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Covalent organic frameworks (COFs) for electrochemical applications

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Covalent organic frameworks are a class of extended crystalline organic materials that possess unique architectures with high surface areas and tuneable pore sizes. Since the first discovery of the topological frameworks in 2005, COFs have been applied as promising materials in diverse areas such as separation and purification, sensing or catalysis. Considering the need for renewable and clean energy production, many research efforts have recently focused on the application of porous materials for electrochemical energy storage and conversion. In this respect, considerable efforts have been devoted to the design and synthesis of COF-based materials for electrochemical applications, including electrodes and membranes for fuel cells, supercapacitors and batteries. This review article highlights the design principles and strategies for the synthesis of COFs with a special focus on their potential for electrochemical applications. Recently suggested hybrid COF materials or COFs with hierarchical porosity will be discussed, which can alleviate the most challenging drawback of COFs for these applications. Finally, the major challenges and future trends of COF materials in electrochemical applications are outlined. **EXAMPLE SERVIEW ARTICLE**
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1. Introduction

The crucial environmental problems related to the increased emission of greenhouse gases and pollutants (CO_x, NO_x, SO_x) and fine particulates) and the growing global energy demand underpin the broad interest in clean energy technologies

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including e.g. electrochemical water splitting for hydrogen production, batteries/supercapacitors for clean energy storage and fuel cells for emission-free electricity generation. $1-6$ To further advance such devices and processes needed for future sustainable energy production, conversion and storage, the development of novel materials is a key task. Chemical reactions occur on the surfaces of electrodes, while transport in membranes occurs through voids and thus it is obvious that porous and high surface area materials can play an important role in the development of such devices. The properties of porous materials have a close relationship with their performances in different electrochemical applications and these properties can vary with the size, structure and arrangement of the pores, the extent of porosity and composition of the materials themselves.^{1,7–9} In particular, porous carbons have been widely investigated as electrode materials in energy conversion and storage devices, as they show high electrical conductivity, can exhibit very large surface areas and porosities to ensure a high number of active sites and sufficient mass transport and are furthermore often cheap and chemically stable.^{7,10} However, as these materials are generally prepared by high temperature treatment, i.e. pyrolysis, their chemical and nano-/microstructure is hardly predictable and controllable. Review Article

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Recently a range of novel porous materials has been developed, whose structures can be precisely manipulated and which are therefore interesting for several electrochemical processes,^{9,11-17} particularly, metal organic frameworks $(MOFs)^{18-24}$ and covalent organic frameworks $(COFs)^{25-33}$ Through the modular nature of these materials, their pore structures and functional groups can be tailored independently, thus enabling the introduction of defined active sites onto the large permanent and accessible surfaces to promote or catalyse targeted electrochemical reactions.³⁴⁻³⁶ Indeed, MOFs have been widely investigated as potential electrocatalysts in recent years. Chemical stability especially in reactions performed in strongly acidic or basic electrolyte solutions as well as low electrical conductivity are however issues that have to be considered when applying MOFs in electrochemical applications.

COFs, a subclass of porous polymers, possess rigid backbones, long-range order and permanent porosity. 2D layered or 3D network structures can be constructed, determined by the geometry of the respective building units and the nature of the covalent linkage between them. $14,37-39$ For the first time, COFs (COF-1 and COF-5) were reported by Yaghi and co-workers, who applied the topological design principles of MOFs in the synthesis and structural determinations of $COFs$ ¹⁶ As they are entirely built from organic building blocks connected by covalent bonds, many COFs show π -conjugated structures and thus at least a modest electronic conductivity. Indeed, many COFs are organic semiconductors with tuneable band gaps. Furthermore, the exclusive covalent connections can yield COFs with high chemical stability. In fact, while the first COFs reported were not even water stable, ongoing development has provided COFs with stability under harsh acidic and/or basic conditions.40,41

Although research efforts on COF applications in electrochemical devices are still in their infancy, these properties together with their large surface area, tuneable pore size, high crystallinity and tailorable architecture have made them promising candidates to promote electrochemical processes (Fig. 1). Several examples for the utilization of COFs in electrochemical applications have been demonstrated recently.^{36,42-44} This review aims to discuss the principles of COF materials design and synthesis to introduce tailor-made properties useful for electrochemical applications. Furthermore, the challenges and future perspectives of COFs for clean energy conversion and storage are described. Finally, also the drawbacks of COFs in electrochemical applications will be discussed, for example their relatively small pores, which inhibit mass transport and

Fig. 1 The most important properties of covalent organic frameworks (COFs), which are decisive for the potential of these materials in electrochemical and other applications.

Arne Thomas

Prof. Arne Thomas studied chemistry in Giessen, Marburg and Edinburgh and received his PhD from the Max Planck Institute of Colloids and Interfaces (MPI-KGF) in Potsdam, Germany. After a postdoctoral stay at the University of California, Santa Barbara, as an AvH fellow, he rejoined the MPI-KGF as a group leader. In 2009, he became a professor at the Technische Universität Berlin, where he is leading the Department of Functional Materials.

their relatively low conductivity, which yield low charge carrier mobility. To alleviate these drawbacks, a range of hybrid and/or hierarchical porous COFs has been suggested recently, which therefore receive a certain attention in this article.

2. Design and fabrication of COF-based materials for electrochemical applications

2.1 Requirements of electrode materials for electrochemical applications

Electrochemical reactions play an important role in the development of novel energy conversion and storage strategies. Some of them are electrocatalytic reactions, including the oxygen reduction reaction (ORR) in fuel cells, $45,46$ oxygen evolution reaction (OER) in electrolytic water splitting and metal–air batteries, $47,48$ hydrogen evolution reaction (HER) in water splitting,^{5,49} and CO_2 reduction reaction (CO_2RR) in pollutant conversion.50,51 Furthermore, challenging electrochemical reactions are found in energy storage devices such as lithiumion batteries^{6,42} and supercapacitors.^{52,53} Thus, the design and synthesis of efficient electrode materials to gain suitable electrochemical performances are crucial for practical applications. There are some general requirements for electrode materials including high conductivity to promote electron transfer between electrode surface and electrolyte solution, large surface areas and pore volumes to ensure a high number of accessible catalytic sites and/or efficient mass transport to the electrodes, and high chemical stability to ensure a long lifetime of the device. Nevertheless, the detailed requirements on structure and properties of electrode materials are of course mainly dependent on the respective electrochemical application in focus. Open Soc Rev

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COFs show some intriguing features, which make them interesting materials for electrochemical applications. There is at first the very good structural control. Indeed, by selection of appropriate linkage motifs and building blocks, the design principles of COFs enable the creation of materials with defined chemical structures in the backbone as well as tuneable pore sizes and pore geometries, which both help to modify the chemical and physical properties of the COFs.

COFs with well-defined active sites for electrocatalytic processes and controllable porosities for efficient mass transport have been developed.⁵⁴ Typically, nitrogen-containing ligands such as bipyridines and porphyrins could act as coordination sites for metal incorporation within COFs, providing single-atom metal catalysts in electrocatalytic reactions. $36,55$ Besides, introduction of redox-active sites into COFs enables enhanced pseudocapacitive performance in supercapacitors as well as the reversable process in rechargeable batteries.^{41,56} On the other hand, COF hybrid materials with controllable porosities for efficient mass transport and conducting materials for good electrical conductivity and mechanical stability have been developed to further improve their electrochemical performance.⁵⁷

2.2 Design principles for COF-based materials for electrochemical applications

As the redox reactions in electrocatalysis and charge–discharge processes are accompanied by electrons, reactant/product and ion transport, several strategies to increase the mass transport and promote the electrical conductivity within the materials can be employed. For example, hierarchical porosities ensuring high surface areas but sufficient mass transport can be introduced in COF electrocatalysts.³⁶ Up to now, various strategies and methodologies have been developed to construct efficient COF-based electrodes (Fig. 2).

2.2.1 Chemical modifications. COFs enable scientists to control the composition and structure of crystalline and porous materials in terms of density and spatial arrangement of chemical functionalities. Depending on the desired application in electrocatalysis and energy storage, the chemical structure and thus number and nature of functional groups and active sites within COFs can be controlled by developing different approaches. Mostly, functional monomers, e.g. bearing redoxactive or metal coordinating groups are directly applied in COF synthesis. If this is not directly possible, backbone modification and post-synthetic modifications are further pathways to tailor the chemical structure and functionality of COFs, which are described in more detail in the following (Fig. 3).

a. Backbone modification. The intrinsic electrocatalytic activity of COFs is related to the presence of functional groups or moieties within the backbone, which are generally introduced through appropriate monomers, called linkers. These linkers bear certain groups allowing them to form robust covalent linkages with other linkers to finally form the COFs. Functional moieties can be either present already in the linker molecules or they are formed during the COF-forming reaction. For the latter, especially triazines,⁵⁸ and imines⁵⁹ have been shown to be suitable for electrochemical reactions. These linkages thus introduce heteroatoms in an otherwise solely sp²-carbon conjugated backbone structure, which can serve as binding sites for metals, basic sites or redox-active sites (Fig. 3a).

In addition to the aforementioned de novo synthesis strategies, the COF backbones can be further modified by postsynthetic modification, for example, by chemical reactions on the linkers or even entire building block exchange for framework-to-framework transformations.60–62

b. Functional group modifications. Constructing functional COFs with redox-active groups is of utmost significance, particularly for supercapacitor, Li-ion battery and proton conductivity applications. $63-65$ However, because of the incompatibility with the building blocks or reaction conditions, it is difficult to directly incorporate such functional groups into COFs via the typical solvothermal and mechanochemical synthetic approaches. In this regard, introduction of functional moieties into COF skeletons through chemical transformation with the preservation of their topological structures enable the possibility of tuning the functional properties of COFs (Fig. 3b).⁶⁶⁻⁶⁹

Fig. 2 Strategies for the design and fabrication of COF-based materials for energy conversion/storage applications.

By applying functional group modifications, it has been demonstrated that functional groups or moieties that cannot be incorporated using direct synthesis can be introduced into COF backbones.⁷⁰

c. Post-synthetic metalation. Post-synthetic modification can also be applied to introduce metal atoms into COFs.⁷¹ Heteroatoms in COF linkages, or functional groups attached to the linkers can be valuable to coordinate and bind metals introduced into the pore system. This can be extended to bind the metals to chelating moieties within the backbone. The presence of periodic and uniform porosity within COFs ensures that the adsorption sites for metals are fully accessible to metals added to the COFs. By applying this strategy, metal–organic complexes, probably already known for homogeneous catalytic reactions, can be immobilized as they act as maintaining part of the COF backbone (Fig. 3c). $36,43,72$ In the meantime, this concept has been exploited many times, especially applying common nitrogen-containing ligands such as bipyridines, $36,55,73$ or macrocyclic structures (e.g. porphyrin). $43,74-76$ Incorporating metals within COFs via controlled bonding also provides a new approach to use single-atom metal catalysts with homogeneous dispersion in electrocatalytic reactions. Besides, the large surface areas and pore sizes in the small nanometre regime also allows the incorporation of small metal

nanoparticles, which can also introduce new functionalities into COFs.

2.2.2 Controlling COF structures on all length scales. As mentioned, an intriguing advantage of COFs in electrochemical applications is that their chemical structure and nanostructure is, at least in principle, entirely known and can be tuned with high precision. However, regarding the microstructure, there are several further issues that have to be addressed. COFs are mostly observed in the form of bulk powders, which cannot be dissolved or melted, and thus can hardly be processed into bulk electrodes or films. Furthermore, reported charge carrier mobility in COFs is still very low, when compared to that of inorganic (semi)conductors and certainly does not get better when COFs are obtained as microcrystalline powders, as mentioned before. Finally, COFs possess mainly pores in the micropore or small mesopore regime, which impede mass transport through the material and thus the accessibility to the actually available abundant active sites on these large surface areas. In this regard, the design of COFs should not only focus on microscopic length scales (i.e. from angstrom to nanometres), but also include the engineering of COF materials on a macroscopic level i.e. from several nanometres to centimetres (Table 1). The challenge is to combine the chemical and crystalline structures of COFs with the possibility to design larger architectures with uniform sizes,

Fig. 3 Strategies for the preparation of functional COFs via post-synthetic modification (PSM) for their applications in electrocatalysis and energy conversion and storage.

well-defined morphologies and/or hybrid components to further enhance the properties of COFs. As an example, combining small and large pores, *i.e.* introducing hierarchical porosity into materials, can enhance ion and mass transfer, which will consequently improve the electrochemical performance, but will also be useful for other applications of COFs. Indeed, within recent years researchers have developed different strategies to control COF structures and morphologies at least on some length scales (Fig. 4).

a. Thin films and membranes. Beside the growth of COF films on typical electrode materials to achieve suitable electron transport and morphological control, growing of COF films can be important to construct other important compounds in electrochemical devices, namely the membranes, which separate the electrodes. Most prominent are probably the proton exchange membranes that are applied to separate the hydrogen and oxygen gases as an electronic insulator and a reactant barrier in fuel cells (Fig. 4a).^{77,78} It has been demonstrated that COFs with well-defined pore structures could provide suitable space to accommodate proton carriers. Moreover, the versatile organic linkers for COF synthesis enable the anchoring of strong acid groups (e.g. sulfonic acid group) on the COF backbone to further improve the proton conductivity.⁷⁹

To synthesize COF-based membranes, COFs were initially blended with polymers to create mixed matrix membranes.^{80,81} With advanced COF synthesis methods, strategies such as in situ growth, 82 layer-by-layer stacking, 83 and interfacial polymerization⁸⁴ have been developed to construct even freestanding COF-based membranes. These membranes show promising applications in proton exchange in fuel cells as well as a huge potential in membrane technology for electrocatalysis.

