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Abstract

Metal organic framework (MOF)-derived nanoporous carbons (NPCs) have been proposed as promising electrode materials for energy storage and conversion devices. However, MOF-derived NPCs typically suffer from poor electrical conductivity due to the lack of connectivity between these particles and a micropore-dominated storage mechanism, which hinder mass and electron transfer, thereby leading to poor electrochemical performance. In recent years, one-dimensional (1D) MOF-derived carbon nanostructures obtained using an electrospinning method have emerged as promising materials for both electrochemical energy storage (EES) and energy conversion applications. In this mini review, the recent progress in the development of MOF-derived 1D porous or hollow carbon nanofibers using the electrospinning method and their application in energy storage (e.g., supercapacitors and rechargeable batteries) and conversion devices (e.g., fuel cells) is presented. The synthetic method, formation mechanism and the structure-activity relationship of such porous or hollow carbon nanofibers are also discussed in detail. Finally, future perspectives on the development of electrospun MOF-derived carbon nanomaterials for energy storage and conversion applications are provided. This review will provide some guidance for future derivations of 1D hollow carbon nanomaterials from MOFs using electrospinning technology.

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Metal-Organic Framework-Derived One-Dimensional Porous or Hollow Carbon-Based Nanofibers for Energy Storage and Conversion

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1 **Abstract**

2 Metal organic framework (MOF)-derived nanoporous carbons (NPCs) have been proposed as promising
3 electrode materials for energy storage and conversion devices. However, MOF-derived NPCs typically suffer
4 from poor electrical conductivity due to the lack of connectivity between these particles and micropore-
5 dominated storage mechanism, which hinder mass and electron transfer, thereby leading to poor electrochemical
6 performance. In recent years, one-dimensional (1D) MOF-derived carbon nanostructures obtained using
7 electrospinning method have emerged as promising materials for both electrochemical energy storage (EES)
8 and energy conversion applications. In this mini review, the recent progress in the development of MOF-derived
9 1D porous or hollow carbon nanofibers using electrospinning method and their applications for energy storage
10 (e.g., supercapacitors and rechargeable batteries) and conversion (e.g., fuel cells) is presented. The synthetic
11 method, formation mechanism and the structure-activity relationship of such porous or hollow carbon
12 nanofibers are also discussed in detail. Finally, the future perspectives on the development of electrospun MOF-
13 derived carbon nanomaterials for energy storage and conversion applications are provided. This review will
14 provide some guidance for future derivations of 1D hollow carbon nanomaterials from MOFs using
15 electrospinning technology.

16
17 **Keywords:** Metal-organic frameworks; hollow carbon nanofibers; nanoporous carbons; energy storage and
18 conversion; electrospinning

19
20

1 **1. Introduction**

2 The rapid increase in world economy has led to a significant increase in global energy requirement, leading
3 to rising concerns of energy shortage and climate change problems for the society.¹⁻⁵ As a result, the
4 development of highly efficient, renewable and clean energy sources is becoming increasingly important. In
5 recent years, many efforts have been devoted to develop various rechargeable and reversible energy storage and
6 conversion devices, such as supercapacitors,⁶⁻⁸ fuel cells,⁹⁻¹² solar cells,¹³⁻¹⁵ and lithium-ion batteries (LIBs).¹⁶⁻¹⁸
7 The results demonstrate that the achievement of EES devices with high energy and power densities require the
8 design and construction of high-performance electrode materials combined with the suitable use of
9 electrolytes.^{19,20}

10 Metal-organic frameworks (MOFs), an emerging but rapidly growing class of porous materials, are
11 assembled by connecting metal ions or clusters with organic linkers. They have attracted considerable interest
12 since firstly reported in 1995.²¹⁻²³ Compared with traditional porous materials, the main advantage of MOFs is
13 their controllable structure and tunable porosity at the molecular level by tuning metal species and organic links.
14 As a result of these advantages, MOFs have been widely applied in adsorption,^{24,25} gas storage/separation,²⁶⁻²⁹
15 catalysis,³⁰⁻³² and drug delivery.³³⁻³⁵ In recent years, the exploration of MOFs in energy storage and conversion
16 applications, such as supercapacitors,³⁶ fuel cells,⁹ and lithium-ion batteries^{37,38} has gained significant attention.
17 However, the poor electrical conductivity and low stability of pure MOFs have impeded their full potential for
18 these applications. More recently, MOFs have received significant attention as novel templates to derive
19 nanoporous carbons (NPCs) for energy storage and conversion applications.³⁹⁻⁴² Since the first report of such
20 derivation by Xu et al.⁴³ in 2008 and a subsequent report by Yamauchi et al.⁴⁴ in 2012, significant advances have
21 been made in the derivation of NPCs from MOFs and the related publications are continuously growing every
22 year (**Fig. 1**).⁴⁵⁻⁴⁹ The development process in the composition and morphology of MOF-derived NPCs is
23 mainly divided into three stages (**Fig. 2**): (i) from single metal MOF to bimetallic MOF;⁵⁰⁻⁵⁴ (ii) from solid
24 structure to hollow structure; the strategies for preparing hollow structures include etching,⁵⁵ rigid
25 inducing/confined pyrolysis,⁵⁶ and templated method;⁵⁷ and (iii) from zero dimension (0D) polyhedral to 1D
26 rods,⁵⁸⁻⁶⁰ 2D sheets,⁶¹⁻⁶³ and 3D arrays.⁶⁴⁻⁶⁶ Compared with 0D, 2D, and 3D nanomaterials, 1D nanostructures,
27 especially those with porous or hollow structures possess many advantages for energy storage and conversion
28 devices. First, porous or hollow structures offer more active sites and reduce the path of transport for both mass
29 and charge. Second, 1D nanomaterials can provide fast pathways for electron transport, and lastly, they can be
30 easily grown on thin films to form self-standing flexible energy storage devices.⁶⁷⁻⁷¹ Due to these advantages,
31 more efforts have been dedicated in recent years to fabricate porous or hollow 1D carbon nanomaterials from
32 MOFs.

