cascaded synergistic effect could effectively inhibit both primary and lung metastasis tumors, which may provide a solution for clinically resolving tumor recurrence and metastasis.

Gene therapy has been receiving considerable attention as an efficient treatment of inherited genetic disorders and a wide range of severe diseases. Not only efficient but also rapid transfection is required since it can maximize the bioavailability of vector-carried genes prior to cellular excretion. However, the “rapid” goal has been less investigated so far in the field of vector-aided transfection research. To address this challenge, Zhang *et al.* synthesized the lysosome-targeting acidity-responsive nanoassembly as gene vectors, which proved the amazing potency to mediate the “superfast” transnuclear gene transport and gene transfection with high efficiency both *in vitro* and *in vivo* (Fig. 57).<sup>408</sup> The nanoassembly was constructed on the pH-reversible covalent boronic acid-diol coupling between 1,3-diol-rich oligoethylenimine (OEI-EHDO) and phenylboronic-acid-modified cholesterol (Chol-PBA). The nanoassembly-mediated transfection at 8 h could afford an outcome that was comparable to that achieved at 48 h by the golden standard of PEI25k; high transfection efficiency could be retained during 48 h. The *in vitro* and *in vivo* results exhibited the low toxicity of this biodecomposable nanoassembly.

With the prominent progress of polymeric delivery systems, novel biomaterials with high biocompatibility and versatile functions are urgently needed. Zhang *et al.* explored the hierarchical micro-/nanostructures of human hair that can be further broken into hierarchical microparticles (HMPS) and

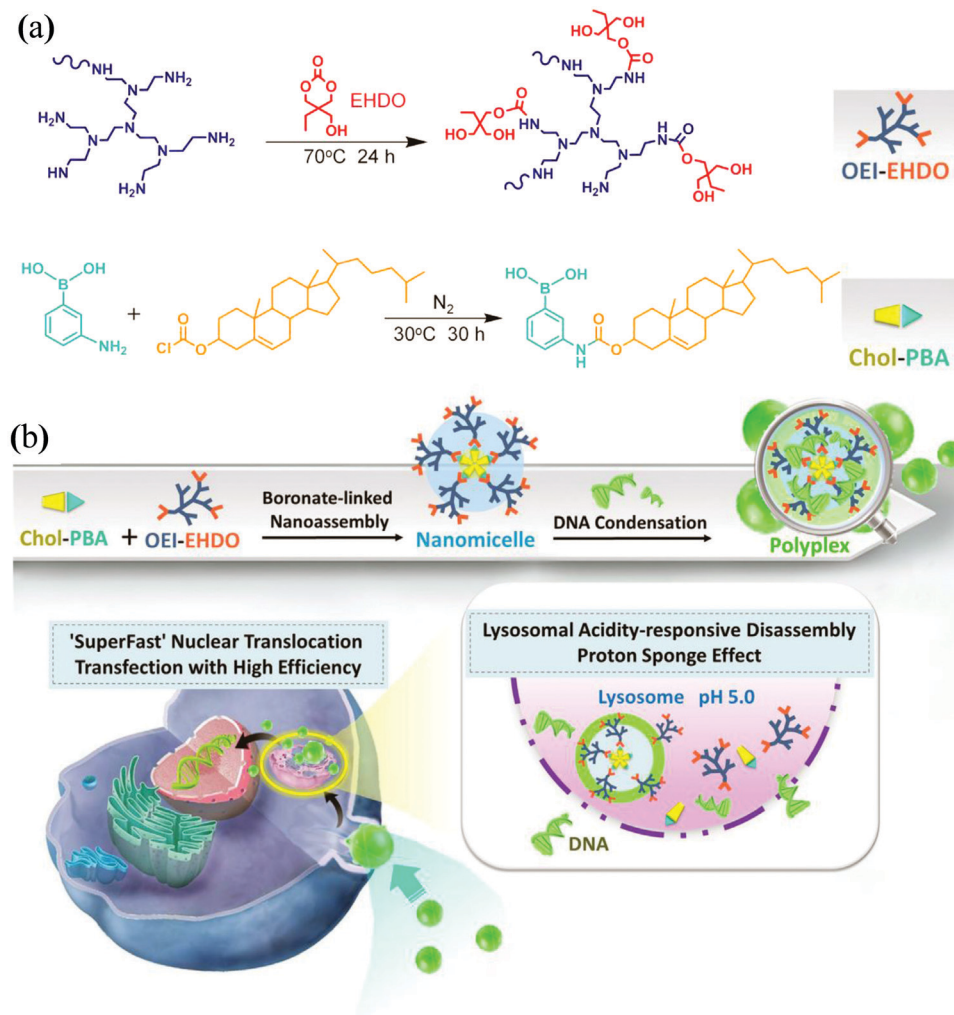


Fig. 57 Acidity-responsive gene delivery for "superfast" nuclear translocation. Reprinted from ref. 408 with permission. Copyright (2016) from Elsevier Ltd.

hierarchical nanoparticles (HNPs) with top-down procedures.<sup>409</sup> When compared with commercialized carriers, such as liposomes or albumin NPs, the obtained particles exhibited high hemocompatibility and negligible immunogenicity. Furthermore, these materials also exhibited peculiar abilities with respect to light absorption and free radical scavenging. Evidently, HMP and HNP could prevent UV-induced damage in skin and relieve symptoms of cataract *in vitro*. Moreover, both HMP and HNP showed satisfactory photothermal conversion ability. By using microcomputed tomography and intravital fluorescence microscopy, it was found that warfarin-loaded HMP could prevent vein thrombosis in mice. In another aspect, HNP modified with tumor-targeted aptamers exhibited dramatic antineoplastic effect and suppressed 96.8% tumor growth *in vivo*. Therefore, the multifaceted materials described here might provide new vectors for addressing biomedical challenges.

Although considerable advances have been achieved in the field of drug and gene delivery, limitations in polymer-based delivery approaches still persist, which have to be overcome before they can be applied as standard practice in clinics.

With regard to drug delivery systems, further developments are certainly needed in this direction, particularly with regard to interactions between the specific microenvironmental conditions of disease and functional design of biomaterials. Determining a clear and stringent understanding of the delivery mechanisms is also necessary to address *in vivo* safety concerns. With regard to gene delivery systems, although polymeric nonviral vectors are generally less efficient in delivering DNA and initiating gene expressions *in vivo* as compared to viral vectors, they have promising potential for producing large quantities under the required GMP conditions. There is certainly higher interest in developing new generations of nonviral vectors, which can not only find the target cells but also put the gene in the correct position on the host genome. Until now, the effective delivery of multiple CRISPR components *in vivo* into host cells still remains a formidable challenge. With the robust development of functional biomaterials, smart polymer-based carriers might become potent vectors for the delivery of CRISPR/Cas9 systems. These developments are very exciting for the drug and gene

delivery fields and could have an enormous impact on pharmaceutical technology and clinical treatments. Numerous challenges in the delivery field need to be addressed, and considerable efforts should be made in this direction before translating them into clinics.

### 6.5 Polyprodrug amphiphiles for drug delivery and theranostics

The use of a polyprodrug strategy for drug delivery provides a plethora of advantages such as customized structural design, ultrahigh drug loading contents, avoidance of premature leakage, and bespoke multifunctional integration.<sup>410</sup> Recently, Liu *et al.* proposed a new concept, namely, polyprodrug amphiphiles, to further enrich the connotation and broaden the vision of polyprodrug strategies. Polyprodrug amphiphiles are a type of specific amphiphilic BCPs, in which one of the blocks is composed of prodrug units. The drug molecules are generally covalently attached to the polymer backbones through stimuli-responsive linkages. The design of stimuli-responsive drug-based monomers may provide the following unique advantages: (1) increasing the water dispersibility and blood circulation stability of water-insoluble and chemically labile drugs; (2) avoiding premature drug release and reducing the systemic cytotoxicity of drugs; (3) optimizing pharmacokinetics and pharmacodynamic performance and elevating the therapeutic efficacy; and (4) integrating diagnostic/imaging functions to develop theranostic nanovectors. In a typical example, an anticancer drug, namely, camptothecin (CPT), was monomerized with reduction-sensitive disulfide linkage, which was subsequently polymerized into amphiphilic PEG-*b*-PCPTM *via* the RAFT polymerization technique by using a PEG-based macroRAFT agent.<sup>411</sup> The CPT loading contents could be delicately tuned and over 50 wt% CPT contents could be facilely loaded. Remarkably, PEG-*b*-PCPTM could be self-assembled into four types of distinct morphologies, namely, spherical micelles, large compound vesicles (LCVs), smooth disks, and unprecedented staggered lamellae with spiked peripheries (Fig. 58). Further studies revealed that, in sharp contrast to the other

three types of nanoassemblies, staggered lamellae possessed extended blood circulation time, unique endocytic pathways, and the fastest cellular internalization rate. Moreover, the four types of nanostructures exhibited drastic CPT release profiles, disassembly kinetics, and *in vitro* cytotoxicities. These observations highlighted the importance of self-assembled morphologies in drug delivery applications and demonstrated that morphology control should be an important consideration in the design of nanomedicines. In this context, Hubbell *et al.* functionalized ibuprofen (IBU)—a well-known antiinflammatory drug—with 2-hydroxyethyl acrylamide *via* the formation of an ester bond.<sup>412</sup> The resultant monomer could be readily polymerized *via* RAFT polymerization to form amphiphilic PEG-*b*-PIBU copolymers by the use of a PEG-based macroRAFT agent. The self-assembly of PEG-*b*-PIBU copolymers led to the formation of both spherical and worm-like micelles, depending on the block ratios of PEG and PIBU. The self-assembly morphology played a critical role in the IBU release behavior and exerted an antiproliferative effect on human cervical carcinoma (HeLa) and murine melanoma (B16-F10) cells.

Indeed, the polyprodrug approach provides a general and robust procedure to fabricate highly efficient nanocarriers with extremely high drug loading contents.<sup>413–417</sup> Moreover, the design of stimuli-responsive drug-based monomers renders it possible to fabricate polyprodrug amphiphiles with varying chain topologies and to integrate other functions. For example, besides linear polyprodrug amphiphiles, hyperbranched polyprodrug amphiphiles were synthesized by the polymerization of CPTM using a methacrylate-monomer-based RAFT agent. Moreover, a magnetic resonance imaging (MRI) contrast agent was initially embedded within the hydrophobic inner cores with suppressed relaxivity, but the MRI signals could be switched on when being exposed to cytosolic-abundant GSH, which also led to the release of the CPT drug by the cleavage of the disulfide linkage (Fig. 59). As such, both the release process of CPT drug and therapy outcomes could be monitored *in situ* and reported on the basis of enhanced MRI signals, which was fairly beneficial for the development of personalized nanomedicines.

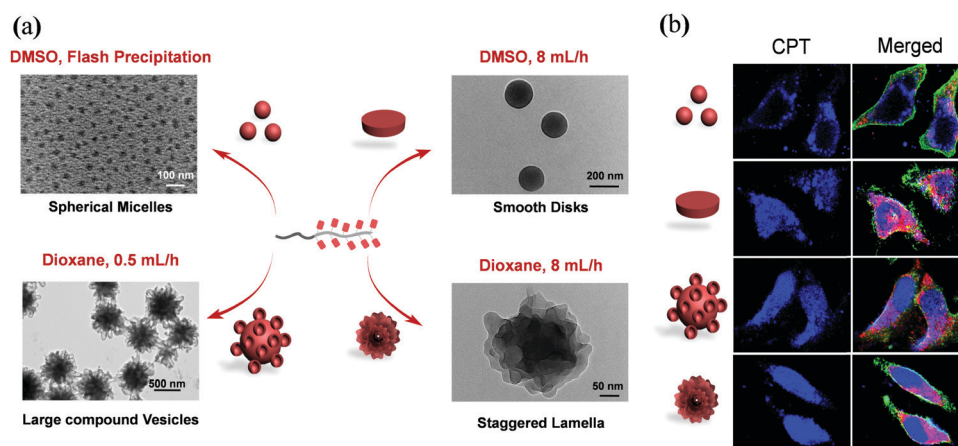


Fig. 58 (a) Self-assembly of PEG-*b*-PCPTM polyprodrug amphiphiles with the formation of four types of nanostructures. (b) Distinct cellular internalization of these four types of nanostructures. Adapted from ref. 411 with permission. Copyright (2013) American Chemical Society.

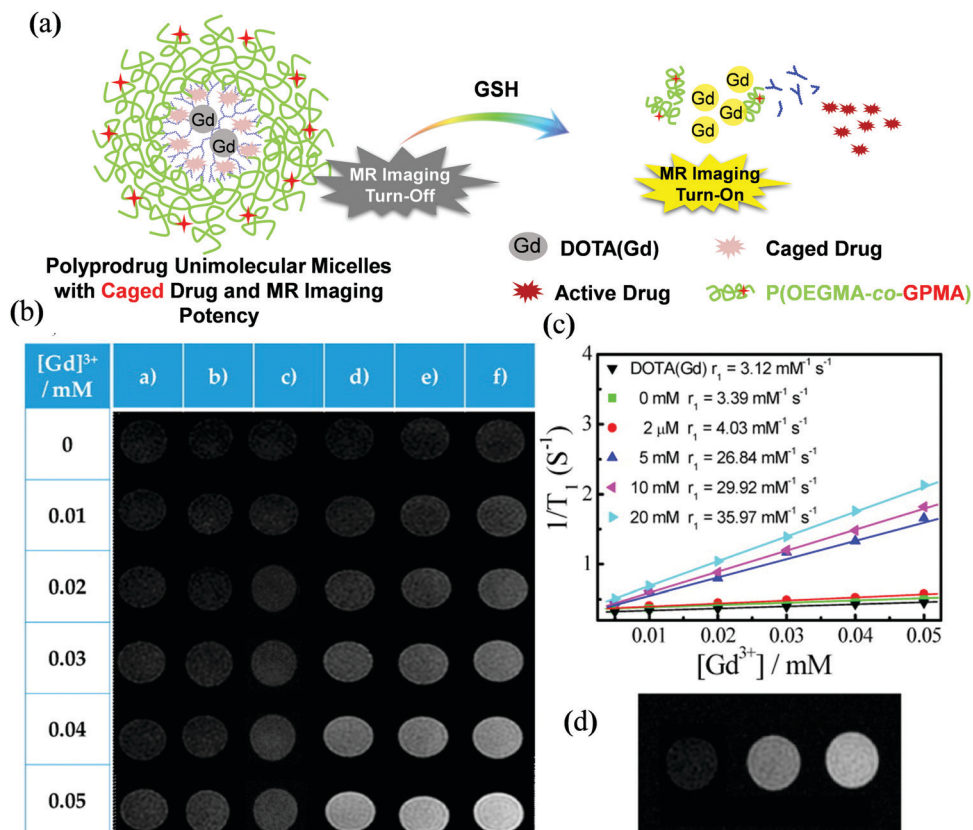


Fig. 59 Hyperbranched polyprodrug amphiphiles with caged CPT prodrug and MRI contrast agents within the inner cores shielded by hydrophilic P(OEGMA-*b*-GPMA) coronas exhibit GSH-triggered intact CPT release and enhanced MRI relaxivity (OEGMA: oligo(ethylene glycol)methyl ether methacrylate; GPMA: *N*-(3-guanidinepropyl) methacrylamide). Adapted from ref. 413 with permission. Copyright (2015) American Chemical Society.

Notably, the self-assembled nanoassemblies of polyprodrug amphiphiles were subjected to extensive dilution and higher shear force in the bloodstream upon administration, disintegrating the assembled nanostructures and compromising the therapeutic efficacy. Hyperbranched polyprodrug amphiphiles can, however, exist in the form of unimolecular micelles even after extensive dilution, providing long-term stability when applied for drug delivery.

The fabrication of polyprodrug amphiphiles represents an emerging but promising approach to design novel therapeutic platforms. The tunable chemical structures, ultrahigh drug loading contents, and versatile self-assembled morphologies can not only provide an opportunity to unravel the interrelation between the assembled nanostructures and therapeutic performance but also shed light on how to overcome the drawbacks encountered in conventional drug carriers. Recently, the direct linkage of hydrophobic and hydrophilic drugs without recourse to other components enabled prodrug amphiphiles to afford drug loading content of 100 wt%.<sup>418</sup> This example suggests that the drug loading contents of polyprodrug amphiphiles could be remarkably elevated by the synergistic introduction of drugs with distinct water solubility. Future direction in this topic can include, but not be limited to, the expansion of drug categories, screening of stimuli-responsive linkages, and introduction of targeting motifs. Meanwhile, the detailed self-assembly

mechanism of polyprodrug amphiphiles and how to precisely control the self-assembled morphologies should be conducted to achieve controlled self-assembly behavior. Moreover, considerable effort should be devoted toward the assessment of the *in vivo* behavior and therapeutic performance of polyprodrug amphiphiles and preclinical evaluations could be carried out at an appropriate time.

## 6.6 Interactions of biomaterial surfaces/interfaces

Biomaterials come into contact with biological environments (*e.g.*, blood, body fluids, tissues) through their surfaces, and the immediately occurring biointerfacial interactions determine the ultimate biocompatibility and the realization of bio-functionality of such biomaterials.<sup>419–421</sup> The communication between biological structures (*e.g.*, cells) and physical objects (*e.g.*, biomaterials) occurs at their interface (called the bio-interface), and the surface of the given biomaterial is one of the most important elements to determine the outcomes of this communication. Biointerface, an interdisciplinary subject among the diverse fields of physics, chemistry, biology, materials science, and biotechnology, is becoming one of the most innovative and expanding areas of science and technology. In the recent decade, considerable efforts have been devoted toward the creation of synthetic biointerfaces either to mimic the biological microenvironment for the fundamental study of cell biology or to

endow the required functions for specific applications.<sup>422,423</sup> For the construction of functional biointerfaces, scientists and engineers have developed a series of surface modification techniques, namely, layer-by-layer (LbL) assembly, surface-initiated polymerization, plasma polymerization, dopamine- or metal-phenolic-networks-assisted deposition, and so on.<sup>424–428</sup> The resultant biointerfaces have been used to investigate cellular behaviors such as adhesion, proliferation, migration, and differentiation in response to surface properties, to improve the blood compatibility and antithrombotic ability for blood-contacting materials and devices, to kill pathogenic bacteria and eliminate biofilm for implants, and to detect analytes and analyze specific biomolecular interactions for biosensors and biochips.<sup>429–435</sup> In addition, the functionalization of the biointerfaces of nanomaterials is also a research hotspot for controlled and targeted drug delivery.<sup>436,437</sup>

Among a series of biointerfacial interactions, protein adsorption has been well recognized as the first significant event, which plays an important role in events such as cell adhesion, platelet activation, and host responses.<sup>438–441</sup> On the positive side, the adsorption of adhesion proteins on cell culture substrates is beneficial for the promotion of cell adhesion and proliferation, which is crucial for regenerative medicine and tissue engineering.<sup>442,443</sup> The selective immobilization of bioactive proteins on the matrix is an important step for the fabrication of biosensors and microarrays.<sup>444,445</sup> On the negative side, nonspecific protein adsorption on synthetic material surfaces can result in several undesired effects such as thrombosis, bacterial infection, and biofouling contamination, leading to the failure of implant materials or the reduction in the sensitivity of diagnostic devices.<sup>446,447</sup> Therefore, understanding and further regulating such interactions between proteins and biomaterial surfaces have become an increasingly important goal for a range of scientific fields and applications.<sup>448–450</sup>

The acquisition of an accurate amount of adsorbed proteins on material surfaces is highly desired for the study of protein–surface interactions and is essential for the evaluation of antifouling properties. However, obtaining protein adsorption in the nanogram range or comparing subtle differences in protein adsorption between various surfaces remains challenging for currently available techniques. Although several methods have been used to determine the quantity of protein adsorbed, the results usually differ. Unfortunately, the reasons leading to such differences are always ignored, which are unfavorable for the rational determination of protein adsorption on a given surface. In this regard, Chen *et al.* compared the three most commonly used quantitative characterization methods, namely, radiolabeling, surface plasma resonance (SPR), and quartz crystal microbalances (QCM), to systematically discuss their applicability for the evaluation of protein adsorption using a series of model gold surfaces with gradient hydrophilicity.<sup>451</sup> They found that when converting SPR signals into protein adsorption, the conversion coefficient was not a fixed value, but it varied with the amount of adsorption. The variation in protein adsorption with wettability measured by QCM was different than that measured by SPR or radiolabeling, possibly because QCM is used to measure

both adsorbed proteins and their associated water molecules. In addition, due to the acoustic principle of QCM, it has performance limitations when working with surfaces modified with thick viscous layers. In order to explore this effect, a series of surfaces modified with self-assembled monolayers or hydrophilic polymer brushes with different thicknesses were prepared, and the protein adsorption on these surfaces measured by QCM and SPR were compared.<sup>452</sup> The experimental results showed that for larger proteins that are expected to adsorb mainly “on top” of the polymer brushes, QCM could not detect a small amount of protein adsorption if the thickness of the polymer brushes exceeded a certain “critical value”; such a critical value increased with polymer viscosity. However, for smaller proteins that are expected to adsorb mainly at the chip substrate and within the polymer brushes, the amount of protein adsorption is detectable by QCM. In conclusion, these findings suggest that QCM data indicating very low protein adsorption on surfaces grafted with polymer brushes should take into account these considerations and should be generally treated with caution.

As mentioned above, the regulation of protein adsorption on biomaterial surfaces (selective binding of favorable proteins and repelling nonspecific proteins) will be beneficial for the promotion of favorable events and the prevention of undesirable events. Taking blood-contacting biomaterials and devices as examples, the initially adsorbed protein layer on the surfaces trigger adverse responses such as plasma coagulation and thrombosis generation. In order to resolve these serious problems, various approaches have been developed to improve the blood compatibility of surfaces by controlling the adsorption of plasma proteins. These modified surfaces can be generally divided into three types: (1) bioinert surfaces that prevent thrombus formation by resisting the adsorption of nonspecific proteins (particularly fibrinogen, which is abundant in plasma and plays a major role in blood coagulation); (2) anticoagulant surfaces that inhibit thrombus formation by suppressing thrombin activity (the key protein in the coagulation process); and (3) fibrinolytic surfaces that destroy the clot (thrombus) once formed by selectively binding essential proteins in the fibrinolytic system. Here, we focus on the recent progresses made in the development of fibrinolytic surfaces by Chen *et al.*; further, the first two strategies are not described in detail due to space limitations; interested readers should refer to other excellent reviews.<sup>431,453,454</sup>

Generally, fibrinolytic surfaces mimic the naturally occurring fibrinolytic processes in the body by the generation of a clot-lysing enzyme called plasmin to break down the nascent clot before it poses any danger.<sup>455</sup> The two main proteins involved in this process were plasminogen (Plg)—a precursor of plasmin—and a plasminogen activator (*e.g.*, tissue plasminogen activator, tPA). Therefore, the main strategy for building fibrinolytic surfaces is to endow the surfaces with the capability to selectively bind Plg and/or tPA in significant quantities when they come into contact with blood. Inspired by natural binding events involved in protein recognition, the bioaffinity ligand for Plg and tPA— $\epsilon$ -lysine with free carboxyl and  $\epsilon$ -amino groups—is screened out and used for the construction of fibrinolytic

surfaces. In the initial works of Chen *et al.*, they attached  $\epsilon$ -lysine to surfaces using hydrophilic polymers (*e.g.*, PEG and poly(2-hydroxy-ethyl methacrylate) (PHEMA)) as spacers; the resultant surfaces could both reduce nonspecific protein adsorption and platelet adhesion as well as capture Plg from plasma with a high degree of selectivity.<sup>456,457</sup> When treated with tPA and incubated in plasma, these surfaces could lyse the clots formed on them or adjacent to them. This clot-lysing ability increased with a decrease in the length of the spacer and increase in the density of  $\epsilon$ -lysine. To simplify the preparation process and avoid multistep surface modification procedures, they designed an  $\epsilon$ -lysine-containing monomer, namely, lysyl methacrylate (LysMA), and used it for either graft copolymerization with HEMA from surfaces containing vinyl groups or copolymerization with HEMA for further blending with polyurethane (PU).<sup>458,459</sup> The Plg-binding capacity of the copolymer films could be simply regulated by varying the compositions of LysMA and HEMA. To further increase the local density of  $\epsilon$ -lysine, Chen *et al.* synthesized a multivalent ligand, namely, a  $\beta$ -cyclodextrin derivative bearing seven  $\epsilon$ -lysine ligands (CD-lysine<sub>7</sub>); they demonstrated that an increase in the local density of  $\epsilon$ -lysine resulted in higher Plg adsorption and higher Plg binding affinity, possibly due to the well-established “multivalent effect.”<sup>460</sup> In particular, using CD-lysine<sub>7</sub> and other bioactive ligands, Chen *et al.* developed a series of dual-functional blood-compatible surfaces *via* a sequential coimmobilization strategy.<sup>461</sup> Surfaces were first modified with a copolymer of poly(2-hydroxy-ethyl methacrylate-*co*-1-adamantan-1-ylmethyl methacrylate) (P(HEMA-*co*-AdaMA)), which served as a linker/spacer for the subsequent sequential attachment of functional peptides (*e.g.*, REDV (Arg-Glu-Asp-Val) peptide with the capability to promote adhesion and proliferation of endothelial cells) *via* covalent bonding and CD-lysine<sub>7</sub> host-guest interactions, respectively.<sup>461</sup> The resultant surfaces exhibited bioactivities toward both the ligands; more importantly, neither of the two functions of peptide and lysine gets compromised by the presence of the other, providing enhanced blood compatibility.

The abovementioned works mainly focused on the selective binding of Plg; evidently, Plg alone is insufficient to generate plasmin as it necessitates the activation of the bound plasminogen. However, the activators of Plg normally exist at extremely low concentrations in bodily fluids. Therefore, surfaces are expected to either release the preloaded tPA or capture tPA from plasma. In this regard, taking advantage of the fact that Plg has higher

affinity than tPA toward surface-bound  $\epsilon$ -lysine, Chen *et al.* developed a new tPA-releasing concept based on protein-protein displacement.<sup>462</sup> This concept was realized by the use of a PU fibrous mat modified with  $\epsilon$ -lysine for the incorporation of tPA *via* specific interaction with lysine residues. As shown in Fig. 60, in contact with the plasma, the preloaded tPA was released from the surface by the displacement of Plg; the amount of released tPA correlated with the amount of adsorbed Plg. This tPA-releasing system provides dual mechanisms for clot lysis, namely, by tPA release from the surface and by plasmin generation on the surface. In addition to using a shared ligand for Plg and tPA, Chen *et al.* also developed a dedicated affinity ligand specific to tPA, *i.e.*, the ARMAPE (Ala-Arg-Met-Ala-Pro-Glu) peptide.<sup>463,464</sup> This peptide is derived from plasminogen activator inhibitor-1 (PAI-1) and is involved in the initial binding of tPA to PAI-1. The binding of tPA through the ARMAPE peptide has a particular advantage in protecting tPA from inactivation by PAI-1 *in vivo*, while the full enzymatic activity of tPA is retained. Taking advantage of  $\epsilon$ -lysine and ARMAPE in exhibiting high affinity separately toward Plg and tPA, a dual-affinity surface formed by immobilizing the two ligands on PU was developed.<sup>465</sup> Differing from the tPA-releasing materials, this dual-affinity surface could continuously enrich Plg from blood and generate plasmin *in situ* to lyse fibrin clots formed on the surface.

Surface modification with bioactive agents capable of combating thrombosis is a widely used strategy for developing antithrombotic biomaterials. However, the exposure of blood to antithrombotic agents on the material surface may cause hemostatic disorders under normal conditions. An ideal blood-contacting material should initiate antithrombotic activity once the thrombus begins to form, while it remains “bioinert” in the normal blood environment. To this end, Chen *et al.* developed a thrombosis-responsive surface coating with the ability to lyse fibrin as it forms.<sup>466,467</sup> Thrombin generation is a key event in blood coagulation/thrombosis and therefore has been used as a physiologic stimulus or trigger in the design of this coating. As shown in Fig. 61, the coating consisted of nanocapsules (NCs) in which the fibrinolysis activator t-PA was encapsulated in a thrombin-degradable hydrogel shell. This t-PA NC surface was stable in normal plasma along with the release of t-PA at a constant rate and efficiently lysing fibrin clots when thrombin was present. In addition, it could release sufficient t-PA for clot lysis in response to endogenous thrombin generated during the clotting process in whole blood.

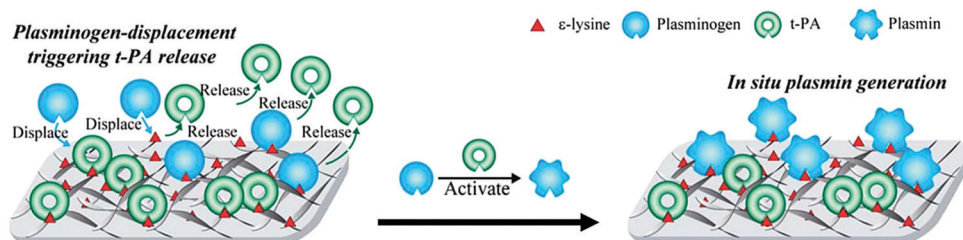


Fig. 60 Schematic illustration of the concept of tPA release based on tPA-Plg displacement. Adapted from ref. 462 with permission. Copyright (2016) Royal Society of Chemistry.