b. Hierarchical porosities. The sole presence of micropores or small mesopores in COFs can lead to diffusion, i.e. mass transport limitations, which is problematic for many applications Table 1 Selected properties of covalent organic frameworks for electrochemical applications. Early reports, controlling macroscopic level and most notable performances

and also for chemical reactions on electrodes. It has been demonstrated that introduction of larger pores (i.e. meso- and macropores) can provide unhindered mass transport to and from the electrode surface. $85,86$ Such a secondary porosity can be useful as the micropores in crystalline COFs already offer large surface areas to enable fast surface reactions. Currently, several synthetic strategies to generate hierarchical porosities (two or more defined pore sizes in one material) in COFs have been explored (Fig. 4b), including the preparation of hollow micro- and meso-tubular porous COFs, ^{87,88} hollow spherical COFs, ⁸⁹ and macro/microporous COFs.36 Such hierarchical porous systems, showing large surface areas, good mass transport and distribution of active sites, should facilitate applications in electrocatalysis, supercapacitors, batteries, etc.⁹⁰

c. Hybrid nanostructures. For electrochemical applications, the low electric conductivity of COFs is one of the major drawbacks. Considering the presence of active sites in the COF backbone, a certain charge-transport pathway within the COF matrix is necessary for good electrocatalytic activities. Generally, the intrinsic bulk conductivity of COFs can be

leveraged by through-bond and through-space charge transport characteristics by applying proper building units and synthetic routes such as full annulation of building blocks through aromatic linkages and maximizing orbital overlap through a strategic choice of building blocks.⁹¹⁻⁹⁴ Nonetheless, the electrical conductivities of COFs are still inferior to those of other typical electrode materials. Intriguingly, formation of COF hybrid materials promise to not only yield morphology control but also to circumvent the conductivity problem, especially when conducting materials such as carbon nanotubes (CNTs), graphene and conductive polymers are applied as a second compound in the composite (Fig. 4c).^{74,95,96} To ensure close contact between COFs and these conducting materials, direct growth of COFs on the substrate as crystalline thin films seems to be the way to go for achieving good electric conductivity and mechanical stability.

d. Terracotta process. Considering the polycrystalline nature of COFs, the processing of COF powders into desired geometrical shapes without using any binder or plasticizer is very difficult. To process the COFs into definite shapes and induce the hierarchical

Fig. 4 (a–e) Strategies for the synthesis of COFs through macroscopically controlled engineering. (f) Types of topologically distinct COF structures. Reproduced from ref. 97 (4d) and ref. 99 (4e) with permission from American Chemical Society, copyright 2017 and copyright 2019.

porosity in the COF matrix, recently, a so-called Terracotta process has been employed.⁹⁷ In this method, grinding of diamine linkers together with p-toluenesulfonic acid (PTSA), water and multitopic aldehydes, followed by heating for a few minutes yielded β ketoenamine COFs in large quantities and with enhanced porosity. As the reagent mixture holds the consistent properties as dough, it is possible to process COFs into various shapes without any additives and binders, thus achieving the construction of different COF sculptures (Fig. 4d). By introducing the use of baking soda during the terracotta process for COF synthesis, even hierarchical porosity with an interconnected pore network has been incorporated in the COF matrix.⁹⁸

e. 3D-printing. To form COF-based macroarchitectures as well as to introduce the macro- and mesoporosity, 3D-printing

methods are another possibility (Fig. 4e). In one example, iminebased polymers were initially co-assembled with Pluronic F127 in an aqueous solution.⁹⁹ By limiting the degree of imine polycondensation, the co-assembled amorphous imine polymer and F127 were used as hydrogels for 3D-printing. Subsequent removal of F127 followed by an amorphous-to-crystalline transformation resulted in the formation of 3D COF monoliths possessing hierarchical porosities and robust mechanical stability. Similarly, graphene oxide (GO) was chosen as a foaming agent, successfully engineering the crystalline COF foams into 3D-printed geometrical forms, for efficient mass transport and adsorption.¹⁰⁰

2.2.3 Construction of COF-based electrodes. To achieve high electrochemical performance, it is critical to construct efficient and durable COF-based electrodes. Since COFs are mostly prepared as the bulk powders that cannot be dissolved

or melted, generally, COF powders are physically dispersed in ethanol solution with additives acting as a binder (e.g. Nafion) and conductor $(e.g.$ activated carbons), to form a homogeneous mixed solution.36,101 Then the resulting ''ink'' can be dropped or spin-coated on different substrates such as a rotating disk electrode (RDE), carbon cloth and nickel foam to form the working electrodes for further applications.⁴⁴ Even though this method enables the construction of COF electrodes, it entails some inherent problems that have to be considered. Thus, partial aggregation or embedding of COF particles, weak interactions between the COF and the substrates or pore blockage by additives can reduce the accessibility of active sites within COFs for substrates or charge carriers. COF electrodes produced in this way are usually compared to bare carbon or other reference electrodes and, when published, logically exhibit an enhanced performance, which is related to the COF material. However, due to the above-mentioned points this kind of electrode manufacturing does not guarantee that the potential of the COF is indeed fully exploited. Review Article

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In situ growth of COF materials on substrates can in contrast yield a uniform distribution of COF films on conductive substrates without any additives, ensuring close contact between COFs and conducting materials. The substrates can be commercial current collectors such as nickel foams, carbon cloths, FTO, ITO, etc.^{102,103} or other conductive substrates such as graphene aerogels.⁵⁷ It can be supposed that such *in situ* grown COF electrodes not only display better mechanical stability and durability than traditional ones prepared by deposition of COF inks, but also exhibit more open and regular pore structures, providing sufficient active sites for electrochemical applications. Furthermore, strategies controlling the COF morphologies at a macroscopic level enable the construction of COFs as selfstanding electrodes. For example, COF membranes have been suggested as flexible electrodes in wearable electronic devices.⁷⁷ On the other hand, considering the low conductivity of selfstanding COF electrodes, the conducting materials (e.g. graphene, $CNTs)^{100}$ can be mixed and processed via 3D printing and Terracotta processes for constructing COF architectures with desirable shapes, sizes and conductivities.

2.2.4 Pyrolysis of COFs to carbon materials. Beside metals, carbonaceous materials are the most used materials in energy conversion and storage because of their large surface area and pore volume, high electrical conductivity and good chemical stability.^{7,8,104-107} A plethora of organic precursors have been applied to synthesize carbonaceous electrode materials. As they are able to create high surface areas or introduce heteroatoms into the carbon materials, such precursors should obviously be (extremely) cheap, easily available and abundant for applications in real-life, i.e., large-scale devices. COFs are hardly fulfilling these latter requirements and seem therefore not to be ideal precursors for carbon materials. Nevertheless, crystalline porous materials have received increasing interest recently as carbon precursors, mainly because it was claimed that their crystalline structure with organized carbon sheets or atoms and metal species in well-defined frameworks could be a promising platform to guide the carbonization process.¹⁰⁸⁻¹¹¹

In this respect, COFs as two dimensional carbon-rich materials with abundant heteroatoms might be used as precursors for heteroatom-doped carbons.^{110,112-115} Furthermore, COFs with implemented coordination sites (e.g. bipyridine and porphyrin structures) for metal ions allow the construction of metal incorporated/heteroatom-doped carbons. The periodical feature of COFs with distributed and separated metal coordination sites is a possible pathway to create single atom catalysts in the final carbon materials.

As the structure of COFs usually collapses when carbonized, which seriously questions the often made statement, that the crystalline structure acts as a scaffold for the carbon materials, templates are normally required for the generation of porous carbonaceous materials from COFs. $110,116$ Indeed, the question can be asked, why should the advantages of a highly ordered material with precise chemical structure be sacrificed to form materials with rather ill-defined structures and unknown active sites, especially if related carbonaceous materials can be prepared from much cheaper precursors. However, it has been shown a number of times that carbon materials prepared from COFs show superior performances compared to carbons prepared from the respective monomers.¹¹⁰ Thus, it seems that the periodic structure of COFs indeed brings benefits as a carbon precursor. In this review, carbon materials from COFs will just be discussed if such a beneficial effect of the defined COF structure on the later carbon materials has been proven.

3. Applications of COF-based materials

As mentioned in the introduction section, COFs exhibit unique properties including large surface areas, ultrahigh porosity, tuneable pore sizes, a relatively high chemical/thermal stability combined with a tailorable architecture. Because of these appealing properties, in recent years, vast research has been witnessed on COF-based materials for various applications, with an increasing interest in their utilization for electrocatalysis as well as electrochemical energy storage.¹¹⁷⁻¹¹⁹ Early reports of COF materials for electrochemical applications as well as some notable performances of later reported COFs in the different electrochemical applications discussed in this review are summarized in Table 1. This review intends to timely summarize the recent progress in this field and to act as an inspiration for further development and utilization of COF materials in electrochemical applications.

3.1 Energy conversion and electrocatalysis

The transformation of energy from one form (chemical energy, heat, electricity, light, motion, etc.) to another is called energy conversion. In history, many devices and systems have been developed to achieve energy conversion. Very early examples of energy conversion systems are applied in windmills, where the kinetic energy of the wind was transformed into mechanical energy for pumping water or grinding grains. Nowadays, windmills are widely disseminated to convert wind into electrical energy, which is the most needed form of energy for our communities.

Indeed, wind and sun energy are the most sustainable sources of energy and consequently their contribution to the energy mix is rising constantly, while resources like fossil or nuclear fuels are slowly but steadily depleting. However, in direct energy conversion devices, such as the named windmills or solar cells, the electrical energy produced must be stored so that it can be used independently from peak wind and sun hours. Electricity can be stored in electrical devices such as batteries or supercapacitors, but can also be converted into chemical energy, for example in an electrolyser, by using the electrical energy to split water into hydrogen and oxygen as storable chemical fuels. When needed, these fuels can be converted back into electrical energy using fuel cells. For all these devices and the processes occurring therein the breaking and formation of chemical bonds take place, which can be achieved by choosing the right electrode materials and electrocatalysts.

Herein, the applications of COFs as electrocatalysts and/or electrode materials in energy-conversion technologies are summarized, highlighting the advantages, limitations and current status of these materials for such applications.

3.1.1 Hydrogen evolution reaction (HER). As one of the most promising renewable energy sources, hydrogen possesses the highest gravimetric energy density compared to other fuels and produces only water as a by-product when used in fuel cells.³ Currently, the world's leading automotive manufactures are engaged in the development of fuel cell vehicles (FCVs) for a sustainable future mobility. It can be expected that the demand for hydrogen will increase sharply, if such hydrogen-powered vehicles enter the market to a large extent. To enable large-scale $H₂$ production in an environmentally-friendly way, water splitting has become an efficient process as the existing methods (e.g. steam reforming) used for hydrogen production are also produce large amounts of carbon dioxide as byproduct.

Water splitting is composed of two half reactions, namely the hydrogen evolution and oxygen evolution reactions, which in many cases are investigated and optimized separately. In principal, a thermodynamic voltage of 1.23 V (25 \degree C) is required to split water into hydrogen and oxygen. However, due to the existing resistances (e.g. solution and contact resistances) and the intrinsic activation barriers present on electrodes, a much higher voltage than the thermodynamic potential is generally required. The excess potential needed is known as the overpotential. The reduction of this overpotential by applying efficient and stable catalysts is the central issue to achieving effective water splitting. Apart from the overpotential, the

indicators to appraise the performance of a given HER electrocatalyst can be readily described by current density, Tafel slope, stability and faradaic efficiency. (i) The current density is defined to determine the electrocatalytic activity of a catalyst. Currents are measured at different applied voltages and normalized to the sample mass (mA mg^{-1}) or geometric electrode area (mA cm^{-2}). In general, the specific overpotential at a current density of 10 mA cm^{-2} (which is the current density expected for a 12.3% efficient solar water-splitting device) in HER is deliberately provided to compare the activities between the electrocatalysts. (ii) The Tafel plot depicts the relationship between current density (j) and overpotential (η) , and the Tafel equation $(\eta = b \log(j) + a)$ is deduced to describe the linear portion of the Tafel plot where the linear slope b is defined as the Tafel slope and the values of b indicate the different catalytic mechanisms for the electrode reaction (Table 2). (iii) Stability testing is conducted to evaluate the practical applicability of a HER catalyst. Two methods are used to characterize the electrocatalytic stability, namely chronoamperometry and chronopotentiometry. (iv) Faradaic efficiency is calculated from the ratio of the experimental hydrogen production to the theoretical hydrogen production, 5 describing the charge efficiency by which electrons are utilized to form hydrogen. Open Soc Rev

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The H_2 evolution process is a multistep reaction on the catalyst surface involving adsorption, reduction and desorption processes of protons/water, dihydrogen and the formed intermediates. As shown in Table 2, the mechanism of HER includes three steps in both acidic and alkaline solutions. The ''Volmer step'' indicates the hydrogen adsorption on the catalyst surface (H*, where * is defined as an active site on the catalyst surface) during the reduction of protons. Molecular hydrogen is then formed through the combination of H^* via either a Heyrovsky or Tafel step. DFT calculations show that the free energy of hydrogen adsorption (ΔG_{H*}) could be a suitable descriptor for the reaction kinetics of HER, where a nearly zero $\Delta G_{\rm H*}$ indicates an optimal HER catalyst. Based on theoretical and experimental studies, the most efficient electrocatalysts for HER are platinum-based materials. Nevertheless, the price and scarcity of platinum hinders its widespread and practical use. Thus, it is quite desirable to develop efficient and affordable HER electrocatalysts composed of earth-abundant elements to achieve costefficient hydrogen production.¹³³⁻¹³⁵ For example, transition metal dichalcogenides,¹³⁶⁻¹³⁸ carbon-based materials,^{139,140} single metal atom-doped materials $141-143$ and polymeric m aterials^{144,145} have been widely studied and applied as candidates for H_2 production. In this section, recent progress in the design of COF-based HER electrocatalysts is described.

where * is defined as an active site on the catalyst surface.