33 Electrospinning, as a versatile and scalable method, has attracted significant interest in both scientific

1 research and practical applications.⁷² Together with carbonization, this method provides a facile and convenient
2 way for preparing 1D porous (PCNFs) or hollow carbon nanofibers (HCNFs).^{4, 19, 73-78} Our group has previously
3 reported the fabrication of electrospun MOF-derived PCNFs and HCNFs for energy storage applications.^{79, 80}
4 MOF-derived PCNFs and HCNFs have emerged as a novel kind of carbon materials for energy storage and
5 conversion applications. In this mini review, we will focus on the recent progress in the fabrication of MOF-
6 derived 1D PCNFs and HCNFs by electrospinning for energy storage and conversion applications (e.g.,
7 supercapacitors, rechargeable batteries, and fuel cells). Herein, we will cover the use of electrospinning method
8 for synthesizing MOF-derived PCNFs and HCNFs and discuss their formation mechanisms, along with their
9 important application in these devices. Finally, the future perspectives of electrospun MOF-derived PCNFs and
10 HCNFs for energy storage and conversion applications are also presented.

11 12 **2. Synthesis and formation mechanism of MOF-derived HCNFs by electrospinning**

13 Electrospinning provides a simple and low-cost method for preparing 1D nanomaterials and it is by far the
14 most commonly used method for fabricating MOF-derived 1D PCNFs or HCNFs.⁸⁰⁻⁸³ Up to now, there are two
15 main approaches for fabricating electrospun MOF-based nanofibers (**Fig. 3**). In route I, MOF nanoparticles are
16 firstly synthesized and then, they are mixed with the polymer to form a homogeneous solution. Next, this
17 precursor solution can be converted into MOF-based nanofibers by electrospinning. In route II, electrospun
18 polymer nanofibers (containing metal ions) are firstly prepared and then, the MOF nanoparticles are grown in-
19 situ on the polymer nanofibers. Finally, the 1D PCNFs or HCNFs are obtained by directly carbonizing the MOF
20 nanofibers under inert atmosphere. To explain the formation mechanism of 1D PCNFs from the electrospinning
21 and subsequent carbonization of MOF/polymer mixtures, zeolitic imidazolate framework-8 (ZIF-
22 8)/polyvinylpyrrolidone (PVP) nanofibers are selected as a typical model.⁸⁴ As depicted in **Fig. 4**, the surface of
23 ZIF-8 polyhedrons possesses abundant Zn²⁺ sites which can coordinate easily with C=O groups present
24 abundantly in PVP. As a result of this coordination effect, the ZIF-8 polyhedrons in the mixture of ZIF-8/PVP
25 are evenly distributed and encapsulated by PVP at a low temperature. Following heat-treatment at 500 °C, the
26 PVP in the mixture decomposed to an amorphous carbon network, while the ZIF-8 polyhedrons retained their
27 crystal structure. Owing to the low calcination temperature, the PVP-derived carbon still possesses numerous
28 C=O bonds which can generate a new and strong interface between ZIF-8 and PVP-derived carbon. With the
29 increase in calcination temperature, the ZIF-8 polyhedrons start to decompose. The strong cohesive interface
30 interaction between PVP-derived carbon and ZIF-8 polyhedrons offers a unique confinement effect which
31 maintains the ZIF-8-derived carbon adjacent to the internal surface of PVP-derived carbon.

32 Another example is the fabrication of 1D HCNFs using the electrospinning and post-synthesis carbonization
33 of a mixture of ZIF-8/PAN nanofibers.⁷⁹ To generate the ZIF-8/PAN nanofibers, ZIF-8 nanoparticles are firstly

1 coated with a layer of PAN. Due to the capability of Zn^{2+} to coordinate with $-\text{C}\equiv\text{N}$ groups existing on the
2 surface of PAN nanofibers, the ZIF-8 particles become tightly encapsulated by the PAN nanofibers. At a low
3 calcination temperature (500 °C), the PAN-derived carbon typically still exhibits many $-\text{C}\equiv\text{N}$ groups which can
4 form a new strong interface between the ZIF-8 surface and the PAN-derived carbon shell. This consequently
5 causes the decomposition of the ZIF-8 particles, which is accompanied by their shrinkage starting from inside to
6 outside, thereby leading to the generation of hollow structure. The formation of such hollow structure can be
7 described as a confinement effect or an interface induction effect, which has been observed in other systems,
8 such as, ZIF-8@SiO_2 .⁵⁶ On the other hand, if no other interaction exists between the surface of the ZIF-8
9 nanoparticles and the PAN layer, inward contraction takes place and a solid structure is achieved. Apart from
10 PAN and PVP, PVA has previously been used to prepare ZnO quantum dot (QD)-decorated carbon nanofibers.⁸⁵
11 This was possible due to the oxygen rich nature of PVA, which enabled it to react with the ZIF-8 particles
12 during carbonization of the ZIF-8/PVA nanofibers to generate ZnO QDs@carbons on the CNF surface.

14 **3. MOF-derived HCNF materials for energy storage and conversion applications**

15 **3.1 Supercapacitors**

16 Supercapacitors (SCs) are considered as promising energy storage devices owing to their unique features,
17 such as high power density, fast charging rate, and long cycling life.^{8, 49} The most significant parameters which
18 govern the capacitive performance of SC electrodes are pore size distribution, conductivity, and surface area.^{8, 86}
19 MOF-derived NPCs with high surface areas and tunable pore size have been widely investigated as potential
20 electrode materials in SCs applications in the past decade.^{47, 87-89} However, the low conductivity of MOF-
21 derived NPCs typically leads to lower specific capacitance, which is often attributed to the poor
22 interconnectivity between these particles. Generally, the specific capacitance of MOF-derived NPCs is around
23 100-300 F g^{-1} in aqueous electrolytes, which is only slightly higher than activated carbons.⁸⁸

24 Recently, electrospinning method combined with carbonization has been proposed as an attractive route to
25 increase the electrical conductivity of MOF-derived NPCs. Previously, our group has fabricated 1D nanoporous
26 carbon fibers (NPCFs) via electrospinning for SC. In our proposed method, ZIF-8 nanoparticles were mixed
27 with PAN solution and converted into ZIF-8/PAN fibers by electrospinning, before being heat-treated at 500 °C
28 under inert atmosphere to obtain NPCFs.⁷⁹ As evident in **Fig. 5a**, the ZIF-8 nanoparticles were transformed into
29 hollow structures after the carbonization process. The CV curves of the NPCFs at various scan rates (**Fig. 5b**)
30 revealed their electric double-layer capacitance and the galvanostatic charge-discharge measurements at various
31 applied currents indicated their favourable conductive behaviors (**Fig. 5c**). The specific capacitances of PAN-
32 derived carbon (PAN-C), ZIF-8-derived NPC (ZIF-8-NPC), and NPCF were measured to be 110, 201, and 332
33 F g^{-1} , respectively, at 1 A g^{-1} (**Fig. 5d**). The enhanced specific capacitance of NPCFs was contributed by their

1 enhanced electrical conductivity, which resulted from the improved interconnectivity between the ZIF-8-derived
2 NPCs. Such improved interconnectivity was possible due to the linking of these particles by the chains of PAN-
3 derived carbon, which provided a pathway to enable smooth charge transport and ionic diffusion during
4 charge/discharge. The Nyquist plots confirmed the higher electrical conductivity of NPCFs compared to PAN-C
5 and ZIF-8-NPC, which was further supported by the four-probe method measurements (**Fig. 5e**).