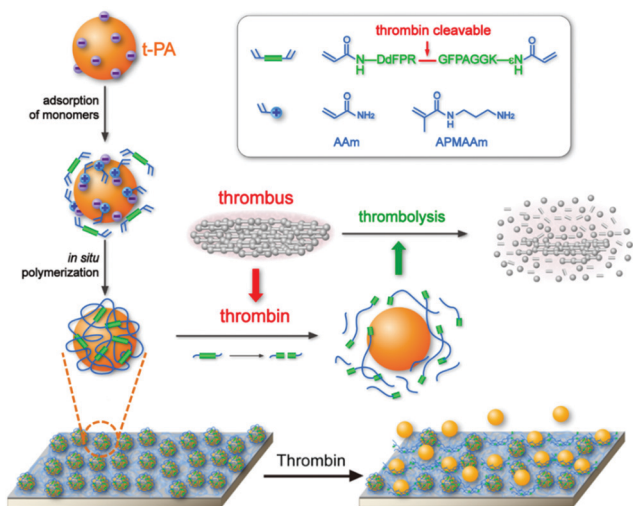


Fig. 61 Schematic illustration of thrombolytic coating based on the thrombin-degradable nanocapsules of t-PA. Adapted from ref. 466 with permission. Copyright (2017) from Wiley-VCH.

### 6.7 Biodegradable polymers for marine antibiofouling

Marine biofouling formed by the settlement and accumulation of microorganisms, plants, and animals on surfaces immersed in seawater has profound effects on the maritime industry. It accelerates biocorrosion, increases hydrodynamic drag (and therefore fuel consumption), and transports invasive species. More than 4000 fouling species live in the sea, and they are fairly different from area to area; they also vary with temperature, salinity, pH, and other conditions. Therefore, marine biofouling is a worldwide problem. Since tributyltin-based coatings have been banned due to their negative effects on the marine environment, it has become imperative to develop eco-friendly antifouling systems.

To combat marine biofouling, Zhang *et al.* proposed the concept of dynamic surface antifouling, where the dynamic surface involves a changeable surface that continuously renews itself in seawater and therefore prevents fouling organisms from landing and adhesion (Fig. 62).<sup>468,469</sup> Dynamic surface antifouling was confirmed by observing the settlement of model microfouling organisms on the surfaces of degradable polymers *via* digital holographic microscopy (DHM), which can reveal the 3D real-time motion of a microorganism. The trajectories of bacteria in the near-surface range with respect to the mean square displacement (MSD) showed that when the polymer degradation rate is sufficiently high, a large portion of bacteria may exhibit superdiffusive motion rather than subdiffusive motion, which represents a transient state before irreversible adhesion. Atomic force microscopy (AFM) measurements demonstrated that adhesion forces between a bacteria-covered colloidal probe and the surface decrease as the degradation rate increases. Accordingly, polymer degradation alters bacterial motion such that they are not able to land on but have to escape from the dynamic surface.

Based on this strategy, Zhang *et al.* developed coatings using PU with a biodegradable backbone: these coatings exhibited

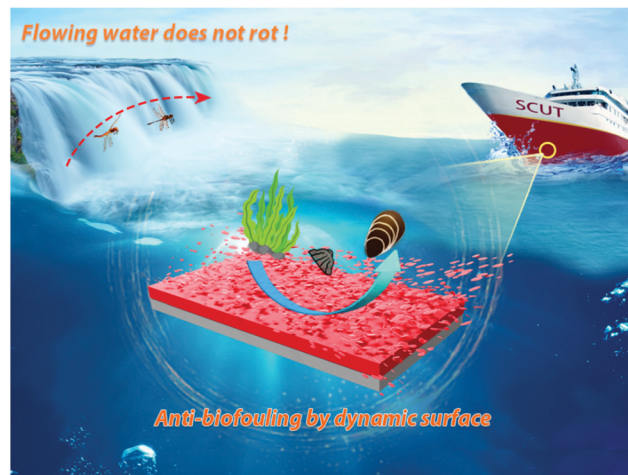


Fig. 62 Schematic of dynamic surface antifouling. Reprinted from ref. 468 with permission. Copyright (2019) Royal Society of Chemistry.

excellent antifouling performance in marine field tests.<sup>470</sup> Furthermore, they synthesized PCL-based PU with triisopropylsilyl acrylate (TIPSA), where the main chains are degradable and the side chains are hydrolyzable in the marine environment. Therefore, the coating surface continuously renews even under the static condition.<sup>471</sup> Marine field tests demonstrated that such a coating has excellent antifouling efficiency over 3 months in natural seawater. The renewal rate of such degradable PU could also be regulated by fouling-resistant side groups, such as *N*-(2,4,6-trichlorophenyl)maleimide or 2-(dimethylamino) ethyl methacrylate.<sup>472</sup> The combination of a dynamic surface and fouling-resistant groups can significantly improve the antifouling efficiency against marine microorganisms (Fig. 63).

To prolong the service life of a coating, a small amount of antifoulant can be added. The system consisting of degradable PU with biobased soft segments such as poly(1,6-hexamethylene adipate) and 4,5-dichloro-2-octyl-isothiazolone (DCOIT, a relatively eco-friendly organic antifoulant) exhibited stable and persistent release of the antifoulant, and the cumulative release linearly increased with time (correlation coefficient > 0.99). Such a coating was free from biofouling after 6 months of

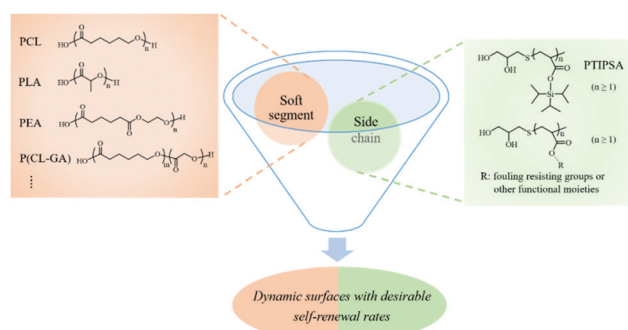


Fig. 63 Dynamic surface formed by PU with polyester as the soft segments and hydrolyzable side chains. Reprinted from ref. 468 with permission. Copyright (2019) Royal Society of Chemistry.

immersion in the sea.<sup>473</sup> DCOIT could be replaced with natural compounds such as butenolide derived from marine bacteria such as *Streptomyces* sp.<sup>474</sup> In the presence of enzymes secreted by a microorganism, PU degraded faster and released more antifoulants, indicating that the environment-adaptive system exhibited higher antifouling ability under heavier fouling pressure.

A dynamic surface can also be constructed by poly(ester-co-acrylate)s synthesized by hybrid copolymerization or ROP. These reactions allow inserting ester bonds in the backbone of polyacrylates with intrinsically remarkable coating performance, generating degradable polyacrylate coatings. A copolymer of methyl methacrylate (MMA) and 2-methylene-1,3-dioxepane (MDO) can enzymatically and hydrolytically degrade because of esters in the main chains, and the degradation rate increased with the ester content.<sup>475</sup> A polymer coating with a low amount of DCOIT exhibited excellent antifouling performance in seawater. Because of the diversity of vinyl monomers, the degradable polyacrylates could be imparted with various functionalities. For instance, the copolymer of MDO, MMA, and tributylsilyl methacrylate with degradable main chains and hydrolyzable side groups exhibits not only controllable surface renewal but also self-smoothing ability, reducing the hydrodynamic drag of ships.<sup>476,477</sup> A coating consisting of such a polymer and DCOIT can effectively resist biofouling in the sea for a long time. Further, degradable polyacrylates with fouling-resistant moieties has excellent antifouling and mechanical performances. For example, copolymer carrying the tertiary carboxybetaine (TCB) ester can be hydrolyzed in seawater to generate zwitterions with resistance to nonspecific protein and bacteria (both marine bacteria and clinically related bacteria).<sup>478</sup> Moreover, these zwitterions were renewed by main-chain degradation, which was favorable to maintain both mechanical robustness and fouling resistance.

In short, biodegradable polymers for marine antibiofouling based on the strategy of dynamic surface antifouling are summarized. Fouling-resistant side groups or antifoulants can be introduced into biodegradable polymers for prolonging the service life of coatings. Moreover, the idea of dynamic surface antifouling was tested to be valid. In the future, environment-adaptive dynamic surfaces that respond to fouling pressure, temperature, salinity, and navigation speed need to be designed, which can yield high efficiency and long service life.

In this section, *in situ* polymerization methods used for protein modification are initially summarized, which offer many potential applications, such as targeted delivery as cancer therapy or BBB penetration as Parkinson's therapy, biosensor, and biofuel. Then, CPs are briefly introduced for their DNA sensing, bioimaging, and therapeutics. After that, various functional polymers used for drug and gene delivery are discussed, namely, stimuli-responsive polymers, polymer vectors, and polyprodrug amphiphiles. Finally, surfaces/interfaces between biomaterials and bioenvironments are discussed, because they play a critical role in biomedical applications. Although significant advances in biomedical polymer materials have been achieved in recent years, there are still many challenges that are needed to be addressed. For example, it is still difficult to

obtain imaging materials with precise imaging performances and good biosafety. The biosafety of polymer materials used in drug delivery is not easy to achieve because they possess the ideal delivery function. Researchers should put a considerable amount of effort to obtain materials with ideal specific functions and biosafety.

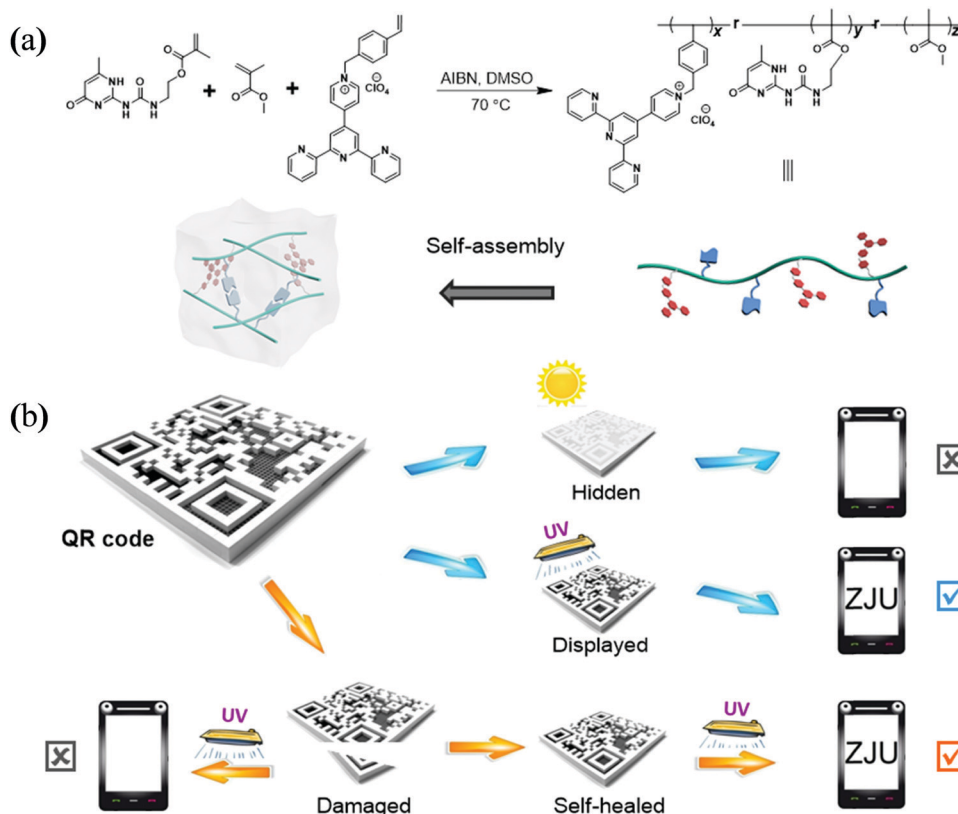
## 7. SP materials

Beyond polymers based on covalent bonds, SPs are produced by a combination of supramolecular chemistry and polymer science.<sup>479–484</sup> They refer to aggregates formed by the noncovalent interactions of low-mass molecular/polymeric building blocks and exhibit polymer properties in solution or in bulk. Noncovalent interactions comprise hydrogen bonding, host-guest interactions, ionic bonding,  $\pi$ - $\pi$  stacking, charge transfer interactions, metal-ligand coordination, hydrophobic interactions, *etc.* Besides conventional polymer properties, SPs exhibit thermal response because they comprise noncovalent bonds with lower bond energy. That is, SPs can be degraded into their building blocks when the temperature is increased, and SPs are reformed when the temperature is decreased; this is a better processing property than that observed in covalent polymers. Moreover, SPs can also show response to other external stimuli, such as light, electricity, and chemicals. The reversibility of noncovalent bonds endows SPs with unique properties, such as self-adaptation and self-healing, which are difficult to be achieved by covalent polymers. In addition, using easy-to-operate noncovalent bonds, various properties and functions can be facilely integrated into SPs and can be reversibly tailored, endowing SPs with potential applications in the robotic industry, biomedicine, and sensors, which has attracted wide attention in this field. In this section, subsequent discussions are divided into four research directions of SPs due to space limitations: responsive fluorescent SP gels, selenium-containing polymer materials, self-assembly of hyperbranched-based SPs, and BCPs.

### 7.1 Responsive fluorescent SP gels

Responsive fluorescent SP gels, a kind of smart gel with a three-dimensional (3D) network structure containing fluorophores, are crosslinked by noncovalent interactions, such as multiple hydrogen bondings, electrostatic interactions,  $\pi$ - $\pi$  interactions, metal-ligand coordination, van der Waals interactions, and host-guest interactions.<sup>485–490</sup> In particular, these noncovalent connections affect the fluorescence properties of the chromophores because their state of aggregation and energy transfer can be regulated by the assembly-disassembly process. Considering these interesting properties, responsive fluorescent SP gels have facilitated the evolution of new materials with applications in fluorescent sensors, biological probes, and imaging agents.<sup>491–493</sup>

For example, Huang *et al.* reported a novel yet facile approach to prepare white-light-emitting fluorescent polymeric materials resulting from the aggregation of a single fluorophore (Fig. 64).<sup>494</sup>



**Fig. 64** (a) Synthesis of the polymer and its self-assembly in solution to form a white-light-emitting fluorescent SP gel *via* intermolecular hydrogen bonding. (b) Schematic representation of data recording, data security protection, and data recovery of a quick response code constructed using this gel. Adapted from ref. 494 with permission. Copyright (2017) Royal Society of Chemistry.

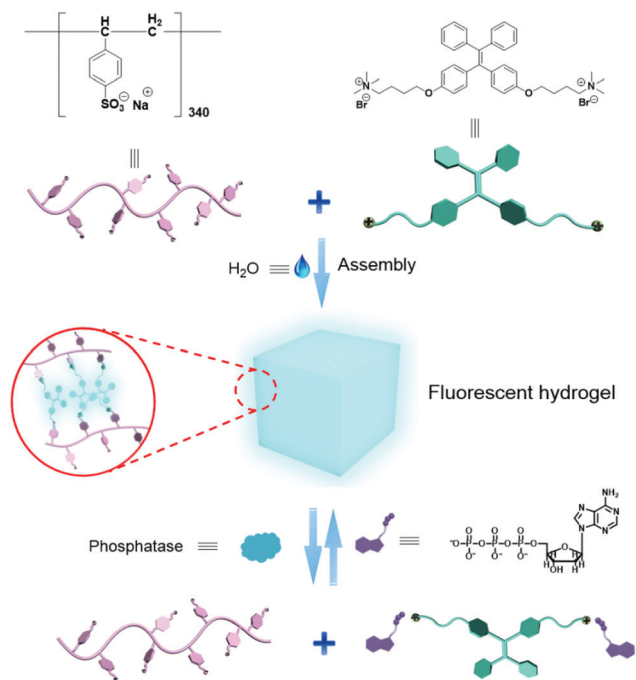
This aggregation resulted from the energetically favorable self-assembly of polymers *via* intermolecular quadruple hydrogen bonding. This advanced material was shown to be an ideal candidate for constructing intelligent information display/storage devices. A protected quick response (QR) code was fabricated based on this white-light-emitting fluorescent supramolecular gel. This code was shown to hide the information under natural light, yet it displayed it under UV light. Therefore, this code was protected and only readable under a specific condition. Furthermore, due to the dynamic nature of hydrogen bonds, the SP gel and the resultant QR code showed self-healing abilities, which is crucial for their applications. Information in an intentionally damaged QR code is incomplete and cannot be read under UV-light irradiation. However, the code can be mended based on the interfacial self-assembly of gels through multiple hydrogen bonding; subsequently, the protected information can be recovered. This novel QR code with improved information security and durability against mechanical damage can find applications in various fields.

Then, Huang *et al.* constructed a novel bioresponsive supramolecular fluorescent hydrogel by electrostatic interactions between poly(sodium *p*-styrenesulfonate) and a TPE derivative containing two quaternary ammonium cations (Fig. 65).<sup>495</sup> This self-assembly process led to the aggregation of TPE, which is a typical luminogen with AIE properties, thereby endowing this hydrogel with AIE properties. Moreover, this supramolecular

fluorescent hydrogel was responsive to biomolecules. When ATP was added to this system, the network structure of the hydrogel was destroyed and the aggregation extent of TPE was decreased, which induced the transition from the gel state to the solution state and decreased the fluorescence intensity of the hydrogel at the same time. Subsequently, after ATPase was added to the solution, ATP was decomposed and the electrostatic interactions between the polymer and TPE were rebuilt, transforming solution into gel and the fluorescence intensity of the hydrogel was recovered. Therefore, the fluorescent supramolecular hydrogel was bioresponsive. Simultaneously, this fluorescent supramolecular hydrogel could cross the cell membrane and enter the cytoplasm. The hydrogel system could be potentially applied to characterize the biooxidation process in the cytoplasm. In addition, it can be further used to prepare supramolecular systems for other bioapplications, such as drug delivery, bioprobes, and biological detection.

## 7.2 Selenium-containing dynamic covalent polymer materials

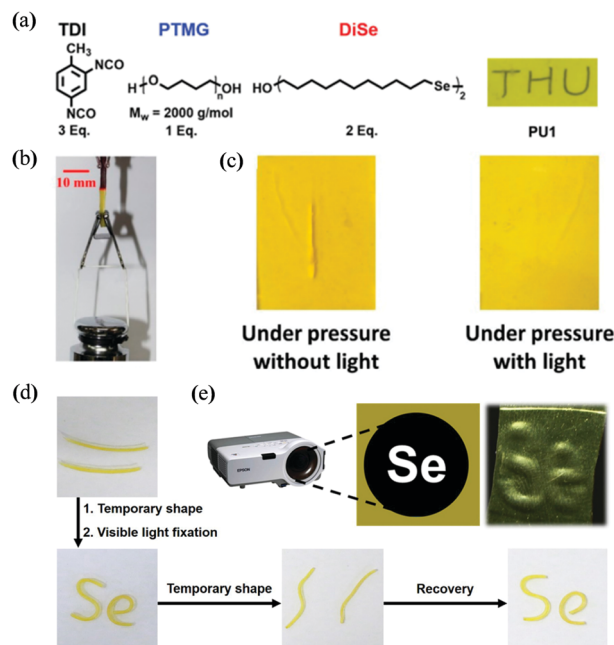
Dynamic covalent bond (DCB), which was first proposed by Jean-Marie Lehn in 1999,<sup>496</sup> refers to a unique chemical bond that can cleave, reform, and exchange under certain stimuli.<sup>497,498</sup> For the past decade, it has been extensively applied in research fields such as adaptive materials and supramolecular chemistry.<sup>499</sup> In 2014, Xu *et al.* reported the discovery of diselenide covalent chemistry.<sup>500–502</sup> Diselenide-containing molecules can



**Fig. 65** Chemical structures of poly(sodium *p*-styrenesulfonate) and TPE derivative, and the schematic representation of the formation and bioresponsive properties of the supramolecular fluorescent hydrogel. Reprinted from ref. 495 with permission. Copyright (2018) Royal Society of Chemistry.

undergo an exchange reaction simply under visible-light irradiation without additional additives *via* a radical mechanism.

The unique visible-light stimuli response behavior of diselenide bonds was applied to produce different types of functional materials. Diselenide bonds containing PU were fabricated,<sup>503</sup> which could self-heal under visible-light stimuli due to the dynamic exchange of diselenide bonds (Fig. 66a–c). Different contents of diselenide bonds could lead to different mechanical properties and self-healing properties. Moreover, employing a laser as the light source not only accelerated the healing process but also offered the potential for remote healing. Next, the crosslinker, glycerol, was added to the earlier recipe, and a thermoset diselenide bond containing PU was produced,<sup>504</sup> which could serve as a shape-memory material. In particular, when the material was heated above 80 °C, it can be bent from its original shape (permanent shape) to another shape (temporary shape). This temporary shape could be memorized by cooling to room temperature. When heating back to 80 °C, the shape could recover from the temporary shape back to the permanent shape (Fig. 66d). It was found that a higher content of diselenide bonds would lead to better shape-memory behavior. Moreover, since the diselenide bonds are light-sensitive, stress relaxation could occur under visible-light stimuli. As a result, it is possible for the material to transfer from a temporary shape to a permanent shape *via* visible-light irradiation in shape-memory cycles. This is a special property as most of the thermoset shape-memory material only has one fixed permanent shape once formed.



**Fig. 66** (a) Composition of the diselenide-bond-containing PU. (b) Material could bear 200 g weight after healing; the red part was stained with Nile red for clarity. (c) The crack disappeared after 24 h of light irradiation, demonstrating the healing property. (d) Typical shape-memory cycle of the material. (e) Material patterning achieved by a normal projector. Reprinted from ref. 503 and 504 with permission. Copyright (2015 and 2017) from Wiley-VCH and American Chemical Society.

As an application demonstration, one can use a commercialized projector to pattern material with designed shapes (Fig. 66e). In a recent study, Xu *et al.* reviewed that Se–S bonds are also a dynamic covalent bond.<sup>505</sup> The dynamic metathesis between diselenide and disulfide could be manipulated by light at different wavelengths. Based on this, a disulfide and a diselenide containing PU were prepared with light-triggered attachable–cleavable behavior. When they are attached together and irradiated by UV light, the Se–S exchange reaction can connect two materials into one piece and could be stretched up to 75% strain. However, for visible light at a wavelength above 410 nm, the welded material would be separated back into two pieces as the Se–S bonds were cleaved back into Se–Se and S–S bonds.

Apart from bulk materials, diselenide bonds could also be incorporated with surface/interface chemistry. Xu *et al.* developed the diselenide dynamic covalent bonds into a versatile surface modification method with fast response (within 30 s) and reversibility.<sup>506</sup> The diselenide bonds could be modified onto different substrates such as PDMS, quartz, and ITO conductive film glass. Different diselenide molecules could then be immobilized onto the surface *via* the diselenide metathesis reaction to achieve a series of functions such as adjusting the wettability and photosurface patterning. Moreover, by using this technique, they successfully achieved liquid motion by visible-light irradiation in a tube under capillary force. Moreover, this technique could also provide an efficient modification method for surface bioconjugation, which has potential applications in clinical usage.

### 7.3 Self-assembly of hyperbranched-based SPs

One important component of the studies on the self-assembly of polymers is to figure out the correlation between the behaviors/properties of self-assemblies and the topological architecture of the involved polymers. Various behaviors and morphologies of the self-assembly of covalent polymers with different architectures have been formulated.<sup>507</sup> Although studies on the same issue of SPs have started, the category and topological structure of SPs, however, are still limited when compared with the large number of covalent polymers. Theoretically, it will be easier to prepare SPs with different architectures, especially complex ones, because modules with different topologies can be separately synthesized and selected to associate *via* specific and strong supramolecular interactions according to the requirement. Moreover, dynamic supramolecular bonds may endow the self-assemblies with interesting properties. In this subsection, the design strategy, self-assembly, and properties of a series of SPs with different architectures are summarized.

As simple architecture, the synthesis and self-assembly of supramolecular linear block copolymers (LBCs) were first reported.<sup>508–510</sup> Two linear building blocks with a recognition unit at one end of each block was mixed, and complexation happened between them. For example, Yuan *et al.* synthesized two end-decorated homopolymers, namely, poly(styrene)- $\beta$ -cyclodextrin (PS- $\beta$ -CD) and poly(ethylene oxide)-ferrocene (PEO-Fc), which could orthogonally self-assemble into a supramolecular diblock copolymer (PS- $\beta$ -CD/PEO-Fc) in aqueous solutions based on terminal host-guest interactions.<sup>508</sup> Supramolecular PS-*b*-PEO further self-assembled into vesicles. The vesicles exhibited voltage-responsive behavior, in which the assembly and disassembly speed could be controlled by the applied voltage strength.

To increase structural complexity, one block of linear polymer can be replaced with dendritic polymers to yield a linear-dendritic block copolymers (LDBC). LDBC refers to polymers comprising a linear chain that is attached to a dendritic block. As a mimic of nature, LDBCs resemble trees in architecture and can be categorized into two types: LDBCs and linear-hyperbranched block copolymers (LHBCs).<sup>511,512</sup> Although intriguing properties were exhibited, the preparation of LHBCs was usually based on covalent polymerization, which involved complicated multistep synthesis, and the covalent nature lacked dynamic controls over their self-assemblies.<sup>513,514</sup> By introducing supramolecular interactions into the polymers, supramolecular LHBCs were successfully synthesized.<sup>515–517</sup> For example, Zhou *et al.* reported a linear-hyperbranched supramolecular amphiphile and studied its self-assembly behaviors.<sup>515</sup> A supramolecular LHBC was constructed *via* the noncovalent coupling of hyperbranched polyglycerol (HPG) grafted from  $\beta$ -cyclodextrin (CD-*g*-HPG) and adamantane-functionalized long alkyl chain (AD- $C_n$ ,  $n = 18$ ), namely, AD- $C_{18}$ , by the specific AD/CD host-guest interactions (Fig. 67). The as-prepared supramolecular  $C_{18}$ -*b*-HPG exhibited amphiphilic characteristics due to the hydrophilic HPG block and hydrophobic alkyl chain. The self-assembly of  $C_{18}$ -*b*-HPG in water resulted in the formation of vesicles with considerable ductility and they were readily deformed by applying

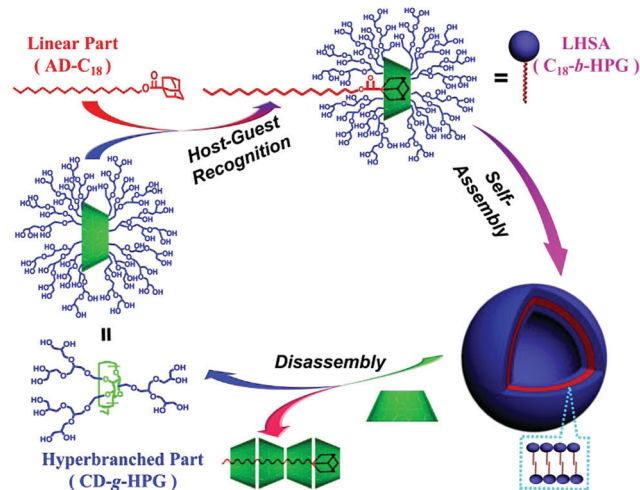


Fig. 67 Preparation, self-assembly, and disassembly processes of  $C_{18}$ -*b*-HPG. Reprinted from ref. 515 with permission. Copyright (2012) American Chemical Society.

force with AFM tips. The addition of a competitive host of  $\beta$ -CD led to the disassembly of vesicles into unimers. The appealing mechanical properties and additive-responsive behaviors of the vesicles were supposed to arise from the supramolecular nature of  $C_{18}$ -*b*-HPG.

The dynamic control over the self-assembly behavior of supramolecular LHBCs could be smarter, more complicated, and hierarchical *via* an appropriate molecular design. Subsequently, Zhou *et al.* prepared a “latent double-amphiphilic” LHBC and studied its morphological transition under light stimulus.<sup>516</sup> Two amphiphiles, namely, linear AZO-PEG (azobenzene-terminated PEG) and hyperbranched CD-*g*-HBPO ( $\beta$ -cyclodextrin-grafted HBPO), were coupled together to form a supramolecular LHBC of PEG-*b*-HBPO through “host-guest” interactions between AZO and CD under visible light. Amphiphilic PEG-*b*-HBPO self-assembled into vesicles in water. Under UV irradiation, however, PEG-*b*-HBPO was conversely dissociated into amphiphilic AZO-PEG and CD-*g*-HBPO. Meanwhile, the vesicles disappeared and turned into a mixture of nanofibers from AZO-PEGs and nanosheets from CD-*g*-HBPOs. This transition was totally reversible: the nanofibers and nanosheets could change back into vesicles again under visible light. The morphological transition, which represented a transition from one regular morphology (vesicle) into two regular ones (fiber and sheet), was reported for the first time and was considered to benefit from the dynamic supramolecular bonds in the polymer.