As discussed above, COFs provide several opportunities to design new HER catalysts. However, the intrinsic properties of polymeric materials, *i.e.* the low conductivity and the moderate stability of COFs in the electrocatalysis process, must also be considered. Starting with an important contribution from Pradhan and co-workers, a crystalline 2D COF (SB-PORPy) was exploited for hydrogen production. The COF was formed through Schiff based condensation between 5,10,15,20-tetrakis- (4-aminophenyl)porphyrin (TAP) and 1,3,6,8-tetrakis(4-formylphenyl)pyrene (TFFPy) (Fig. 5a and b).¹⁰¹ The synthesized COF with a surface area of 869 $\mathrm{m^2~g^{-1}}$ (Fig. 5c) was mechanically ground with carbon black in a 1:1 weight ratio. When investigated as a HER electrocatalyst in 0.5 M H_2SO_4 aqueous solution, the SB-PORPy COF exhibits a lower onset potential of 50 mV compared to that of Vulcan and bare glassy carbon, demonstrating its electrocatalytic activity for HER (Fig. 5d). As no metals were added or involved during synthesis, it was speculated that the imine nitrogens $(-N)=$ were protonated as free docking sites for HER, forming hyperporphyrinic structures with positive charge. On the other hand, by taking advantage of the conducting channels derived from the π electronic conjugation between pyrene and porphyrin, the H–H combination could occur with the help of other adjacent imine nitrogen sites, which indicates a Volmer–Heyrovsky mechanism on the SB-PORPy COF surface for the HER. As the first of its kind, this work demonstrated the potential application of COFs for hydrogen production, even though carbon Review Article

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black was applied as a conductive additive to promote the conductivity of the system.

Inspired by the catalytic activities of porphyrin and metalloporphyrins in biological systems and chemical catalysis,^{146,147} similarly, Bhaumik's group reported a metal-free porphyrinbased microporous organic polymer (TpPAM) as a HER electrocatalyst for hydrogen production.¹⁴⁴ Even though the as-synthesized covalent organic polymer was found to be amorphous, its HER performance underlined the potential applications of porphyrinbased polymers for HER catalysis, with a low overpotential $(250 \text{ mV at 10 mA cm}^{-2})$ and a small Tafel slope $(106 \text{ mV dev}^{-1}).$ However, based on DFT calculations, the authors concluded that the origin of HER activity of TpPAM was derived from the porphyrin cores, which is contradictory to the above explanation. Thus there is certainly an interest to create more COF-based electrocatalysts with varying molecular architectures to clarify the HER mechanism of such metal-free systems.

Furthermore, metal atoms have been incorporated into porphyrin-based networks to promote the performance for HER. For example, a cobalt porphyrin-based COF was reported by Villagrán and co-workers for electrocatalytic hydrogen generation in strong acidic media.¹⁴⁸ The crystalline polymer exhibits a large surface area and its catalytic HER performance (deposited on FTO glass slides by using silver glue) can be ascribed to the existence of porous channels and exposed active sites derived from metalloporphyrins. In addition, the enhanced HER performance in comparison to the cobalt porphyrin monomer gives

Fig. 5 (a) Topological structure of SB-PORPy COF; (b) simulated structure of SB-PORPy COF; (c) N₂ sorption isotherm at 77 K of SB-PORPy COF; (d) linear sweep voltammetry (LSV) profile of the SB-PORPy COF electrode compared to those of the bare glassy carbon electrode (GCE) and Vulcan. Reproduced from ref. 101 with permission from the American Chemical Society, copyright 2017.

strong evidence that the construction of an extended conjugated system can promote the electrocatalytic efficiency and confirms the role of crystallinity and well-ordered channels present in COFs for enhancing the HER performance.

Furthermore, COFs can be used as support to immobilize and stabilize small nanoparticles with HER catalytic activity.^{127,149} Covalent triazine frameworks (CTFs) are a chemically stable class of COFs synthesized using triazine (C_3N_3) units as crosslinkers.^{58,127,150–152} Hu and co-workers reported a CTF@MoS₂ composite by in situ growing $MoS₂$ nanoparticles into the 1D channel arrays of CTFs (Fig. 6a).¹²⁷ The influence of the amount of $MoS₂$ was studied, and it was demonstrated that the CTF@MoS₂-5 (where 5 represents the mass ratio of $CTFs:MoS₂$) exhibited catalytic kinetics with an overpotential of 93 mV at 10 mA cm^{-2} , which is close to that of the benchmark catalyst (20% Pt/C) and lower than those of the pure CTFs or $MoS₂$ and CTFs@MoS₂-X with different mass ratios (Fig. 6b). It was suggested that the crystalline 1D channel arrays formed by the stacking of 2D in-plane porous layers could promote the electron transmission, mass diffusion as well as the accessibility of $MoS₂$ active sites in the H_2 evolution process (Fig. 6c). While these are promising examples for COF-based, non-noble metal or even metal-free HER catalysts, the so far presented electrocatalytic performances are still lower than those for Pt-based systems but also for many other non-noble metal HER catalysts. So far, the poor conductivity of COFs and their lower stability, especially under a strong electrolyte environment seem to be the major bottlenecks to achieving practical application for large-scale H_2 production. Future work should therefore also concentrate on the development of COFs with enhanced stability and new approaches to ensure an intimate contact with conducting supports to promote the HER performance of COF-based electrocatalysts. Chem Soc Rev

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3.1.2 Oxygen evolution reaction (OER). The other half reaction in water splitting, the OER, involves a 4e⁻ transfer step and suffers from sluggish kinetics and a high overpotential, thus is actually considered as the main bottleneck of overall water splitting and consequently sustainable hydrogen production. The proposed mechanisms of OER include several intermediate species (OH $*$, OOH $*$ and O $*$) that are produced in the process of OER and subsequently react to give O_2 (Table 3).¹⁵³ Their binding energies exhibit a strong correlation, and thus form a scaling relation between OOH* and OH*. As a result, in OER more critical steps have to be considered than in HER either in acidic or alkaline solution, involving the breaking of O–H bonds and the subsequent formation of $O=O$ double bonds.

Similar to HER, indicators such as overpotential, Tafel slope, faradaic efficiency and stability are normally applied for elucidating the electrocatalytic performance of a given OER catalyst, which are also important to gain insight into the OER mechanism.⁴⁷ For example, the overpotential at a catalytic current density of 10 mA $\rm cm^{-2}$ is commonly compared to judge the performance of different electrocatalysts. As for HER, the Tafel slope is another crucial indicator for an OER catalyst in characterizing the relationship between current density and overpotential. Typically, a smaller Tafel slope is expected to imply good electrocatalytic kinetics of an OER catalyst, which means that the current density can increase faster with a smaller change in overpotential. As OER is operated at a high applied voltage (V), the chemical stability is another important factor to evaluate the degradation of OER catalysts. Descriptors such as the chemical composition, crystallinity, micro- and macro-scale morphology and of course electrochemical performance over time are usually considered to evaluate the stability of electrocatalysts. Currently, noble metal oxides

Fig. 6 (a) The formation process of CTF@MoS₂; (b) polarization curve of CTF@MoS₂ for HER catalysis in 0.5 M H₂SO₄, compared to those of other materials such as Pt/C (green), pure CTFs (black), MoS₂ (rose red) and CTF@MoS₂-X with different mass ratios; (c) schematic structure of crystal 1D channel arrays derived from 2D in-plane porous layer stacking. Reproduced from ref. 127 with permission from John Wiley and Sons, copyright 2019.

(e.g. $RuO₂$ and IrO₂) are still considered as the best OER catalysts, even though they have drawbacks such as high-cost, and scarcity. In this regard, many efforts have been devoted to search for active and low-cost alternatives, including transition metal oxides and their derivatives,¹⁵⁴⁻¹⁵⁸ carbon-based materials^{159,160} and single-atom systems.^{161,162}

To gain a better understanding of the mechanism of this multielectron reaction, OER catalysts with well-defined active sites and structures are preferentially designed, making COFs an interesting option. Probably the most often used approach is the incorporation of metal-coordinating units (e.g. bipyridine and porphyrin structures) into the COF skeletons to finally achieve the immobilization of metal-complexes within the framework. Such metalated COFs might be promising OER catalysts by taking advantage of a high amount of accessible metal active sites on the high surface areas and the masstransfer channels provided by the pores. For example, Banerjee's group synthesized a bipyridine-COF by reacting 1,3,5-trifomylphloroglucinol (Tp) and 2,2'-bipyridine-5,5'-diamine (Bpy) via

the Schiff base reaction.⁵⁵ Bipyridine moieties were used for cobalt ion coordination, yielding abundant Co–N active sites, mimicking metal-bipyridine molecular systems (Fig. 7a).^{163,164} The obtained bipyridine-COF with Co incorporation (Co-TpBpy) possesses an accessible surface area of \sim 450 $\mathrm{m}^2\,\mathrm{g}^{-1}$ (Fig. 7b) and a reasonable OER performance with an overpotential of 400 mV at 1 mA cm^{-2} (a calculated overpotential of \sim 520 mV at 10 mA cm-2) under neutral pH conditions (Fig. 7c). In addition, Co-TpBpy exhibits good stability with a similar polarization curve retention after 1000 scans from 0.6 V to 1.8 V (vs. RHE). It should be noted that acetylene black was applied as a conductive additive to enhance the conductivity of the system with a mass ratio of 9 : 1 for Co-TpBpy to acetylene black.

Metal porphyrins with their four-coordinated (metal– N_4) structures were one of the first molecules tested for noble metal-free OER.^{165,166} In particular, the porphyrin structure with high/low-valent metal ion incorporation was preferentially designed for oxidizing/reducing catalytic cycles.^{167,168} Consequently, porphyrin-based COFs have been designed as

Fig. 7 (a) Schematic illustration of TpBpy and Co-TpBpy with Co incorporation; (b) N₂ sorption isotherms of TpBpy (black) and Co-TpBpy (red); (c) LSV profiles of Co-TpBpy before and after 1000 cycles from 0.6 V to 1.8 V in a phosphate buffer (pH 7.0). Inset: An enlarged view of the LSV profile Reproduced from ref. 55 with permission from the American Chemical Society, copyright 2016.

an OER electrocatalyst. For instance, Chen's group described a metalloporphyrin-based graphdiyne analogue (Co-PDY) by using Co-coordinated 5,10,15,20-(tetra-4-ethynylphenyl)porphyrin (Co-TEPP) as the monomer to fabricate a novel graphdiyne analogue (Fig. 8a).126 Notably, graphdiyne is composed of two acetylene (butadiyne) linkages ($-c \equiv c-c \equiv c$) between the aromatic rings and possesses a unique electronic structure.^{169,170} Consequently, the formed Co-PDY, even though just low crystallinity was reported, shows a uniform distribution of $Co-N₄$ moieties, serving as active sites for OER. Meanwhile, the conjugated structure of porphyrin and the butadiyne linkage could promote the electron transfer in the OER process, achieving the consecutive activity. To further improve the OER electrocatalytic activity and stability, copper foam (CF) was applied as a substrate for polymer growth, serving as a robust 3D porous conductive network (Fig. 8b and c). When used as a working electrode in an alkaline electrolyte (1.0 M KOH), Co-PDY/CF provided an enhanced OER activity with an overpotential of 270 mV at 10 mA cm^{-2} , in comparison to those of bare CF substrate

(442 mV) and PDY/CF (381 mV) (Fig. 8d). A chronopotentiometry measurement was further performed to confirm the chemical stability of Co-PDY/CF for OER (Fig. 8e).

Mass transport (of electrolyte molecules, ions, gas molecules etc.) is a crucial factor for any electrocatalytic process. The micro- or small mesopores of COFs on one hand increase their surface area but can be too small to ensure unhindered mass transport throughout the pore system. The construction of hierarchical porosities, which is the introduction of additional mesopores or macropores into COFs is supposed to be an effective way to facilitate mass transport through COFs. COFs with hierarchical pore structures have been prepared by applying polystyrene spheres (PSs) as a hard template (Fig. 8f).³⁶ The macroporous COFs preserve their crystalline structure and high surface area (Fig. 8g and h). The benefit of additional macroporosity was proven by applying macro/microporous co-coordinated bipyridine-COFs (macro-TpBpy-Co) as OER catalysts. A low overpotential (380 mV) was required to reach a current density of 10 mA cm^{-2} , which is superior to that of a

Fig. 8 (a) Schematic representation of the synthesis of Co-PDY. (b) SEM image of Co-PDY/CF, inset: SEM image of copper foam (CF); (c) TEM image of Co-PDY/CF; (d) LSV profiles of Co-PDY/CF, PDY/CF, and CF; (e) chronopotentiometry curve of Co-PDY/CF for OER at 1.50 V (vs. RHE); (f) schematic illustration of the synthesis of macro-TpBpy COF with macroporous structure; (g) SEM image of macro-TpBpy; (h) N₂ sorption isotherms of macro-TpBpy and TpBpy; (i) polarization curves of macro-TpBpy-Co. Reproduced from ref. 126 with permission from The Royal Society of Chemistry, copyright 2019; and ref. 36 with permission from the American Chemical Society, copyright 2019.

COF without hierarchical porosity (TpBpy-Co) (Fig. 8i), thus demonstrating the improvement regarding mass diffusion and accessibility of active sites in the electrocatalytic process.

3.1.3 Oxygen reduction reaction (ORR). Oxygen reduction is a fundamental reaction related to the energy conversion devices including fuel cells and metal–air batteries. As for OER, the oxygen reduction process is composed of multistep reactions, involving multiple adsorbed intermediate species such as O^* , OH^* and OOH^* .¹⁷¹ Depending on the number of electrons transferred, the oxygen reduction process could be divided into two pathways: the direct $4e^-$ transfer pathway from O_2 to H₂O; and the 2e⁻ transfer pathway from O_2 to H₂O₂ (Table 4).