6 Apart from PAN, 1D conductive polyaniline (PANI) nanofibers have also attracted considerable attention as
7 electrode materials for SCs, owing to their simple and low-cost fabrication, high chemical stability, and good
8 electrical conductivity.⁹⁰ Previously, 3D cross-linked PANI/N-doped porous carbon fibers (3D-CPC) have been
9 obtained from the in-situ polymerization of ZIF-8-derived N-doped porous carbon fibers with aniline.⁹¹ The
10 galvanostatic charge-discharge measurements revealed the higher specific capacitance of the 3D-CPC (75 F g⁻¹)
11 relative to 3D cross-linked PANI (504 F g⁻¹) and N-doped porous carbon (325 F g⁻¹) at a current density of 1 A
12 g⁻¹. Furthermore, the specific capacitance of the 3D-CPC only slightly decreased from 755 to 618 F g⁻¹, with the
13 increase in current density from 1 to 20 A g⁻¹, indicating its superior rate performance. In addition, the 3D-CPC
14 electrode also showed good cycling performance with high capacitance retention of 91.2% after 1000 cycles at
15 a high current density of 20 A g⁻¹. The excellent electrochemical properties of the 3D-CPC for SC were ascribed
16 to the N-doped porous carbon which could offer more active sites for the fast transfer of ions and electrons and
17 the high structural stability of the 3D cross-linked PANI fiber which could prevent the electrode material from
18 swelling and shrinking.

19 The electrochemical performance of MOF-derived CNFs for SCs can be further enhanced by combining
20 them with pseudocapacitive materials.^{85, 92} For instance, hierarchical nanofibers composed of numerous ZnO
21 QDs@carbons and CNFs prepared via the carbonization of electrospun ZIF-8/poly(vinyl alcohol) (PVA)
22 nanofibers displayed improved specific capacitance (346 F g⁻¹ at 0.5 A g⁻¹) and enhanced rate capability (79%
23 capacitance retention at 8 A g⁻¹) compared to ZIF-8-derived carbon and pure CNFs.⁸⁵ The outstanding
24 electrochemical performance of the ZnO QDs-decorated CNFs (ZPCNFs) was attributed to several reasons.
25 First, the uniformly dispersed ZnO QDs throughout the entire CNFs would increase the total capacitance of the
26 composite by offering pseudocapacitance. Second, the 1D CNFs could provide highly conductive pathways for
27 the transport of electrons during the charge/discharge process. Lastly, the hierarchical pores of the ZPCNFs
28 offered many electroactive sites and electrolyte transfer channels to enhance the transportation of ions
29 throughout the architecture, leading to improved capacitance and rate capability.

30 An integrated self-charging power system combining flexible fiber-shaped coaxial asymmetric
31 supercapacitors (FASCs) with flexible triboelectric nanogenerators (FTENGs) can enable simultaneous
32 harvesting and storage of energy for providing power for flexible electronics.⁹³ Uniform carbon nanotubes
33 fibers (CNTFs)@ZnCo₂O₄@Zn-Co-S hybrid array (HA) derived from bimetallic Zn-Co-ZIF have previously

1 been utilized to fabricate such self-charged power system.⁹⁴ To obtain the hybrid array, the CNTFs were initially
2 immersed in a mixture solution containing Zn^{2+} and Co^{2+} solution under hydrothermal conditions and the
3 resulting product was heat-treated at 350 °C under inert atmosphere to obtain the CNTFs@ZnCo₂O₄ nanowire
4 array (NA). The grown ZnCo₂O₄ NA was subsequently used as a source of both Zn^{2+} and Co^{2+} ions as well as a
5 skeleton for the in-situ growth of the bimetallic Zn/Co-ZIF (CNTFs@ZnCo₂O₄@Zn/Co-ZIF). The final
6 conversion to CNTFs@ZnCo₂O₄@Zn-Co-S HA was achieved by heat-treating the CNTFs@ZnCo₂O₄@Zn/Co-
7 ZIF HA in air at 350 °C to convert it to CNTFs@ZnCo₂O₄@ZnCo₂O₄ HA, followed by a secondary reaction
8 with thioacetamide at 70 °C. The flexible FASC device was assembled by utilizing the CNTFs@ZnCo₂O₄@Zn-
9 Co-S as the cathode and the CNTFs@H-Co₃O₄@CoNC HA as the anode. This FASC device showed a high
10 specific capacitance of 117.18 mF cm⁻² at a current density of 1 mA cm⁻², which slightly decreased to 67.5 mF
11 cm⁻² with the increase of current density to 10 mA cm⁻². The excellent electrochemical performance of this
12 FASC device was further highlighted by its high areal energy density of 32.01 μW h cm⁻² at an areal power
13 density of 698.42 μW cm⁻². Moreover, when the power density was increased to 7000 μW cm⁻², the energy
14 density of the device still remained as high as 12.38 μW h cm⁻², suggesting the synergistic effect of the
15 CNTFs@ZnCo₂O₄@Zn-Co-S HA. The excellent electrochemical performance of the assembled FASC device
16 was attributed to the unique hierarchical and porous structure which provided many electroactive sites and a
17 facile ion diffusion path.