To further increase the structural complexity, one can replace both linear blocks with dendritic blocks to finally obtain a Janus macromolecule. Janus particles have attracted extensive attention owing to their unique and fascinating properties correlated to their asymmetric structure and functionalization as well as their promising potential to be used as new building blocks in self-assembly.<sup>518,519</sup> Janus dendritic polymers, which possess a 3D structure, were regarded as molecular Janus particles. Although covalent Janus dendrimers were successfully obtained through complicated multiple synthesis steps, the preparation of Janus

hyperbranched polymers (JHBPs) was difficult because the apex in HBP may be destroyed by intramolecular cyclization during the “one-pot” synthesis process. Zhou *et al.* overcame the obstacle by introducing supramolecular bonds into HBPs, and they reported the first example of JHBPs.<sup>520</sup> The JHBP of HBPO-*b*-HPG was obtained by complexation between CD-*g*-HPG and hyperbranched HBPO with an azobenzene group (AZO-*g*-HBPO) at the apex *via* specific AZO/CD host-guest interactions (Fig. 68). HBPO-*b*-HPG self-assembled into narrowly distributed vesicles in water under visible light; on the contrary, the vesicles disassembled into unimers under UV-light irradiation. In addition, data from dissipative particle dynamics (DPD) simulations matched very well with the experimental data and revealed the dynamics of the self-assembly process from randomly distributed molecules into the final vesicles and detailed structures of the vesicles.

By adding one more block, Zhou *et al.* further increased the structural complexity and obtained a “hyperbranched-linear-hyperbranched” triblock copolymer that resembled a dumbbell in shape.<sup>521</sup> The dumbbell-like supramolecular triblock copolymer (DSTC) was constructed *via* noncovalent host-guest coupling between the monotelechelic HPG grafted from  $\beta$ -cyclodextrin (CD-*g*-HPG) and ditelechelic azobenzenes-encapped linear polystyrene (AZO-PS-AZO). The resulting DSTC was amphiphilic and self-assembled into vesicles with a monolayer structure in water under visible light, and the vesicle size depended on the molecular weight of the PS block in the DSTC. In addition, the vesicles could disassemble into unimers under

UV-light irradiation, revealing light-responsive self-assembly behavior.

Later on, the same group reported an even more complexed dandelion-like SP (DSP) with a “sphere-star-parachute” structure.<sup>522</sup> The DSP was prepared by the noncovalent coupling of  $\beta$ -CD-encapped HBPO-*star*-PEO and adamantane-1-3,4,5-tris(*n*-dodecan-1-yloxy)benzamide (AD-*t*C<sub>12</sub>) comprising an adamantane head and three dodecyl tails *via* specific AD/CD host-guest interactions (Fig. 69). The DSPs exhibited novel hierarchical self-assembly behavior from vesicles to nanotubes through a fusion process in water. As a proof of concept, the DSPs were used to build an artificial light-harvesting system in an aqueous environment, taking advantage of the supramolecular nature and the self-assembled nanotube structure. The system was constructed by incorporating hydrophobic donors in the hydrophobic hyperbranched cores inside the nanotubes and the hydrophilic acceptors on the nanotube surfaces. This system could effectively prevent  $\pi$ - $\pi$  stacking between the donors and could provide a suitable distance between the donors and acceptors within the Förster radius, thereby yielding high energy transfer efficiency of over 90% in water. This work provides a solution to the low energy transfer efficiency in aqueous light-harvesting systems due to the poor solubility of fluorophores in water.

The introduction of supramolecular chemistry into polymers considerably simplified the preparation of polymers with complex architectures. Different “modules” can be separately synthesized

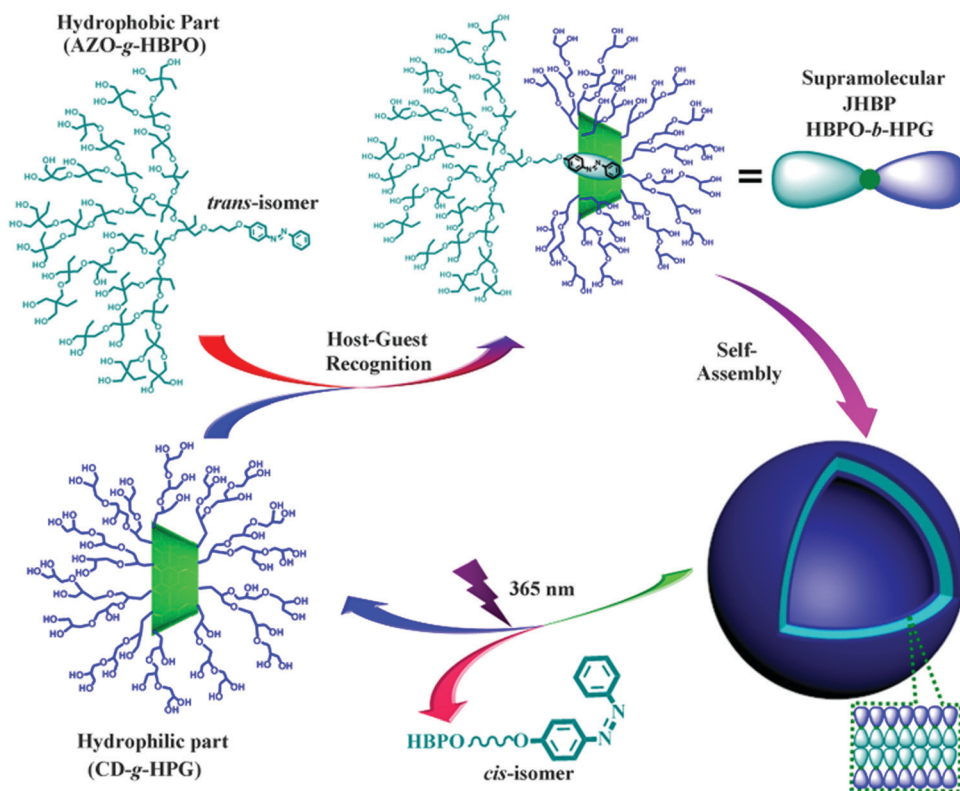


Fig. 68 Preparation, self-assembly, and disassembly processes of the supramolecular JHBP of HBPO-*b*-HPG. Reprinted from ref. 520 with permission. Copyright (2013) American Chemical Society.

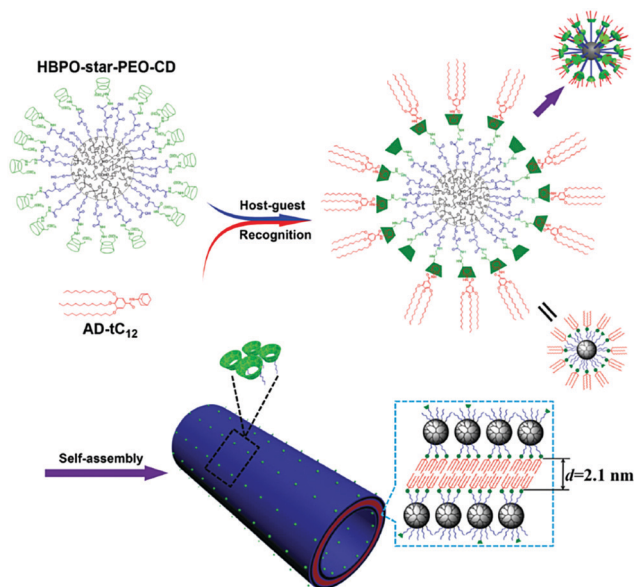


Fig. 69 Preparation and self-assembly processes of DSPs. Reprinted from ref. 522 with permission. Copyright (2016) from Wiley-VCH.

and assembled together through supramolecular bonds to finally get SPs with various topological structures. Following this design strategy, more and more SPs with different architectures and components are expected to emerge in the future, which can certainly extend the properties of polymers as well as their applications. Moreover, the complex structure and dynamic nature of supramolecular bonds in SPs may reveal more intriguing behaviors and properties of their self-assemblies.

#### 7.4 Self-assembly of BCPs under confinement

Self-assembly of BCPs has attracted considerable attention for many decades since it can generate well-ordered structures in a wide range of morphologies.<sup>523,524</sup> The preparation and use of

polymer materials usually suffers because of the external environment. For example, strong shear, elevated temperature, high pressure, and limited space are required in the process of injection molding. These extreme external environments have a significant impact on the microstructure and properties of polymer materials, endowing them with unique functionalities. When the self-assembly of BCPs takes place within confined spaces, the interfacial interactions, symmetry breaking, structural frustration, and confinement-induced entropy loss of the polymer chains play a dominant role in the final self-assembled structure, thereby resulting in some novel nanostructures that are not available in bulk and solution states.<sup>524–528</sup> Moreover, functional inorganic NPs are usually introduced to coassemble with BCPs or directly assembled with polymer-tethered NPs, which can effectively combine the respective functions of the polymers and NPs.<sup>529,530</sup>

Despite the rapidly increasing number of publications related to the self-assembly of BCPs or polymer-tethered NPs under confinement, there are still many open questions. For instance, the mechanisms of the confined assembly of BCPs are still under careful investigation. Control over the uniformity and stability of the assemblies and to get functional assemblies under confinement are still challenging. Due to their broad potential applications, the assembly behavior of BCPs and NPs under confinement has attracted considerable attention and has been the main research topic of Zhu *et al.* in the past decade. In this subsection, recent progresses made with regard to the assemblies of BCPs and inorganic NPs under confinement are addressed.

**Under 3D confinement.** Emulsion droplets were selected by Zhu *et al.* as soft and deformable confined space to systematically investigate the 3D confined assembly of BCPs.<sup>524–526</sup> By controlling the interfacial interactions, degree of confinement, and polymer–polymer interactions, BCP colloidal particles with tunable shapes and internal structures were created by confining BCPs into 3D droplets (Fig. 70). Unique shapes (*e.g.*, onion-, bud-, and pupa-like NPs with alternately stacked layers) can

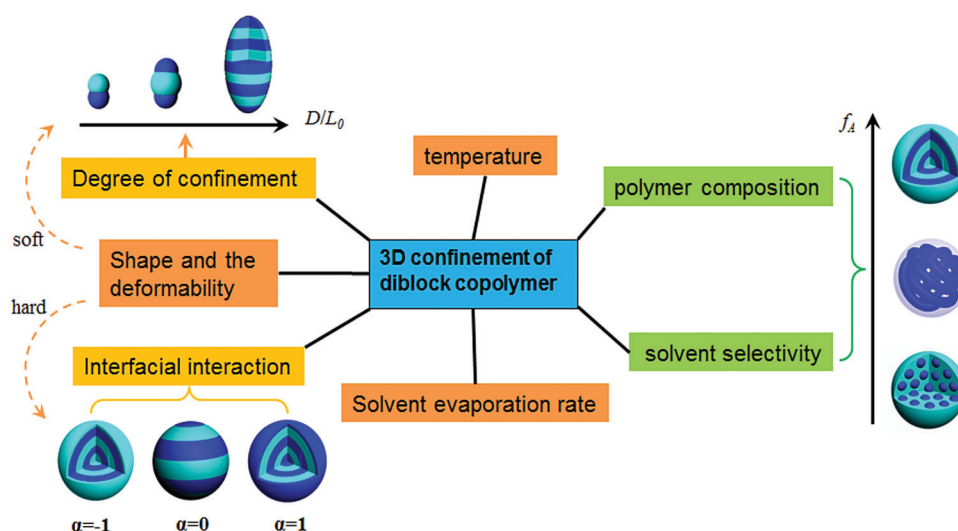


Fig. 70 Factors affecting the assembly behavior of BCPs in 3D confinement.

be obtained by tailoring the properties of the surfactants. In particular, neutral interfaces were created by blending surfactants with different affinities to achieve pupa-like NPs comprising alternately stacked layers. Interestingly, they succeeded in simultaneously tailoring both shape and internal structures of the NPs by solvent-absorption annealing. Moreover, the reversible transformation of nanostructured BCP particles could be achieved by switching the interfacial properties from neutral to selective interface.<sup>525</sup> This functional particle could have applications in drug delivery, catalysis, and smart material formation.

Using the concept of confinement, Zhu *et al.* constructed stimuli-responsive photonic crystal microspheres of BCPs.<sup>527</sup> Recently, the polystyrene-*block*-poly(2-vinyl pyridine) (PS-*b*-P2VP) particles with tunable periodic onion-like microstructures (Fig. 71) have been prepared *via* confined self-assembly in emulsion droplets, which exhibited angle-independent structural color after swelling in an ethanol/water mixed solution.<sup>528</sup> Interestingly, the structural color can be effectively tuned by varying the ethanol concentration because of the selective swelling of P2VP domains. These photonic microspheres showed potential applications in bio-diagnosis, colorimetric sensors, encoding, and active displays. This concept of confinement could be extended to construct confined BCP shells (*e.g.*, capsules). For example, hollow microcapsules with tunable shell structures have been fabricated under confinement, and the release behavior of encapsulated cargoes in these structured capsules was found to be strongly dependent on the shell structure.<sup>528</sup>

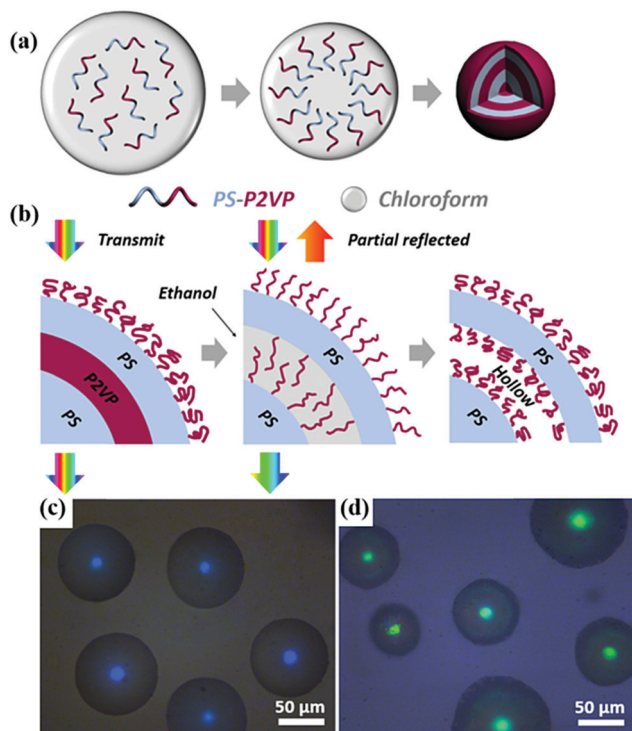


Fig. 71 Schematic illustration showing the preparation of PS-*b*-P2VP photonic microspheres through confined self-assembly in emulsion droplets. Adapted from ref. 527 with permission. Copyright (2018) from Wiley-VCH.

The above results indicate that confined assembly is a facile yet robust approach to control the shape and internal structure of BCP colloidal particles. Yet, despite the significant progress achieved in recent years, the assembly of BCPs in 3D confined space is still an emerging field. A better understanding of the control parameters that govern the confined assembly and nanostructures formation of BCPs should be explored in the future, which can certainly facilitate the design and fabrication of BCP materials with well-controlled structures and unique properties.

**Polymer-tethered inorganic NPs under confinement.** The self-assembly of polymer-tethered inorganic NPs under confinement is of great interest due to their unique structures and built-in functionalities. Zhu *et al.* reported the 2D confined assembly of PS-Au NPs in anodic aluminum oxide channels. Several unique assemblies (*e.g.*, linear chain, zigzag, two-NP layer, three-NP layer, and hexagonally packed NP structures) were obtained by manipulating the pore size and molecular weight of PS ligands.<sup>529</sup> Moreover, the electric field was applied to align the polymer-tethered Au nanorods under cylindrical confinement.<sup>530</sup> Various interesting hybrid assemblies (*e.g.*, single-, double-, triple-, or quadruple-helix; linear; and hexagonally packed structures) were obtained by adjusting the pore size of the channels, ligand length, and electric field orientation. Consequently, the surface plasmonic property of the assemblies can be tuned. The concept of the assembly of polymer-tethered inorganic NPs can be used for generating photonic crystal elastomers.<sup>531</sup> The photonic elastomers were prepared by incorporating isotropically arranged monodisperse SiO<sub>2</sub> NPs within a supramolecular elastomeric matrix based on metal coordination interaction between amino-terminated poly(dimethylsiloxane) and cerium trichloride. The photonic elastomers exhibited excellent tunable mechanical strength, angle-independent structural color, and self-healing capability, which may be useful in wearable devices, optical coating, and visualized force sensing.

Furthermore, a “rapid liquid-liquid interface confined assembly” strategy was proposed, which has been used to generate a non-close-packed PS-tethered Au NPs superlattice monolayer (AuNPs@PS SM) on the centimeter scale.<sup>532</sup> In this work, it was shown that a small amount (one drop, ~1 μL) of PS-tethered Au NPs solution could spontaneously spread on the water surface to form a monolayer thin film. During this process, AuNPs@PS self-assembled into centimeter-sized free-standing thin film with a non-close-packing structure (Fig. 72). The generated film has been successfully used to fabricate high-performance pentacene-based organic nano-floating-gate memory (NFGM). The *d* space of NPs (*e.g.*, interparticle distance) and the surface plasmon resonance spectra of the film can be tailored by adjusting the molecular weight of the tethered polymers. Precise control over the *d* value allows the successful fabrication of photosensitive NFGM devices with highly tunable performances from short-term memory to non-volatile data storage. More interestingly, after incisively revealing the assembly mechanism, it was found that this strategy is general and can be applied to NPs with various sizes. This strategy is significantly meaningful to microelectronic, photovoltaic devices,

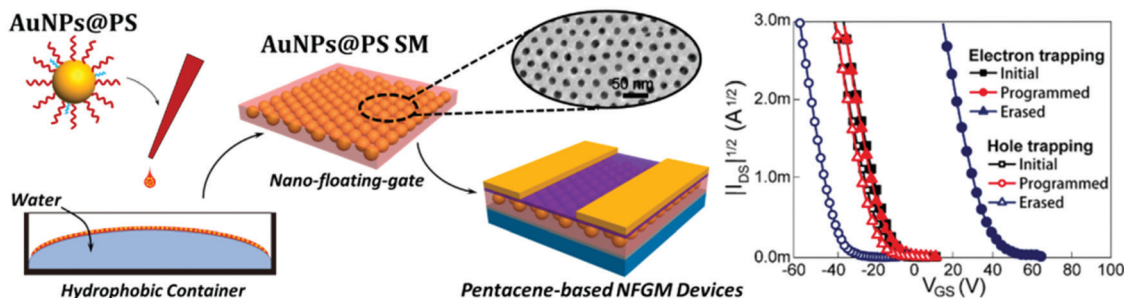


Fig. 72 Schematic illustration showing the strategy for fabricating AuNPs@PS SM through rapid liquid–liquid interface assembly on a convex water surface, and the design for NFGM devices based on a superlattice monomer. Adapted from ref. 532 with permission. Copyright (2018) from Wiley-VCH.

and biochemical sensors. These results indicate that the confined assembly of a polymer–NP system is a promising approach for tuning the ordered structure and functionality of hybrid materials.

The confined assembly of BCPs and inorganic NPs is a facile yet effective strategy to generate polymer and hybrid materials with well-ordered structures. Polymer particles with switchable shapes and internal structures can offer unique properties for this particular material. In addition, versatile hybrid nanomaterials can be designed on the basis of complementary advantages of functional inorganic NPs and polymeric materials *via* the confined assembly of polymer-tethered inorganic NPs.

However, despite the great progress made in recent years, the confined self-assemblies of BCPs and inorganic NPs still require more studies for a better understanding of the control parameters that govern the confined self-assembly and nanostructure formation. It still remains a challenge to utilize the well-defined BCPs or NP nanostructures for versatile applications (*e.g.*, solar cells, sensors, targeted drug delivery, metal catalysis, and magnetic storage devices) by taking advantage of the integrated and unique properties of these hybrid materials.

We first discuss responsive fluorescent SP gels that have many potential applications as fluorescent sensors, biological probes, imaging agents, drug delivery, bioprobes, biological detection, *etc.* Polymer materials containing dynamic diselenide bonds are then introduced, because they can be endowed with many functionalities, such as self-healing properties, adjusting wettability of the surface, and photosurface patterning. The various topological nanostructures of hyperbranched SPs and BCPs can be self-assembled by synthesizing different hyperbranched BCPs or under different confinements, which is summarized in the last part of this section. In general, the recent developments of SPs have been briefly addressed, which may provide some clues to readers for designing more abundant and higher-level SPs to meet the needs of various fields. Many challenges still remain for SPs, such as the *in situ* monitoring of the assembly process, exposing the rules of electrons, energy and mass transfer, and realizing the construction and practical applications of smart devices made of SPs.

## 8. Stimuli-responsive polymer materials

Stimuli-responsive polymers are smart materials whose chemical or physical properties can undergo reversible or irreversible

changes when they are exposed to external stimuli, such as temperature, pH, light, electricity, magnetic field, mechanical force, and biochemical molecules. They have drawn a great deal of research attention due to their applications in nanomedicine, diagnostics, flexible devices, smart actuation, optical storage, energy materials, and environmental governance. The existence, form, and mechanism of responsive polymer materials can be embodied in molecular chains, colloidal particles, self-assembled ordered aggregates, (hydro)gels, thin films, liquid crystal (LC) elastomers, and bulk materials. The response modes of such materials are different, including geometric dimension (shape) changes and recovery, changes in microenvironment polarity, cleavage or rebonding of covalent bonds, color and luminescence, surfaces infiltration, self-healing behavior, and so on. These different response modes are accompanied by their applications, such as shape changes that can mimic the action of muscles, cleavage or rebonding of covalent bonds that can control drug delivery and release, and self-healing behavior that can heal the polymers themselves. In this section, we only discuss electrochromic polymer materials and photoresponsive liquid crystal polymers (LCPs). SMPs, a type of stimuli-responsive polymers, will be addressed in the next section because of their unique mechanism and properties.

### 8.1 Electrochromic polymer materials

Electrochromism is the phenomenon in which materials can reversibly change their optical properties such as transmittance, reflectance, or absorbance in response to external potential.<sup>533,534</sup> Since the discovery of the electrochromic effect in transition-metal oxides by Deb *et al.* at the end of the 1960s, it has been applied in many fields such as smart windows, electronic displays, and mirror devices; therefore, state-of-the-art studies on electrochromism can have a significant impact on the environment, energy, and economy.<sup>535–537</sup>

The main classes of electrochromic materials are CPs (polythiophene (PTH), polyaniline (PANI), polypyrrole (PPy), and their derivatives), viologens and their derivatives, and transition-metal oxides ( $\text{WO}_3$ ,  $\text{NiO}$ , and  $\text{V}_2\text{O}_5$ ).<sup>538–540</sup> All the above-mentioned materials have gained popularity due to their ease of processability, high optical contrast, rapid switching times, and high coloration efficiency. In addition, many researchers have not only focused on improving the EC properties of the materials but

also attempted to combine other special functions. Hence, we provide examples of instances in which such materials have been constructed.

In terms of CPs, PTH and their derivatives are widely studied due to their modifiable structures to obtain multifunctional materials. In 2016, Xu's group designed a novel polymer with AIE and electrochromic properties, which was realized by introducing the TPE group into thiophene main chains (Fig. 73).<sup>541</sup> The polymer exhibited weak photoluminescence in THF, but its corresponding film prepared by spray-coating exhibited intense yellow-green fluorescent light at 540 nm. The color of the polymer film could be switched from bright yellow to navy blue by applying a relatively low voltage. An electrochromic device (ECD) was fabricated by using this polymer, which differs from common ECDs because both its color and fluorescent state could be synchronously switched by applying different potential level; therefore, the polymer is a unique candidate for electrofluorochromic and electrochromic applications.

AIEgens are strongly emissive in their aggregate states but weakly emissive in dilute solutions. The dual-state emission (DSE) polymers have strong emissions in both solution and solid states, broadening the scope of fluorescent displays. Xu *et al.* reported two kinds of PTH derivatives containing *trans*-stilbene (P1) and fumaronitrile (P2) groups, respectively.<sup>542</sup> These polymers emitted intense fluorescence in both solution and film states, attributable to the restriction of intramolecular rotation caused by multiple C-H $\cdots\pi$  bonds existing in the neighboring polymer backbones. Driven by different applied potential levels, the polymer films showed reversible electrochemical oxidation reduction reactions with colors changing from yellow-green to sky-blue and red to rufous. Furthermore, electrofluorochromism between the fluorescent state and nonfluorescent state happened synchronously (Fig. 74). This unique property makes the polymers a special candidate for optical applications.

With regard to composite electrochromic materials, Xu *et al.* investigated a quantum dots–polymer composite film, achieving the triple functions of electrochromism, electrofluorescence, and light-induced coloration effects (Fig. 75).<sup>543</sup> Combining the advantages of both cadmium selenide (CdSe) quantum dots and conjugated electrochromic polymers, the composite film

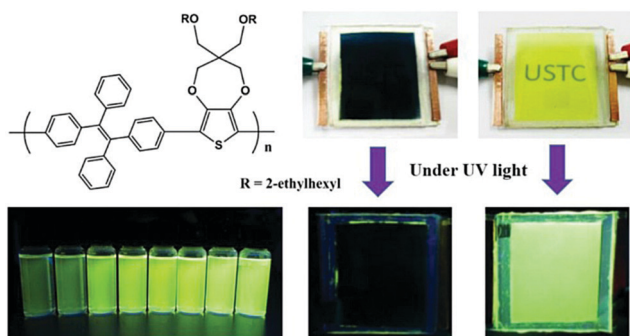


Fig. 73 Photographs of polymer-films-based ECDs and their corresponding transmittance and emission modulation. Reprinted from ref. 541 with permission. Copyright (2015) American Chemical Society.

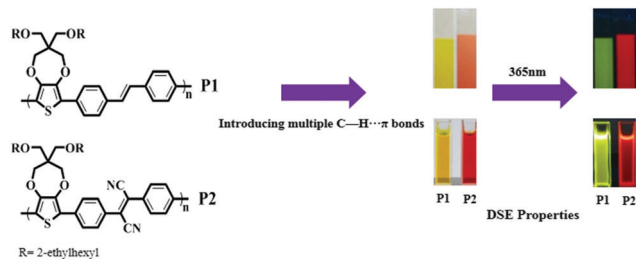


Fig. 74 Electrofluorochromic and electrochromic bifunctional polymers with DSE by introducing multiple C-H $\cdots\pi$  bonds. Adapted from ref. 542 with permission. Copyright (2018) from Elsevier Ltd.

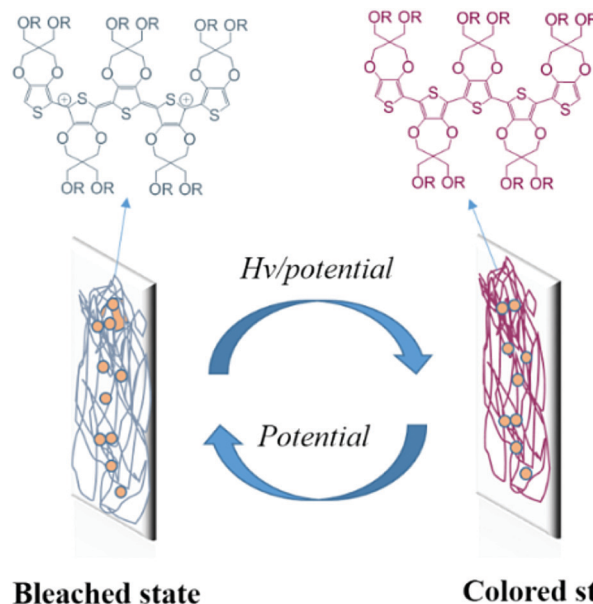


Fig. 75 Trifunctional CdSe quantum dots–polymer composite film with electrochromic, electrofluorescent, and light-induced coloration effects. Adapted from ref. 543 with permission. Copyright (2018) from Elsevier Ltd.

exhibited excellent electrochromic properties with low redox potential, high optical contrast, and fast response time. In addition, because of its enhanced performance, the composite film could switch between the nonfluorescent and fluorescent states, where the photoluminescence intensity was more than twice that of the pristine polymer film. Furthermore, the material could gradually become self-colored from the bleached state under sunlight irradiation.