Typically, to obtain a high efficiency in the oxygen reduction process, the 4e⁻ transfer pathway is required. For an ORR catalyst, the selectivity of the $2e^-$ or $4e^-$ pathway is determined by the adsorption energetics of the aforementioned intermediates $(O^*, OH^*$ and $OOH^*)$ and the reaction barriers on the catalyst surface. Currently, the most suitable ORR electrocatalysts are based on the precious Pt metal and its alloys, as evidenced by the volcano plot that counts experimental ORR activity as a function of hydroxyl binding energy.^{46,172} However, also many non-noble metal ORR electrocatalysts, including carbon-based catalysts,^{173,174} MOFs and their derivatives,^{175,176} as well as transition metal oxides 177 have been reported to promote the ORR performance.

To evaluate the performance of ORR catalysts, the most frequently used technique is steady-state polarization on a rotating disk electrode (RDE) or a rotating ring-disk electrode (RRDE). Typically, on an RDE, the indicators such as onset potential, half-wave potential and limiting current density could be calculated as the standard criteria to evaluate the ORR performance for different electrocatalysts. Besides, the total electron transfer number (n) is another important indicator to demonstrate the progress of ORR, which can be calculated by the Koutecky–Levich (K–L) equation on an RDE or directly measured on an RRDE through the equation.

It is known that a 2e⁻ transfer pathway of ORR can produce H_2O_2 (in acidic medium) or HO_2 ⁻ (in alkaline medium), whereas it is still unclear whether to define it as a key intermediate of the dominant ORR mechanism or a side product.⁴⁶ It is essential to further study the interaction of intermediates with catalyst surfaces to understand the ORR mechanism and develop advanced catalysts to promote the ORR performance. Currently, COFs in their pristine state exhibit promising potential for ORR.178–180 In particular, inspired by the naturally available biocatalysts such as cytochrome c oxidase in mitochondria, which contain as active center an iron (n) -porphyrin structure serving as the ''ORR'' catalyst in cellular organelles, COFs have been prepared using macrocyclic clusters such as porphyrin, phthalocyanine and tetraazannulene. This provides a biomimetic strategy for designing ORR catalysts by mimicking their homogeneous metal-complex counterparts.¹⁸¹ The activity of metalloporphyrin COFs for ORR has been evaluated by DFT calculations. Similar to the results of metalloporphyrin COFs for OER, metal–porphyrin structures with Co, Rh and Ir incorporation could be potential candidates for ORR catalysis considering their intermediate free energies close to those of an ideal catalyst.⁷⁵

Recently, Xiang's group synthesized a 2D COF material (COF_{BTC}) where the phthalocyanine iron centres were arranged into rigid networks and endowed the COF_{BTC} with positively charged centres (Fig. 9a).¹²⁵ The XRD pattern confirms the crystal structure of the as-synthesized COF_{BTC} (Fig. 9b). Notably, at alkaline conditions, the COF_{BTC} layers can be exfoliated and solvated to form a stable dispersion, which is ascribed to the insertion and absorption of negative hydroxyl groups onto the positively charged centres within the COF_{BTC} stacks (Fig. 9c). When applied for ORR with carbon paper as the working electrode in 0.1 M KOH aqueous electrolyte, the COF_{BTC} catalyst with ordered N-coordinated Fe centres and conjugated structures provides a half-wave potential of 0.9 V vs. RHE and an onset potential of 0.965 V vs. RHE at a COF_{BTC} concentration of 1.25 g mL⁻¹ (Fig. 9d), which compare well to those of some reported metal–nitrogen–carbon based catalysts.¹⁸² When further applied in a zinc-air battery with O_2/OH^- and Zn^{2+}/Zn as redox couples, this COF can be directly used as the catalyst for ORR, due to its solubility in the electrolyte. As such, the COF_{BTC} could provide more accessible active sites compared with the conventional air electrodes such as catalyst-coated carbon paper. Expectedly, the COF_{BTC} catalyst exhibits practical applications in zinc–air flow batteries with voltage gaps (0.77 V at 5 mA $\rm cm^{-2}$, 0.83 V at 10 mA $\rm cm^{-2}$, and 0.96 V at 20 mA cm^{-2}) and high energy efficiencies (60.7% at 5 mA cm⁻², 59.0% at 10 mA cm⁻², and 52.6% at 20 mA cm⁻²) (Fig. 9e and f). Review Article

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> To improve the electron transport during the electrocatalytic process, the hybridization of COFs with conducting additives such as graphene and CNTs has been achieved by in situ polymerization.95,183 For instance, Xiang's group reported a hybridized covalent organic polymer/graphene (COP/rGO) by self-assembling highly ordered porphyrin COPs with graphene.⁹⁵

Fig. 9 (a) Schematic representation of the synthesis process for COF_{BTC}; (b) XRD pattern of COF_{BTC}; (c) schematic illustration of the exfoliation and dissolution processes of COF_{BTC} in an alkaline solution; (d) LSV profile of the COF_{BTC} in O₂-saturated 0.1 M KOH. Inset: Scheme for the detailed operation of the measurement with carbon paper as electrodes. (e) Schematic configuration of rechargeable Zn–air batteries; (f) galvanostatic discharge–charge cycling curves of Zn–air batteries at different current densities of 5, 10, and 20 mA cm $^{-2}$. Reproduced from ref. 125 with permission from the American Chemical Society, copyright 2019.

As a result, the conductivity showed a significant increase (from 3.06×10^{-9} to 2.56×10^{-1} S m⁻¹) compared to pure COPs. It is assumed that the synergetic effects between active COP and high electrical conductivity of rGO are responsible for the high ORR activities and long-term stability.

COF materials have been recently used as precursors to construct carbon-based electrocatalysts for ORR, such as metal-free carbon,¹¹² single-atom doped carbon¹¹⁰ as well as metal–carbon composites.184–187 Nevertheless, the direct pyrolysis of COFs destroys their inherent porous structure forming bulk carbons with hardly controllable structures. With this in consideration, Jiang's group applied phytic acid (PA) as a template to guide the pyrolysis, which successfully converted a traditional COF (TAPT-DHTA-COF) into nanosized 2D carbons (Fig. 10a).¹¹² It was demonstrated that PA can enter the 1D channels of COFs, acting as templating and directing agents during the pyrolysis process of COFs at 1000 $^{\circ}$ C under N₂ (Fig. 10b). As a result, the resulting 2D COF (PA@TAPT–DHTA– COF₁₀₀₀) possesses good conductivity, hierarchical porosity and abundant catalytic edges with phosphorus and nitrogen doping.

Later on, the PA@TAPT-DHTA-COF₁₀₀₀ was pyrolyzed under $NH₃$ at 900 \degree C for 0.5 h, to further improve its specific surface area (1160 m^2 g⁻¹), pore volume (0.59 m^3 g⁻¹) and ORR catalytic activity (Fig. 10c and d). PA@TAPT-DHTA- $\text{COF}_{1000\text{NH}}3$ as a metal-free ORR electrocatalyst achieved a half-wave potential of 0.831 V (vs. RHE), an onset potential of 0.941 V (vs. RHE), and a limiting current density of 7.2 mA cm^{-2} , superior to those of Pt/C (Fig. 10e).

Recently, our group applied silica nanoparticles as a hard template to convert a bipyridine-COF (TpBpy) into mesoporous carbon via carbonization.¹¹⁰ A molecular organizer (p-toluenesulfonic acid, PTSA) was used for COF growth and enabled the formation of $SiO₂(Q)$ TpBpy composites via a grinding approach. Bipyridine moieties (2,2'-bipyridine-5,5'-diamine, Bpy) were chosen and incorporated within the COF backbone to coordinate metal, here Fe ions. When taken as a carbon precursor with a pyrolysis process followed by suitable treatment with HCl and NaOH, the bipyridine-COF with Fe incorporation was successfully converted into mesoporous carbon (mC-TpBpy-Fe) with a surface area of 362 $m^2 g^{-1}$ and abundant Fe–N_x active sites.

Fig. 10 (a) Structure of TAPT-DHTA-COF; (b) PA-templated synthesis of PA@TAPT–DHTA–COF; (c) N2 sorption isotherms of PA@TAPT–DHTA– COF_{1000NH3} (red), PA@TAPT-DHTA-COF₁₀₀₀ (blue) and TAPT-DHTA-COF₁₀₀₀ (black); (d) pore size distribution of PA@TAPT-DHTA-COF_{1000NH3}; (e) ORR performances of PA@TAPT–DHTA–COF_{1000NH3} (red), PA@TAPT–DHTA–COF₁₀₀₀ (blue), TAPT–DHTA–COF₁₀₀₀ (green), and Pt/C (black) in 0.1 M O₂-saturated KOH aqueous solutions at 1600 rpm. Reproduced from ref. 112 with permission from John Wiley and Sons, copyright 2018

Electrocatalytic studies show that mC-TpBpy-Fe exhibits an onset potential of 0.92 V (vs. RHE) and half-wave potential of 0.845 V (vs. RHE), which are comparable to those of commercial Pt/C in oxygen-saturated 0.1 M KOH. In addition, the performance was superior to carbonaceous materials derived from the carbonisation of the pure monomers or an amorphous polymer network with similar chemical structure. It can thus be suggested that the periodicity of the COF precursor yield a more homogenous distribution of functional groups within the carbon framework, i.e. that the formation of a crystalline COF precursor is indeed beneficial for the preparation of highly active and porous Fe–N–C catalysts.

3.1.4 Bifunctional OER/ORR catalysts. As mentioned above, OER and ORR are key processes in renewable energy conversion. To design efficient bifunctional electrocatalysts both for ORR and OER is of significance in the commercialization of renewable energy devices such as Zn–air batteries. Typically, the rechargeable Zn–air battery consists of a Zn anode, a separator soaked in an aqueous alkaline electrolyte and an air cathode. In a discharge process, the Zn anode is oxidized, while $O₂$ from air diffuses through the cathode electrode and is then reduced by electrons to form OH⁻. This process reverses during charging, where the Zn recovers at the anode electrode and $O₂$ is regenerated, evolving at the cathode electrode. For a rechargeable battery, indicators including discharge/charge potentials, power density and galvanostatic discharge–charge cycling are generally considered to evaluate its performance such as rechargeability and energy efficiency.

Recently, many research efforts have been dedicated to the development of efficient bifunctional electrocatalysts for the OER and ORR.188,189 Based on density functional theory (DFT) calculations, Promarak and co-workers studied the free energy changes of intermediates both for ORR and OER on metalloporphyrins in a 2D framework.⁷⁵ The metalloporphyrin frameworks with 14 different metal sites were applied to investigate the free energies of the reaction intermediates, including OH*, OOH^* and O^* . As a result, especially the metalloporphyrin frameworks with Co, Ir and Rh incorporation could be potential candidates for both the OER and ORR. In this regard, Zhang's group reported a porphyrin framework (''POF'') through dehydration polymerization between pyrrole and benzene-1,4 dialdehyde (BDA) (Fig. 11a).¹⁹⁰ Molecular dynamic (MD) simulations and XRD measurements were performed to confirm that the constructed POFs possess a slightly ordered structures distinguished from those of fully amorphous polymers. After adding cobalt ions, the Co-coordinated POF exhibited electrocatalytic performances both for OER and ORR, demonstrating its potential application as a bifunctional electrocatalyst.

Considering the limitation of electronic transport of such crystalline materials, the authors applied carbon nanotubes (CNTs) as scaffolds to fabricate a CNT@POF hybrid material and explored its practical application in zinc–air batteries. It was found that the POF was uniformly coated on CNTs as thin layers with a calculated thickness of \sim 4 nm, thus fully exposing the porphyrin active sites to reactants (Fig. 11b).⁷⁴ When applied as the cathode in a Zn–air battery (Fig. 11c), the CNT@POF hybrid exhibits a superior current density than that of a reference $Pt/C + Ir/C$ cathode, while the discharge and charge potentials for CNT@POF are 1.21 V and 1.99 V at a required current density of 20.0 mA $\rm cm^{-2}$ (0.95 V and 2.02 V for

Fig. 11 (a) Schematic illustration of the synthesis of POF; (b) Photographs and SEM image of CNT@POF. (Inset shows a schemtic drawing of the POF-CNT composite); (c) left: chemical structure of POF with Co incorporation. The hydrogen (white), carbon (green), nitrogen (blue) and cobalt (pink) atoms are marked in different colours. Right: Schematic illustration of a flexible Zn-air battery; (d) polarization curves of CNT@POF in the rechargeable Zn-air battery; (e) charge/discharge cycling of CNT@POF and Pt/C + Ir/C at 2.0 mA cm⁻². Reproduced from ref. 190 with permission from John Wiley and Sons, copyright 2019; and ref. 74 with permission from the Royal Society of Chemistry, copyright 2018.

Pt/C + Ir/C) (Fig. 11d). The peak power density of CNT@POF (237 mW cm^{-2}) is much higher than that of the reference Pt/C + Ir/C cathode (78 mW cm^{-2}). Besides, the galvanostatic discharge–charge cycling curves were conducted at 2.0 mA $\rm cm^{-2}$ to investigate the cycling performance of CNT@POF in a rechargeable Zn–air battery, where the CNT@POF hybrid shows a smaller voltage gap of 0.71 V than that of $Pt/C + Ir/C$ (0.77 V), demonstrating its impressive performance in rechargeable flexible Zn–air batteries (Fig. 11e).