18 From the above reports, it is evident that electrospinning is a feasible and attractive method for preparing
19 1D MOF-derived PCNFs or HCNFs with high electrochemical performance for SC applications. In general, the
20 benefits of MOF-derived PCNFs or HCNFs for SC applications are:

- 21 • *High structural stability* which prevents significant swelling and shrinkage of the electrode.
- 22 • *Hierarchical porosity* which provides many electroactive sites for ions and numerous electrolyte transfer
23 channels to enhance ionic transport throughout the structure.
- 24 • *1D fiber-like structure* which offers highly conductive pathways for the transport of electrons during
25 charge/discharge.
- 26 • *Improved interconnection between the MOF-derived NPCs* which provided a ‘pathway’ to enable
27 efficient charge transport and ionic diffusion during charge/discharge.

28 Despite the significant progress in this field, many previous reports tend to focus on ZIF-derived carbons and
29 more studies are needed on the preparation of 1D PCNFs or HCNFs from other MOFs in the future.

31 **3.2 Rechargeable batteries**

32 Rechargeable batteries, such as LIBs and sodium-ion batteries (SIBs) are among the most popular energy
33 storage systems in our daily lives, playing important roles in personal computers, smart grids, and electrical

vehicles.⁹⁵⁻⁹⁷ One of the key factors which determine the performance of rechargeable batteries is the electrode materials. Therefore, many studies have been carried out to develop advanced electrode materials with high specific capacity, good rate performance, and excellent cycling stability. Electrospun MOF-derived PCNFs or HCNFs possess high surface area, good electrical conductivity, and hierarchically porous structures, which make them promising electrode materials for rechargeable batteries. For instance, Lou et al. have demonstrated the excellent electrochemical properties of hierarchical Co₃O₄/CNT microtubes derived from the in-situ growth of ZIF-67 on electrospun polyacrylonitrile (PAN) nanofibers for LIBs (**Fig. 6a**).⁹⁸ The hollow nature of the ZIF-67/PAN-derived Co₃O₄/CNT microtubes can be clearly observed in **Fig 6b-d**, which was attributed to the Kirkendall effect during the annealing process. Due to their hierarchical hollow architectures, the volume variations which occurred during charge-discharge processes could be effectively minimized. As a result, the ZIF-67/PAN-derived Co₃O₄/CNT microtubes exhibited good rate performance, with the reversible capacity being decreased only slightly from 832 mAh g⁻¹ at 0.75 A g⁻¹ to 643 mAh g⁻¹ at 3 A g⁻¹ (**Fig. 6e**) and good cycling stability with no obvious capacity fading after 200 cycles at 1 A g⁻¹ (**Fig. 6f**). Goodenough and co-workers reported the fabrication of nitrogen-doped hollow porous carbon nanofibers (HPCNFs-N) through the in-situ growth of bimetallic ZIF (Zn-Co-ZIF, BMZIF) on PAN nanofibers for SIBs (**Fig. 6g**).⁹⁹ The BMZIF can combine the advantages of each individual ZIF material (ZIF-8 affords the porous structure and nitrogen doping; ZIF-67 provides highly graphitized carbon). As a result, the synthesized N-doped CNFs showed excellent electrochemical performance with high reversible capacities of up to 346 mAh g⁻¹ and excellent cycling life of over 10,000 cycles with no visible decrease in capacity when employed as an anode material for SIBs (**Fig. 6h**).

Lithium-sulfur (Li-S) batteries have gained increasing attention in recent years due to their lightweight nature and high energy density. Theoretically, Li-S batteries can deliver a large specific capacity as high as 1670 mA h g⁻¹, and hence, they are capable of exhibiting high energy density of up to 2600 W h kg⁻¹.^{94, 100} In addition, Li-sulfur batteries show other advantages compared to LIBs, such as lower fabrication cost, environmental friendliness, and wide availability of the electrode-active sulfur material. However, there exist some challenges which need to be solved before they can be commercialized for use in electronic devices and hybrid vehicles. These challenges include low electrode utilization, poor high-rate performance, and rapid fade in capacity, due to the poor ionic or electronic conductivity of sulfur and dissolution of the intermediate polysulfides in the electrolyte, also known as the ‘shuttle effect’. Porous carbon materials, e.g., microporous, mesoporous, and macroporous carbon, have been widely investigated as cathode hosts for Li-S batteries. In general, macroporous carbon is not so effective in activating enclosed sulfur and restricting soluble polysulfides, which causes poor long-term stability. In comparison, mesoporous carbon materials have been successfully used to some extent to prepare sulfur cathodes with better long-term cycling stability, however, the

1 electrochemistry of cyclo-S₈ still remains in the systems, and the dissolution and shuttle effect of polysulfides
2 remain a major problem.¹⁰¹ In recent years, microporous carbon materials have been proposed as an ideal
3 cathode host for sulfur.^{102, 103} For instance, the loading of sulfur into microporous carbon fibers with an average
4 pore size of around 1 nm, the catalytic activity and cycling stability of sulfur could be significantly enhanced.¹⁰²
5 Nonetheless, it is still difficult to completely solve the problems arising from the electrochemistry of cyclo-S₈.
6 As such, it is desirable to load sulfur into carbon substrates with hierarchical porosity to enable optimized
7 confinement of sulfur. In such hierarchical porous carbon materials, the micropores will enable the storage of
8 sulfur as chain-like sulfur molecules to ensure high specific capacity and good cycling performance, whereas
9 the presence of mesopores will increase the pore volume of the carbon materials to achieve a high amount of
10 sulfur loading and ensure the smooth transfer of Li ions to promote high rate capability.¹⁰¹

11 MOF-derived microporous carbon polyhedrons (MCPs) encapsulated PAN nanofibers have previously been
12 proposed as effective sulfur immobilizer for Li-S batteries, owing to the possibility to disperse sulfur uniformly
13 inside the nanofibers and in the micropores of MCPs.⁹⁴ The encapsulation process was achieved by
14 electrospinning a mixture of MCPs and PAN, followed by the loading of sulfur via a two-step reaction at 155 °C
15 and at 300 °C, respectively. The optimized S/MCPs-PAN composite (containing 52 wt % sulfur) exhibited an
16 initial large reversible capacity of around 790 mAh g⁻¹, which only slightly decreased to 789.7 mAh g⁻¹ in the
17 second cycle, and the discharge capacity was maintained at 666.2 mAh g⁻¹ after 200 cycles, indicating both high
18 sulfur utilization (90.7%), and good capacity retention (84.4%). The good electrochemical performance of the
19 S/MCPs-PAN-52 composite was attributed to its high structural stability which offers 3D ionic or electronic
20 transport paths and its effective sulfur utilization owing to the existence of micropores. However, when the
21 sulfur content was increased to 65%, the resulting composite showed low utilization of sulfur and poor rate
22 performance due to the formation of a thick solid electrolyte interphase (SEI) layer. These results indicate that
23 the state and content of sulfur can influence the utilization and reversible capacity of sulfur cathode in
24 carbonate-based electrolytes greatly. Specifically, when a low amount of amorphous sulfur is distributed
25 uniformly inside MCPs-PAN nanofibers, the thin SEI layer formed during the discharge process can be reverted
26 back after charging and this is critical for ensuring effective sulfur utilization and good cycling stability of the
27 composite. On the other hand, the presence of a large amount of crystalline sulfur active material can lead to the
28 generation of thick and dense SEI layer on the surface of the nanofibers which consequently leads to poor sulfur
29 utilization and capacity retention.