Insolubility is a major limitation of conducting electrochromic polymers. To allow materials to be incorporated into large-area device architectures of interest, considerable efforts have been devoted over the last few decades. Water processability offers a green alternative to conventional organic-solvent-based processing methods that can facilitate the utilization of electrochromic polymers in industrial-scale device applications. Xu *et al.* reported a highly regiosymmetric water-processable blue-to-transparent electrochromic film based on diethylmalonate-derivatized PTH (Fig. 76).<sup>544</sup> The polymer exhibited high electrochromic contrast ( $\Delta T_{\text{max}}$ ) of 56% at 580 nm and a

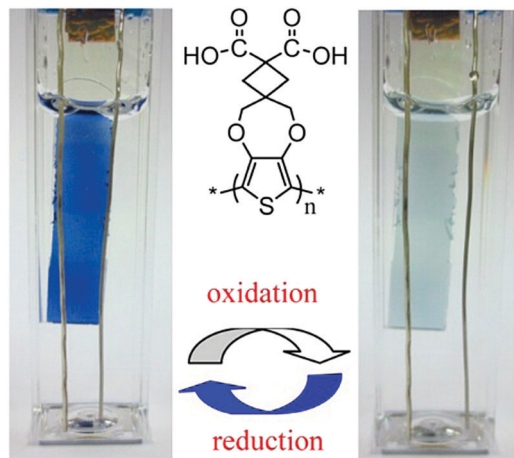


Fig. 76 A highly regiosymmetric, water-processable blue-to-transparent PTH-derivatized EC film. Reprinted from ref. 544 with permission. Copyright (2015) from American Chemical Society.

relatively fast switching speed of 1.8 s; further, it exhibited contrast loss of only 11% (from 56% to 45%) for a squarewave potential step of 5 s over 11 000 switching cycles, making it a desirable candidate for large-area electrochromic applications such as windows and displays, which have been long-lived aspirations, as reported by Xu *et al.*

Viologen is another widely studied organic electrochromic material. In an electrochromic system, viologen and its derivatives undergo reversible reduction under applied potential to form a viologen radical cation, which is considered to be one of the most stable radical molecules. The viologen radical cation exhibits a deep color because of its intense absorption of visible

light. Xu *et al.* reported a novel electrofluorochromic dual-function devices based on aryl-substituted viologen derivatives.<sup>545</sup> These devices comprise four viologen derivatives, namely, 4,4'-(1,4-phenylene)bis(1-butylpyridinium)hexafluorophosphate (PV), 4,4'-(naphthalene-1,4-diyl)bis(1-butylpyridinium)hexafluorophosphate (NV), 4,4'-(anthracene-9,10-diyl)bis(1-butylpyridinium)hexafluorophosphate (AV), and 4,4'-(benzo[*c*][1,2,5]thiadiazole-4,7-diyl)bis(1-butylpyridinium)hexafluorophosphate (BV). They were designed and functionally synthesized by introducing benzene, naphthalene, anthracene, and benzothiadiazole rings, respectively, between the two pyridine rings, which endowed these viologen derivatives with electrochromic and fluorescent switching properties. The devices comprising benzene viologen, naphthalene viologen, and benzothiadiazole viologen exhibited exceptional electrochromic properties with vivid color changes, high optical contrast, and fast response time under applied bias. In addition, owing to the increased conjugated p-framework degree of the designed compounds as compared to those of conventional viologens, all the fabricated devices could achieve intense fluorescence emissions (Fig. 77).

## 8.2 Photoresponsive LCs

As a kind of intelligent material, stimuli-responsive crosslinked liquid crystal polymers (CLCPs) combine the advantages of polymer networks and liquid crystal molecules, such as elasticity, anisotropy, molecular cooperation effect, and stimuli responsiveness.<sup>546</sup> Upon the application of appropriate external stimulation, the change in the mesogen arrangement in CLCPs could be transmitted to various macroscopic shape changes through the liquid crystal network. The incorporation of photochromic moieties, such as azobenzene, into CLCPs enables photoinduced

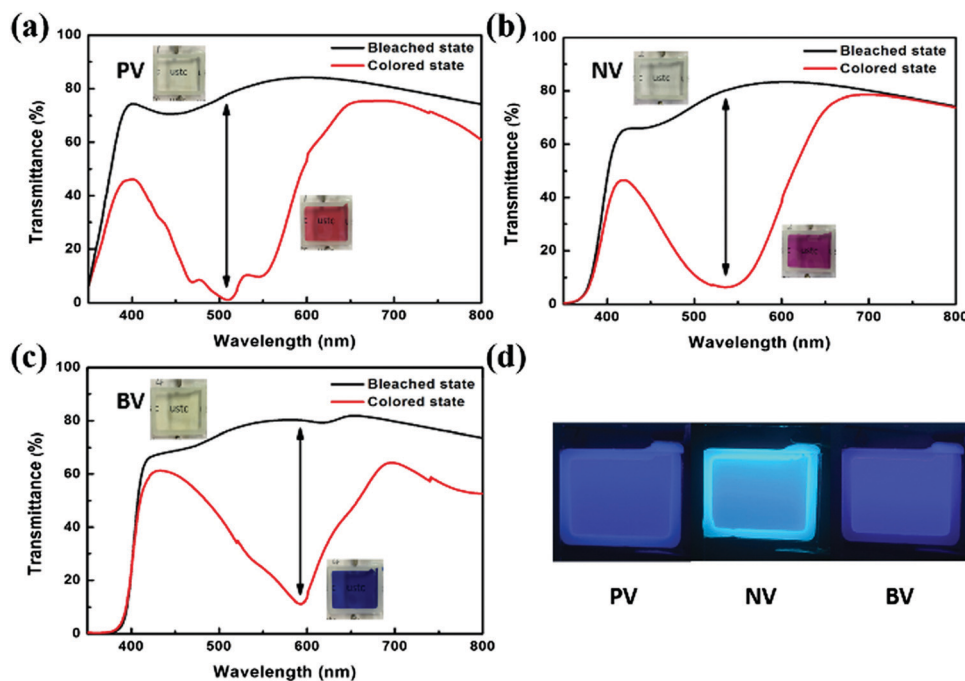


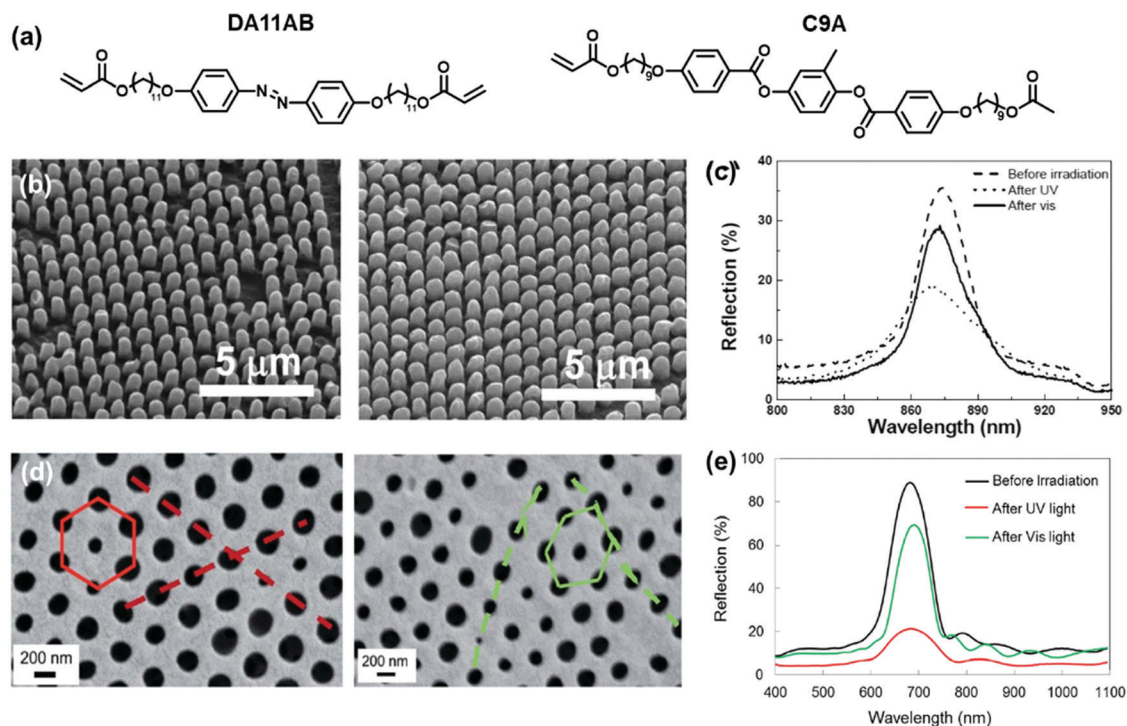
Fig. 77 UV-vis transmittance spectra of ECDs: (a) PV, (b) NV, (c) BV, and (d) photographs of their emissive states. Adapted from ref. 545 with permission. Copyright (2018) from Elsevier Ltd.

deformation in a remote and precise manner. The reversible *trans-cis* isomerization of azobenzene upon specific light irradiation induces a change in the LC order and consequently leads to quick shape variations in the CLCP samples.<sup>547</sup> The deformation process is based on the nanoscale molecular motion of the inner LC mesogens and requires neither complex architectural design nor precise assembly; therefore, photoresponsive CLCPs are appropriate materials to use for fabricating microscale actuators instead of traditional mechanical structures.<sup>548</sup> This subsection mainly describes the recent research by Yu *et al.* regarding the fabrication of photoresponsive microstructured CLCP actuators for photonic crystals, superhydrophobic surfaces, and microfluidics.

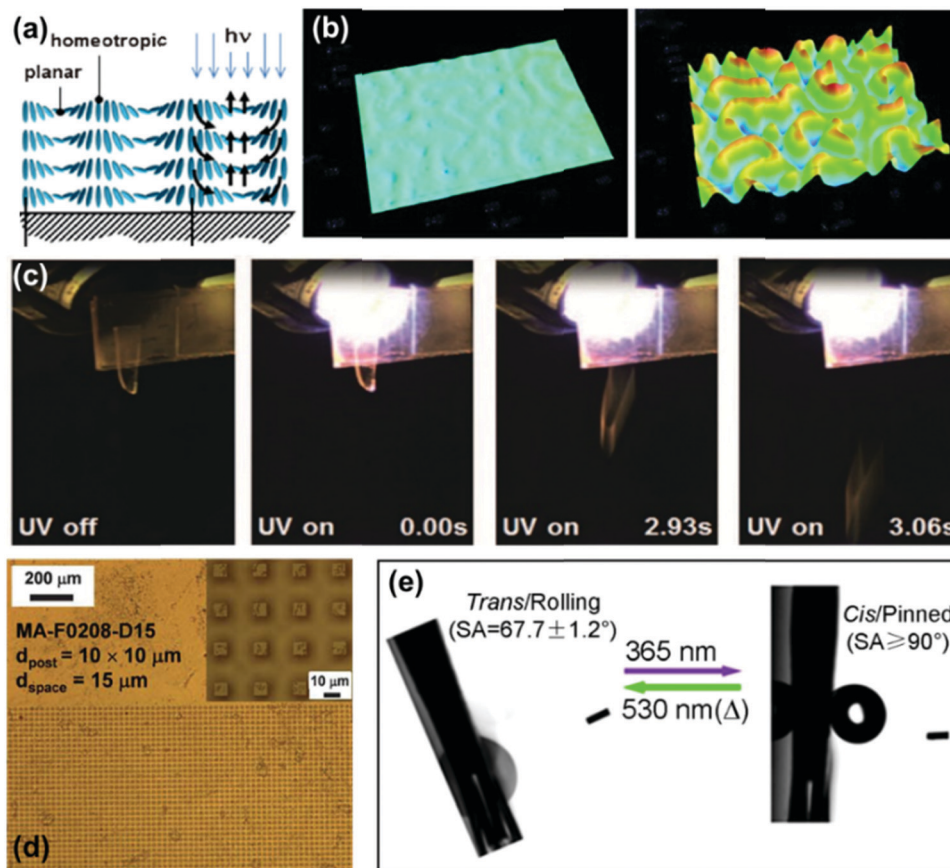
Photonic crystals contain periodic microstructures that reflect light of a particular wavelength. Adjustment of their microstructures can induce related reflection changes and have potential applications in display, communication, and anticounterfeiting technologies.<sup>549</sup> Most of the existing phototunable photonic crystals comprise small LC molecules that can facilitate a change in their arrangement. Yu *et al.* firstly reported phototunable CLCP photonic crystals that exploited the advantage of photoinduced deformation (Fig. 78a).<sup>550</sup> A two-dimensional (2D) periodic microarray was fabricated with azobenzene-containing CLCP *via* the replica molding technique. In accordance with geometry changes in the CLCP microarray induced by the photoisomerization of

azobenzene groups, the ON/OFF switching behavior on the reflection spectra of the microarray was observed when it was alternately irradiated with UV-vis light (Fig. 78b and c). Subsequently, they prepared a photo and thermal dual-responsive CLCP inverse opal, which was the smallest CLCP photonic structure.<sup>551</sup> Upon alternative irradiation of UV and visible light or temperature variations, the hole size of the inverse opal film changed (Fig. 78d), leading to rise and fall in the intensity of the reflection spectra (Fig. 78e), which could be ascribed to the deformation of CLCP induced by the photochemical reactions of azobenzene moieties or thermal-induced phase transition.

Liu and Broer *et al.* designed a kind of novel azobenzene-containing CLCP coating with the ability to change its surface topology in the microscale upon UV-light irradiation (Fig. 79a).<sup>552</sup> Due to the presence of chiral molecules in the LC mixture, LC mesogens spontaneously formed helical structures (the helix axis was parallel to the substrate), and the surface of the polymerized CLCP film was almost flat. Upon UV-light irradiation, protrusions were formed on the surface at planar positions, while wells were generated in homeotropic areas. As a result, the topological morphology of the CLCP surface was changed into a fingerprints-like structure (Fig. 79b). The mechanism of this remarkable photomechanical property is based on the reversible change in the order parameters by a photoinduced conformational change in the azobenzene units. Photoactivated generation of disorder in a



**Fig. 78** (a) Chemical structures of the LC monomers DA11AB and C9A. (b) SEM images of CLCP microarray before (left) and after (right) the irradiation of UV light. After UV-light irradiation, the mean diameter of the pillars increases from 0.587 to 0.695 μm, yielding a diameter increase of about 18%. (c) Reflection spectra of CLCP microarray under UV-light irradiation and subsequent visible-light irradiation. (d) SEM images of the inverse opal film (left) before and (right) after UV-light irradiation. The red and green regular hexagon and straight lines represent the arrangement of holes before and after UV-light irradiation. The distance between every two holes of the inverse opal film is 350 nm. (e) Reflection spectra of CLCP inverse opal film under UV-light irradiation and subsequent visible-light irradiation. Adapted from ref. 550 and 551 with permission. Copyright (2012 and 2014) from Wiley-VCH and Royal Society of Chemistry.



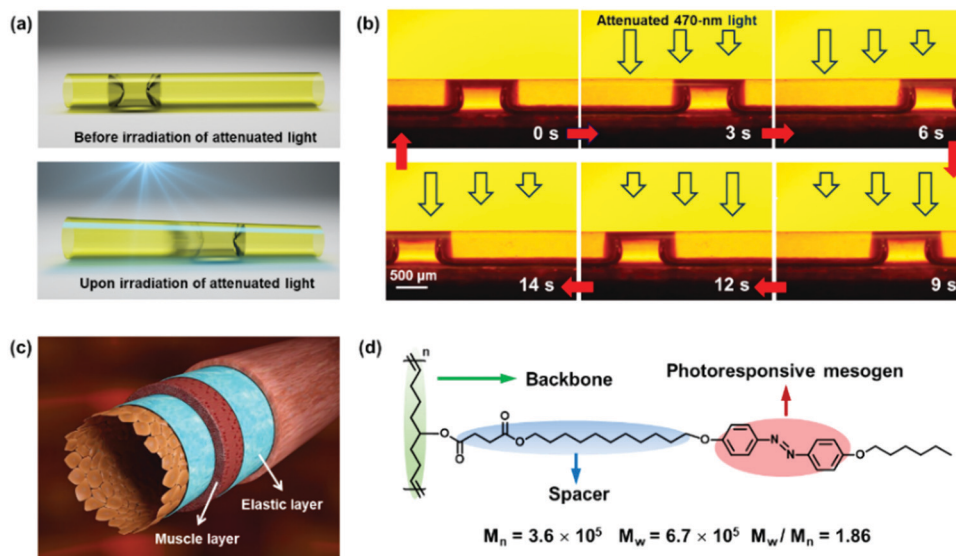
**Fig. 79** (a) Schematic representation of mesogen alignment in dynamic fingerprints. (b) 3D image of an initial flat state (left) and surface topographies under UV exposure (right). (c) Snapshots of a gripper that releases an object upon UV illumination. (d) Optical images of a microarrayed azobenzene CLCP film. (e) Light-controlled quick and reversible switching of superhydrophobic adhesion between rolling and pinning on microarrayed CLCP with a 2  $\mu\text{L}$  water droplet. Adapted from ref. 552 and 555 with permission. Copyright (2014 and 2012) from Wiley-VCH and Royal Society of Chemistry.

LC network produces free volume that can lead to the controlled formation of dynamic corrugations at its surface. In addition to the use of dynamic friction in controlling the grip and release (Fig. 79c), this dynamic surface microstructure can yield versatile applications, such as tracking light in a solar energy system, liquid mixing, or flow regulation in microfluidic channels.<sup>553,554</sup>

Superhydrophobic surfaces play wide-ranging roles in self-cleaning materials, lossless droplet transport, and open micro-reactors. When the adhesive force of a superhydrophobic surface is adjustable, it can work as a switch to control the movements of water droplets. In most of the existing works, the wettability and adhesive force of a superhydrophobic surface could be adjusted at the same time by changing the surface microstructures or chemical composition. It was challenging to regulate the adhesive force while maintaining the superhydrophobicity of a surface. In order to achieve the noncontact regulation of superhydrophobic adhesion, a CLCP microarrayed surface was fabricated by means of a secondary replicating method (Fig. 79d). The water sliding angle of the surface could be rapidly and reversibly adjusted by light irradiation while maintaining its large contact angle (Fig. 79e).<sup>555</sup> Recently, Yu *et al.* fabricated two different submicroarrayed LCP surfaces using the high-throughput etching technique of colloidal lithography.<sup>556</sup> These two surfaces

were easily fabricated by modulating different types of etching masks, which were uniform with average diameters of sub-250 nm and sub-400 nm. Owing to the surface chemistry and surface morphology designed to induce substantial hydrophobicity, the two surfaces exhibited completely different wetting behaviors with regard to water adhesion, mimicking a rose petal and lotus leaf, respectively. Here, azotolane was used as the photoresponsive moiety instead of azobenzene to achieve all-visible-light control; therefore, the wettability of these films could be reversibly transferred when alternately irradiated by light at 470 and 530 nm. This kind of noncontact real-time photocontrollable switching of microdroplet adhesion might inspire and facilitate the designs and fabrication of novel microfluidic devices and lossless transportation.

In 2016, Yu *et al.* demonstrated a conceptually novel way to manipulate liquids in a micro-LCP channel by exploiting the asymmetric capillary force arising from photoinduced deformation (Fig. 80).<sup>557</sup> Inspired by the lamellar structure of artery walls, they judiciously designed and synthesized a novel azobenzene-based linear liquid crystalline polymer (LLCP), where the flexible backbones and azobenzene mesogens self-assembled into ordered nanoscale lamellar structures due to the molecular cooperation effect. The non-crosslinked LLCP



**Fig. 80** (a) Schematics showing the motion of a slug of fully wetting liquid confined in a tubular microactuator (TMA) driven by photodeformation. (b) Lateral photographs of the light-induced motion of a silicone oil slug in a TMA fixed on a substrate that was taken through an optical filter to remove light with wavelengths below 530 nm. (c) Schematic illustration of the structure of artery walls. The middle coat of an artery, called the tunica media, comprising alternating muscle layers and elastic layers, which are responsible for stimuli-responsive deformation and mechanical robustness, respectively. (d) Molecular structure of LLCPC. Reprinted from ref. 557 with permission. Copyright (2016) from Nature Publishing Group.

was firstly fabricated into 3D microtubes, which exhibited geometry changes from cylindrical to conical upon light irradiation with gradient intensity, inducing asymmetric capillary force to move the inner water slugs. When compared with existing photocontrollable water transportation technologies based on wettability gradients or the Marangoni effect, a tubular LLCPC microactuator could transfer more types of liquids with faster speed. The transportation of multiphase liquids and even a different mix of liquids were achieved with remote photocontrol. Furthermore, the great advantage of water transportation in LLCPC microtubes is that when compared with traditional microfluidics based on nonresponsive materials, air pumps and complex pipelines are not needed to drive the liquid movement. Hence, the technology developed by Yu's group has great potential to simplify the microfluidic system and create small-volume micro-reactors, laboratory-on-a-chip contexts, and microoptomechanical systems.

In recent years, several burgeoning technologies, such as colloidal lithography, inkjet printing, molding, and electrospinning, have evolved to facilitate the fabrication of actuators at the micrometer and even nanometer scales, bringing photoresponsive LCP materials closer to real applications. It is anticipated that the preparation of microscale actuators will become the mainstream for the future development of photo-deformable LCPs, which requires not only the guidance of new concepts and theories but also creative material design and application of novel processing methods.

Electrochromic polymers with other functions, such as electrofluorochromism, electrofluorescence, light-induced coloration effects, water processability, and fluorescent switching property, are briefly reviewed. The design and fabrication of photoresponsive LCP actuators for photonic crystals, superhydrophobic surfaces,

and microfluidics are also addressed. However, further effort is needed for integrating stimuli-responsive polymers with other functions to fully demonstrate their broad potential applications. The future development of stimuli-responsive polymer materials is to further improve their specificity, sensitivity, and selectivity, as well as to promote their biomimetic construction and functional integration, broadening the potential applications.

## 9. SMP materials

Shape-memory effect (SME) for polymers<sup>16,17</sup> has been known for over 60 years. Products based on SMPs have also received commercial success since the early days, well-known examples of which are shrinkable tubing and shrinkable labels. However, this research field has only started to gain significant attention in this century, spurred by the realization of its enormous potential for use as deployable medical devices and aerospace structures.<sup>17</sup> In general, SMPs fall into two categories: one-way and two-way SMPs (2W-SMPs). SMPs that can recover their permanent shape from a temporary shape when exposed to an external stimulus but cannot change back to its temporary shape unless they are reprogrammed by applying an external force are called one-way SMPs. SMPs that can reversibly change between two independent shapes under two various external stimuli are called 2W-SMPs. For re-imparting the SME to the former structure after recovery, a new programming step has to be applied, which is not suitable for advanced smart applications. In addition, new applications beyond the widely pursued medical devices (particularly in advanced manufacturing) have emerged. Other advances<sup>16,17</sup> have also been made in extending shape recovery triggering mechanisms beyond direct heating. In this

section, we briefly highlight the most notable recent developments in extending fundamental behaviors/mechanisms for one-way and 2W-SMPs.

### 9.1 One-way SMPs

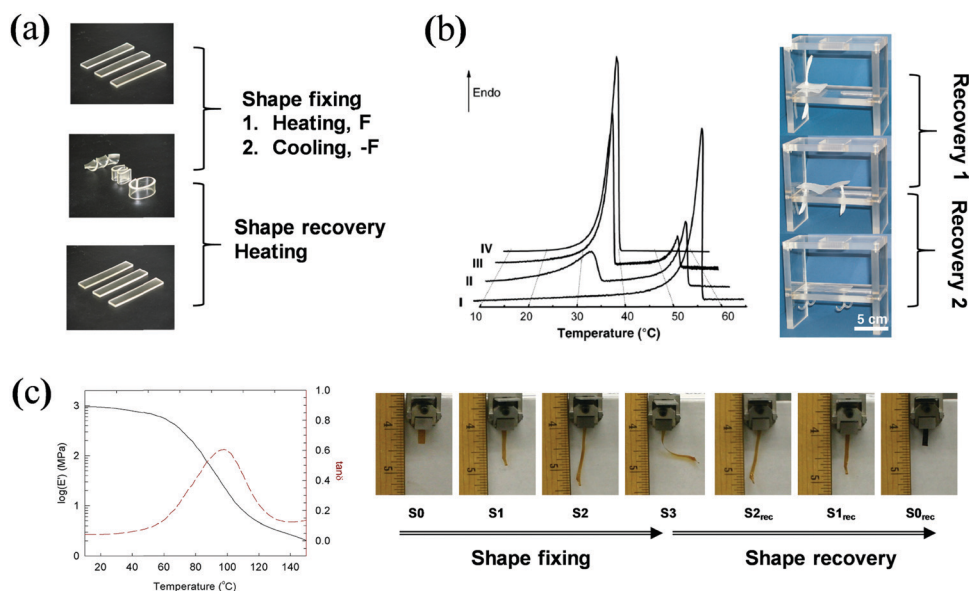
Depending on the number of memorized shapes, one-way SMPs can also be divided into two types: dual and multiple SMPs, according to their capability to memorize two or more shapes, respectively. Recently emergent thermadapt SMPs comprising dynamic covalent bonds are discussed in this subsection, because most examples of such SMPs are irreversible.

**Dual SMPs.** In its most common form, an SMP relies on one thermal transition to activate the polymer chain mobility by heating and freezing the chains by cooling. Chain freezing and activation allow temporary shape fixing and permanent shape recovery, respectively.<sup>17</sup> The corresponding thermal transition (typically melting or glass transition) is called shape-memory transition ( $T_{\text{trans}}$ ). The simplest and most classical dual SMPs rely on one thermal transition to fix one temporary shape, which can be recovered upon heating (Fig. 81a). Counting the permanent shape, each shape-memory cycle, therefore, involves two shapes. Although each shape-memory cycle can be repeatedly conducted, the need for reapplying an external force for reprogramming is required. Otherwise, the permanent shape cannot be switched back into the temporary shape. This forms the basis for dual SMPs. Progresses beyond this behavior are outlined hereafter.

**Multiple SMPs.** In 2006, Lendlein *et al.* revealed a triple SME (TSME), namely, two thermal transitions in an SMP network can be utilized for fixing two independent temporary shapes (Fig. 81b).<sup>558</sup> Although materials with two distinct transitions are not uncommon, the need to control each of them independently both in terms of transition temperature and their ratio is

not easily achievable from the material synthesis standpoint. This is further constrained by other properties (*e.g.*, surface characteristics) that should be designed into the material for potential device applications. These molecular design constraints stand in sharp contrast to the versatile designability of dual SMPs. In 2010, Xie *et al.* unraveled a more versatile physical mechanism that allowed achieving multishape-memory behavior beyond the triple version.<sup>559</sup> Instead of relying on two distinct thermal transitions for the triple-shape effect, a single transition in a broad temperature span of 75 °C was found to be sufficient for at least quadruple-shape-memory behavior (Fig. 81c). The underlying principle of this multi-shape effect is that a single broad transition is equivalent to the accumulation of an unlimited number of narrow transitions (called elemental memory units). Therefore, the number of temporary shapes is theoretically unlimited; however, in practice, it is constrained by the need to distinguish the temporary shapes by fixing them at sufficiently different temperatures. Nevertheless, the mechanism behind the multiple SME effect considerably extends the capability of SMPs. Critically, materials with a single broad transition are much more accessible than materials with two distinct thermal transitions.

**Thermadapt SMPs.** The geometry of the permanent shape plays a critical role in the device applications of SMP. Classically, SMP can be categorized as thermoplastic and thermoset SMP, where the permanent shapes of both of them are constrained by the molding modality. Here, 4D printing offers an alternative way to fabricate more complex permanent shapes,<sup>560</sup> but satisfying the printability characteristic limits the choices of materials. On this front, SMP networks containing dynamic covalent linkages provide unique opportunities.<sup>561,562</sup> Covalent bond exchange permits network topological rearrangement that can be utilized to reconfigure the permanent shape without being constrained by molding. The underlying solid-state



**Fig. 81** One-way SMEs. (a) dual-shape-memory cycle; (b) DSC curves for polymers with two distinct phase transitions and the associated triple-shape-memory recovery; (c) DMA curve for a polymer with a single broad transition and the associated multishape-memory cycle. Adapted from ref. 17, 558 and 559 with permission. Copyright (2015, 2006, and 2010) from Elsevier Ltd., National Academy of Sciences, U.S.A., and Nature Publishing Group.