3.1.5 Carbon dioxide reduction reaction (CO_2RR) . Carbon dioxide $(CO₂)$ emission from the utilization of fossil fuels constitutes a potential threat to the global climate. Transforming $CO₂$ into a valuable carbon feedstock is therefore an important goal for a sustainable economy.191–193 However, because of the competitive side reactions (for example: HER) and the stable $C=O$ double bond, the $CO₂RR$ is kinetically challenging. The reaction process of $CO₂RR$ mainly involves three steps: (1) adsorption of $CO₂$ molecules on a catalyst surface; (2) the breaking of C–O bonds and/or the forming of C–H bonds; (3) the rearrangement and desorption processes of products. In particular, the second step is a complex process with multielectron transfer pathways producing different products. Depending on the number of migrated protons and electrons, $CO₂$ can be converted into different products, including formic acid, methane, methanol, ethanol, ethylene, carbon monoxide and many others.¹⁹¹ In most cases, the products are multiple species rather than a single product. The type and amount of species mainly depend on the surface properties of electrocatalysts and the applied electrode potentials. Consequently, electrocatalysts with various sizes, morphologies, crystal structures and chemical states have been designed to increase the $CO₂RR$ kinetics and selectivity.194,195 To evaluate the performance of a CO2RR electrocatalyst, the indicators generally include onset potential, overpotential and current density. Besides, due to the formation of multiple products in the $CO₂RR$ process, the

faradaic efficiency is required to evaluate the conversion efficiency and selectivity of catalysts. Typically, faradaic efficiency (FE) is defined through the following equation for a certain product:

$$
\text{FE} = \frac{\alpha nF}{Q}
$$

where α is the electron transfer number to form a certain product; F is the Faraday constant; Q is the total charge during the reduction process; n is the number of moles of the product produced.

Currently, the most widely employed electrocatalysts for $CO₂$ conversion are traditional bulk metals and metal-containing inorganic materials.^{51,196,197} However, these metal-based catalysts are generally poisoned by impurities or some strongly bound intermediates, showing limited active sites. In this regard, various advanced catalysts have been studied to further enhance CO_2RR ,^{194,198,199} and recently COFs have also been applied to this reaction.^{44,73,200,201}

Starting with an important contribution from Yaghi and coworkers, two Co-based COFs (i.e. COF-366-Co and COF-367-Co) were synthesized through the imine condensation of 5,10,15,20 tetrakis(4-aminophenyl)porphinato cobalt [Co(TAP)] with 1,4 benzenedicarboxaldehyde (BDA) and biphenyl-4,4'-dicarboxaldehyde (BPDA), respectively, serving as electrocatalysts to convert $CO₂$ into carbon monoxide (CO) (Fig. 12a).⁴³ The as-synthesized Co-COFs exhibit high crystallinity and large surface areas (1360 $\mathrm{m^2~g^{-1}}$ for COF-366-Co and 1470 $\mathrm{m^2~g^{-1}}$ for COF-367-Co), enabling abundant active sites for $CO₂RR$. The faradaic efficiency (FE $_{\rm CO}$) of the COF-366-Co is 90% at -0.67 V, showing a superior catalytic performance compared to that of the molecular cobalt complex Co-(TAP). At standard temperature and pressure (STP), COF-366-Co could yield more than 36 mL of carbon monoxide per mg of COF during a 24 hour period. When the framework is expanded through the replacement of BDA with

Fig. 12 (a) Schematic illustration of metalloporphyrin-derived COFs (COF-366-Co and COF-367-Co); (b) cyclic voltammograms (CVs) of COF-366-Co and COF-367-Co in a CO₂-or N₂-saturated medium. A bare carbon electrode was applied as the background (black solid line). (c) Long-term bulk electrolysis at 0.67 V (vs. RHE), showing the volumes of CO₂ (H₂) produced by COF-367-Co, COF-366-Co, and Co(TAP). Reproduced from ref. 43 with permission from AAAS, copyright 2015.

BPDA, the expanded COF-367-Co with larger pore size provides a greater number of active cobalt-porphyrin moieties and thus possesses an improved capacity of $CO₂$ adsorption within the framework. Consequently, the COF-367-Co exhibits enhanced catalytic performance compared to that of the COF-366-Co (Fig. 12b and c), where the faradaic efficiency of COF-367-Co is 91% and the production of carbon monoxide per milligram is more than 100 mL at -0.67 V during a 24 hour period. X-ray absorption spectra show that the COF structure is of importance in modulating the electronic properties of molecular centers. This demonstrates that because of the interaction between the COF lattice and cobalt atoms, the cobalt centers within COFs could possess a more delocalized electronic structure, which is responsible for the high activity and selectivity of the reduction process from $CO₂$ to $CO₂$. This work thus clearly proves the advantages of ordered and porous COF materials for carbon dioxide reduction.

Given the unique features such as a fully conjugated skeleton, high chemical stability and the abundant presence of heteroatoms, CTFs exhibit strong affinity to $CO₂$ and can be considered as promising candidates for CO_2RR ²⁰² Zhuang and co-workers designed a Ni-coordinated porphyrin-based covalent triazine framework (NiPor-CTF).²⁰³ The formed NiN₄ centres are uniformly dispersed within the frameworks. When applied as electrocatalyst for $CO₂RR$, the NiPor-CTF possesses a superior selectivity for CO conversion with a faradaic efficiency of $>90\%$. DFT calculations indicated that the NiN₄ active sites could decrease the kinetic energy barriers of the transition from $*CO₂$ to $*COOH$, thus improving the performance of CO2RR. Similarly, Nakanishi's group reported CTF-based materials with incorporation of 3d metal atoms (Co, Ni or Cu)

as electrocatalysts for CO_2RR ²⁰⁴ They proved that the CTFs with Co and Ni incorporation (Co-CTF and Ni-CTF) could effectively provide reductive activity from $\rm CO_2$ to $\rm CO$ at -0.5 vs. RHE. Particularly, the Ni-CTF possesses a superior faradaic efficiency of 90% at -0.8 V during CO formation, in contrast to the corresponding Ni–porphyrin (using tetraphenylporphyrin, TPP). It is evidenced by the aid of first principles calculation that because of the low-coordination environment in the Ni-CTF, the intermediates could be stabilized on the metal atoms, which means the free energy barriers for the formation of intermediate species (adsorbed COOH) on the metal atoms in CTF is lower than those on the TPP supports.

Due to the opposite properties of the strength of a linkage and its reversibility required for the crystallization process of COFs, it is a challenge to synthesize COF materials with high chemical stability in general and of course also for electrocatalysis application. In this regard, the attempts to build chemically strong COFs via keto-enol tautomerism enabled the synthetic possibilities of imine-based COFs.⁴⁰ From such an imine COF, Deng's group synthesized an amine COF (COF-300-AR, ''after reduction'') by direct reduction of the corresponding imine COF (COF-300) (Fig. 13a).¹²⁸ The amine COF maintains the crystalline structure of the parent COF-300 and exhibits a satisfactory chemical stability both in acidic and alkaline solutions. Notably, the COF-300-AR here is not the electrocatalyst but is deposited on a silver electrode to form a molecularly defined interface for the catalytic $CO₂RR$. It is demonstrated that the COF with amine linkages provides chemically sorptive sites to concentrate $CO₂$ molecules to form carbamates close to the electrode surface, and then the silver

Fig. 13 (a) Reduction process of COF-300 to COF-300-AR; (b) schematic representation of the molecularly defined interface between the COF-300- AR and silver electrode. Reproduced from ref. 128 with permission from Elsevier Inc., copyright 2018.

surface with sufficient electrons could activate the carbamates, leading to the selective generation of CO (Fig. 13b). This work provides a novel approach by using COFs to achieve the construction of ordered porous structures on the surface of silver electrodes. Forming the molecularly defined interface achieves the favourable diffusion of $CO₂$ molecules and good CO₂RR selectivity.

3.2 Energy storage

The process of capturing energy produced at one time for later use is called energy storage. Devices used for energy storage are supercapacitors, accumulators or batteries. With the discovery of electricity, there was a search for effective approaches for storing the electricity for use on demand. In the past decades, the change in renewable energy requirements and the development of nanotechnology has enabled the energy storage industry to continue to develop, adapt and transform. To build more sustainable energy infrastructures and achieve the economic utilization of electric energy, there is a need for sustainable and large-scale energy storage devices or systems.

3.2.1 Supercapacitors. Supercapacitors with high power density and long cycling life are considered as promising energy storage devices. Depending on the energy storage mechanism and the nature of the electrode material, supercapacitors exhibit two distinct processes for storing electricity: the nonfaradaic processes related to the electrochemical double layer and the faradaic processes derived from reversible redox processes on the electrode (i.e. pseudocapacitance).^{52,205,206} Supercapacitors are generally related to the electrochemical process occurring at the interface of the electrolyte and electrode. In the non-faradaic process, found in electrochemical double layer supercapacitors, the electrical energy is stored through the adsorption of ions onto electrode surfaces. Consequently, electrodes with high surface areas (e.g. porous carbons) provide maximum energy storage.^{207,208} Meanwhile, the faradaic processes in pseudocapacitors are characterised by reversible chemical reactions, i.e. oxidation/reduction reactions occurring on the electrode surfaces, which usually provide

higher supercapacitor performance in contrast to those of electric double layer formation only.²⁰⁶ Electrochemical parameters, such as capacitance, power density, energy density and cycling stability, are applied to characterize the performance of a supercapacitor. 209 The capacitance is one of the key parameters, which is defined by the ratio of the change in electrical charge to the corresponding change in electrical potential. In particular, the specific capacitance is calculated as the most significant indicator to compare the ability of charge storage for supercapacitor materials. When normalized by the surface area, volume, and mass of the active materials, the obtained specific capacitance is correspondingly defined as the areal capacitance (F cm⁻²), volumetric capacitance (F cm⁻³) and gravimetric capacitance (F g^{-1}), respectively. Similarly, the power density and energy density of a supercapacitor can be gravimetrical or volumetrical, where the power density indicates the efficiency of energy delivery and uptake by W kg⁻¹ or W m^{-3} and the energy density demonstrates the amount of stored and deliverable electrical energy by W h kg^{-1} or W h m⁻³. Cycling stability is another important parameter, which is usually estimated in the form of capacitance retention rate, which is obtained in a galvanostatic charge–discharge (GCD) test through comparison of the capacitance before and after thousands of cycles.

For electrochemical double layer supercapacitors, the most widely used electrode materials are porous carbons with high surface area because of their low cost and enormous availability, whereas the electrode materials for pseudocapacitors depend on metal oxides, redox-active polymers and functionalized porous carbons that conduct the electrostatic and pseudocapacitive processes simultaneously. Even though pseudocapacitors possess superior specific capacitance compared to electrochemical double layer supercapacitors, the fast-redox reactions in pseudocapacitors usually lead to swelling and shrinkage of the electrodes, resulting in poor mechanical stability and inaccessible active sites. In this regard, the design of advanced electrode materials with stable and controllable porous structures, high surface areas and integrated redox-active groups, is crucially required to achieve an efficient redox process in supercapacitors.

COFs can be promising electrode materials for electrochemical capacitors due to their unique advantages such as large surface areas, tuneable pore sizes and shapes as well as the adjustable redox-active groups within the frameworks. $210-217$ It can be supposed that the long-range ordered structure with high surface area could promote the ion transport at the interface of the electrode and electrolyte. In addition, the tuneable building units enable COFs to be functionalized by the incorporation of redox-active sites, suitable for pseudocapacitive energy storage.²¹⁸ In this section, the development of COF-based electrode materials applied in supercapacitors is summarized.

In 2013, Dichtel's group firstly applied COFs as electrode materials for pseudocapacitors by introducing redox-active anthraquinone moieties from 2,6-diaminoanthraquinone (DAAQ) into the crystalline framework (DAAQ–TFP COF) that was synthesized by condensing DAAQ and 1,3,5-triformylphloroglucinol (TFP).⁶³ However, due to the poor conductivity, the DAAQ–TFP COF exhibited only 40 F g^{-1} capacitance after 5000 chargedischarge cycles. To address the conductivity issues, DAAQ–TFP COF thin films were fabricated on Au substrates through the slow reaction of the TFP monomer with DAAQ (Fig. 14a and b). 219 The thin films enable many more redox-active sites to be accessible and an order of magnitude higher capacitance was obtained compared to the bulk COFs that were synthesized as randomly oriented microcrystalline powder. Thus, this work not only demonstrated the possibility to apply redox-active COFs in pseudocapacitors but also the impact of regulating the morphology and microstructure of the COFs to enhance their electrochemical performance.

Furthermore, to enhance the electrical conductivity of redoxactive COF films, conductive polymers were integrated into the ordered channels within the COF matrix. Dichtel and

co-workers applied the above DAAQ–TFP COF as a scaffold to achieve the infiltration of the conductive poly(3,4-ethylenedioxythiophene) (PEDOT) into the channels of COFs (Fig. 15a).⁹⁶ The as-fabricated PEDOT-infiltrated COF (PEDOT-modified DAAQ–TFP) exhibited greatly enhanced electrochemical activities with good accessibility of the redox-active groups within DAAQ– TFP COF. It is supposed that the PEDOT chains could act as electron transfer channels to the redox-active groups, achieving a high efficiency of electron transfer between electrodes and redoxactive groups. Consequently, the galvanostatic charge–discharge process was conducted to evaluate the capacitance of samples at different charging rates, where the PEDOT-modified DAAQ–TFP COF film exhibited superior capacitance in comparison to the DAAQ-TFP COF film, with a maximum capacitance of 350 F $\rm cm^{-3}$ at a charging rate of 10C and 50% of the capacitance was maintained even at 1600C (Fig. 15b). Moreover, the PEDOTinfiltrated COF shows good chemical stability after 10 000 cycles, where the capacitance shows no significant changes at the charging rates of 10C and 100C (Fig. 15c). Review Article

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Subsequently, a chemically stable and redox active COF (TpOMe–DAQ COF) was developed by Banerjee's group through the condensation of 2,4,6-trimethoxy-1,3,5-benzenetricarbaldehyde (TpOMe) and 2,6-diaminoanthraquinone (DAQ) (Fig. 16a). 41 The chemical stability of TpOMe–DAQ COF was derived from the effect of both steric and hydrophobic protection of imine $(C=N)$ bonds by the neighbouring-OCH₃ functionalities and the interlayer C-H \cdots N H-bonding (Fig. 16b). By applying a mechanochemical grinding approach, TpOMe–DAQ COF was processed as thin film and applied as a free-standing electrode material for supercapacitors in 3.0 M $H₂SO₄$ solution. Given the good porosity of the COF and redox activity of the quinone/hydroquinone couple, the as-synthesized TpOMe–DAQ

Fig. 14 (a) Structure of DAAQ–TFP COF with random orientation on the electrode; (b) schematic illustration of the DAAQ–TFP COF as an oriented thin film on an electrode surface. Reproduced from ref. 219 with permission from the American Chemical Society, copyright 2015.