30 Lithium-selenium (Li-Se) batteries have been considered as alternatives to Li-S batteries with some
31 additional advantages, including higher electrical conductivity, higher energy density, and better cyclability. Li-
32 Se batteries exhibit lower theoretical specific capacity (678 mA h g⁻¹) compared to that of Li-S batteries (1670
33 mA h g⁻¹) due to the smaller mass of selenium compared to sulfur.¹⁰⁴ However, due to the much higher density

1 of Se compared to S, Li-Se batteries can generate a nearly similar volumetric capacity ($3253 \text{ mA h cm}^{-3}$) as Li-S
2 batteries ($3467 \text{ mA h cm}^{-3}$), and therefore, similar energy and power densities. Furthermore, the electrical
3 conductivity of selenium ($1 \times 10^{-3} \text{ S m}^{-1}$) is significantly higher than that of sulfur ($5 \times 10^{-28} \text{ S m}^{-1}$), thus
4 highlighting another advantage of Li-Se batteries over Li-S batteries.⁷² Similar to Li-S batteries, the Se cathode
5 in Li-Se batteries also undergoes volume expansion upon reaction with lithium due to the incapability of the Se
6 lattice structure to accommodate the insertion of Li^+ as result of insufficient space. In addition, the Se cathode
7 also suffers from the dissolution issue of high-order lithium selenides (Li_2Se_x , $x > 4$) in electrolytes, which often
8 leads to low Se utilization, inferior capacity, and poor cycling stability. To mitigate this issue, it is desirable to
9 distribute the electroactive Se materials uniformly in a conductive porous matrix in which there is a sufficient
10 space to accommodate the expansion of the Se cathode. However, at present, it is still challenging to optimize
11 the size of the micropores because very small micropores will lead to poor accommodation of the electroactive
12 material and sluggish Li^+ diffusion.¹⁰⁴ In contrast, larger micropores can facilitate improved Se dissolution
13 during cycling and decrease capacity fading, thereby leading to enhanced electrochemical performance. Ideally,
14 the size of the micropores needs to be larger than 1 nm to enable accommodation of Se_8 molecules. Apart from
15 microporous carbon, mesoporous carbon also can also offer a suitable matrix for accommodating the Se
16 electroactive material to enable stable cycling performance.

17 In order to combine the advantages of micropores and mesopores, activated bimodal porous carbon
18 nanofibers (BP-CNFs) have been prepared by the carbonization of ZIF-8/PAN nanofibers followed by KOH
19 activation for Li-Se batteries.¹⁰⁵ **Fig. 7a** depicts the formation of BP-CNFs homogeneously filled with chain-
20 like Se from the combined electrospinning and carbonization of ZIF-8/PAN nanofibers. In the first step, the
21 PAN/ZIF-8 nanofibers were prepared via electrospinning using a solution containing ZIF-8 nanoparticles and
22 PAN. Following heat treatment at $800 \text{ }^\circ\text{C}$ under argon atmosphere, mesoporous carbon nanofibers (M-CNFs)
23 (**Fig. 7b and 7c**) were achieved from the carbonization of organic linkers in ZIF-8 and PAN, and removal of the
24 remaining Zn metal by etching with hydrochloric acid. To activate the M-CNFs, the etched M-CNFs were
25 reacted with KOH and calcined at $800 \text{ }^\circ\text{C}$ in an argon atmosphere. Finally, the Se-loaded BP-CNFs (BP-
26 CNFs/Se) were synthesized through a melt-diffusion process by mixing the BP-CNFs with Se powder followed
27 by final heating of the resulting mixture under argon atmosphere. The confinement of chain-like Se within the
28 mesopores of BP-CNFs generated by the KOH activation and the presence of graphitic carbon are evident in
29 **Fig. 7d-h**. When evaluated as a cathode host for Li-Se batteries, the BP-CNFs/Se electrode exhibited excellent
30 cycling stability with a high capacity retention of 79.2% after 300 cycles (588 mA h g^{-1}) at a current density of
31 0.5 C (**Fig. 7i**), along with high rate performance. Specifically, the discharge capacities of the BP-CNFs/Se
32 cathode at current densities of 0.5, 1.0, 2.0, 5.0, and 10.0 C were 696, 670, 645, 607, and 568 mAh g^{-1} ,
33 respectively (**Fig. 7j**). In comparison, the discharge capacities of M-CNF/Se greatly decreased from 60 to 1 mA

h g⁻¹ with the increase in current density from 0.5 to 10.0 C. The high discharge capacities and excellent rate performance of the BM-CNFs/Se electrode for Li-Se batteries were attributed to the presence of bimodal pores (mesopores and micropores) generated by the carbonization of ZIF-8 nanoparticles and KOH activation, which promoted high utilization of chain-like selenium in the micropores and ensured easy access of electrolyte through the micropores.

3.3 Fuel cells

In the past few years, polymer electrolyte membrane fuel cells (PEMFC), a new type of sustainable energy conversion technologies, have gained significant attention due to their excellent efficiency, high energy density and negligible emission of harmful gases.¹⁰⁶ The performance of PEMFC mainly depends on several important electrocatalytic reactions, such as oxygen reduction reaction (ORR), hydrogen evolution reaction (HER), and oxygen evolution reaction (OER).¹⁰⁶⁻¹⁰⁸ At present, the major challenge associated with PEMFC is the slow reaction kinetics of ORR at the cathode, in which a large quantity of catalysts is needed to produce sufficient energy output. Currently, Pt-based catalysts are the most commonly used catalysts for ORR.¹⁰⁹⁻¹¹³ However, Pt catalysts exhibit several disadvantages, such as high cost, poor stability, the crossover effect, and CO poisoning.¹¹⁴ These drawbacks have limited the wider commercialization of PEMFCs. As such, it is highly desirable to replace expensive Pt-based catalysts with non-noble metal electrocatalysts which exhibit high catalytic activity, low cost, and excellent long-term cycling stability.