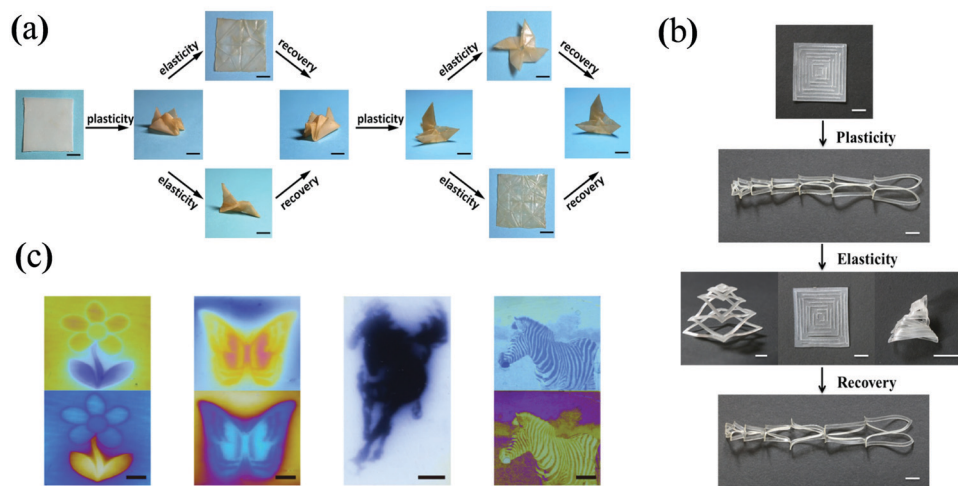


Fig. 82 Thermadapt SMP. (a) Origami-enabled shape memory; (b) kirigami-enabled shape memory; (c) digitally defined stress map visualized as the birefringent color under polarized light. Adapted from ref. 561, 562 and 563 with permission. Copyright (2016, 2016, and 2018) from Wiley-VCH, AAAS, and Nature Publishing Group.

plasticity offers a mechanism to manipulate the permanent shapes *via* origami<sup>561</sup> or kirigami<sup>562</sup> techniques. Therefore, geometrically complex shape morphing (as shown in Fig. 82a and b) could be realized. Although many complex permanent shapes can also be fabricated by 3D printing, solid-state plasticity uniquely and repeatedly allows permanent shape reconfiguration (Fig. 82a) in a cumulative fashion.<sup>561</sup> Importantly, a wide variety of covalent reversible bonds are accessible to design this class of materials, including the most common building units for classical SMP, ester, and urethane bonds.<sup>561,562</sup> Given the unique features of this type of SMP, Xie *et al.* called them thermadapt SMP in distinction to thermoplastic and thermoset SMPs. Besides the mold-free permanent shape reconfiguration, it is believed that there are many unique benefits that have yet to be explored. In a recent effort along this direction, Xie *et al.* illustrated that the synergetic manipulation of plasticity and elasticity (shape memory) resulted in an unparalleled opportunity to digitally program the stress map in a polymer (Fig. 82c).<sup>563</sup>

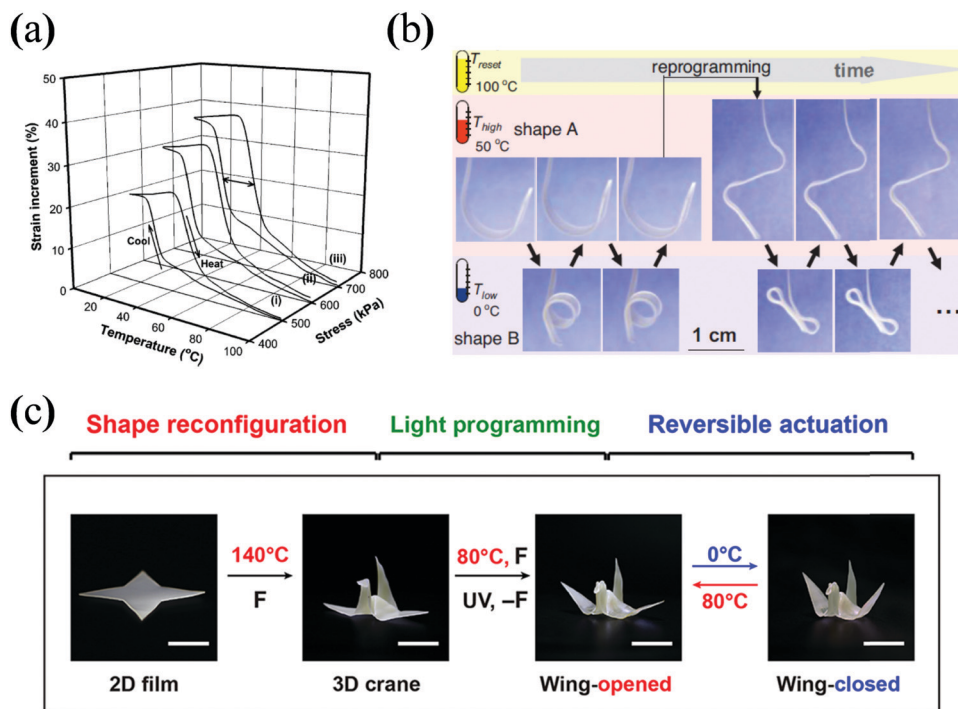
## 9.2 2W-SMPs

Although the one-way effect is sufficient for numerous applications, it is highly desirable to formulate two-way SME (2W-SME). Until now, 2W-SMPs have been further classified into quasi and true 2W-SMPs, depending on whether they need applied force to take effect or not. The feasibility of quasi 2W-SME was demonstrated by Mather *et al.* in 2008.<sup>564</sup> Their study demonstrated that a crystalline SMP, under a constant external load (Fig. 83a), could undergo reversible shape-shifting with temperature variations (below its crystallization and above its melting transition). Mechanistically, network anisotropy due to an external load causes directional crystallization/melting in the polymer, resulting in heating-induced contraction and cooling-induced elongation. Nevertheless, the versatility of this two-way shape-shifting behavior is severely constrained by the need to maintain an external load in the process. This limitation was elegantly addressed by Lendlein *et al.* in 2013 and Sheiko *et al.* in 2014.<sup>565,566</sup> In a network system

with two crystalline phases, the phase with higher melting transition was utilized to lock in internal stress, which permitted reversible shape-shifting *via* the other phase with the lower melting transition. Forgoing the external load, *i.e.*, true 2W-SME, versatile reversible shape-shifting in a programmable manner can be realized (Fig. 83b).<sup>565</sup>

In view of practical usability, true 2W-SMPs are obviously superior to the quasi versions. The available 2W-SMPs of this type include the following: (i) polymer laminates,<sup>567–570</sup> which are made by stacking layers of crosslinked semicrystalline polymers or crosslinked semicrystalline polymers/elastomers; (ii) liquid crystalline elastomers (LCEs),<sup>571–573</sup> and (iii) cross-linked semicrystalline polymers.<sup>574–582</sup> Laminated SMPs are difficult to be fabricated because of macroscopic heterogeneity. Moreover, they can only implement two-way bending deformation. With respect to LCEs, which operate because of the reversible anisotropic phase transition of mesogens, the synthesis routes are complicated and the experimental conditions are rigorous. Comparatively, 2W-SMPs derived from crosslinked semicrystalline polymers are rather promising in consideration of the accessibility of raw materials as well as manufacturing and usage of the end products.

Lendlein *et al.* made use of a specific semicrystalline polymer with a broad melting temperature range, which was divided into an upper melting temperature ( $T_m$ ) range (as the geometry-determining domain) and a lower  $T_m$  range (as the actuator domain).<sup>583</sup> The material realized 2W-SME by repeatedly melting the lower  $T_m$  portion. Similarly, the method was also adopted by other researchers to enable 2W-SME *via* the partial melting of crystalline regions.<sup>566,584</sup> Moreover, Wu *et al.* synthesized a two-way shape-memory IPN consisting of crosslinked elastomeric and crystalline components.<sup>585</sup> Meng *et al.* reported a two-way shape actuator with double chemical networks.<sup>586</sup> Xie *et al.* demonstrated much more complex 2W-SME in a polymer network with a single crystalline phase.<sup>587</sup> In particular, the network comprised both thermo- and photoreversible bonds. Thermoreversible bonds

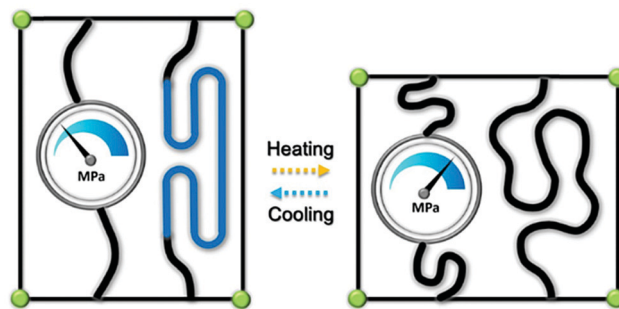


**Fig. 83** 2W-SME. (a) Quasi two-way shape-memory cycles under various constant stresses; (b) stress-free programmable two-way shape memory; (c) soft robot based on light-programmable two-way shape memory. Adapted from ref. 564, 565 and 587 with permission. Copyright (2008, 2013, and 2018) from American Chemical Society, Wiley-VCH, and AAAS.

were utilized to fabricate a complex 3D shape *via* solid-state plasticity. More importantly, photoreversible bonds allowed locking the network anisotropy in a spatioselective manner. Consequently, a flat sheet of polymer could be programmed into a 3D robotic crane that could undergo reversible wing-flapping (Fig. 83c). The rapid development in 2W-SME technology can open up future opportunities in unique device design.

By carefully analyzing the principles of documented systems, it is evident that an internal stress provider is imperative for building up a free-standing semicrystalline 2W-SMP regardless of its appearance and the way of interaction with the crystalline phases (Fig. 84), besides the chemical or physical crosslinkages that define the permanent shape.<sup>588</sup> The internal stress provider (*e.g.*, the higher  $T_m$  portion) plays the same role as external stress used for semicrystalline quasi 2W-SMPs and closely cooperates with the crystalline phase when temperature changes, inducing oriented recrystallization of the melted phase during cooling and restoring mechanical energy during heating.

Inspired by this finding, a series of facile methods are proposed by Zhang *et al.* to prepare 2W-SMPs by introducing new internal stress providers. Crosslinked elastomeric styrene-butadiene-styrene block copolymer (SBS),<sup>589</sup> for example, was successfully converted into an internal expansion stress provider in the crosslinked PU/SBS blend after appropriate programming. The temperature-induced reversible opposite directional deformability of the crystalline phase of PU, compressed SBS accompanying melting, and orientated recrystallization of PU led to the formation of the desired 2W-SME. Moreover, alkoxyamine moieties were introduced into crystalline polycaprolactone



**Fig. 84** Design principle of a semicrystalline 2W-SMP with an internal stress provider.

(PCL)-based crosslinked PU.<sup>590</sup> Owing to the synchronous bond fission/radical recombination of C–ON bonds in alkoxyamine, the stretched PU network underwent uneven partial stress relaxation depending on the triggering status of C–ON bonds. Accordingly, the considerably relaxed portions tended to maintain their extended status, while the nearly unrelaxed portions tended to recover their original undeformed shape. These two opposite tendencies generated an internal tensional force, which worked with the bidirectional deformation driven by the melting/recrystallization of the crystalline phase such that 2W-SME could be detected.

In fact, PCL-based PU can be coupled with 2W-SME in a simpler way. As revealed by the recent exploration by Zhang *et al.*,<sup>588</sup> proof-of-concept crosslinked crystalline PU was trained only by stretching and thermal treatment without using any

additional chemicals. The reversible melting/recrystallization of the crystalline phases elaborately collaborated with the tensed hydrogen bond network, because the inherent hydrogen bond network could function as an internal stress provider. The programmed crosslinked PU exhibited reversible shrinkage and extension with changing temperatures.

It is worth noting that SME-aided self-healing has been extensively investigated with the rapid development of self-healing polymers.<sup>591</sup> Contraction strain during the operation of SME can initially close cracks, facilitating intimate interfacial contact for the subsequent reestablishment of bonding. The method is particularly suited to the restoration of wider cracks; however, the existing solution mostly lies in one-way SME coupled with repeated self-healability. It is believed that an ideal case should be the cooperation between reversible SME and reversible self-healability. When reversible intrinsic self-healing ability is elaborately combined with 2W-SME in a single polymer, a larger crack can be repeatedly closed autonomously and then rebonded whenever it appears. In this context, no additional programming is needed to regain the SME after each healing event, which would give full play to multiple intrinsic self-healing events.

To verify this design, the abovementioned external stress-free two-way shape memory crosslinked PU/SBS<sup>589</sup> was modified by including alkoxyamine into the skeleton (Fig. 85).<sup>592</sup> The homolysis temperature of the included C–ON bond, *i.e.*, self-healing temperature of the material, was tuned to be higher than the triggering temperature of the 2W-SME. The thermal retractability offered by the SME coupled with the reversible C–ON bonds proved to be able to indeed successively close and repair a wider crack (width: ~250 μm). These works broaden the scope of SMP and intrinsic self-healing strategy.

In summary, significant progress has been recently made in the field of SMPs. Expansion of the shape-memory behavior beyond classical dual SMPs is particularly noteworthy, which includes multi-SMPs, thermadapt SMPs, and 2W-SMPs. These developments demonstrate the rich designability of SMPs, a key

advantage over its metallic counterpart, namely, a shape-memory alloy (SMA). These advances at the fundamental material level can lead to practical benefits at the device level in the long run.

## 10. Polymer membranes for separation

In recent years, polymer membranes have become emerging materials for separation due to their obvious merits when compared with conventional separation techniques, such as distillation, extraction, crystallization, and filtration. In particular, polymer membranes possess low carbon footprint, low operating costs, energy efficiency, high product quality, small spatial requirements, and so on. The increasing concentration of CO<sub>2</sub>, oily wastewater, shortage of freshwater, and water pollution have promoted the development of separation membranes, particularly polymer membranes.<sup>593</sup> Polymer membrane science is also an interdisciplinary subject, which involves polymer chemistry, polymer physics, materials science, material processing, and modification.<sup>594</sup> The increasing demand for water resources, comprehensive treatment of drinking water, desalination of seawater and brackish water, and recycling of wastewater have become more serious problems than the energy issue. Therefore, the fabrication of separation polymer membranes with excellent properties has become imperative, yielding high efficiency, flux, and selectivity; good antifouling performance; and high chemical stability. Generally, the applications of polymer membranes include four main subgroups: reverse osmosis, pervaporation, gas separation, and electrodialysis. Each subgroup has its own applications. For example, the main industrial application of reverse osmosis is the desalination of brackish water and seawater. The principal application of pervaporation is the dehydration of organic solvents. In this section, we present the recent advances in polymer membranes for oil/water separation, blood purification and water treatment, and water desalination.

### 10.1 Polymer membranes for oil/water separation

Polymer membranes have been widely used in diverse separation applications and have advantages in energy consumption, operation process, and secondary pollution problems over other separation technologies such as distillation, sublimation, crystallization, extraction, and ion exchange.<sup>595–598</sup> However, besides these advantages, surface wettability is a major drawback of traditional polymer membranes. Most polymer membranes such as polysulfone (PS), polypropylene (PP), and polyvinylidene fluoride (PVDF), are not superwetting (superlyophilic or superlyophobic) and suffer from serious fouling, particularly with oil/water separation.<sup>599–605</sup> The membrane flux and separation efficiency quickly decline due to fouling, which impedes the long-term use of such membranes and causes high cost for treating large amounts of oil/water mixtures. In contrast, superwetting polymer membranes have distinctly different wetting and dewetting behaviors with respect to oil

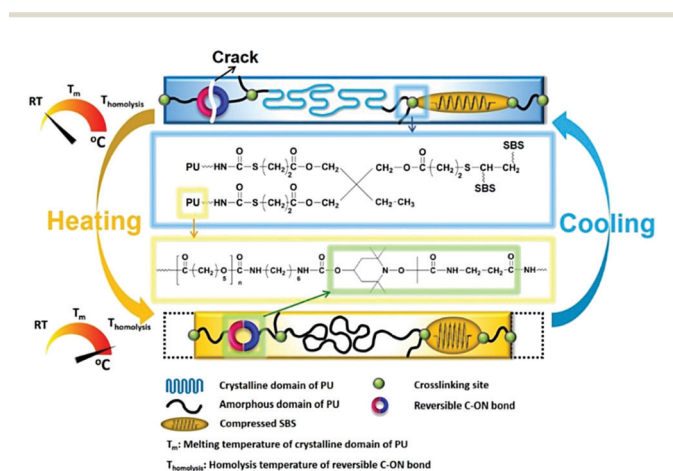


Fig. 85 Schematic of a crosslinked PU containing reversible C–ON bond for the self-healing of wider cracks. Reprinted from ref. 592 with permission. Copyright (2018) American Chemical Society.

and water. For instance, a superhydrophilic membrane is superoleophobic in water, and a superoleophilic membrane is superhydrophobic in oil. Therefore, superwetting polymer membranes are capable of more effective separation of oil from water or water from oil by the selective permeance and repellence and simultaneously control the fouling issue to the minimum.

In order to improve the antifouling performance of polymer membranes, blending modifications that involve directly blending lyophilic or lyophobic additives in a polymer matrix during the membrane-manufacturing process have been widely adopted because of their ease of operation and scaling-up feasibility for industrialization.<sup>601–603</sup> However, only a few blending additives can endow polymer membranes with superwetting property, particularly superhydrophilicity. An effective method for fabricating superwetting polymer membranes is surface modification that involves grafting or coating superlyophilic or superlyophobic modifiers onto the membrane surface. Another brand-new strategy for fabricating superwetting polymer membranes is to construct artificial hierarchical micro-/nanostructures on the membrane surface and accomplish superwettability *via* a combination of surface chemical composition and surface roughness. Very recently, nanomaterial-constructed membranes with the advantages of ultrathin thickness and tunable pore size at the nanometer scale have been comprehensively investigated. Polymer-decorated superwetting nanocomposite membranes were also tentatively applied for oil/water separation and exhibited both high efficiency and ultrahigh flux far beyond traditional polymer membranes.

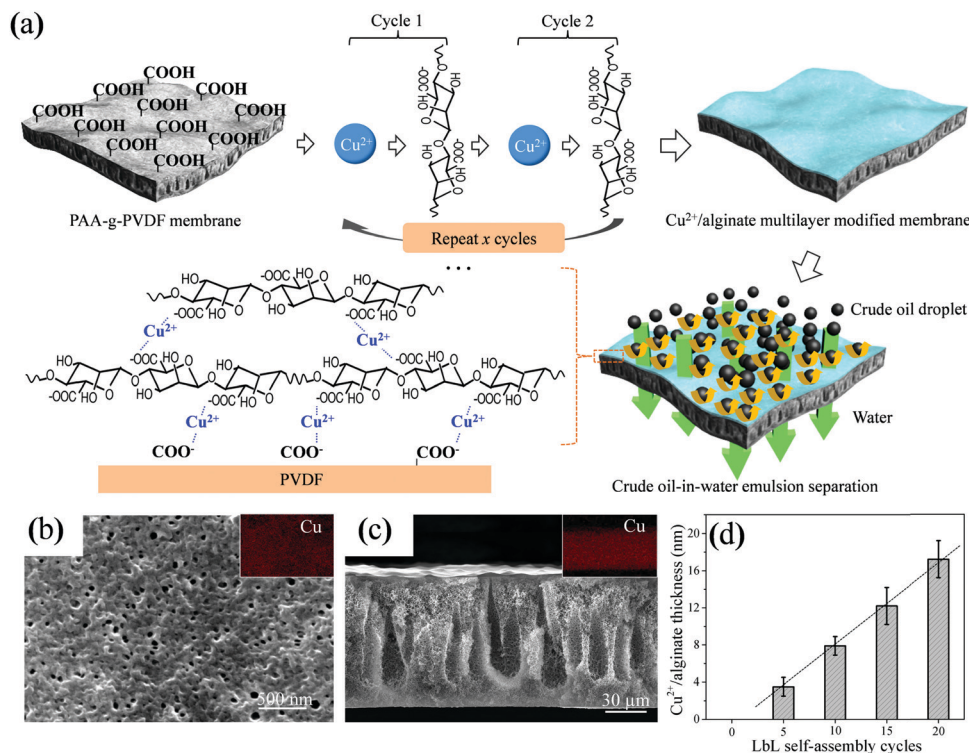
**Fabrication of superwetting membranes *via* surface modification.** When compared with blending modification that usually causes insufficient and uneven coverage of additives on the membrane surface, surface modification is a more effective method to ensure that the modifiers sufficiently and evenly distribute on the membrane surface and therefore endow polymer membranes with the desired superwetting property. A variety of surface modification methods, such as surface grafting, biomimetic adhesion, surface mineralization, and atomic layer deposition, have been adopted to fabricate superwetting polymer membranes *via* chemical bonding or physical absorption.<sup>606–611</sup>

Researchers usually adopt the surface grafting method in which superlyophilic polymer chains are immobilized onto membrane surfaces and function like “nanobrushes” to endow polymer membranes with superlyophilicity and antifouling property. In general, reactive groups need to be introduced onto the membrane surface in advance by introducing initiator sites or exposing the membrane to low-temperature plasma, ultraviolet,  $\gamma$ -ray, electron beam radiation, *etc.*<sup>612–614</sup> The grafted polymers are usually grown *via* “grafting to” or “grafting from” approaches.<sup>615</sup> Jin *et al.* synthesized the zwitterionic polyelectrolyte, poly(3-(*N*-2-methacryloxyethyl-*N,N*-dimethyl) ammoniopropanesulfonate), and grafted it onto a commercially available PVDF microfiltration membrane *via* a surface-initiated ATRP process.<sup>616</sup> After grafting the zwitterionic “nanobrushes,” the wettability of the PVDF membrane transferred from hydrophobic to superhydrophilic and superoleophobic in the presence of

water. Because of the superwetting property, this membrane could thoroughly separate a series of oils from oil-in-water emulsions with extremely high separation efficiency of above 99.999%, simultaneously with excellent anti-oil-fouling and cyclic performances. However, it has intrinsic drawbacks of time consumption and harsh chemical environment requirement during membrane fabrication, which considerably impedes the commercialization of this method to manufacture superwetting polymeric membranes.

Surface coating is another widely adopted method to fabricate superlyophilic polymer membranes and has promising potential for industrial mass production of membranes. PDA and polyethyleneimine (PEI) are a common coating combination for fabricating superhydrophilic polymer membranes, since PDA can be easily and robustly deposited on various solid substrates and superhydrophilic PEI is easy to be grafted on PDA *via* a Michael addition reaction. Xu *et al.* reported the fabrication of a superhydrophilic and underwater superoleophilic nanosilica-decorated PP microfiltration membrane with the assistance of a PDA/PEI intermediate layer.<sup>607,609</sup> In this case, PEI played an important role in the polycondensation of silicic acid *via* the so-called biomineralization process. Benefiting from the robust superwetting property, this nanosilica-decorated PP membrane exhibited very high permeation flux and very high oil rejection for surfactant-stabilized oil-in-water emulsions, as well as outstanding flux recovery after simply rinsing with water. It is worth noting that PDA, a well-known “bio-glue,” is widely used to coat not only polymer membranes but also inorganic membranes, and it can act as an intermediate layer before the formation of the desired separation layer. Other coating materials, such as metallic oxide NPs, metallic carbonate NPs, and even 10 nm-thick metallic oxide nanolayers, have been adopted to fabricate superwetting and antifouling polymer membranes.<sup>610,611,617</sup> Meanwhile, a collaborative mechanism exhibiting concomitant fouling resistance and fouling release during oil-in-water emulsion separation was proposed by Jiang *et al.* to explain the outstanding anti-oil-fouling performance of these membranes with both inorganic hydrophilic microdomains and organic hydrophobic microdomains. A drawback of the superwetting membranes obtained *via* surface coating is the instability of the coating layer, which is problematic for long-term use, particularly in acidic and alkaline environments.

A challenge in oil/water separation is separating highly viscous crude oil from oily water, where crude oil is easy to adhere onto the membrane surface and cause the fouling of membranes, even for those polymer membranes with underwater superoleophobicity toward light oil. A widely recognized theory is that the superoleophobicity and anti-oil-fouling property of a membrane is directly affected by its surface hydratability that can result in the formation of a sufficient hydrated layer to protect the membrane surface from oil contact and adhesion.<sup>618,619</sup> Therefore, hydrogels with strong hydratability are ideal materials for modifying polymer membranes and achieve low-adhesive superoleophobicity for the antifouling separation of crude oil. The key point in modifying polymeric membranes by hydrogels is to form sufficiently small versions on



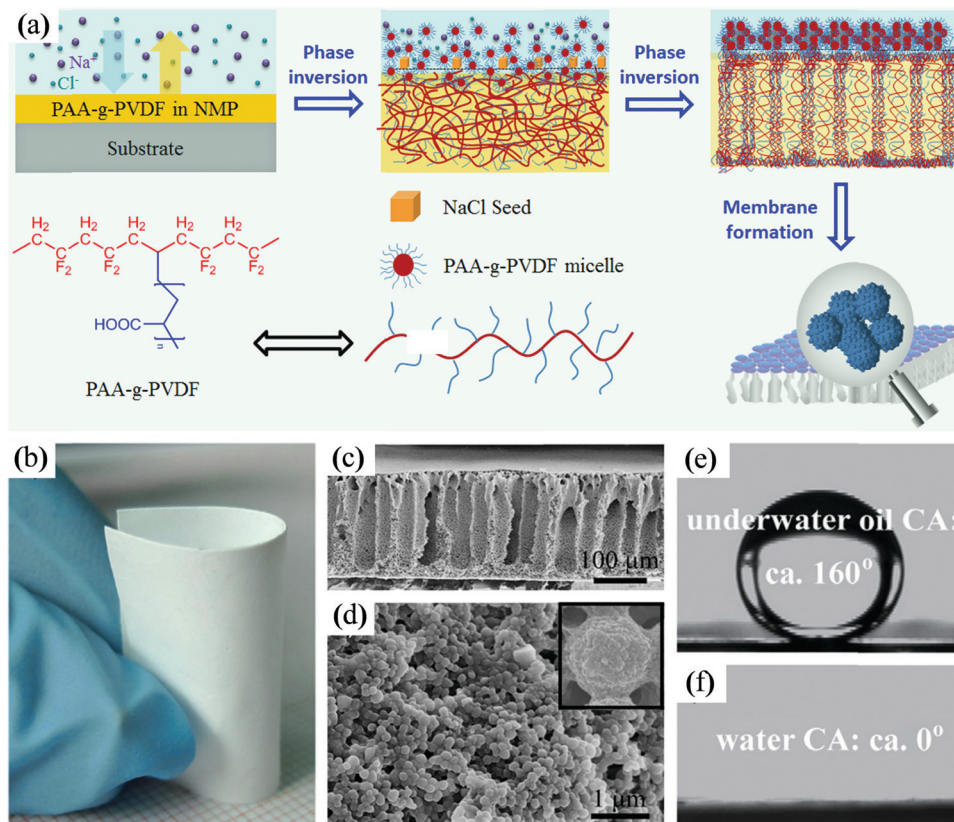
**Fig. 86** Ultrathin alginate-hydrogel-modified PAA-g-PVDF membrane. (a) Schematic showing the formation of alginate hydrogel on the membrane surface via LbL self-assembly and the antifouling separation of crude-oil-in-water emulsion. (b) Top-view and (c) cross-sectional SEM images of the membrane after 10 LbL cycles. Insets show the corresponding EDX images of Cu element distribution. (d) Thickness of the alginate hydrogel as a function of LbL cycles. Reproduced with permission from ref. 620. Copyright (2018) Wiley-VCH.

the membrane surface such that the micro- and nanosized membrane pores are unblocked. Jin *et al.* used a LbL self-assembly method with molecular-scale controllability to coat an ultrathin alginate hydrogel layer on a poly(acrylic acid)-grafted polyvinylidene fluoride (PAA-g-PVDF) ultrafiltration membrane (Fig. 86).<sup>620</sup> The thickness of the hydrogel layer was highly controllable from 3 to 17 nm, and the nanosized membrane pores were effectively maintained. An ionic bond was designed between the hydrogel layer and membrane substrate to ensure membrane stability. The hydrogel-modified membrane was superhydrophilic and exhibited underwater superoleophobic properties with nearly zero oil adhesion for crude oil and was capable of separating crude-oil-in-water emulsions with ultrahigh efficiency and outstanding antifouling ability. Another approach is to synthesize nanohydrogels and then decorate the nanohydrogel on polymer membranes. For instance, a sulfobetaine zwitterionic nanohydrogel (diameter: 50 nm) was synthesized via an inverse microemulsion polymerization process and then grafted onto a polymer microfiltration membrane.<sup>621</sup> The resulting membrane was also endowed with superwetting and anti-oil-fouling properties. It should be noted that the size of the nanohydrogels should match with the surface pore size of the polymeric membrane.

**Membranes with micro-/nanohierarchical surface structures.** According to classic Wenzel and Cassie models, the wetting behavior of a liquid on a solid surface is determined by the chemical composition (lyophilicity with high surface energy or lyophobicity with low surface energy) and surface roughness.<sup>622,623</sup>

Micro-/nanohierarchical surface roughness on polymeric membranes can provide considerably enhanced lyophilicity or lyophobicity and even generate superlyophilic or superlyophobic membranes.

The artificial construction of micro-/nanohierarchical surface structure with sufficient roughness during the membrane-forming process is crucial to obtain superwetting polymer membranes without subsequent surface modification. Jin *et al.* reported the construction of a spherical structure with the size of several hundreds of nanometers on a PAA-g-PVDF membrane via a salt-induced phase-inversion method (Fig. 87).<sup>624</sup> Distinguished from the general approach of adding salt as a pore-forming additive to change the pore structure of the membrane, the salt that was added into the coagulation bath at a nearly saturated concentration can function as nucleates to induce the aggregation of polymer micelles. These micelle aggregates on the membrane surface grow additionally in the subsequent phase-inversion process into the desired spherical particles. Synergistically combined with the artificial micro-/nanohierarchical surface structure and the hydrophobicity of PAA-g-PVDF, the resulting PAA-g-PVDF membrane has intrinsic superhydrophilicity and underwater low-adhesive superoleophobicity. Meanwhile, this membrane could effectively separate both surfactant-free and surfactant-stabilized oil-in-water emulsions, either under low applied pressure or solely driven by gravity. An inert solvent-induced phase-inversion method was also used to generate a spherical structure on PVDF membranes.<sup>625</sup> The added ammonia water was prone to induce



**Fig. 87** PAA-g-PVDF membrane with a micro-/nanohierarchical surface structure. (a) Schematic showing the fabrication of the membrane by a salt-induced phase-inversion process. (b) Photograph of the membrane. (c) Cross-section and (d) top-view SEM images of the membrane. Photographs of (e) an oil droplet on the membrane in water and (f) a water droplet on the membrane in air. Reproduced from ref. 624 with permission. Copyright (2014) Wiley-VCH.

localized microphase separation and caused the formation of “incipient precipitation” in the form of PVDF clusters, which functioned as a growing point to generate spherical microparticles during the subsequent phase-inversion process. As a result, the obtained PVDF membrane was uniformly skinless and comprised spherical microparticles with hierarchical micro-/nanostructures, which resulted in the superhydrophobicity and superoleophilicity of the membrane. Based on the oil-favoring and water-repellent properties, this PVDF membrane can also be applied for separating emulsified water from oil and organic solvents.