Fig. 15 (a) Schematic representation of the synthesis of PEDOT-modified DAAQ–TFP COF film; (b) average capacitance of PEDOT-modified DAAQ– TFP and DAAQ–TFP at different charging rates; (c) extended cycling of PEDOT-modified DAAQ–TFP film showing stability over 10 000 cycles. Reproduced from ref. 96 with permission from the American Chemical Society, copyright 2016.

displays an areal capacitance of 1600 mF cm^{-2} (gravimetric capacitance 169 F g^{-1}) and good cycling stability (>100000) with no significant changes in capacitance retention and coulombic efficiency (Fig. 16c and d).

Recently, Banerjee and co-workers reported two kinds of redox-active COFs, that is TpPa– $(OH)_2$ and TpBD– $(OH)_2$, by introducing redox-active hydroquinone moieties into COF backbones through Schiff base condensation of 1,3,5-triformylpholoroglucinol (TFP) with 2,5-dihydroxy-1,4-phenyldiamine [Pa-(OH)₂] and 3,3'-dihydroxybenzidine [BD-(OH)₂], respectively.²¹⁸ The reversible proton-coupled electron transfer $(2H^+/2e^-)$ of hydroquinone/benzoquinone moieties are responsible for the specific capacitance of these redox-active COFs.

The hybridization of COFs with conducting materials has been developed to further improve the electrical conductivity and subsequently the electron transport during the charge– discharge process. For example, Sun's group reported a COF/ NH2-rGO composite as a supercapacitor electrode through the condensation of 1,3,5-triformylbenzene and 1,4-diaminobenzene, and a further composite with amine-modified reduced graphene oxide (NH₂-rGO).²²⁰ It was described that the synergistic effect of the active COFs and conductive graphene supports increased the electrochemical performance. Similarly, 3D graphene was modified by growing COFs on its surfaces through an interfacial polymerization method, where redox-active moieties (anthraquinone) were deliberately chosen to increase the electrochemical activities.²²¹ The as-prepared electrode material had a good conductivity and provided more accessible electrochemically active moieties. As a result, the COF/graphene composite displays improved specific capacitance in alkaline media in comparison to that of the powder 2D COFs.

Very recently, ultralight COF/rGO aerogels with hierarchical porosity were reported. The COFs were produced in the presence of graphene oxide using hydrothermal treatment. Notably, during this treatment, the COF is not only formed as a thin layer on the graphene sheets, but GO is also reduced to rGO, which additionally stacks up in a 3D fashion to finally form a COF/rGO gel. This gel can be transformed into an ultralow density COF/rGO aerogel by freeze-drying (Fig. 17a).⁵⁷ Diaminoanthraquinone (Dq) was selected as a redox-active unit,

Fig. 16 (a) Schematic representation of the synthesis of TpOMe–DAQ, and the reversible transformation between quinone and hydroquinone; (b) structural illustration of the interlayer C–H \cdots NH-bonding in TpOMe–DAQ; (c) galvanostatic charge–discharge and (d) cycling stability performance (10 mA cm⁻²) of TpOMe–DAQ thin sheets. Reproduced from ref. 41 with permission from the American Chemical Society, copyright 2018.

Fig. 17 (a) Schematic illustration of the fabrication of COF/rGO aerogels; (b) N₂ sorption isotherms of COF/rGO, COF and rGO; (c) a photograph of a COF/rGO aerogel held by a leaf; (d) the galvanostatic charge–discharge curves of COF/rGO in 0.5 M H₂SO₄ at 0.5, 1, 2, 3 and 4 A g⁻¹; (e) the cycling stability of COF/rGO at 8 A g⁻¹. The inset image shows the charge/discharge profiles for 1–5 and 4996–5000 cycles. Reproduced from ref. 57 with permission from Springer Nature, copyright 2020.

enabling COFs with pseudocapacitive performance. The obtained COF/rGO provides a specific surface area of \sim 246 m² g⁻¹, lower than that of the pure COF (498 $\mathrm{m^2\,g^{-1}}$) due to the existence of rGO (Fig. 17b). By taking advantage of the morphology of the rGO aerogel and the surface-grown COFs, the COF/rGO aerogel exhibits a low density of \sim 7.0 mg cm⁻³, which is light enough to

be held by a leaf (Fig. 17c). In addition, the COF/rGO composite shows good mechanical strength without breaking when undergoing repeated compression and expansion. As a result of the good electron transport within graphene networks, the COF/rGO aerogel could be applied as an electrode directly for supercapacitors without the addition of conducting binders or additives.

In comparison to rGO and pristine COFs, the COF/rGO showed a superior capacitance (269 F $\rm g^{-1}$ at 0.5 A $\rm g^{-1})$ and enhanced cycling stability (Fig. 17d and e), which were higher than most of the reported COF-based electrodes.

Similar to graphene, CNTs have also been used to fabricate hybrid composites with COF materials.²²² Very recently, Zhang's group has reported an olefin-linked COF by introducing 1,3,5-triazine units and 3,5-dicyano-2,4,6-trimethylpyridine into the molecular framework. 223 The obtained conjugated COF was assembled with CNTs to form a flexible thin-film as the supercapacitor electrode, which exhibited an areal capacitance of \sim 15.2 mF cm⁻² and an energy density of \sim 7.3 mW h cm⁻³. It was demonstrated that the synergetic effect of the electron delocalization of olefin-linkages as well as the conductive CNTs could facilitate the electron transport via a strong π - π affinity between them, leading to an improved supercapacitor performance. Furthermore, the introduction of CNTs enabled the construction of free-standing COF/CNT thin films, which possess good capability and flexibility, showing promising application in the development of flexible and wearable devices as electrode materials.

As mentioned before, carbon-based materials with suitable pore size distribution and high specific surface area have been applied as electrode materials for supercapacitors. COF-derived carbons have also been investigated for this application as heteroatom-doped carbons,^{114,224-227} hierarchically porous carbons, 228 porous carbon with a core-shell structure¹¹⁵ and carbon nanoparticles.²²⁹ It is supposed that the regular arrangement of heteroatoms and ordered structures make COF materials promising precursors to construct heteroatomdoped carbons with tuneable porosities, controllable morphologies and high surface areas. Xu and co-workers reported a hierarchically porous B-doped carbon with meso/microporous structures using COFs as precursors, where improved porosities were achieved by applying $ZnCl₂$ as a template via a molten-salt approach.²²⁸ The obtained hierarchically porous carbons exhibited a specific surface area of 1828 $\mathrm{m^2\,g^{-1}}$ and supercapacitive performance with specific capacitance of 160 F $\rm g^{-1}.$ However, it can be of course questioned if COF-derived carbons are really competitive electrode materials compared to traditional carbons (e.g. activated carbons) considering their sophisticated synthesis and consequently higher cost.

3.2.2 Rechargeable batteries. Rechargeable batteries (including Li-ion batteries, Li–S batteries, Na-ion batteries, etc.) are currently intensively discussed and applied to enable emission free electric vehicles.^{6,230} In rechargeable batteries, the metal ions transfer between an anode and cathode during the charging/discharging process. Currently, a significant challenge in rechargeable batteries is to design high energy electrode materials, with high electrochemical performance without scarifying their chemical and mechanical stability during long-term cycles. The close relationship between electrode materials and battery performances including specific capacity, charge–discharge ability and recycle stability could provide insight into the design principle of efficient electrode materials. In this section, we concentrate on the COF-based

electrode materials and discuss their potential applications in Li-ion batteries, Li–S batteries and Na-ion batteries.

As the most widely utilized member in the rechargeable battery family, Li-ion batteries (LIBs) have many advantages regarding their high energy density, low self-discharge and large voltage window, and are therefore applied successfully in many everyday applications. However, the slow kinetics, poor power density, concerns of safety, cost, Li scarcity and low longevity create major challenges to their practical applications in electric vehicles. 231 Currently, the rational design of advanced electrode materials is applied to address the above concerns. In particular, the performance of anodes provide large room for improvement in terms of higher power density and better cycling stability. $232-234$ Currently, commercial LIB anodes are built from carbon materials because of the good electric conductivity, hierarchical structures for lithium ion intercalation and low cost. $235,236$ However, the graphite anodes generally face the problems of low specific capacity and safety issues. Thus, a vast amount of research work is being conducted to study alternatives, including metal anodes (such as Sn, Sb, Al, Mg, Ag and their alloys),^{232,237,238} transition metal oxides²³⁹⁻²⁴¹ and silicon.^{242,243} Chem Soc Rev

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Recently, COF-based materials have also been explored as potential anode materials in LIBs due to their layered and porous structures, which should allow facile charge carrier transport. In particular, the 2D structures of COFs provide sufficient internal channels for lithium atom insertion/adsorption and short-ended paths for fast lithium ion diffusion. It was demonstrated by theoretical calculations that 2D COFs preferred to adsorb lithium atoms in a regular symmetry, and lithium atoms tended to cross the 1D pores efficiently due to the low energy barrier, enhancing Li storage capacity.²⁴⁴ Zhao's group reported two conjugated COFs as anodes for Li-ion batteries by the condensation of aldehydes and amines under solvothermal conditions (N2-COF and N3-COF), 245 where the numbers indicate the amino groups in the amino derivatives (Fig. 18a and b). When applied to a Li-ion battery as an anode material, the as-fabricated N2-COF exhibited a charge capacity of 600 mA h g^{-1} with a capacity retention of 82% after 500 cycles. For the battery with N3-COF as the anode, a similar performance was observed (593 mA h g^{-1}) with a capacity retention of 81% (Fig. 18c and d). Moreover, the cycling stability of N2-COF and N3-COF were conducted at different current densities $(0.2 - 5 \text{ A g}^{-1})$ (Fig. 18e). The charge capacity of N2-COF shows a slight decrease from ~ 806 to 497 mA h g⁻¹ with increase of the current density from 0.2 to 5C. When the current density decreased from 5 to 2C, the N2-COF exhibits a recovered capacity of 607 mA h g^{-1} , proving its stability over multiple cycles and various current densities. A similar result can be observed with the N3-COF (Fig. 18f). The performance of N-COFs in LIBs was attributed to the uniform porosity, conjugated structures and considerable chemical stability, showing the promising application of COFs for Li-ion batteries.

Liu and co-workers have designed a conductive 2D COF (TThPP), where polyporphyrin moieties were connected by 4-thiophenephenyl groups via in situ oxidative polymerization

Fig. 18 Structures of (a) N2-COF and (b) N3-COF; charge–discharge cycles at 1 A g⁻¹ of (c) N2-COF and (d) N3-COF; chemical stability tests at different current densities (0.2–5 A g⁻¹) of (e) N2-COF and (f) N3-COF. Reproduced from ref. 245 with permission from the Royal Society of Chemistry, copyright 2016.

where porphyrin and thiophene serve as the core and linking bridge, respectively.²⁴⁶ When the TThPP was grown on the surface of a copper foil to be used as an anode material in a LIB, the COF-based polyporphyrin film exhibits good electrical conductivity with a reversible capacity of \sim 666 mA h g⁻¹. The high conductivity and capacity of TThPP was attributed to an eclipsed alignment of the polyporphyrin sheets, which can promote the process of charge transfer within the frameworks. To further increase the conductivity and stability, carbon-based conductive materials such as CNTs and graphene were also utilized for the preparation of composites with COFs, forming hybrid electrode materials for Li-ion batteries. For instance, Wang and co-workers synthesized a hybrid COF@CNT composite as the anode material. It was observed that the CNTs could affect the growth process of COFs, and consequently were coated by few-layered COFs via π - π interaction (Fig. 19).⁴² The as-fabricated COF@CNT composite exhibits a high reversible capacity of 1536 mA h g^{-1} with good retention after 500 cycles at 100 mA g^{-1} . The long-term stability can be ascribed to the few-layered morphology of the COFs and the good conductivity of CNTs within the hybrid composite.

Recently, Li–S batteries have received much attention due to their high theoretical energy density, gravimetric capacity and low production costs.²⁴⁷ Compared with traditional cathode materials such as LiFePO₄ and LiCoO₂, the usage of a sulfur cathode endows Li–S batteries with a relatively high theoretical gravimetric capacity of 1672 mA h $g^{-1}.^{248}$ However, before applying it into practical applications, some issues need to be considered, including the formation of soluble lithium polysulfides (Li₂S_x, 3 \leq x \leq 8) and insoluble sulfides (Li₂S₂/Li₂S) in the charge-discharge process. 248 It is supposed that the

dissolved polysulfides could lead to the low efficiency and poor stability of Li–S batteries due to the shuttle effect in the charging process between cathode and anode. Thus, development of advanced materials as a sulfur cathode, which can effectively inhibit the loss of soluble intermediates, would have a significant impact in this field. Since COF materials possess high porosity, low density and robustness, and their open pores and rigid networks can promote the transport of electrolyte ions, it is proposed that COFs could become potential cathode materials for hosting sulphur in Li-S batteries.²⁴⁹⁻²⁵¹ In this regard, CTFs, a distinct class of nitrogen-rich COF materials, were applied for the first time as a novel host material for the insertion of sulphur to capture dissolved polysulfides, thus enhancing the stability of Li-S batteries.¹²² This work paved the way to applying COF materials as cathode materials for Li–S batteries, but it should be noticed that more advanced methods and strategies still need to be developed to improve the performance of the COF-based sulfur cathodes.