MOF-derived NPCs, especially heteroatom-doped NPCs, with high surface area, high chemical stability, and tunable structure have shown some promise for electrocatalytic reactions.^{39, 41} However, due to the inherent presence of micropores in MOFs, MOF-derived NPCs usually have a micropore-dominated structure which hinders electron and mass transport and therefore, lowers the overall reaction kinetics. 1D porous or hollow carbon nanostructures, such as carbon nanotubes (CNTs) and CNFs have attracted considerable interest for ORR owing to their large surface area, high strength to weight ratio, excellent electrical conductivity and good stability, characteristics which are highly desired for ORR catalysts.⁸⁰ However, pure CNTs or CNFs do not possess sufficiently high catalytic activity toward ORR for fuel cell applications. Therefore, by combining the advantages of MOF-derived NPCs and porous/hollow CNFs, CNFs with hierarchical porosity (mesopores and micropores) and heteroatomic doping, novel catalysts with high electrocatalytic performance for fuel cell reactions can be achieved. In such catalysts, the hierarchical pores are beneficial for enhancing the effective surface area and facilitating mass transportation, while the uniform multiheteroatom doping can enhance the overall conductivity and catalytic performance, such as N-doping which can promote redistribution of atomic charge density and spin density, in nanocarbons, which in turn promotes better oxygen adsorption and reduction.¹⁰⁶

1 For example, nitrogen-enriched carbon fibers with hierarchical micro-mesoporous structures (NEMCF)
2 were previously synthesized through the in-situ confinement pyrolysis of electrospun ZIF-8/PVP fibers at 900
3 °C, which showed high catalytic activity for ORR.⁸⁴ The origin of the mesopores in the fabricated NEMCF was
4 attributed to the strong cohesive interface interaction between PVP-derived carbon and ZIF-8 polyhedrons,
5 which provided a significant confinement effect to maintain the ZIF-8-derived carbon adjacent to the internal
6 surface of PVP-derived carbon during the pyrolysis process, thereby shrinking the ZIF-8-based framework from
7 center to outside, leading to the formation of uniform mesoporous structure. Benefiting for the hierarchical
8 meso-microporous structure and the rich nitrogen doping (6.06 at. %), the 1D NEMCF showed slightly higher
9 catalytic activity together with enhanced stability and methanol resistance compared to commercial 30 wt%
10 Pt/C catalyst in 0.1 M KOH. Specifically, the hierarchical porous structure allowed for rapid mass transfer and
11 persistent exposure of active sites, avoiding the temporary or permanent deactivation of these sites due to the
12 blocking or collapse of the catalyst pores. In addition, the relatively high proportions of pyridinic and graphitic
13 N sites on the surface of the NEMCF resulted in high electron conductivity. In general, the graphitic N
14 determines the limiting current density, whereas the pyridinic N is likely to improve the onset potential due to
15 the conversion of the reaction route from a $2e^-$ -dominated process to a $4e^-$ -dominated process. The study further
16 revealed that the types of the active sites and their space distribution are more important in determining the
17 ORR activity and selectivity of the NEMCF than the total nitrogen content.

18 In order to enhance the electrocatalytic activity of MOF-derived 1D PCNFs or HCNFs for ORR,
19 multiheteroatomic doping (e.g., N, Co, P, B, S, etc.) is often implemented to modify their surface polarity and
20 electronic properties of nanocarbons. For instance, Jia and co-workers reported a simple one-step method for
21 the tunable synthesis of Co–N doped carbon microsphere/nanofiber hybrids (CSFHs) via electrospinning of
22 ZIF-67 and PAN and demonstrated the possible morphology control from a pearl necklace-like nanofiber to a
23 microsphere/nanofibers 3D structure by controlling the ratio of ZIF-67/PAN.¹¹⁵ When evaluated for ORR, the
24 CSFHs obtained with ZIF-67/PAN ratio of 8:1 and calcined at 800 °C showed comparable catalytic activity to
25 commercial Pt/C electrode, as indicated by onset potential (E_{onset}) of 0.845 V and half-wave potential ($E_{1/2}$) of
26 0.794 V. Furthermore, the CSFH electrode exhibited a Tafel slope of 150 mV per decade, better than the Pt/C
27 electrode (161 mV per decade), indicating its high kinetic activity for ORR. Meanwhile, the H_2O_2 yield and
28 electron transfer number (n) measured using the CSFH electrode were below 3.0% and *ca.* 3.99 at all potentials
29 in 0.5 M H_2SO_4 , which are relatively similar to those of the Pt/C electrode. The excellent ORR activity of the
30 CSFHs was attributed to the synergistic effects of the high surface area, the large quantity of micropores, and
31 Co-N doping.

32 N, Co-containing MOF-based hierarchical carbon fibers have previously been synthesized by incorporating
33 bimetallic Zn, Co-ZIF with electrospun Co^{2+} /PAN fibers (**Fig. 8a and 8b**). The resulting Zn, Co-ZIF based

1 carbon fibers (ZCP-CFs) inherited the merits of carbons from both MOFs and PAN fibers and they exhibited
2 several desirable characteristics for ORR, including large surface areas, high degree of graphitization, highly
3 dispersed N and Co active species and excellent electrical conductivity. As a result of these advantageous
4 features, the ZCP-CFs displayed high activity for ORR in 0.1 M KOH, including more positive half-wave
5 potential (0.135 V vs. Ag/AgCl), higher diffusion-limited current density (5.95 mA cm⁻²) and kinetic-limiting
6 currents, lower Tafel slope (62 mV dec⁻¹) higher selectivity (number of electron transferred, $n = 3.97$) compared
7 to Zn, Co-ZIF derived carbon (ZIF-C). and Zn, Co-ZIF free carbon fiber (CP-CFs) (**Fig. 8c and 8d**). More
8 notably, the ZCP-CFs catalyst exhibited very close ORR activity to the commercial Pt/C catalyst, but with
9 improved durability and better tolerance to methanol crossover. The above studies clearly indicate that high
10 surface area, high electrical conductivity by Co and N-doping, and hierarchical porosity are essential for
11 enhancing the ORR activity of MOF-derived carbon materials.