Generally, oil has much lower surface energy than water and wets a membrane more easily, so that the most hydrophilic or superhydrophilic membranes with high surface energy are simultaneously superoleophilic. Liu *et al.* reported this phenomenon with a PVDF-blend membrane. This membrane is superhydrophilic and underwater superoleophobic because of the inner hydrophilic additive and its spherulites-petaloid surface structure.<sup>626</sup> On the other hand, the membrane is superoleophilic and underoil superhydrophobic. Therefore, such a type of superhydrophilic polymeric membranes can achieve both oil separation from water and water separation from oil. When compared with surface modification, the artificial construction of a micro-/nanohierarchical surface structure yields membranes with intrinsic surface components and

omits the component-releasing problem for surface-modified membranes. However, it is still difficult to construct controllable micro-/nanohierarchical structures on the polymeric membrane surface by traditional methods. Innovative methodologies on this strategy are highly desired.

**Nanomaterial-constructed composite membranes.** The membrane with ultrathin thickness at the nanometer scale while retaining desirable pore structure and high mechanical strength is an ideal membrane for separation but hard to be accomplished with traditional polymeric or ceramic membranes. In recent years, the development of ultrathin membranes composed of one-dimensional (1D) nanomaterials (such as CNTs, metal hydroxide wires, and metal oxide wires) and 2D nanomaterials (such as graphene nanosheets, graphene oxide nanosheets, and metal sulfide nanosheets) with unprecedented separation performances have been reported. These nanomaterial-constructed membranes have been extensively explored for devising separation applications, including oil/water separation, molecule separation, ion sieving, and gas separation.

Single-walled carbon nanotube (SWCNT), a chemically stable and high-strength 1D nanomaterial, is a good choice to construct ultrathin network membranes with tunable thickness and pore size at the nanometer scale. Jin *et al.* constructed SWCNT membranes with nanopores and intrinsic superoleophilic-hydrophobic

property, and they applied this membrane for the separation of water from water-in-oil emulsions for the first time.<sup>627</sup> However, it was impossible for a pristine SWCNT membrane to achieve oil separation from oily water because of its wettability. Therefore, surface modification of the SWCNT membrane becomes imperative. The challenge in modifying an inert SWCNT membrane is to guarantee the uniform distribution of the modifier. In order to overcome this problem, a PDA layer with a thickness of several nanometers needs to be uniformly coated on every SWCNT in advance, which functions as an intermediate layer for the further grafting of PEI (Fig. 88).<sup>628</sup> The fabricated SWCNT/PDA/PEI composite membrane exhibited superhydrophilicity and underwater superoleophobic property with ultralow oil adhesion, and it could effectively separate nanosized oil from oil-in-water nanoemulsions. According to the classic Hagen–Poiseuille fluid dynamic theory, membrane flux is inversely proportional to the total distance of the liquid running through the membrane.<sup>629</sup> Hence, a thinner membrane corresponds to higher membrane flux. This nanomaterial-constructed composite membrane exhibited extremely high flux far beyond traditional polymeric and ceramic ultrafiltration membranes benefiting from its ultrathin thickness. It is believed that this new type of nanomaterial-constructed membrane is a promising alternative for traditional membranes in large-scale applications in the near future. A critical problem for the application of nanomaterial-constructed membranes is to achieve scalable and controllable membrane fabrication.

## 10.2 Polyethersulfone (PES) membranes for blood purification and water treatment

Over the past several decades, polymeric membranes have been widely used for separation and purification requirements, such as molecular separation, water purification, plasma collection, and blood purification. Among the polymeric membranes used

for blood purification and water treatment, PES membranes have shown good thermal and chemical resistance. However, some drawbacks associated with PES (such as its hydrophobic nature and adhesion of proteins and organisms) exist, which may yield undesirable results.

**Blood purification.** Undesired protein adsorption and platelet adhesion can lead to the formation of a thrombus and even cause the death of patients during blood purification. To overcome these problems, various approaches have been developed and they can be divided into bulk blending, surface grafting, surface coating, and their combinations. Amongst these methods, the bulk blending of functional amphiphilic polymers is the traditional way to modify PES membranes by the simple incorporation of functional additives into the casting solution. For instance, to combine various functions in PES membranes, 4,4'-diphenylmethane diisocyanate has been used as the hard segment to enhance the miscibility between PU and PES, and *N*-methyl-diethanolamine (containing a tertiary amino group) has been used as the soft segment.<sup>630</sup> After blending tertiary amino-terminated PU with PES, the composite membrane exhibits outstanding adsorption capability of toxic bilirubin (over 20 mg g<sup>-1</sup>). By simply functionalizing the tertiary amino group, the membrane bearing zwitterionic PU exhibited excellent blood compatibility; with quaternary ammonium salts, it exhibited antibacterial property. After incorporating Ag NPs into the carboxybetaine-functional composite membrane, dual functions of antibacterial and antifouling properties could be achieved. Different from adding macromolecules with -OH, -NH<sub>2</sub>, or -COOH, various functionalities could be obtained by easily converting the tertiary amino group into zwitterions or quaternary ammonium salt, and the developed PUs have promising potential as readily available, multifunctional, and easy-to-use materials for biological applications.

Zhao *et al.* also developed a new method to modify PES membranes by “*in situ* crosslinking polymerization.” That is,

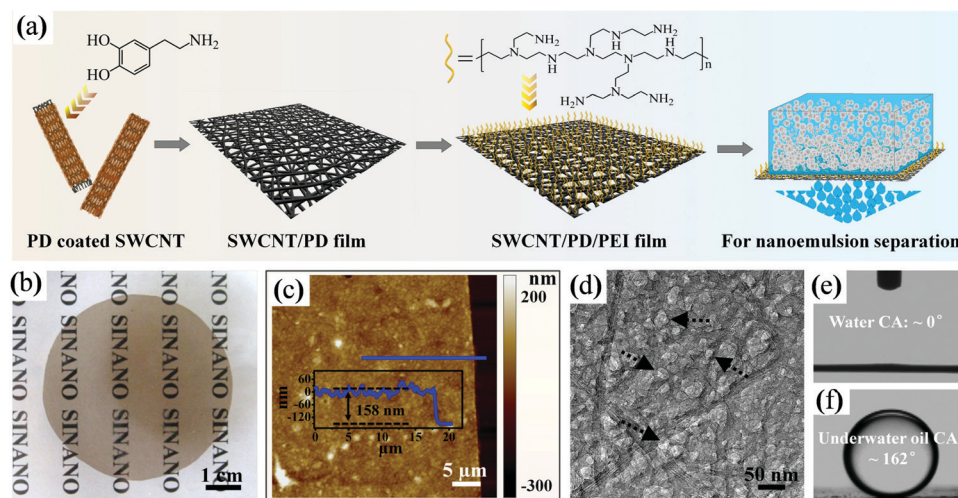


Fig. 88 SWCNT/PD/PEI composite membrane. (a) Schematic showing the fabrication of a membrane and the separation of oil-in-water nanoemulsion. (b) Digital photograph of a free-standing membrane. (c) AFM image and height profile of the membrane. (d) TEM image of the membrane. Photographs of (e) a water droplet on the membrane in air and (f) an oil droplet on the membrane underwater. Reproduced from ref. 628 with permission. Copyright (2015) Royal Society of Chemistry.

functional monomers are introduced into a PES casting solution followed by a thermally induced crosslinking reaction, and the obtained solution is then directly converted into the membrane by a phase-inversion technique. For instance, PES was selected as the membrane matrix, and sodium acrylate (AANa) and sodium p-styrene sulfonate (SSNa) were selected as the functional monomers.<sup>631</sup> After the *in situ* crosslinking polymerization of AANa and/or SSNa, a series of functional membranes were obtained, as shown in Fig. 89a. These functional membranes showed lower protein adsorption, suppressed platelet adhesion by 93%, and prolonged clotting times by 60 s for activated partial thromboplastin time (APTT) when compared with those obtainable from pristine PES membranes. Moreover, the membranes showed good antifouling property. These results confirmed that the highly efficient and convenient *in situ* crosslinking polymerization endowed PES membranes with excellent biocompatibility, thereby having great potential for use in blood purification.

By a combination of *in situ* crosslinking polymerization and post-functionalization, Zhao *et al.* could also integrate thin hydrogel films on the surface of PES membranes. For instance, hydroxyl groups were introduced onto the PES membrane by the use of *in situ* crosslinking polymerization of HEMA; then, double bonds were also introduced *via* the reaction between hydroxyl groups and acryloyl chloride. Sulfobetaine methacrylate (SBMA) and AANa were selected as the functional monomers to fabricate thin hydrogel layers onto the surfaces of PES membranes by UV-light-initiated polymerization.<sup>632</sup> Ag ions were absorbed into the hydrogel layers by the carboxylate groups from AANa, as shown in Fig. 89b. The excellent hemocompatibility of the modified membranes was confirmed by undetectable plasma protein adsorption, suppressed platelet

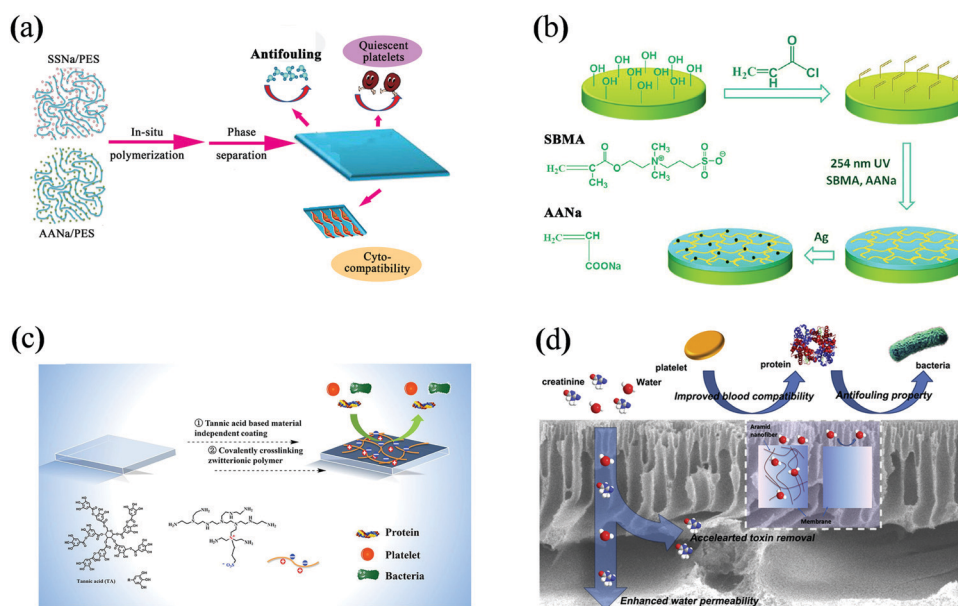
adhesion, prolonged clotting times, low hemolysis ratio, and suppressed blood-related complement activation. In addition, the strong antibacterial ability could last for over 5 weeks.

The *in situ* crosslinking reaction could also be performed between different polymers. Zhao *et al.* straightforwardly prepared a thin layer within a few minutes *via* the coordination reaction between tannic acid (TA) and Fe<sup>III</sup>. Thereafter, a reactive zwitterionic polymer (PEI-S) was post-crosslinked with the matrix layer *via* a dip-coating process at room temperature, as shown in Fig. 89c. Combining TA with an active zwitterionic polymer was an effective approach for constructing a universal, stable, and low-cost antifouling material surface.<sup>633</sup>

In spite of the addition of organic polymers, organic nano-materials could also enhance the filtration and biological performances of polymeric membranes. For instance, Zhao *et al.* reported the usage of CNTs and aramid nanofibers (ANFs) to enhance the ultrafiltration and biological performances of PES membranes.<sup>634</sup> The flux recovery ratio, protein adsorption, and bacterial adhesion tests validated that the modified membranes exhibited improved antifouling, blood compatibility, and efficient adsorption of small molecular toxins during dialysis applications of the composite membranes (Fig. 89d).

**Water treatment.** As mentioned above, PES has some drawbacks, such as its hydrophobic nature and adhesion of proteins and organisms. For water treatment, the modification of PES necessitates finding a compromise between hydrophobicity and hydrophilicity and to localize a positive effect on the flux and fouling resistance but without sacrificing mechanical stability. Recently, Zhao *et al.* fabricated a series of ion-exchange and smart membranes.

By simply blending stimuli-responsive materials into PES, a membrane with redox-responsive hydraulic permeability was



**Fig. 89** (a) *In situ* crosslinking polymerization of monomers; (b) *in situ* crosslinking polymerization combined with post-crosslinking of monomers; (c) surface coating followed by *in situ* crosslinking of polymers; (d) direct blending of nanofibers and nanotubes. Adapted from ref. 631–634 with permission. Copyright (2016, 2017, 2018, and 2017) Elsevier Ltd and American Chemical Society.

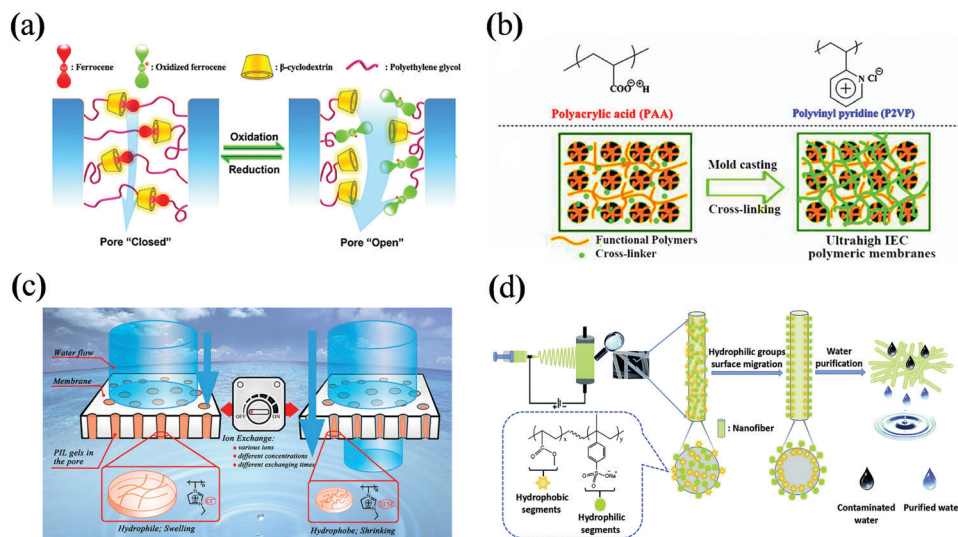


Fig. 90 (a) Direct blending of redox-responsive materials; (b) *in situ* crosslinking copolymerization of ion-exchange monomers; (c) *in situ* crosslinking copolymerization of salt-responsive monomers; (d) direct blending of amphiphilic polymers coupled with electrospinning technology. Adapted from ref. 635, 636, 638, and 639 with permission. Copyright (2015, 2016, 2017, and 2018) Elsevier Ltd, American Chemical Society, and Royal Society of Chemistry.

fabricated for the first time by introducing a reversible redox-responsive system based on the host-guest complex between ferrocene and  $\beta$ -cyclodextrin.<sup>635</sup> As a result, the water flux of the membrane with 18.3 wt% redox-responsive materials was  $270 \text{ mL m}^{-2} \text{ mmHg}^{-1} \text{ h}^{-1}$  in the original state, which increased up to  $1022 \text{ mL m}^{-2} \text{ mmHg}^{-1} \text{ h}^{-1}$  in the oxidation state and decreased to  $538 \text{ mL m}^{-2} \text{ mmHg}^{-1} \text{ h}^{-1}$  in the reduction state. Changes in the pore size and porosity of the modified membrane has outstanding potential for application in chemical detection and biological treatments, as shown in Fig. 90a.

*In situ* crosslinking copolymerization method is also a good choice to fabricate functional PES membranes for water treatment. To develop PES membranes with high ion-exchange capacities (IECs) but low swelling degree, the universal mold casting approach was performed by *in situ* crosslinking negatively/positively charged monomers in the presence of PES, as shown in Fig. 90b. As a result, crosslinking monomers provided high IECs along with restricting the swelling of these hydrophilic monomers with high crosslinking density and hydrophobic polymers. For instance, Zhao *et al.* chose AA for fabricating cationic-exchange membranes (CEM) and 2-vinylpyridine (2VP) for fabricating anionic-exchange membranes (AEM).<sup>636</sup> The new method yielded ultrahigh IECs of  $7.88 \text{ mequiv g}^{-1}$  for CEM and  $6.27 \text{ mequiv g}^{-1}$  for AEM, which are 8.8 and 7.0 times higher than that of Nafion 117 ( $0.89 \text{ mequiv g}^{-1}$ ), respectively.

Salt-responsive PES membranes were prepared by the use of *in situ* crosslinking copolymerization method with methacrylateoethyl trimethyl ammonium chloride ([MTA][Cl]) as the functional monomer.<sup>637</sup> The anion strength response of the membranes was positive, and the responsive coefficient of the membranes could be 26 times in response to an aqueous solution of NaCl (as shown in Fig. 90c). The cation species

and concentrations had a minimal influence on the responsive behavior. Meanwhile, the membranes responded to both anion species ( $\text{PF}_6^-$ ,  $\text{BF}_4^-$ , and  $\text{SCN}^-$ ) and their strengths. After slightly modifying the *in situ* crosslinking copolymerization method for a liquid system, novel anion-responsive smart PES membranes could be fabricated by filling PES microporous membranes (solid system) with poly(ionic liquid)s (PILs) gels.<sup>638</sup> The wetting properties of these PILs could be controlled by changing their counteranions (CAs), and therefore, the filled PIL gel gates in the membrane pores could spontaneously switch from the “closed” state to the “open” state by recognizing the hydrophilic CAs in the environment and *vice versa*. As a result, the flux values of the smart membranes could be adjusted from a very low level ( $0 \text{ mL m}^{-2} \text{ mmHg}^{-1}$  for  $\text{Cl}^-$ ,  $\text{Br}^-$ , and  $\text{BF}_4^-$ ) to a relatively high one ( $430 \text{ mL m}^{-2} \text{ mmHg}^{-1}$  for TFSI). The anion-responsive gating behavior of PILs-filled membranes is fast, reversible, and reproducible. The proposed smart membranes are highly attractive for ion-recognizable chemical/biomedical purifications.

To rapidly remove toxins from wastewater, the simple blending of amphiphilic polymers coupled with the electrospinning method was developed. For instance, a novel ultrafast adsorption nanofibrous membrane (UFAM) for organic dyes was prepared *via* simple functional group surface migration.<sup>639</sup> An amphiphilic copolymer of MMA and SSNa (mass ratio of 3 : 2) was synthesized by free radical polymerization and further blended into PES followed by electrospinning. Only by soaking in water and lyophilization, the hydrophilic and negatively charged groups migrated on the PES nanofibrous membrane surface, and the obtained UFAM was found to be exceptionally efficient in the removal of methylene blue (MB) from an aqueous solution, as shown in Fig. 90d. In particular, an ultrafast adsorption rate for MB with an adsorption equilibrium time of less than 5 min was reported.

In addition, the UFAM showed excellent performance with respect to recyclability, dynamic filtration–purification, and selective adsorption of cationic dyes from mixed dye solutions.

As a short summary, for either blood purification or water treatment, the *in situ* crosslinking copolymerization method can avoid multistep purification, while the crosslinking degree should be controlled at a certain level. Otherwise, homogenous gels are formed in the case of a high crosslinking degree of monomers. In contrast to *in situ* crosslinking polymerization, the direct blending of amphiphilic polymers or small hydrophobic molecules may necessitate more steps for synthesis and/or purification, while the surface/interface migration of amphiphilic polymers may induce some dramatical merits, such as the fast adsorption of toxins. Different from *in situ* crosslinking polymerization and direct blending, the post-crosslinking of thin hydrogel films (even by several nanometers) could result in significant surface modification. However, precise control over the thickness of thin hydrogel films is desirable, because a thick hydrogel film can result in sacrificing the permeability of PES membranes. Notably, the abovementioned strategies for modifying PES membranes are also applicable to improve the performance of other polymeric matrixes.

### 10.3 Nanofiltration membranes (NFMs) for seawater desalination

Nanofiltration is a widely used pressure-driven membrane forming technique with low cost and high efficiency; it is used in separating organic dyes and multivalent ions, and it offers a promising solution to alleviate the formidable issue of freshwater scarcity.<sup>640</sup> Diverse manipulation strategies have been developed to meet various expectations of NFMs, such as interfacial polymerization, mussel-inspired codeposition, and LbL assembly, for constructing selective layers on the top surfaces of porous substrates. Although these NFMs have made tremendous progress in the past few decades, they still suffer from a tradeoff between the water permeation flux and salt rejection. Therefore, efficient strategies are required to break the tradeoff effect and to simultaneously achieve high water permeation flux and high salt rejection. In recent years, a series of appealing approaches have been developed to improve the performances of NFMs, such as substrate optimization, sacrificial-layer application, interlayer introduction, and nano-materials incorporation.

Livingston *et al.* reported the formation of sub-10 nm polyamide nanofilms by controlling interfacial polymerization on a sacrificial layer of cadmium hydroxide nanostrands.<sup>641</sup> However, these ultrathin nanofilms were only suggested as separating layers in NFMs that were used in processing organic solvents. Metal–organic framework NPs (ZIF-8) were also used as a sacrificial layer by Jin *et al.* to facilitate the formation of a rough polyamide selective layer with crumpled nanostructures.<sup>642</sup> These crumpled nanostructures were believed to increase the effective area of the selective layer for water permeation. The NFMs thus obtained exhibited high water permeability (up to  $53.5 \text{ L m}^{-2} \text{ h}^{-1} \text{ bar}^{-1}$ ) with salt rejection of above 95% for  $\text{Na}_2\text{SO}_4$ . Nevertheless, it should be noted that

heavy metal compounds such as cadmium hydroxide and ZIF-8 were used in the fabrication processes of NFMs, which are harmful to human health if their ions are left in the produced water. Therefore, Livingston *et al.* tried to fabricate ultrathin selective layers at the free aqueous–organic interface.<sup>643</sup> Smooth sub-8 nm polyamide nanofilms were synthesized to provide fast water transport through the nanofilm composite membranes. Manipulating the intrinsic nanofilm thickness from  $\sim 15$  down to 8 nm revealed that water permeance proportionally increased with a decrease in the thickness, after which it increased nonlinearly to  $2.7 \text{ L m}^{-2} \text{ h}^{-1} \text{ bar}^{-1}$  as the thickness was further reduced to  $\sim 6$  nm. However, this method is limited for practical applications because of the difficulty in fabricating ultrathin films in a large area at the free aqueous–organic interface. On the other hand, Tan *et al.* modulated the interfacial polymerization by “controlling” monomer diffusion with water-soluble polymers to facilitate the formation of crumpled nanostructures in the polyamide selective layer.<sup>644</sup> The thickness of the crumpled polyamide films (the authors called them nanoscale Turing structures) was about 20 nm or less, which was much thinner than those fabricated by traditional methods. Therefore, the as-prepared NFMs showed both high water permeability and high salt rejection, which could break the tradeoff effect.

Apart from polyamide ultrathin films obtained by interfacial polymerization, Xu *et al.* developed mussel-inspired chemistry to fabricate selective layers of NFMs. For example, as schematically shown in Fig. 91,  $\text{CuSO}_4/\text{H}_2\text{O}_2$  was found to trigger the rapid deposition of PDA-based ultrathin coatings with high uniformity and enhanced stability.<sup>645–648</sup> These mussel-inspired coatings exhibit two kinds of nanopores (around 0.6 and 1.0 nm) for versatile molecular separation. The corresponding NFMs exhibited promising potential for water desalination as well as organic solvent separation.<sup>649</sup> Polyamines such as PEI were used to promote the formation of mussel-inspired ultrathin selective layers *via* Michael addition and Schiff base reactions between a series of polyphenols and polyamines.<sup>650–655</sup> At the same time, the surface potential and surface hydrophilicity were remarkably modulated because the used PEI was a type of positively charged polyelectrolyte.<sup>650,651</sup> Therefore, the as-prepared NFMs showed different salt rejection properties *via* the Donnan effect as compared to those exhibited by negatively charged polyamide-based

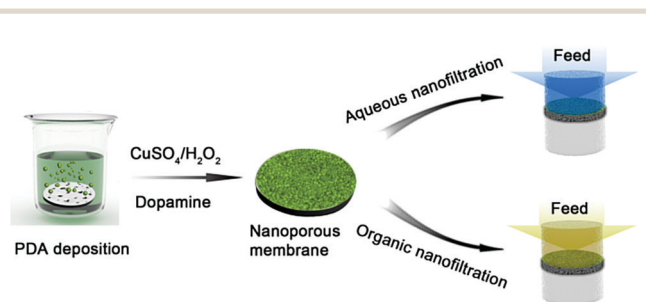


Fig. 91  $\text{CuSO}_4/\text{H}_2\text{O}_2$  triggered the rapid formation of PDA-based ultrathin selective layers with high uniformity and enhanced stability for both aqueous nanofiltration and organic nanofiltration. Reprinted from ref. 649 with permission. Copyright (2017) American Chemical Society.

membranes. Their surface properties could be further tuned by the addition of inorganic NPs or the biomineralization of inorganic nanofilms and nanorods.<sup>656–659</sup> The resulting organic–inorganic composite NFMs could be applied for long-term salt rejection or dye removal from water due to the enhanced structural stability or the self-cleaning property endowed *via* photocatalysis. It should be pointed out that the mussel-inspired selective layers can be formed on the porous substrates *via* direct deposition<sup>649–659</sup> or through the contradiffusion approach.<sup>660–662</sup> In the latter case, the fabricated mussel-inspired films usually showed lower thickness with narrow pore size distribution for achieving precise separation.

On the other hand, nanostructured interlayers were suggested by Xu *et al.* to optimize the substrate surfaces for modulating interfacial polymerization. Nanomaterials, such as CNTs, cellulose nanocrystals, and graphene oxide nanosheets or mussel-inspired coatings, were used to fabricate these interlayers.<sup>663–667</sup> Taking cellulose nanocrystals as an example, the nanomaterials could be vacuum-filtrated on the porous substrate to form a uniform and hydrophilic thin coating. This coating adsorbed and distributed the aqueous solution of diamine on the substrate surface for interfacial polymerization. Therefore, the polymerization process could be controlled to a certain extent for facilitating the formation of ultrathin polyamide films with homogeneous characteristics. The thickness of the selective layer was normally lower than 50 nm and the corresponding NFMs exhibited overall desalination performance superior to that of state-of-the-art materials reported so far (Fig. 92). This work proposes a simple

way to fabricate advanced NFMs with outstanding performance for practical applications.

Recent progresses made in the fabrication of polymer membranes for use in various applications, such as oil/water separation, blood purification, water treatment, and seawater desalination, are introduced. Moreover, design strategies and preparation methods of polymer membranes for obtaining good performances, such as good antifouling, high efficiency, ultrahigh flux, high mechanical strength, low protein adsorption, platelet adhesion abilities, good blood compatibility, high IECs, high salt rejection, *etc.*, in the above applications are also presented. Although great advances in polymer membrane technology has been achieved, and a few of them have been industrialized, they still have their respective problems that need to be resolved. As an example, fabricating polymer membranes capable of good antifouling and chemical stability while maintaining high permeation flux and very high oil/salt rejection is still a challenge for researchers and engineers.