For utilizing the long-range ordering and high porosity present in COFs, the same group further explored the applications of COFs for Li–S batteries by designing a 2D porphyrinbased COF (Por-COF) with large pore volume and narrow pore size distribution, which was used as a host for sulphur storage (Fig. 20a and b). 252 The galvanostatic charge/discharge test of Por-COF/S was conducted at a current rate of 0.5C in the voltage range of 1.8–2.7 V (Fig. 20c). Two plateaus were found both in discharge curves and charge curves (2.1 and 2.3 V for discharge curves; 2.25 and 2.37 V for charge curves), indicating the reversible redox reaction between sulfur and lithium polysulfides ($Li₂S_x$, $2 < x < 8$). The discharge capacity of the Por-COF/S was calculated after three cycles, which was almost

Fig. 20 (a) Synthesis and structure of Por-COF; (b) schematic representation of the charge and discharge process of the Por-COF/S; (c) galvanostatic charge and discharge profiles of the Por-COF/S at a rate of 0.5C; (d) cycling stability and coulombic efficiency of the Por-COF/S at a rate of 0.5C. Reproduced from ref. 252 with permission from the Royal Society of Chemistry, copyright 2016.

unchanged and maintained at 870 mA h $\rm g^{-1}.$ In addition, the cycling performance was evaluated at a charge/discharge rate of 0.5C where the Por-COF/S possessed a capacity of 633 mA h g^{-1} after 200 cycles, with a capacity decay rate of 0.16% per cycle, indicating its stable cycling performance. Moreover, the coulombic efficiency of the Por-COF/S was calculated to be 96% in

the cycling process, demonstrating the alleviation of the shuttle effect from dissolved polysulfides (Fig. 20d). In comparison to their previous study, the loaded sulphur within the large pore volume of Por-COF is responsible for the improved cycling performance.

It is supposed that physically isolated sulphur in the COF channels cannot satisfy the prevention of the shuttle effect

from Li–S intermediates, thus leading to an unsatisfactory energy storage performance. To suppress the shuttle effect, recently, the immobilization of polysulfide chains on the pore walls of COFs was demonstrated, and a series of functionalized COFs has been designed by grafting polysulfide chains on the pore walls through covalent coupling.^{69,253,254} For example, Jiang's group designed an imine-linked COF by the condensation of 1,3,6,8-tetrakis(4-formylphenyl) pyrene (TFPPy) and 4,4',4",4"'-(ethene-1,1,2,2-tetrayl)tetraaniline (ETTA) via a solvothermal method (Fig. 21a).²⁵⁴ The imine linkages (C=N) present in the COF (TFPPy–ETTA-COF) can induce the polymerization of sulphur to form polysulfide chains at elevated temperatures, which can be anchored covalently on the channel walls via C-S bonds. As the impact of polysulfide chains grow within COF channels, the as-synthesized polysulfide@TFPPy–ETTA-COF exhibits a capacity of 1069 mA h g^{-1} with a long discharge/ charge plateau, which is much higher than that of S@TFPPY– ETTA-COF (723 mA h g^{-1}), where sulfur is just incorporated in

the pores (Fig. 21b). Considering the remarkable enhancement of performance, it has been confirmed that the polysulfide chains can both promote the capacity and the efficiency of sulfur utilization. In addition, the cycling performances conducted at 0.1C showed the superior capacity of polysulfide@TFPPy–ETTA-COF to S@TFPPy–ETTA-COF, while the coulombic efficiency did not change during cycling (Fig. 21c). It is supposed that the anchored polysulfide chains are the redox-active sites and suppress the shuttle effect. Apart from the immobilization of polysulfide chains, morphology-controlled engineering was developed for the preparation of porphyrin COFs with hollow spherical structures (POF-HSs).²⁵⁵ The POF-HSs show two effects, namely physical confinement and chemical adsorption of polysulfides, thus alleviating the shuttle effect of polysulfides.

In addition to LIBs and Li–S batteries, all-solid-state Li-ion batteries, as next-generation energy storage devices, have attracted much attention in recent years. To achieve high-performance of

Fig. 21 (a) Structure of TFPPy–ETTA-COF and schematic representation of polysulfide@TFPPy–ETTA-COF; (b) galvanostatic charge–discharge tests of polysulfide@TFPPy–ETTA-COF (red) and S@TFPPy–ETTA-COF (blue) at a rate of 0.1C; (c) chemical stability and coulombic efficiency of polysulfide@TFPPy–ETTA-COF (red) and S@TFPPy–ETTA-COF (blue) at a rate of 0.1C. Reproduced from ref. 254 with permission from the Royal Society of Chemistry, copyright 2019.

an all-solid-state Li-ion batteries, the development of advanced solid electrolytes could be the key determinant. Key materials for the all-solid-state lithium batteries are solid electrolytes, and are generally oxide- and sulfide-based materials. Although several materials have been applied as solid electrolytes such as oxideand sulfide-based materials, some problems still remain for practical applications.256 A big challenge could be the formation of an internal short-circuit in the charging–discharging process. To solve this problem, organic materials with good ion conductivity could be an option. In this regard, ionic COFs have been explored as a Li-conducting solid electrolyte in Li-ion batteries due to their high ion conductivity. For example, Zhang's group synthesized a novel type of ionic COFs containing hybridized boron anionic centres and tuneable counter cations $\left[\mathrm{(NMe}_2\right)^+$ for ICOF-1 and Li⁺ for ICOF-2].²⁵⁷ By taking advantage of the high accessibility of Li ions, ICOF-2 exhibited a high Li-ion conductivity $(3.05 \times 10^{-5} \text{ S cm}^{-1})$ and a Li⁺ transfer number value of \sim 0.80, achieving the efficient transportation of Li ion at roomtemperature. In similar efforts, a 3D anionic COF was synthesized through the condensation of trimethyl borate and hydroxy groups in γ -cyclodextrin (γ -CD), where the formed tetrakis(spiroborate) tetrahedra needed to balance their negative charge with counterions.132 Because of the insertion of flexible moieties, the formed COFs (CD-COFs) provide a high Li-ion conductivity of \sim 2.7 mS cm⁻¹ and long-term Li-ion stripping/plating stability. In addition, polyelectrolyte COFs were reported by Jiang's group by introducing flexible oligo(ethylene oxide) chains into the COF pores.258 When complexed with Li ions, the polyelectrolyte COF shows enhanced ion conductivity, cycle life and thermal stability Chem Soc Rev

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that extend the potential applications of COF materials as solidstate ion conductors in Li-ion batteries.

Similar to Li-ion batteries, the sodium-ion battery (NIB) is another type of rechargeable battery that uses sodium ions (Na⁺) as the charge carriers, which are much more abundant than Li-ions. Considering the larger size of Na⁺, COF materials having soft and flexible structures can help in buffering the large volume expansion during sodiation. In addition, the insertion of redox-active sites into COF skeletons, which is desirable to construct COF-based anode materials for Na-ion batteries.56,259 As mentioned above, DAAQ–TFP COFs with redox-active anthraquinone moieties have been applied as electrode materials for pseudocapacitors (Fig. 14). Recently, Lu and co-workers applied this redox-active COF (DAAQ–TFP COF) also as an anode for Na-ion batteries (Fig. 22a).⁵⁶ With different exfoliation processes for the DAAQ-COF, four samples were obtained with thicknesses of 4–12 nm, 50–85 nm, 100– 180 nm and 100–250 nm (Fig. 22b). Consequently, the cycling performance of samples with different thicknesses was tested at a rate of 100 mA g^{-1} showing that the 4-12 nm sample exhibit superior capacity (420 mA h g^{-1}) in comparison to the thicker samples. This was attributed to the weak stacking interaction of 2D-COFs which may enhance the capacity and stability by restraining the self-exchange behaviour of the interlayer electrons and creating more accessible active sites. In addition, the 4–12 nm sample also showed a good cycling stability with 99% retention at 5 A g^{-1} over 10 000 cycles (Fig. 22c). Moreover, the authors demonstrated that radical intermediates (C – $O[•]$ and α - C) are of importance in the sodiation–desodiation

Fig. 22 (a) Synthetic route and structure of DAAQ-COF; (b) simulated structure of DAAQ-COF and the schematic representation of the stacked 2D-COF with different thicknesses; (c) cycling performance of 2D-COFs with different thicknesses and the coulombic efficiency of the 4-12 nm thick COF at 100 mA g⁻¹; (d) cycling performance of the COF with a thickness of 4–12 nm at 5 A g⁻¹. Reproduced from ref. 56 with permission from the American Chemical Society, copyright 2019.

process, and their redox activity and stability could be determined by the stacking behaviour of COFs.

In summary, COFs have shown to efficiently enhance the energy storage capacity in rechargeable batteries as a combined result of the presence of ordered channels, high porosities and possible integration of redox moieties. Even though the application of COFs in this field is still in an initial stage, their comparable performances to those of other reported materials show their promising applications in energy storage devices.

3.2.3 Fuel cells. In comparison to traditional power sources, fuel cells possess many advantages including high conversion efficiency, low maintenance, portability, and environmental friendliness. Among them, proton exchange membrane fuel cells (PEMFCs) are operated at a low temperature range in which COFs can be applied. As a critical component of PEMFCs, the proton exchange membrane (PEM) takes charge of the transport of protons from the anode to cathode, thus recombining with oxygen to form water. Thus, designing the PEMs is referred to as the heart of the PEMFC, and the proton conduction materials have subsequently spurred tremendous interest among researchers. Since the production of the perfluorosulfonic acid membrane with polytetrafluoroethylene as a support, which is called 'Nafion', it soon became a standard for PEMFC and remains so to date.²⁶⁰ However, the upper operation temperature of Nafion is low (only 80-100 $^{\circ}$ C) and it has been observed that its proton conductivity significantly drops after 100 $^{\circ}$ C.²⁶¹ Currently, many other types of PEMs have been developed, including polymeric,^{262,263} ceramic²⁶⁴ and inorganic–organic composite membranes, aiming at designing a stable PEM with high proton conductivity.²⁶⁵

COF materials are relatively lightweight, provide a variety of functionalities and have membrane processability like polymers through chemical or physical modifications. The high degree of internal ordering within COFs can also facilitate loading and transport of proton conducting substrates.²⁶⁶ As a proof of concept, Banerjee's group reported an azo $(-N=N-)$ -functionalized COF (Tp–Azo COF) through the reaction between triformylphloroglucinol (Tp) and 4,4'-azodianiline (Azo) (Fig. 23a). 267 The as-synthesized COFs are comprised of b-ketoenamine-linkages that yield enhanced chemical stability against acids, ensuring the sustainability under harsh conditions in fuel cells. Because of the introduction of phosphoric acid (H_3PO_4) into the framework, the azo units were protonated and then formed hydrogen bonds with $H_2PO_4^-$. Consequently, a hydrogen-bonding network was formed in the 1D channels with further interaction with free H_3PO_4 molecules (Fig. 23b). The reported proton conductivity values could reach 6.7 \times 10^{-5} S cm⁻¹ under anhydrous conditions and 9.9×10^{-4} S cm⁻¹ under 98% relative humidity (Fig. 23d and e). These results indicate the potential of COF-based functional proton conducting materials for applications in fuel cells. As such, in the following years, Banerjee's group reported several proton conductive COF materials based on b-ketoenamine-linkage COFs, such as sulfonic-acidbased COF (TpPa-SO₃H) with intrinsic proton conductivity under anhydrous conditions⁷⁹ and mechanochemically synthesized bipyridine-COF with H_3PO_4 incorporation.²⁶⁸

Beyond the doping of porous materials with acids, it has been shown that heterocyclic bases can be utilized as proton conducting materials. In this case, the strength of hydrogen bonding interactions could be weak, and the rotation of acid and base molecules could become possible. As expected, a high level of proton conductivity can be achieved via the Grotthuss mechanism. In order to achieve this level of proton conductivity in COFs, Jiang's group reported a mesoporous COF (TPB–DMTP-COF) through the reaction between 2,5-dimethoxyterephthalaldehyde (DMTP) and 1,3,5-tri(4-aminophenyl)benzene (TPB) via a solvothermal method (Fig. $23f$).⁶⁵ The authors studied the proton conduction across mesoporous channels in this framework, which could be applied as a host material for loading proton carriers. In this work, triazole and imidazole were chosen as N-heterocyclic proton carriers, providing trz@TPB–DMTP-COF and im@TPB–DMTP-COF with high loading capacities of 180 and 155 wt%, respectively. The frameworks can achieve satisfying proton conductivities $(4.37 \times 10^{-3} \text{ S cm}^{-1}$ at 130 °C) that are 2–4 orders of magnitude higher than those of microporous and non-porous polymers, implying the importance of doping N-heterocyclic proton carriers for enhanced proton conductivity. Another work about proton conductive COF was reported by Zhu's group, who introduced a cationic COF that was synthesized by combining a cationic monomer, ethidium bromide (EB) (3,8-diamino-5-ethyl-6-phenylphenanthridinium bromide), with 1,3,5-triformylphloroglucinol (TFP) via Schiff base reactions. By immobilizing $PW_{12}O_{40}^3$ ⁻ into this porous cationic framework, the proton conductivity was found to be enhanced to 3.32 \times 10^{-3} S $\rm cm^{-1}$ at 97% relative humidity. These contributions have shown the promising applications of COF materials in fuel cells and will probably encourage researchers to further design functional COF-based proton conducting materials. Although moderate values have been achieved using COFs as a proton conductor, most of the reported materials show conductivities only after doping with acids or N-heterocyclic proton carriers, limiting the utility of COFs for real time applications. On the other hand, because of their robust covalent backbone structures, COFs are generally synthesized as fluffy microcrystalline powders with low processability. It is practically difficult to post-fabricate COF powders for the demands of membrane separation technology. Even though additives and binders are generally used to achieve the fabrication of COF membranes, the possible pore blocking effect from the additives can decrease the accessibility of well-defined pores and functional ligands.²⁶⁹ Review Article
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3.3 Electrochemical sensors

Electrochemical sensors have gained extensive attention for the analysis of species of biological, environmental and industrial interest. 270 Typically, an electrochemical sensor consists of a working electrode and a counter electrode connected by a certain electrolyte. Through the reaction between working electrode and analytes, electric signals can be recorded, which are proportional to the analyte concentrations. In comparison with the applications in electrocatalysis and energy storage, just a few COFs have been applied as electrochemical sensors.