12 In another report, multiheteroatom-doped (N, B, P, and S heteroatom-doped) hierarchical porous carbon
13 nanofibers (denoted as carbon NBPS) synthesized using ZIF-8 coated worm-like micelles have been
14 investigated as catalyst for ORR.¹¹⁶ To fabricate such complex multiheteroatom-doped porous CNFs, polymeric
15 worm-like micelles were initially prepared through the self-assembly of poly(ethylene glycol)-*block*-poly(4-
16 vinylpyridine) (PEG₁₁₃-*b*-P4VP₁₁₈) micelles and DNA followed by cross-linking with 1, 4-dibromobutane. In
17 this case, the polymeric worm-like micelles serve as a source of multiheteroatomic doping and hierarchical
18 porosity, which contributed the excellent textural characteristics of the porous CNFs following carbonization.
19 Subsequently, 3D network of ZIF-8 nanofibers was obtained through the in-situ crystallization of ZIF-8 on the
20 surface of the polymeric worm-like micelles, which exhibited hierarchical porosity, including macropores,
21 mesopores, and micropores. Such network formation was promoted by the likelihood of the coated ZIF-8
22 particles to fuse and interconnect together to form cross-linking points. The doping of the hierarchical porous
23 CNFs with B, P, and S was achieved by dispersing in solutions containing B₄O₇²⁻, HPO₄²⁻, and SO₄²⁻,
24 respectively, and the absorption rate of these ions could be controlled to allow concentration tuning of these
25 dopants. When evaluated as a catalyst for ORR, the carbon NBPS showed an onset potential of -0.086 V,
26 comparable to that of commercial 20% Pt/C catalyst (-0.074 V), but with superior methanol tolerance and long-
27 term durability. Specifically, the order of ORR activity was revealed to be: 20% Pt/C > carbon NBPS > carbon
28 NBP > carbon NBS = carbon NPS > carbon NS > carbon NB > carbon NP > carbon N. The excellent catalytic
29 activity of the carbon NPBS was largely contributed by the existence of multiheteroatomic doping and the
30 hierarchical pores with abundant micropores, mesopores, and macropores which facilitated the fast mass
31 transportation of reactants and provided extensive contact between reactants and active sites as well as the 1D
32 carbon nanofibrous network which promoted improved electron transfer.

33 In recent years, research on the development of bifunctional catalysts for water splitting is rapidly

1 increasing. Molybdenum disulfide (MoS_2) is an efficient catalyst for HER, yet its low electrical conductivity
2 restricts its charge transfer rate and electrocatalytic performance. To address this issue, MoS_2 nanosheets were
3 grown on ZIF-67/PAN-derived carbon fibers (CoNC@MoS_2) to improve their electrical conductivity, as
4 schematically illustrated in **Fig. 8e**.¹¹⁷ In order to obtain CoNC@MoS_2 , uniform PAN nanofiber films were
5 initially prepared through electrospinning technique. Next, leaf-like ZIFs (ZIF-L) were directly grown on PAN
6 nanofibers through a simple liquid-phase deposition method at room temperature. Then, the MOF particles on
7 the nanofibers were carbonized at 800 °C under nitrogen atmosphere to form Co, N-doped carbon flakes grafted
8 on CNF films (CoNC/CNF). Finally, MoS_2 nanostructures were grown on the surface of the leaf-like carbon
9 flakes through a facile solvothermal treatment with ammonium thiomolybdate ($(\text{NH}_4)_2\text{MoS}_4$) at 200 °C, leading
10 to the formation of $\text{CoNC@MoS}_2/\text{CNF}$ films. The resulting $\text{CoNC@MoS}_2/\text{CNF}$ films exhibited unique
11 hierarchical architectures with interconnected vine-like CNFs, which imparted good flexibility and excellent
12 electrical conductivity for both HER and OER.

13 For HER measured in 1 M KOH electrolyte, the $\text{CoNC@MoS}_2/\text{CNFs}$ displayed remarkable HER catalytic
14 activity, in which the overpotential required to generate a current density of 10 mA cm^{-2} (η_{10}) was 143 mV,
15 superior than those of CNF, CoNC/CNF , and MoS_2/CNF (673 mV, 425 mV and 257 mV, respectively).
16 Furthermore, the $\text{CoNC@MoS}_2/\text{CNF}$ electrode delivered a low Tafel slope of 68 mV, which was superior to
17 those of CoNC/CNF (79 mV^{-1}) and MoS_2/CNF (89 mV^{-1}), indicating its fast HER catalytic kinetics.
18 Furthermore, when employed in OER in 1 M KOH electrolyte, the $\text{CoNC@MoS}_2/\text{CNFs}$ displayed superior
19 catalytic activity with a low η_{10} value of 350 mV, which was superior to those of MoS_2/CNF and CoNC/CNF
20 (570 mV and 430 mV, respectively) (**Fig. 8f**). In addition, the $\text{CoNC@MoS}_2/\text{CNFs}$ also exhibited a low Tafel
21 slope of 51.9 mV^{-1} , which was significantly lower than those of CoNC/CNF (89.7 mV^{-1}), MoS_2/CNF (119.3
22 mV^{-1}), and RuO_2 (98.1 mV^{-1}) (**Fig. 8g**), thereby indicating the enhanced catalytic kinetics and high catalytic
23 activity of $\text{CoNC@MoS}_2/\text{CNFs}$. Moreover, an increase of only 17 mV in the value of η_{100} was observed after
24 1500 continuous cycles (**Fig. 8h**), indicating the excellent OER stability of $\text{CoNC@MoS}_2/\text{CNFs}$ in a basic
25 medium. The enhanced HER and OER activities of the $\text{CoNC@MoS}_2/\text{CNFs}$ were attributed to the presence of
26 hierarchically porous carbon fibers which not only allowed rapid mass transport but also provide a ‘highway’
27 for fast electron transfer, thereby confirming the superiority of MOF-derived HCNFs. The above studies clearly
28 highlight the potential of hybrid nanostructures assembled from MOFs and CNFs as efficient self-supported
29 electrocatalysts, thus opening new possibilities into the development of functional free-standing materials for
30 fuel cell applications.