## 11. Energy storage polymers

The two worst crises for humans in the 21st century are energy shortage and environmental pollution. This is because fossil fuels—the dominant energy supplier—will eventually be depleted in the near future and they are also the cause of environmental pollution. Therefore, it is highly desirable to develop green and renewable sources of energy to replace fossil fuels as the main energy source. Energy storage is one of these solutions. Hence, dramatic progress in the field of energy storage materials and devices has been witnessed during the last decade. In the quest for high energy and power densities, fast electrode kinetics, long life cycle, and stable energy storage systems, various kinds of materials and different device configurations have been explored. The two main energy storage systems are supercapacitors and batteries. Although energy storage in both these systems depends on electrochemical processes, different mechanisms lead to different energy and power densities in such systems. Supercapacitors have been previously considered to have low energy density and high power density as well as long cycle life as compared to those obtainable from batteries, while the recent progresses made in both these fields have narrowed down this gap.<sup>668,669</sup> Even though much progress has been made in the field of electrode materials, including various forms of carbons, nanostructured materials, transition metals, and metal oxides, the energy density of supercapacitors and the power density of batteries are still limited. The charge storage mechanism in such energy storage systems is mainly responsible for these limitations.<sup>670</sup> This problem can be overcome either by inventing a new device with a different and effective charge storage mechanism or by selectively designing materials with superior properties. However, most energy storage devices, such as LIBs and supercapacitors, have some shortcomings, such as being heavy, rigid, and bulky, which cannot meet the demands of flexible and portable devices. In recent years, stretchable/portable devices

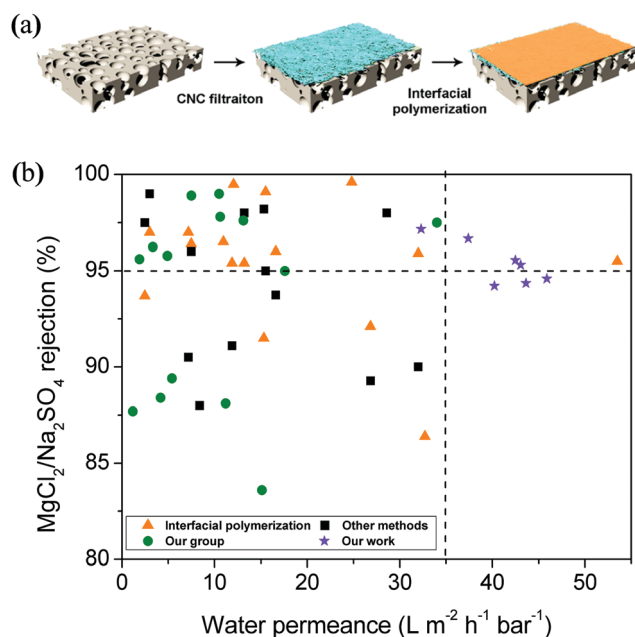
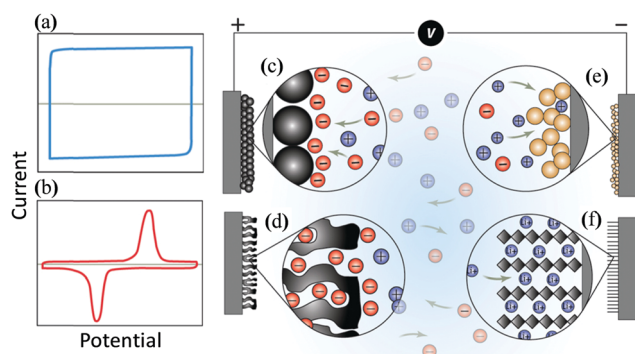


Fig. 92 (a) Schematic illustration of the formation of a nanostructured interlayer from cellulose nanocrystals (CNC) and ultrathin polyamide selective layer by interfacial polymerization; (b) comparison of typical performances of the as-fabricated NFMs by mussel-inspired chemistry (●) and interlayer strategy (★) with those reported in the literature. Reprinted from ref. 664 with permission. Copyright (2017) Royal Society of Chemistry.

have attracted considerable attention, such as wearable devices, artificial skin sensors, and bendable mobile phones.<sup>671,672</sup> Organic molecules are considered to be promising materials for flexible energy storage devices due to their flexibility, diversity, relative abundance, low cost, *etc.* However, they still face challenges with regard to stability and low cycle life due to solubility in liquid electrolytes. This problem can be considerably resolved by the use of small organic molecules to build functional polymers. Therefore, we present the advances in energy storage devices in this section, such as LIBs, supercapacitors, and fiber-shaped energy devices made of polymers.

### 11.1 Supercapacitors

Supercapacitors offer several advantages over batteries, such as high power density and long cycle life. It is always advantageous to have a device that can be charged at an ultrafast rate to its full capacity with a longer lifetime. In contrast to the ion diffusion mechanism in batteries that can reduce the overall kinetics, the ion adsorption mechanism on the surface of electrodes in supercapacitors can facilitate the fabrication of devices that can be charged within minutes.<sup>673</sup> Supercapacitors are also expected to bridge the gap between batteries and capacitors. A typical example of the cyclic voltammetry (CV) curves of a supercapacitor and battery is shown in Fig. 93a and b, respectively. Based on the charge storage mechanisms, supercapacitors can be divided into two categories: electrostatic double-layer supercapacitors (EDLCs) and pseudocapacitors. Conventional materials with a high surface area have been explored in the field of capacitors, but interest regarding them has been rising in pseudocapacitors. Several new materials, such as polymers, are emerging as potential candidates for next-generation supercapacitors. Redox-active polymers and conducting polymers capable of pseudocapacitive energy storage are directly used as electrode materials. Porous polymers and heteroatom polymers are used to produce porous carbon and heteroatom-doped carbon, respectively. In this subsection, the applications of polymers in both types of supercapacitors are briefly discussed.



**Fig. 93** Typical CV curves of a (a) supercapacitor and (b) battery, and the charge storage mechanism for different materials: (c) surface charge accumulation in carbon and (d) porous carbon; pseudocapacitive behavior for (e) redox reaction and (f) ion intercalation. Reprinted from ref. 669 with permission. Copyright (2014) from AAAS.

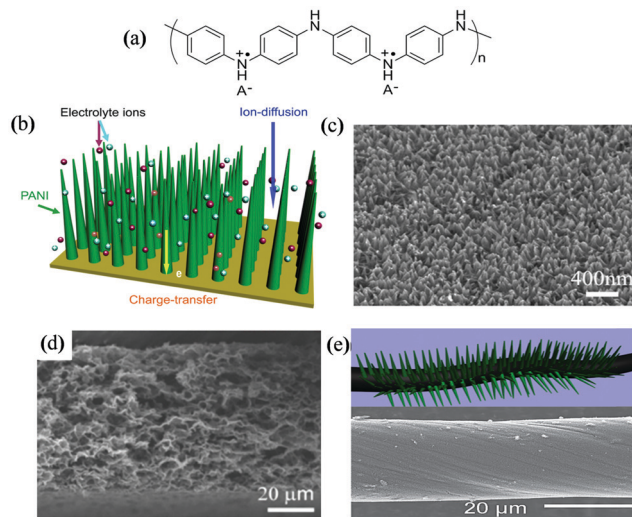
EDLCs primarily work through the interfacial accumulation of charges at the electrode–electrolyte surface.<sup>674</sup> When a charged object is placed in a liquid, the countercharge from the liquid starts to accumulate near the surface, giving rise to an electric double layer.<sup>675</sup> Therefore, surface-dependent charge storage mechanism in EDLCs needs materials with high specific surface, large porosity, and uniform pore distribution.<sup>676</sup> Different types of carbons are the material of choice in this regard (Fig. 93c–f). A comprehensive report in this regard is out of the scope of this review and can be found elsewhere.<sup>677,678</sup> For the sake of general understanding, we would like to introduce a few examples where polymers have been employed either to get heteroatom-doped porous carbon or composites with carbon for superior properties.

Heteroatom-doped carbon-based materials show superior properties in energy storage applications, particularly supercapacitors. There are various strategies to obtain heteroatom-doped carbon-based materials. Nitrogen- and oxygen-doped hierarchical porous carbon structures were fabricated *via* the carbonization of a quinone amine polymer precursor on the surface of MgO as a template.<sup>679</sup> A high content of heteroatoms along with a large surface area were obtained, which ensures high pseudocapacitance as well as pathways for faster ion diffusion, leading to excellent charge storage properties. In another report, a template-free three-step carbonization process was developed to achieve nitrogen-doped 3D porous carbon-based materials with interconnected pores.<sup>680</sup> The nitrogen content was found to be 9.71%, and the capacitor based on two-electrode systems delivered capacity of 221 F g<sup>-1</sup> and showed excellent cycle life.

Although the stability of EDLCs is very high, they suffer from low gravimetric and volumetric energy densities.<sup>670</sup> Research is being conducted for resolving this issue. One of the emerging trends is to investigate materials with pseudocapacitance.

Based on the mechanism of faradaic redox reaction, pseudo-supercapacitors usually offer higher theoretical capacity than EDLCs. Reversible redox reactions, similar to that in batteries, take place when potential is applied across the electrodes. Redox-active conducting polymers are one of the most promising candidates because of their low cost, structural diversity, high capacitance, and reasonable electronic conductivity. Various conducting polymers, such as PANI, PPy, PEDOT, and their derivatives, have been extensively studied.<sup>676,681,682</sup> A few examples in this field will be summarized in the subsequent subsection.

Wei *et al.* synthesized large arrays of vertically aligned PANI through a one-step template-free approach on various conducting substrates (Fig. 94a–c).<sup>683</sup> The vertical growth of nanowires on a conducting substrate with a narrow diameter was beneficial to the ion diffusion process. The highest specific capacitance of 950 F g<sup>-1</sup> was obtained, and specific capacitance of 780 F g<sup>-1</sup> was maintained at a high current density of 40 A g<sup>-1</sup>. The capacitance behaviors in the different electrolytes, such as 1 M HClO<sub>4</sub>, 1 M lithium bis(trifluoromethanesulfonyl), and nonsolvent electrolyte ionic liquid (IL), were also investigated. This study revealed that the oriented nanostructures could dramatically improve the electrochemical properties. Subsequently, the same group



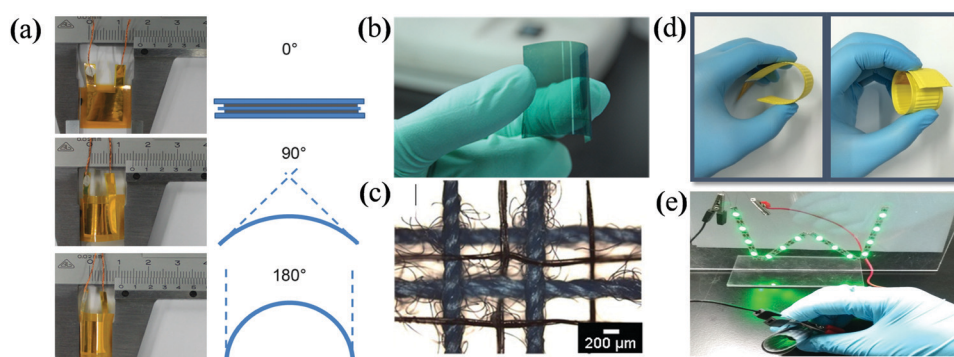
**Fig. 94** Application of conducting polymers in supercapacitors: (a) structure of PANI, (b) schematic mechanism of faster ion diffusion in nanowire arrays of PANI, (c) morphology of PANI nanowire array, (d) interconnected porous network of 3D graphene-based flexible film, and (e) schematic of the growth of PANI nanowires on CNT yarn and SEM image of actual yarn. Reproduced from ref. 683, 685 and 686 with permission. Copyright (2010 and 2013) American Chemical Society and Wiley-VCH.

investigated the synergistic effect of PANI and graphene oxide by growing vertically aligned PANI nanowire array on the surface of graphene oxide sheets.<sup>684</sup> Electrochemical capacitance and stability of the obtained nanocomposite were higher than those of both the individual components. Different dimensional materials can be composited to obtain functional nanomaterials with superior properties.

Wei *et al.* synthesized a free-standing porous 3D reduced graphene oxide (3D-RGO) film using a template method for application in flexible supercapacitors.<sup>685</sup> High charge-discharge rate performance was observed due to the presence of an interconnected porous structure, ensuring the diffusion of electrolyte ions into the inner surface electrodes (Fig. 94d). In addition, the growth of PANI nanowires on 3D-RGO resulted in

excellent charge storage properties. Supercapacitors based on the 3D-RGO PANI nanocomposite could retain 90% of its initial capacity after 5000 charge–discharge cycles. In another study, PANI nanowires were grown on CNT yarn through an *in situ* process to assemble a wire-like supercapacitor (Fig. 94e).<sup>686</sup> CNT yarn acted as both active material and current collector because of its worthwhile conductive properties. A thread-like supercapacitor exhibited excellent energy storage properties both in terms of rate performance and cycling stability. Capacity was retained under different bending states and angles, showing remarkable flexible properties. The free-standing flexible film composed of 3D-RGO PANI was also employed to assemble a flexible supercapacitor, which exhibited remarkable energy storage properties even under applied stress (Fig. 95a). The wire or fiber-shaped devices can be further woven into textiles to fabricate smart clothes (Fig. 95b). In a rather interesting study, a multifunctional flexible energy storage smart (ESS) window was fabricated by combining PANI nanowires with a transparent substrate (Fig. 95c).<sup>687</sup> Energy storage and electrochromic functions were, therefore, integrated into a single device, leading to its potential application as a smart window. The device showed reasonable charge storage properties, which could be further improved by adjusting the structure and designing new materials. This ESS window can be further integrated with solar cells to combine energy production together with the storage function. Schon *et al.* joined fullerene monomers by a cyclobutene ring to synthesize a branched chain fullerene polymer.<sup>688</sup> Based on the negative-charge-accepting pseudocapacitive mechanism, an n-type supercapacitor was fabricated, which exhibits good capacity and cycling stability.

These results suggest that polymers are potential candidates for next-generation commercial supercapacitors (either EDLCs or pseudocapacitors). Although significant progress has been made in the recent decade, there are still several challenges that need to be addressed. For example, energy density is still not sufficient. This may be achieved by designing new materials with high capacity or developing new electrolytes and processes. For example, lithium-ion-based supercapacitors can operate at higher voltages, leading to higher energy densities.<sup>668,670</sup>



**Fig. 95** Different flexible energy storage devices: (a) flexible supercapacitor and schematic of bending at different angles, (b) photograph of a single flexible and transparent electrode, (c) yarn supercapacitor integrated into conventional cotton yarn, (d) wave-like flexible battery bent into different shapes, and (e) illumination of LEDs by a bent flexible battery. Reproduced from ref. 685, 686, 687 and 689 with permission. Copyright (2012, 2013, and 2018) from Royal Society of Chemistry and Wiley-VCH.

## 11.2 LIBs

Batteries are another major area where polymers are being explored either as active material for energy storage or as additive for different purposes (Fig. 95d and e). In this subsection, we will summarize the recent examples elaborating the applications of functional polymers as active electrode materials. Electroactive small molecules are usually polymerized to improve stability in liquid electrolytes. For example, a novel hyper-crosslinked polypillar[5]quinone (poly-P5Q) was used as the electrode material in LIBs.<sup>690</sup> It offers very low practical capacity, but it provides new insights into the reduction of solubility of electroactive small molecules by crosslinking them. Solubility in the electrolyte is one of the major problems for organic materials. The stability of poly-P5Q in the liquid electrolyte was improved by this method. Two isomeric pairs of poly(anthraquinonyl imide)s were prepared, and their electrochemical properties as LIB cathodes were studied.<sup>691</sup> It was shown that isomerism played an important role in electrochemical performance. An all-solid-state Li-organic battery with quinone-based polymer cathode and composite polymer electrolyte was reported.<sup>692</sup> The ionic conductivity of the solid electrolyte was excellent when the temperature was 343 K. Although this performance was good, there is still a need to develop effective systems capable of the best performance at room temperature. An anode material for LIBs derived from aromatic imides with high capacity was reported due to its low potential.<sup>693</sup> Based on the density functional theory calculation and natural bond orbital charge analysis for electron transfer processes and reaction free energies, they proposed a 12-electron transfer process to accommodate the high experimental capacity of aromatic amides. A quinone-based polymer accommodating 4 lithium ions was reported with high charge storage capacity.<sup>694</sup> Lithiated poly(dihydroxyanthraquinonyl sulfide) (LiDHAQS) was selected as the material of choice because of its high degree of carbonyl groups. The experimental capacity of 3.5 Li<sup>+</sup> per monomer was obtained, which is less than the theoretical value but higher than all the other quinone-based electrodes reported so far.

An *in situ* polymerization method was adopted to synthesize 3,4,9,10-perylenetetracarboxylic dianhydride (PTCDA)-based polyimide/carbon nanotube (polyimide/CNT) nanocomposite electrode, which could be used in LIBs.<sup>695</sup> Polymerization on the surface of CNTs was found to increase the capacity as well as cycling stability. Here, 93% of the initial capacity was found to be retained when the polyimide/CNT nanocomposite electrode was used after 300 cycles, while only 74% of the initial capacity was retained for PTCDA/CNTs. Pyromellitic dianhydride (PMDA) was polymerized with ethylene diamine in the presence of a 3D-RGO film to fabricate a flexible electrode for LIBs.<sup>250</sup> The composition of the electrode was found to contain 80% of the weight of the active materials and 20% of 3D-RGO without any other additive or binder. 3D-RGO/polyimide flexible cathode exhibited initial capacity of 175 mA h g<sup>-1</sup> as compared to 120 mA h g<sup>-1</sup> when polyimide was bound with conductive carbon in the presence of a binder and a conventional electrode structure. Similarly, significant enhancement in the cycling performance

was also reported. Through electrochemical impedance spectroscopy (EIS) studies, it was concluded that an effective 3D-RGO network could lead to excellent conductivity, which would eventually yield better rate and cycling performances. Polyimide suffers from poor conductivity, leading to poor performance and low material utilization. To overcome this issue, PMDA was *in situ* polymerized in the presence of CNTs to obtain composite materials with much better electronic properties.<sup>696</sup> The vacuum filtration technique was used to fabricate a binder and metal-current-collector-free electrode for LIBs. In this case, the inter-connective network of CNTs acted as the current collector, which is much lighter than the commercially available aluminum current collector, thereby effectively reducing the dead mass. Significant enhancement in electrochemical properties was reported in this case, where the electrode was still able to deliver high capacity of 120 mA h g<sup>-1</sup> even at a very high rate of 20 C. The influence of monomer concentration on morphology was also studied in this research: it was observed that large aggregates start to grow instead of flakes on the CNT surface at a certain amount of monomer concentration. Therefore, the optimum monomer concentration is also important to obtain good results. Subsequently, the same group reported another interesting strategy to design large-area flexible cathodes for LIBs.<sup>697</sup> A simple rolling method was used to fabricate the electrode film, which was mounted on a free-standing CNTs film by applying pressure. The CNT film acted as a mechanical support and current collector in this system (Fig. 96a). The polyimide obtained from the polymerization of PMDA and triethyl amine was used as the active material. The large-area flexible cathode showed superior electrochemical properties over a conventional electrode and retained 80% of its initial capacity after bending for 1000 times.

In another example, PMDA and anthraquinone were polymerized together to obtain high-capacity polyimides.<sup>698</sup> A simple vacuum filtration method together with CNTs was employed to obtain a flexible, free-standing, binder-free electrode for LIBs (Fig. 96b). High capacity of 190 mA h g<sup>-1</sup> at 0.1 C was obtained due to the three-electron transfer system. Capacity of 120 mA h g<sup>-1</sup> (~63% of that at 0.1 C) was maintained at a high current density of 20 C. The cathode retained more than 95% of its initial capacity after 300 charge-discharge cycles. Excellent rate performance was assumed due to the inter-connective network of CNTs in the cathode architecture. EIS results also showed that the charge transfer resistance decreased significantly with an increase in the concentration of CNTs, which has an influence on the electrochemical performance. The composite cathode showed optimal performance when the CNT content was 30%. A flexible cathode film exhibited good mechanical properties under stress. The tensile strength of the composite was comparable to that of CNTs. A sulfur-linked quinone-based polymer was reported for long-cycle-life LIBs.<sup>689</sup> A free-standing electrode was assembled together with CNTs acting as a binder and current collector. When applied as the cathode, it was able to retain about 89% of its initial capacity after 500 charge-discharge cycles. Excellent rate performance was also observed. Furthermore, a flexible

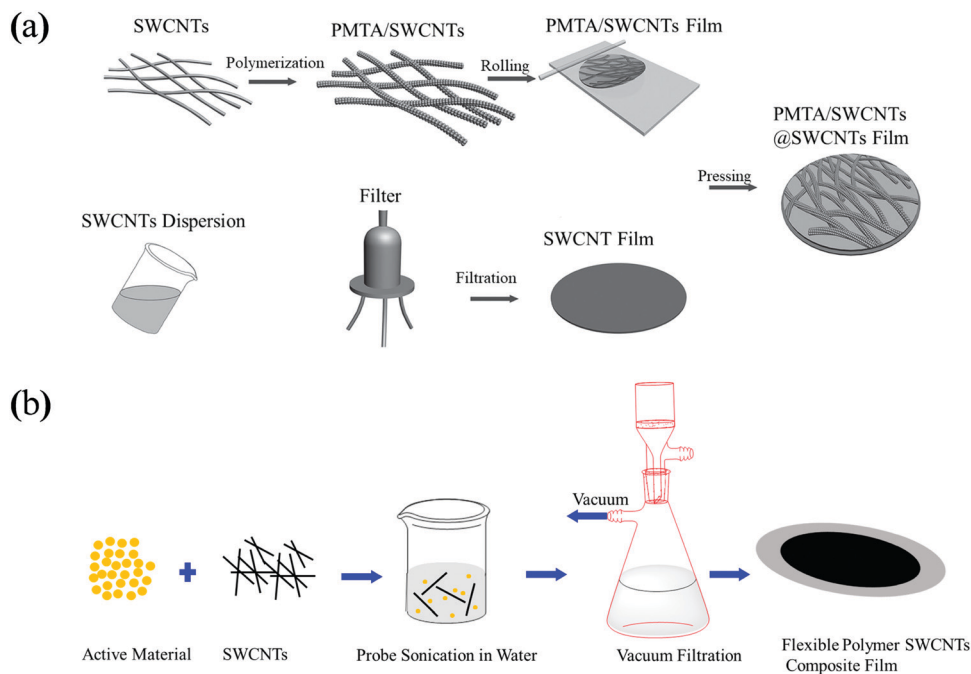


Fig. 96 Schematic of fabricating a flexible electrode: (a) embedding a conventional cathode film into a CNT film acting as a flexible current collector and (b) fabrication of a free-standing film. Reprinted from ref. 689 and 697 with permission. Copyright (2015 and 2018) Wiley-VCH.

battery was assembled with a lithium metal anode, which could retain almost 88% of its initial capacity after 2000 bending and unbending cycles (Fig. 95d and e). In short, the CNTs played a crucial role in supporting the free-standing film due to their excellent mechanical properties and enhancing the electronic conductivity of the free-standing film.

A PMDA-based polyimide was synthesized *via in situ* polymerization on the surface of few-layer exfoliated graphene (FLEG) sheets as the cathode for LIBs.<sup>699</sup> The utilization of the active material was improved by employing this composite material in the cathode, and polyimide-FLEG cathode delivered initial capacity of  $177 \text{ mA h g}^{-1}$  at 0.1 C. The cathode was able to retain almost 80% of its initial capacity after 200 charge-discharge cycles. The charge transfer resistance was significantly reduced by employing this cathode with a composite structure.

So far, polymer composites with CNTs or graphene are discussed. However, these two types of materials are expensive for commercial applications. A recent work reported an *in situ* polymerization strategy for synthesizing a polyimide composite with carbon black (CB).<sup>700</sup> In this case, insulating polyimide and conductive additive can be tightly contacted, and the crystallization of polyimides can also be inhibited. By optimizing the synthesis conditions, a composite that could deliver extraordinary capacity at high current densities could be obtained, which yielded excellent rate performance. In comparison to the *in situ* polymerized composite, mechanically mixed composite electrodes exhibited much poorer performance. Consequently, CB can be an alternative to synthesize *in situ* polymerized composite electrode materials.

Based on the above discussion, it can be concluded that free-standing composite electrodes with CNTs or graphene

offer various advantages over the conventional ones. These conductive additives can significantly improve the electronic conductivity. Because of good conductivity, free-standing films not only can act as a current collector but also offer excellent electron transport kinetics. Getting rid of the metal current collector in these films can lead to a reduction in the dead mass of the electrode, thereby effectively increasing the volumetric energy density. The microporous structure in these films is beneficial for the penetration of the electrolyte as well as fast diffusion of lithium ions in the electrode structure. These films offer reasonable mechanical properties with inherent flexibility, which is very beneficial for flexible LIBs.

Researchers are looking forward to some options other than LIBs. As alternatives, sodium-ion batteries (SIBs) are getting increasingly popular owing to their advantages over LIBs.<sup>701</sup> In principle, polymers are promising candidates for next-generation SIBs since they follow the conversion-reaction-based energy storage mechanism, making them electroactive toward most metal-ion batteries. A PMDA-based polyimide with microflower-like morphology was tested as the anode in SIBs.<sup>702</sup> A two-step enolization reaction with redox potential below 1.5 was observed during the electrochemical performance, suggesting its suitability as the anode for SIBs. Sodium terephthalate is one of the most promising anode materials for SIBs due to its low redox potential, while the performance is not stable due to its undesirable dissolution in the electrolyte. To overcome this problem, a polymer was reported bearing sodium terephthalate as the side group acting as the redox-active unit; here, the polymer backbone ensured stability.<sup>703</sup> By suitably adjusting the structure with an electron-donating group, researchers were able to obtain lower potential and much better cycling performance with almost 100%

coulombic efficiency for over 100 cycles at 30 mA g<sup>-1</sup>. Metal-ion batteries are going to become increasingly popular, and polymer materials can be the material of choice in the future.<sup>704,705</sup>

### 11.3 Fiber-shaped energy devices

Due to the high flexibility and easy fabrication typically derived from the solution process, a variety of functional polymers are explored for soft electronic devices. The use of polymers for thin-film electronic devices has been extensively discussed by many review articles, while it is rare to summarize their applications in a type of new soft electronic devices, *i.e.*, fiber-shaped electronic devices that attract increasing attention with the rapid advance of information technology and artificial intelligence.<sup>706-711</sup> When compared with the generally bulky or thin-film counterparts, fiber-shaped electronic devices demonstrate some unique and promising properties. For instance, they can accommodate complex deformations such as bending in various directions, twisting, and stretching. Furthermore, they can also be woven into soft, breathable, and comfortable electronic fabrics to satisfy the requirements of portable and wearable electronics.

Functional polymers are the key components for the realization and high performance of fiber-shaped electronics devices. In this subsection, a fiber electrode is first presented; then, we summarize the different types of fiber-shaped electronic devices, and the future direction is highlighted at the end. Different from earlier review articles, in this paper, we focus more on discussing them from a viewpoint of practical applications.

Currently available fiber-shaped electronic devices include solar cells, triboelectric and piezoelectric generators, supercapacitors, LIBs, lithium-sulfur batteries, and metal-air batteries.<sup>712</sup> With the help of a timeline, different types of fiber-shaped electronic devices are summarized in Fig. 97. It starts from sensors, solar cells, and light-emitting devices, and then extended to supercapacitors and batteries, almost according to the degree of difficulty in making them into a fiber

shape in the lab. With further increasing interest from the industry, the main efforts are focused on the continuous fabrication of fiber-shaped energy-harvesting (*e.g.*, solar cells) and storage (*e.g.*, LIBs) devices, which may soon find large-scale applications in many fields.

The electrode has always played a key role in electronics, and it is even more important for fiber-shaped electronic devices. It is necessary to carefully control the composition and microstructure of fiber electrodes because, when compared with traditional plates or film substrates, the curved surface of a fiber electrode makes it much more difficult to deposit thin active layers, particularly under a continuous fabrication process. In addition, the fiber electrodes should be simultaneously flexible, strong, and conducting. This flexibility guarantees to meet the production process necessary in the industry, such as roll-to-roll fabrication. The fiber electrodes also serve as substrates; therefore, they must be sufficiently strong to bear various deformations without breaking during use. Of course, charges would transport along the fiber electrode, and the pathways are much longer in fiber-shaped electronic devices as compared to their planar counterparts. Therefore, they should have higher electrical conductivities to provide satisfactory performances at lengths ranging into meters or even kilometers for real-world applications.

Polymers, metals, and carbon nanomaterials represent the three main categories of candidates for fiber electrodes. Polymers have been mostly explored as they are efficient for both moiety control and surface modification. A spectrum of conducting polymers such as PANI, PPy, and PEDOT have been mainly explored. The resultant fibers were flexible but typically with relatively low conductivities (<10<sup>2</sup> S cm<sup>-1</sup>). A second phase with much higher conductivity, *e.g.*, CNTs, is therefore added to increase the conductivities up to 10<sup>3</sup> S cm<sup>-1</sup>. However, it still needs to be further enhanced to meet the industrial production requirements of the resulting fiber-shaped electronic devices. They are expected to show conductivities on the level of 10<sup>4</sup> S cm<sup>-1</sup> at room temperature. Recently, bare CNTs have also

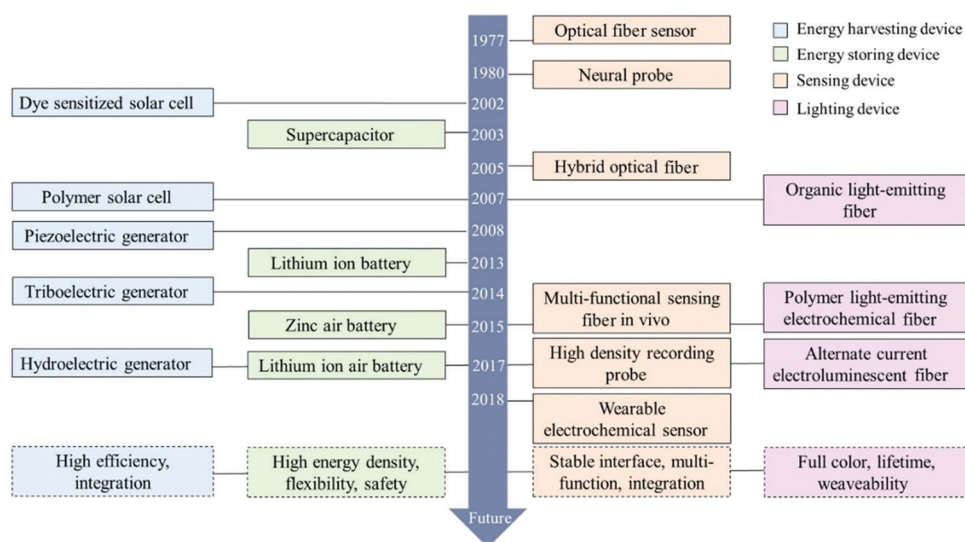


Fig. 97 Advances in fiber electronic devices with four main functions: energy harvesting, energy storing, sensing, and lighting according to the timeline.

been assembled or coassembled with polymers into flexible and continuous fiber electrodes based on chemical vapor deposition processes, and the CNTs are highly aligned to provide high conductivities of  $10^4 \text{ S cm}^{-1}$ .<sup>713–715</sup> However, the use of chemical vapor deposition techniques makes the synthesis much more expensive when compared with those for polymers, and large-scale production remains difficult.