Fig. 23 (a) Synthesis and structure of Tp–Azo; (b) the formation of hydrogen bonding between H₂PO₄⁻ anion and azo units; (c) schematic illustration of H₃PO₄ doping in COFs; (d) proton conductivities of PA@Tp–Azo under anhydrous conditions. Inset image shows a distinct color change (red to black) in Tp-Azo COF powder upon H₃PO₄ treatment; (e) proton conductivities of PA@Tp-Azo under hydrous conditions; (f) proton conduction across 1D mesopores of crystalline COFs with different proton carriers. Reproduced from ref. 267 with permission from the American Chemical Society, copyright 2014; and ref. 65 with permission from Springer Nature, copyright 2016.

The low density and high surface areas of COFs can be considered as an ideal platform for adsorption/separation and detection/sensing. $271,272$ In addition, depending on the modular nature of COFs, their pore structures and functional groups can be tailored independently, thus enabling the introduction of electroactive sites/groups into COF channels to potentially bind and detect the specific species. For example, Zhu and co-workers have recently applied DAAQ–TFP COF (DQ-COF) as a substrate to study its capability for the removal and electrochemical detection of hydrazine from water solutions.²⁷³ By virtue of the high surface area and the presence of abundant anthraquinone sites, DQ-COF exhibits a hydrazine removal capability as high as 1108 mg $\mathrm{g}^{-1}.$ Consequently, the DQ-COF was dropped onto a nickel matrix to form the DQ-COF/Ni composite (DQ-COF/Ni), which was further applied for the detection of hydrazine by using an amperometry technique.274 As revealed, DQ-COF/Ni presents a wider linear range (0.5-1223 μ M) and lower detection limit (0.07 μ M) towards

hydrazine detection than those of reported hydrazine sensors (e.g. NiCo₂O₄²⁷⁵ and MnO₂²⁷⁶). Its high detection performance was attributed to the synergistic effect of the electroactive DQ-COF and Ni matrix accompanied by the enhanced electron transfer. Apart from hydrazine, environmental pollutants, such as heavy metals (e.g. lead²⁷⁷), nitrophenol²⁷⁸ and bisphenol A^{279} have also been studied as detection species by using COF-based electrochemical sensors.

Biomedical analysis is another prominent area for COF applications via electrochemical sensing. Analytes such as dopamine,²⁸⁰ glucose,²⁸¹ H₂O₂,²⁸² ascorbic acid (AA)²⁸³ and prostate specific antigen (PSA)²⁸⁴ have shown positive electrochemical responses to COF-based materials. To further improve the detection performance, COFs are generally constructed with other functional materials to form COF-based composites, to enhance their selectivity or sensitivity for detection molecules. For example, metal nanoparticles (NPs) are widely applied in

Fig. 24 SEM images of (a) 3D-KSC; (b) 3D-KSC/COFTAPB–PDA; (c) 3D-KSC/COFTAPB–PDA/CuNPs and (d) 3D-KSC/COFTAPB–PDA/PtNPs; (e) amperometric response of 3D-KSC/COFTAPB–PDA/CuNPs under stirring in 0.1 M NaOH solution with continuous addition of glucose; (f) linear relationship between current density and glucose concentration; (g) amperometric response of 3D-KSC/COFTAPB–PDA/PtNPs under stirring in 0.2 M PBS (pH = 7.0) solution with continuous addition of H₂O₂. Inset: Magnification part of the curve; (h) linear relationship between current density and H₂O₂ concentration. Reproduced from ref. 281 with permission from Elsevier Inc, copyright 2019.

detecting targeted compounds such as glucose (Cu NPs) and H2O2 (Pt NPs). To avoid the aggregation of metal NPs during the deposition process, Wang and co-workers designed a functionalized 3D porous carbon-COF hybrid (3D-KSC/COFTAPB-PDA), by growing a COF thin film $(COF_{TAPB-PDA}, TAPB: 1,3,5$ tris(4-aminophenyl)benzene, PDA: 1,4-phthalaldehyde) on pore walls of $3D-KSC.²⁸¹$ By taking advantage of the exposed active sites in the COF film, the spherical Cu and Pt nanoparticles were deposited uniformly on the $3D-KSC/COF_{TAPB-PDA}$ electrode by electrostatic potential deposition, named 3D-KSC/ COF_{TAPB-PDA}/CuNPs and 3D-KSC/COF_{TAPB-PDA}/PtNPs, respectively (Fig. 24a–d). After optimization, the amperometric response of 3D-KSC/COF_{TAPB-PDA}/CuNPs was recorded by continuously adding glucose solution at 0.5 V (Fig. 24e). As shown in Fig. 24f, the linear range of glucose detection by 3D-KSC/ COF_{TAPB–PDA}/CuNPs was 4.69 μ M–1.57 mM and 1.57–7.07 mM with a detection limit of 1.54 μ M. The result was much better than those of 3D-KSC/CuNPs $(6.33 \mu M - 4.07 \text{ mM})$, and detection limit of 2.08 μ M) as well as some of the Cu/Cu_xO related electrochemical sensors.²⁸⁵ Similarly, the 3D-KSC/COF_{TAPB-PDA}/ PtNPs also provided a comparable performance in the detection of $H₂O₂$ (Fig. 24g and h).

4. Chemical stability of COFs and their impact on electrochemical operations

The periodicity in COF structures is achieved *via* dynamic covalent chemistry. In this regard, a high crystallinity in COFs is always then observed when good reversibility of the newly formed covalent bonds is ensured. This naturally means that the bonds can be easily opened again, consequently resulting in limited chemical stability. Indeed, crystallinity and stability seem to be contradictory properties of COFs. Thus, the first reported boroxine and boronate-ester COFs have been found to be unstable in water and even under humid conditions, as the linkages are susceptible to attack by nucleophiles (e.g., water molecules) at the electron-deficient boron sites.¹⁶ As such, boroxine and boronate-ester COFs have several limitations for real applications, especially those involving aqueous solutions for electrocatalysis (HER, OER, ORR, etc.). To circumvent these limitations, various novel linkages have been explored for the construction of COFs in recent years (Fig. 25). Covalent triazine frameworks (CTFs) synthesized by the cyano-trimerization are found to be extremely robust towards strong acid or base environments as well as in the presence of strong oxidizing and reducing agents, and thus were considered as promising candidates for a range of electrochemical applications.^{286,287} Nevertheless, to ensure reversibility, the triazine formation was carried out under harsh conditions in salt melts, which limits the amount of applicable monomers and also yielded defectrich materials with lower crystallinity.²⁶⁹

Imine,²⁸⁸ azine²⁸⁹ and hydrazone²⁹⁰ COFs linked by C=N bonds exhibit superior hydrolytic stability compared to that of the first reported boron-based COFs. The chemical stability of most imine, azine and hydrazone COFs is however still unsatisfactory since they undergo hydrolysis under acidic conditions, resulting in some limitations for their electrochemical applications in acidic media.²⁹¹ Notably, β -ketoenaminelinked COFs offer a much enhanced stability in aqueous, acidic and basic media, due to the irreversible tautomerization of their initially formed imines. Indeed β -ketoenamine-linked COFs with

varying porosity and diverse functionalities have been tested for various electrochemical applications such as OER, ORR, supercapacitors, batteries and proton conductivity.41,55,269

Polyimide COFs show good framework retention in pure water and acid solutions; however, alkali aqueous solutions significantly affect the framework stability due to the nucleophilic substitution in the presence of a base.²⁹² Due to their limited chemical stability, silicate organic frameworks (Si-COFs) have not been explored for electrocatalysis.³³ However, cyanovinylene and vinylene-linked COFs possessing fully conjugated backbones are very promising, and are significantly less prone to hydrolysis compared to boroxine,¹⁶ boronate ester,²⁹³ or imine²⁸⁸ linkages, while still showing high crystallinity. Indeed, shortly after their discovery, cyanovinylene and vinylene-linked COFs have already been used for electrocatalytic applications.²⁹⁴⁻²⁹⁶ Similarly, dioxin(polyarylether)- and ester-linked COFs show considerable chemical stability, which might be useful for exploring their electrochemical applications.²⁹⁷

Nonetheless, beside the increasing stability of COFs in acid or basic media, there are still some other issues, which have to be considered before applying the COFs for practical applications, especially when a voltage is applied over a long-term operation. Concerns regarding COFs in long-term applications are the evolution of morphology/structure of COFs and of initially well-defined active sites in electrolytes under applied voltages. For example, when a Co-bipyridine-based COF was applied as an electrocatalyst for oxygen production, the conversion of $Co(\mu)$ to $Co(\mu)$ was detected, to form cobalt hydroxide/ oxyhydroxide on the surface after long-term OER operation, both of which are also OER active.36

Besides, attributing electrochemical performance to the morphology/structure of COFs is valid only if the COF structure remains intact throughout the electrochemical process. So, it is crucial to analyse and discuss their structures before and after long-term operation, to identify the real active sites as well as the operational stability of constructed COF-based electrodes. Generally, ex situ characterizations of spent catalysts (e.g., XRD, XPS, XANES, SEM and TEM) are a facile solution to illustrate structural changes of COFs. However, this can be problematic if an additive (Nafion, carbon, etc.) has to be added to the COF for

the electrochemical testing. Furthermore, in situ techniques can be applied to detect formed intermediates and conversion of active centres during operation. Indeed, techniques like operando ambient pressure XPS,¹⁵⁸ in situ Raman spectroscopy²⁹⁸ and *in situ* TEM²⁹⁹ are commonly used nowadays in electrocatalytic/chemical processes and have enabled a detailed insight into reaction mechanisms and evolution of active sites.

5. Summary and outlook

Since the discovery of covalent organic frameworks in 2005 by Yaghi and co-workers, a series of COFs has been designed and synthesized targeting versatile applications. Given their controllable and well-defined structure, high porosity and surface area, COFs are promising materials for energy conversion and storage applications. Indeed, COFs feature certain advantages, which distinguish them from other typical materials used in this field. Compared to carbon- and graphene-based materials, there is at first very good structural control. By selection of appropriate linkage motifs and building blocks, materials with defined chemical structures, tuneable pore sizes and structures and low densities can be prepared, which are not seen in other typical carbon-based electrode materials. For example, metalcoordinating moieties in COFs can act as well-defined active sites for electrocatalysis or redox-active sites can enable an enhanced pseudocapacitive performance in supercapacitors or reversable processes in rechargeable batteries. On the other hand, the exclusive covalent connections can yield fully π -conjugated COFs enabling enhanced charge transport, which is often problematic in MOFs.

To achieve high and robust electrocatalytic performance, increasing the number of active sites and intrinsic activity of each individual active site is required, which is enabled by the high surface area of COFs. Meanwhile, as the redox reactions in electrocatalysis generally happen between electrolyte and electrodes along with ion transport and electron exchange, novel synthetic approaches of COF-based electrocatalysts focus on facilitating mass transport and promoting the electric conductivity during electrocatalytic processes. The introduction of hierarchically porous structures and conducting materials into COFs is thus a promising option to create more advanced electrocatalysts.

In general, to design and fabricate efficient COF-based materials for electrochemical applications, different strategies have been developed, including:

(i) Chemical modifications: depending on the desired application in electrocatalysis, COF structures can be controlled by developing different strategies including backbone modification, functional group modifications and post-synthetic modifications.

(ii) Controlling COF structures on all length scales: COFs at the macroscopic length scale can combine their crystalline structures with the possibility to design architectures with well-defined morphologies and hybrid components, endowing them with good mass transfer and high conductivity.

COF-based materials have been applied as electrocatalysts, for HER, OER, ORR and CO2RR, as electrodes in supercapacitors and rechargeable batteries and as solid electrolytes or proton conducting membranes in batteries and fuel cells. However, several challenges for COFs in energy applications still exist. Primarily, the low stability or limited durability largely limits the practical realization of COF materials in energy-related devices. Under strong aqueous acidic/alkaline electrolytes, the collapse of COF structures could cause that active centres are getting inaccessible and that mass as well as charge transport is impaired. In addition, the poor processability of COFs could be another main issue for their applications in electrocatalysis as most synthetic methods for COFs involve solvothermal reactions of precursors in sealed glass tubes at elevated temperatures, yielding completely insoluble powders comprised of randomly aggregated crystallites. This makes the integration of COF materials into energy-related devices challenging. Review Article

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Therefore, it is desirable to develop synthetic and processing methodologies for construction of COFs with enhanced stability and processability that allow shaping, placing and orienting COFs as per necessity, while integrating them with other components of the devices for stable operation. For improving the electrocatalytic performance and energy storage performance in COFs, it seems furthermore necessary to improve their electrical conductivity. Growing of oriented COF thin films on a conducting support or the preparation of composites has been described as an option to achieve the same. Finally, using the new stable and fully conjugated COFs composed of cyanovinylene- or olefin-linkages will not only be advantageous for enhancing the recyclability (robustness), but also for improving charge transport properties that are mandatory for electrochemical applications.

Diverse building units and flexible molecular design are available for scientists to design COF-based electrocatalysts at the molecular level with precisely controllable active sites. COFs with their defined structures indeed offer the possibility to study optimal structures of active sites, and to understand the underlying chemical and electrochemical reactions. The combination of advanced in situ/in operando characterization techniques and computational tools, both of which can be more suitably applied to a crystalline material than to an amorphous or a rather ill-defined material, can indeed yield novel insight into structure–function relationships in electrochemical devices.

Conflicts of interest

There are no conflicts to declare.

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