Generally, positive and negative fiber electrodes are twisted into fiber-shaped electronic devices.<sup>712</sup> For instance, a fiber-shaped dye-sensitized solar cell was wound from a Ti wire coated with active materials and a CNT fiber (Fig. 98a), followed by the introduction of liquid electrolytes. However, the use of liquid electrolytes necessitated very strict sealing treatment that remains challenging, while the replacement with polymer gel electrolytes largely decreased the PCE. To this end, a fiber-shaped polymer solar cell that was completely solid had been invented by the use of CPs as active materials (Fig. 98b). Besides the above twisted structure, a lot of effort has also been devoted toward fabricating solar cells with a coaxial structure. In other words, a thin and transparent conducting layer was coated or wound on an active-layer-modified fiber electrode. For instance, a fiber-shaped PVSC was firstly produced by depositing active perovskite materials on the fiber electrode and then winding the aligned CNT sheet as the second electrode (Fig. 98c).<sup>711</sup> The twisted structure favors continuous production, but the counter-electrode fiber may shield light with low PCE. In contrast, it is not

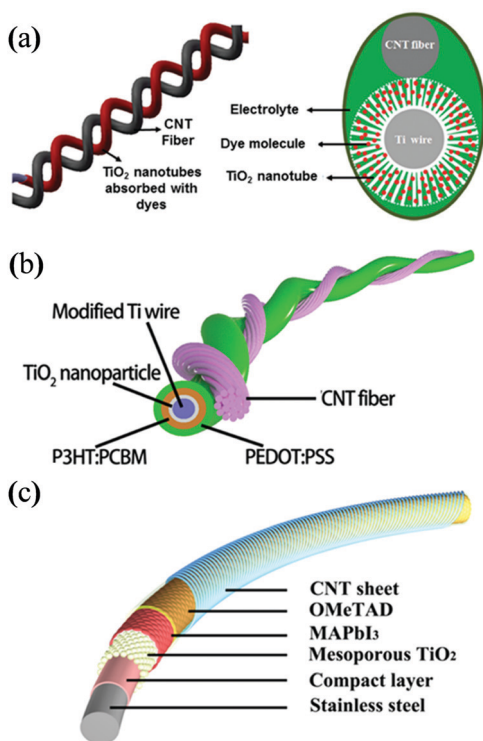
easy to continuously prepare a thin-film electrode for a coaxial structure.

It should be emphasized that the interface in a fiber device exhibits several unique features as compared to its 3D counterparts. For instance, for a fiber-shaped polymer solar cell with a twisted structure, the interface between the highly curved charge and hole transport layer are different from conventional flat interfaces. They are concentrated at the contact points of the two twisted fiber electrodes while uniformly distributed through the entire interface of the planar structure. In the case of coaxial structures, external electrodes are required to be stably and uniformly attached to the outer surface; therefore, high mechanical and electrical properties are desirable for achieving high PCEs.

When photoactive materials are replaced by electrochemically active materials, fiber-shaped supercapacitors, batteries, and other energy storage devices could be obtained.<sup>709–711</sup> Like solar cells, the fiber-shaped energy storage devices are also made with both twisted and coaxial structures. Currently, fiber-shaped supercapacitors have been mostly explored possibly due to their easy fabrication, although they have found few applications. Recently, fiber-shaped batteries have started to attract increasing attention; now, they have become the primary candidate for real applications. Studies involving fiber-shaped LIBs are being conducted by a few companies for enabling their industrial production.

Fiber-shaped energy-harvesting and storage functionalities have also been integrated into a single fiber device. They may be further woven into flexible and even stretchable power textiles toward portable and wearable applications. Although it seems easy to assemble the above fiber devices into textiles based on the weaving process, it remains challenging to effectively connect so many fiber electrodes in the power textile. There exist many reports that have discussed solar cells or battery textiles, but they are also limited to relatively small sizes (in the range of centimeters). There is still a long way to go for fiber-shaped energy-harvesting and storage devices from the lab to the market. Scientifically speaking, further investigations into a distinctive interface are necessary to guide performance enhancement. Technically speaking, the emphasis will be made on generic and effective strategies for the continuous preparation of fiber electrodes and reliable connection of fiber-shaped devices.

The two main energy storage systems (supercapacitors and batteries) made of polymers are first introduced. However, such energy storage devices are usually rigid, heavy, and bulky. Subsequently, flexible and portable fiber-shaped energy devices are presented from the viewpoint of practical applications. From the above contents, it is evident that polymers have achieved considerable progress in basic research and practical applications in the field of energy storage. The main challenges needed to be addressed in the future are as follows: (1) fabricating supercapacitors with high energy density and batteries with high power density; (2) designing fiber-shaped energy devices with excellent electrochemical performance and mechanical stretchability.



**Fig. 98** Fiber-shaped solar cells with different active materials and architectures. (a) Fiber-shaped dye-sensitized solar cell twisted from Ti wire and CNT fiber. (b) Fiber-shaped polymer solar cell with a twisted structure. (c) Fiber-shaped PVSC with a coaxial structure. Reproduced from ref. 712 with permission. Copyright (2016) Wiley-VCH.

## 12. COFs

COFs are a class of crystalline porous polymers assembled from organic building blocks by strong covalent bonds.<sup>716–722</sup> Different from traditional polymers, they allow the precise and atomically controllable integration of struts into long-ranged ordered networks. COFs feature extended porous structures with high surface area, large cavity, good thermal stability, and low density. Moreover, the tunable pore size, shape, and environment, as well as the backbone functionalization of the network realized by pre- and/or post-synthesis modification, further expands their applications in gas storage and separation, optoelectronics, catalysis, energy storage, and electronic devices.<sup>723–730</sup>

In COF chemistry, the molecular units can be accurately arranged into extended frameworks under the guidance of reticular chemistry,<sup>731</sup> where the functionality and pore metrics can be further tuned. Since the first report on COF-1 and COF-5 by Yaghi *et al.*,<sup>732</sup> COFs with various structures and characteristics have been designed and synthesized. The structures of COFs are largely determined by the geometry and dimensions of the building units and linkages, which can be classified into 2D and 3D frameworks. In 2D COFs, the structural units are covalently aligned in 2D atomic layers that further stack *via* noncovalent forces to form eclipsed or staggered configurations.<sup>733,734</sup> For 3D COFs, the building blocks are covalently and periodically oriented to form 3D architectures, and the entire crystalline structure gets retained by covalent bonds.

The syntheses of most organic polymeric materials are kinetically controlled reactions, where the covalent bonds are irreversibly formed, resulting in amorphous products. To afford extended crystalline structures, the chemical reactions should be reversible and allow for “error-checking” and “proofreading” to give thermodynamically stable and highly ordered networks. In this way, the structural defects can be repaired by a reverse reaction; consequently, the bonds can be reformed to achieve crystalline architectures. Several reversible reactions have been employed to prepare COFs, and they can be grouped into B–O, C=N, C–O, Si–O, C–N, N–N, C–C, *etc.* (Fig. 99) based on the linkage type. The first two COFs (COF-1 and COF-5) were synthesized by the self-condensation of boronic acid and the cocondensation of boronic acid and catechol to give boroxine- and boronate-ester-linked structures, respectively.<sup>732</sup> Borosilicate and spiroborate were incorporated to prepare boron-based COFs, and most of them exhibited good thermal stability but poor stability when exposed to water or acid, because water promoted the reverse reaction and led to the decomposition of COFs.<sup>735,736</sup> To improve the chemical stability of COFs in water and even strong acid or basic aqueous solutions, more chemical reactions, *e.g.*, phenazine- and  $\beta$ -ketoenamine-based COFs, were further explored to fabricate COFs.<sup>737–739</sup> Post-synthesis modification was also utilized to stabilize the COF structures. Further, it was possible to employ reversible–irreversible cascade reactions or even irreversible nucleophilic aromatic substitution reactions to synthesize COFs, and the

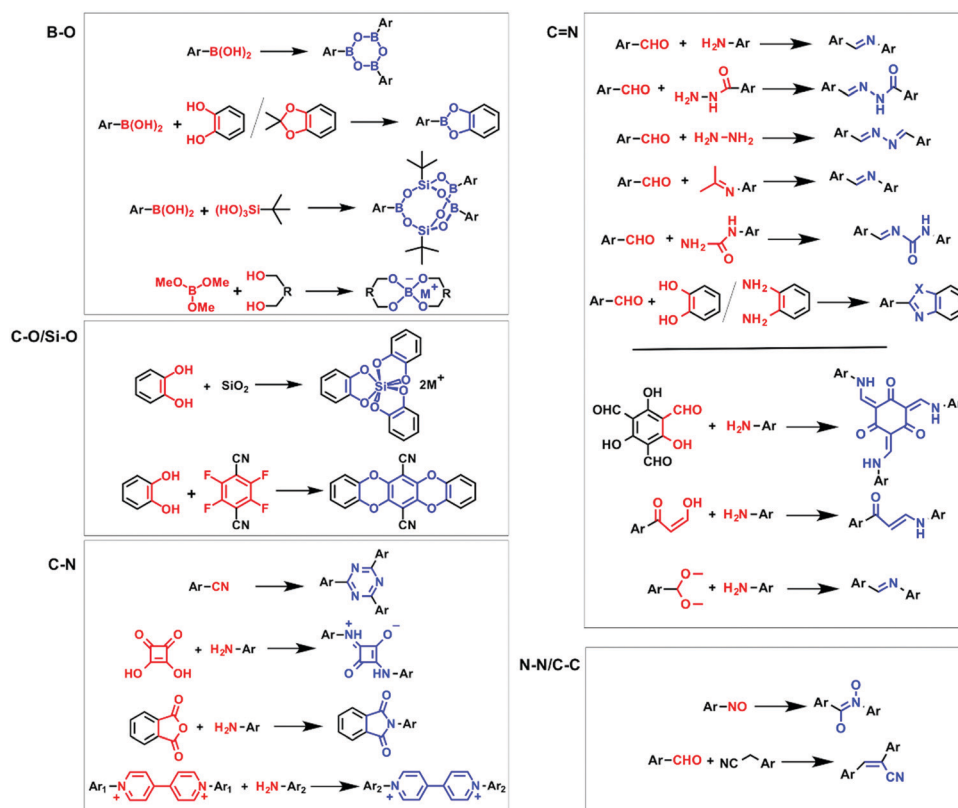


Fig. 99 Chemical reactions for the preparation of COFs.

irreversibility of the reaction imparts the resulting COFs with good chemical stability.<sup>737,740,741</sup>

Under the topology direction, COFs with targeted structures could be fabricated by an appropriate selection of the building units. In the target topology net, the structure can be deconstructed into basic geometric units, and further substitution with building molecules of certain connectivity and geometry would yield a COF possessing the same underlying net. To ensure that the conformation of the building blocks remains unchanged during the assembly process, rigid units with discrete directionality are preferred to construct well-defined networks. A wide range of molecules, such as simple arenes to extended  $\pi$  aromatics with heteroatoms, coordinated strings, and large macrocycles, have been employed to construct COFs. In 2D COFs, topologies such as *hcb*, *sql*, *kgm*, and *hxl* have been realized in COF structures, where there is only one type of edge linking the vertices.<sup>732,742–744</sup> To improve the structural complexity, Zhao *et al.* reported the heterostructural mixed linker strategy to construct COFs with three types of pores.<sup>745</sup> Employing the condensation reaction of a tetraamine ( $D_{2h}$  symmetry) and two dialdehydes ( $C_2$  symmetry) of different lengths, they successfully synthesized a COF with hierarchical pores of inequilateral hexagon, as well as small and big triangular shapes. Other strategies such as desymmetrized vertex design and utilizing macrocycles as building blocks have also proven to be capable of giving COFs with periodically heterogeneous pore structures.<sup>746,747</sup> With regard to 3D COFs, due to the limited choices of building units and synthesis difficulties, the reported topologies of 3D structures in COFs are only *bor*, *ctn*, *pts*, *dia*, *rra*, *srs*, and *lon* (Fig. 100).<sup>748–753</sup> Feng *et al.* introduced the flexible building block of  $\gamma$ -cyclodextrin ( $\gamma$ -CD) into 3D COFs for the first time.<sup>751</sup> Utilizing the transesterification reaction between trimethyl borate and hydroxyl groups, they could construct CD-COFs possessing an anionic framework with spiroborate linkage and tunable counter ions identified as the *rra* topology.

Once the organic building units and chemical reactions are judiciously determined, finding the appropriate synthesis

conditions is also important to obtain crystalline porous frameworks. COFs are generally prepared by solvothermal synthesis, and the reaction vessels charged with precursor solutions are usually vacuumed to low pressure before being sealed. After heating for several days, polycrystalline products with high surface areas can be obtained. The preparation of single crystals has always been a challenge in COF chemistry, and the products always end up as polycrystalline or even amorphous solids. Recently, Yaghi *et al.* reported the first examples of COF single crystals suitable for single-crystal X-ray diffraction data collection.<sup>753</sup> The strategy here is to add excess aniline as a nucleation inhibitor and competitive modulator during the synthesis of imine-based COFs (COF-300, COF-303, LZU-79, and LZU-111) at ambient temperature, which improves the reversibility during the course of bond formation. Dichtel and coworkers developed a seeded-growth method to prepare single-crystal 2D boronate ester-linked COFs.<sup>754</sup>

However, solvothermal synthesis is not applicable for industrial production processes because of the long reaction time, involvement of a large number of solvents, and harsh reaction conditions. To shorten the reaction time, microwave heating is also employed to assist the synthesis of COFs with higher production yields and purity.<sup>755</sup> Solid-state reaction is a good choice for the fast and scale-up production of COFs. Banerjee *et al.* successfully synthesized  $\beta$ -ketoenamine-linked COFs by using a solvent-free mechanochemical method and further improved the crystallinity and porosity of COFs by adding the external coagent of *p*-toluene sulfonic acid that helps to control the covalent bond formation during solid-state preparation.<sup>756,757</sup> This solid-state synthesis was further applied with a twin-screw extruder for large-scale COF synthesis, showing promising potential for industrial production.<sup>758</sup> Another solvent-free synthesis method is ionothermal synthesis, where molten salts or ILs were used as solvents.<sup>759</sup> Fang *et al.* reported the successful preparation of a series of 3D COFs utilizing ILs as the green solvent at ambient temperature and pressure in a short reaction time, which opens up a new way for green and massive production of COFs.<sup>760</sup> Other methods such as continuous flow synthesis, sonochemical method, and photochemical synthesis have also been reported for synthesizing COF powders with high production rates.<sup>761–763</sup>

For practical applications such as catalysis, electronic devices, and sensing, these powder-form materials need to be processed into certain shapes, particularly thin films. The most direct and common way to synthesize COF films is solvothermal synthesis. Here, the substrates are immersed in the precursor solutions during the reaction, resulting in the growth of COF coating layers on the substrate surface. Various substrates have been used to prepare COF films such as graphene, indium-doped tin oxide, gold, and silicon dioxide, and the type of substrates influences the quality and orientation of the films.<sup>764–767</sup> However, it is difficult to control the morphology and achieve large-area films by using this method. Interfacial synthesis is also employed to prepare COF films with good control to be cast and evaporated giving free-standing membranes with preferential orientation.<sup>768</sup> For the large-scale synthesis of COF membranes, the continuous flow method

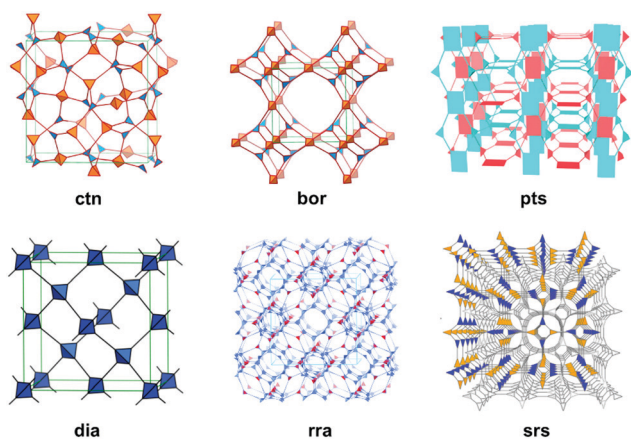


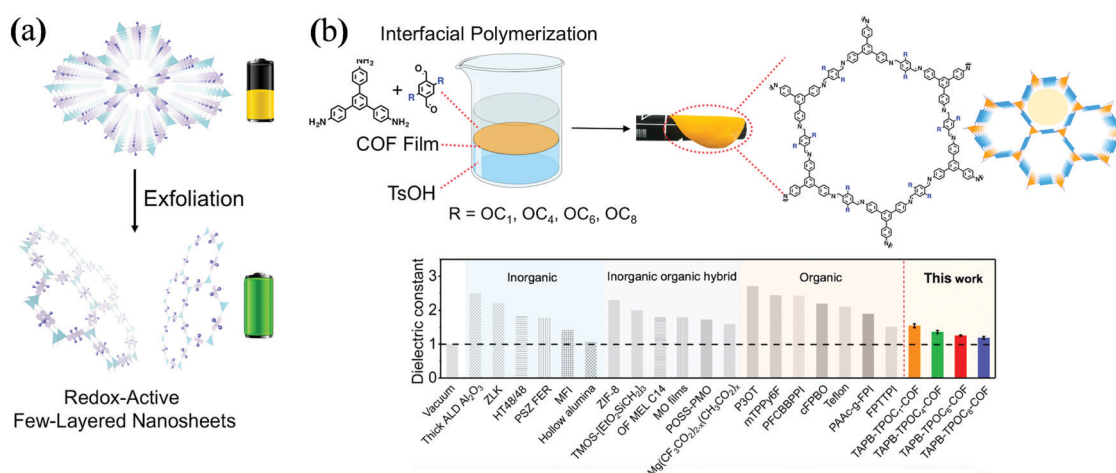
Fig. 100 3D topological diagrams of COFs. Adapted from ref. 716, 748, 750, 751 and 752 with permission. Copyright (2017, 2007, 2009, 2016, 2017, and 2018) AAAS, American Chemical Society, and Wiley-VCH.

and molecular baking strategy show promise. The former technique developed by Dichtel and coworkers enables the growth of COFs under the continuous flow condition, achieving good control over the film thickness and avoiding contamination from the powdered precipitates.<sup>769</sup> Standalone and defect-free COF membranes with large areas can be obtained by the molecular baking method, and the concept was proven in the cases of  $\beta$ -ketoenamine and imine COF membranes.<sup>770</sup> Interfacial synthesis is also employed to prepare COF films with good control over the nucleation process and film thickness, and the growth of COFs can be on either the air-liquid interface or the liquid-liquid interface.<sup>726</sup> Using this method, Feng *et al.* prepared large-area crystalline monolayers and multi-layer 2D polymers with controlled thickness by the Schiff-base condensation reaction at the air-water and water-chloroform interfaces, respectively.<sup>771</sup> The obtained monolayer film ( $\sim 0.7$  nm) on a wafer (lateral size: 4 inch) showed high crystallinity and exceptional mechanical robustness. Another mild way to prepare COF films is room-temperature vapor-assisted conversion synthesis, which was first reported by Medina *et al.*<sup>772</sup> Basically, the precursors were first dropped onto the substrate and then exposed to organic solvent vapor. The thickness of the resulting film could be precisely controlled by tuning the volume and concentration of the precursor droplet. This strategy was effective for preparing BDT-COF and COF-5. Later, colloidal suspension of COF was directly used.

Bulk 2D COFs are layered structures whose atomic layers are stacked *via* weak interactions such as van der Waals force and/or hydrogen bonding, and the breakage of the interlayer  $\pi$ - $\pi$  interactions can lead to single- or multilayered COF thin films, namely, COF nanosheets (CONs). The force to exfoliate the bulk materials can be either internal forces (which is the case for the self-exfoliation method) or external forces (used in solvent-assisted exfoliation, mechanical delamination, and chemical exfoliation strategies).<sup>773-776</sup> The solvent-assisted method is the most commonly used method due to its simplicity and low cost.

Further, the mechanical delamination method usually involves grinding or ball milling, which is an effective and powerful way to exfoliate. The self-exfoliation and chemical exfoliation methods often require a special chemical structure designed by the building block. Gu *et al.* employed ionic building blocks to construct COFs with electrostatic repulsion that can resist interlayer interactions. As a result, the afforded framework can self-exfoliate into monolayers or multilayers in solutions, and the electrophoretic deposition was further utilized to prepare COF films with controllable thickness.<sup>777</sup> Wang *et al.* reported a self-limiting solid-vapor interface reaction strategy to directly prepare large-scale crystalline COF sheets with one-atom thickness *via* a Schiff base reaction, avoiding the need for further exfoliation processes.<sup>778</sup>

COFs have been widely studied for gas storage and separation due to their high porosity and low density. Furthermore, their designable structure and diverse functionality make them highly promising for many other applications. In the field of energy storage and electronic devices, Wang *et al.* mechanically delaminated redox-active COFs into few-layer nanosheets and applied them as a battery cathode that yielded considerably improved performance because of the shortened  $\text{Li}^+$  transportation path in the exfoliated COFs (Fig. 101a).<sup>725</sup> Furthermore, they prepared a series of COF films by interfacial polymerization, and these films showed ultralow dielectric constants under high humidity due to the porous and symmetrical structure as well as the incorporated alkoxy chains (Fig. 101b).<sup>726</sup> In the field of photoluminescence, Wang *et al.* designed and synthesized an AIE-luminogens-based 3D COFs, which could emit yellow fluorescence upon excitation; their further coating on a blue LED afforded a white LED with long working life.<sup>779</sup> As demonstrated by Feng *et al.*, COFs are a good platform for phosphorescence study, and the phosphorescence behavior can be tuned by adjusting the interlayer distances in a crystallization-induced phosphorescence-phosphor-based COF.<sup>780</sup> In the field of catalysis, the first example was reported by Wang *et al.*, and the COF-based



**Fig. 101** (a) Schematic illustration for the exfoliation of 2D redox-active COFs into exfoliated COFs as cathodes for LIBs. (b) COF films prepared by interfacial polymerization, and their dielectric constants compared with those of typical commercial and representative ultralow- $k$  materials. Adapted from ref. 725 and 726 with permission. Copyright (2017 and 2018) American Chemical Society and Wiley-VCH.

catalyst was prepared by the metalation of an imine-based 2D COF with Pd(II), showing excellent yields, high stability, and facile recyclability for the Suzuki–Miyaura coupling reaction. This can be attributed to the highly exposed active sites and the rapid diffusion of products in a porous scaffold.<sup>723</sup> Cui *et al.* successfully prepared a series of multivariate chiral COFs with organocatalyst-modified channels, serving as efficient asymmetric catalysts with stereoselectivity and diastereoselectivity.<sup>724</sup>

In summary, COFs have shown promise in diverse applications due to their unique characteristics. They are ideal platforms for various fundamental studies because of their highly ordered and porous structures. To further expand their structural complexity and diversity, multicomponent networks with hierarchical pores should be developed. The crystallinity of COFs still remains a challenge, and new reactions should be explored to construct highly crystalline materials with controllable stability and functionality. For functional exploration, COF-based composites are desirable. For example, well-aligned channels with tunable size and environment in COFs could be utilized to accommodate functional guest molecules. The final goal is to apply these porous materials in real-life applications. Therefore, low processability and scalability issues should be resolved. Researchers have proposed many promising strategies to deal with these problems and have made significant progress in this field. It is believed that the continued study on this emerging direction can yield practical applications of COFs in the future.

### 13. Conclusion and outlook

Advanced functional polymers are very interesting materials for broad applications requiring specific functions in their corresponding fields, such as biomedicine, aerospace, artificial intelligence, bionics, robotics, energy, electronics, separation, *etc.* During the past few decades, tremendous efforts have been devoted toward the advancement and development of this field. In this review, we addressed the notable recent progresses made in functional polymers, including their concepts, classifications, design strategies, preparation methodologies, performances, structure–property relationships, potential applications, and challenges. From these discussions, we can conclude that functional polymers play a critical role in the progress and development of our society and will continue to do the same in the future.

Although great advances in functional polymeric materials have been achieved, numerous challenges still persist and should be seriously considered. A few examples are considered for each type of functional polymer as follows:

(1) Although ATRPs with organic photocatalysts have been rapidly developed in recent years, the synthesis is still limited to the polymerization of methyl acrylate in the presence of metal catalysts. Therefore, developing ATRP in the presence of a metal-free catalyst still needs to be determined.

(2) For luminescent polymers, pure and stable white-light emissions and highly efficient deep-blue-light emissions from

phosphorescent materials have been rarely reported, which remain to be further developed.

(3) PCEs of OSCs are still much lower than those of PVSCs. Researchers should focus on the following aspects: (1) designing more efficient donors/acceptors, (2) controlling the aggregate structures of materials and the morphology of blend films, and (3) realizing the application of multifunctional organic photovoltaic devices.

(4) An ultralow-*k* polyimide still needs to be designed and prepared without sacrificing other properties. A simple synthesis method of NLO materials should be formulated.

(5) Many challenges in the drug delivery field need to be tackled and more efforts should be focused before translating them into clinical applications. For example, overcoming biological barriers and realizing highly efficient drug delivery systems need to be addressed. Because of the complexity of the ocean environment, marine antibiofouling polymer coatings with environmental adaptation need to be developed in the future along with maintaining high efficiency and long service life.

(6) The main challenge of SPs is to design and fabricate materials that possess both dynamic capabilities and strong mechanical properties. The self-healing property can be facilitated in the soft state, while the mechanical strength of SPs is usually low. Materials with good mechanical properties usually have poor dynamic capabilities due to the presence of rigid polymer chains and restricted molecular mobility. This problem may be addressed by incorporating dynamic covalent bonds into SPs that can be cleaved and recombined when triggered by external stimuli, such as light or temperature.

(7) Electrochromic polymers have problems of trivial film processing, uneven electrochromism, ultraviolet degradation, *etc.*, which can seriously limit large-scale preparation and cycling stability of the ECD.

(8) Integrating multifunctions, *e.g.*, shape-memory and color-changing abilities, into a single SMP should be investigated for extending its potential applications. That is to say, shape and color may be synergistically changed when the SMP is triggered by an external stimulus. Moreover, SMPs are still investigated in the basic research phase; it is highly demanded to promote their development in practical applications.

(9) In the process of oil/water separation, seawater desalination, and blood purification, impurities, salts, sticky molecules, and organic compounds can be easily deposited on the surface of a membrane, promoting membrane fouling and therefore reducing membrane life. Antifouling polymer materials are highly desirable for membrane separation industries.

(10) Design of energy storage devices with high performances, such as high energy density, high power density, and long cycle life, is still challenging. Although flexible, bendable, twistable, and stretchable energy storage devices, *i.e.*, fiber-shaped energy devices, have been investigated for several years, they are still in the early stages of lab-scale research. Therefore, the development of novel strategies for the fabrication of such devices in both academia and industry has become necessary.

(11) Although many strategies have been developed to improve the crystallinity of COFs, obtaining larger single crystals is fraught with difficulties.

## Contributions

K. W. wrote sections of Abstract, Introduction, Conclusion and outlook, and transition sections. Z. A., D. H., A. Q., R. H., Y. H., J. Z., W. F., J. S., Z. C., B. T., Y. D., and Z. L. wrote the section of polymerization methodologies. J. G., A. Q., B. Y., and L. W. wrote the section of luminescent polymer materials. T. L., X. Z., H. C., J. L., Y. H., and F. H. wrote the section of photovoltaic polymer materials. Y. Z., Q. L., and Z. L. wrote the section of other electrical and optical polymer materials. X. J., X. Z., S. W., D. H., J. L., Z. Z., Y. S., X. Z., X. Z., S. L., H. C., and G. Z. wrote the section of bio-related polymer materials. H. W., F. H., H. X., Y. Z., and J. Z. wrote the section of supramolecular polymer materials. C. X., X. Z., X. Q., and Y. Y. wrote the section of stimuli-responsive polymer materials. T. X. and M. Q. Z. wrote the section of shape memory polymer materials. J. J., W. Z., C. Z., and Z. K. X. wrote the section of polymer membranes for separation. K. A., Z. W., X. S., and H. P. wrote the section of Energy storage polymers. Y. Z., X. F., and B. W. wrote the section of covalent organic frameworks. B. Z. T. proposed and corrected the paper.

## Conflicts of interest

There are no conflicts to declare.

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