4. Conclusions and prospects

Electrospun MOF-derived 1D hollow carbon nanofibers (HCNFs) have emerged as promising materials for energy storage and conversion applications. HCNFs can overcome the limitations of MOF-derived NPCs, so that long cycle life, high capacity/capacitance and/or high catalytic activity can be realized. In this mini review, the preparation method, the formation mechanism, and the structure-activity relationship of MOF-derived HCNFs have been summarized. From our survey of the literatures, it is deduced that the composition of the MOFs, the thermal treatment conditions, and post-treatment are critical factors which need to be considered to obtain HCNFs with large surface area, high electrical conductivity and hierarchical porosity, which are beneficial for enhancing their electrochemical performance for EES applications. Despite the rapid progress on MOF-derived HCNFs for EES applications, this research field requires further developments. More efforts should be devoted to realizing the practical applications of HCNFs by focusing on the following points: (a) Many previous works were focused mainly on the synthesis of MOF-derived HCNFs and investigation of their supercapacitive or electrocatalytic performance, thus more theoretical studies are needed to provide more insights into the structure-activity relationship; (b) Previously reported studies were focused mainly on ZIF-derived HCNFs, whereas other common MOFs, such as MIL and Prussian blue (PB) series are far less explored, hence future studies should attempt to explore the potential of these MOFs as precursors for the derivation of HCNFs; (c) Future works should address the challenges in assembling MOF-derived HCNFs into flexible or wearable energy storage devices. Overall, the electrospinning method has provided a new and exciting avenue for fabricating functional 1D hollow carbon materials from MOFs, especially for clean energy utilization. It is expected that the electrochemical performance of HCNFs for energy storage and conversion applications may be further improved in the future by doping with several active elements or forming heterostructures with other conductive inorganic or organic nanomaterials. We anticipate that this mini review may provide researchers with some guidance for developing new MOF-based electrochemical devices in the future.

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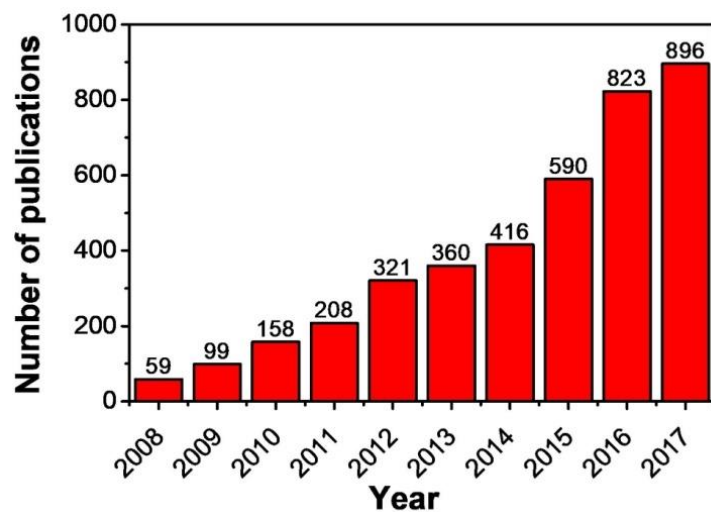
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1 **LIST OF FIGURES**

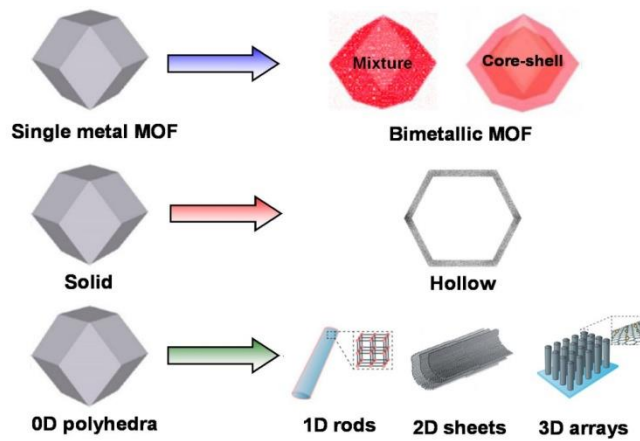
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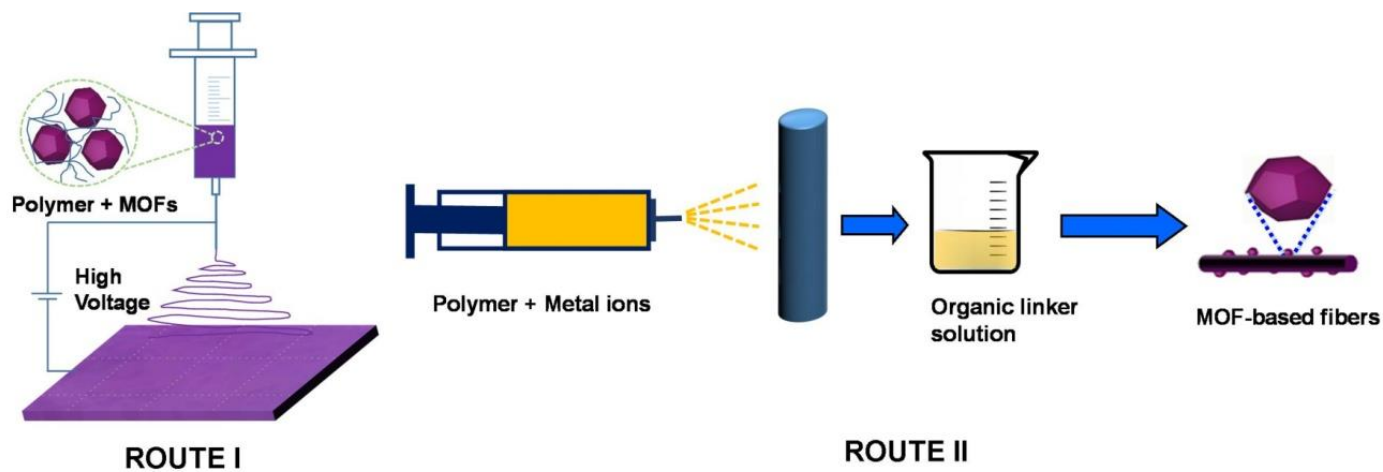
4 **Fig. 1.** Number of articles published on MOF-derived carbon materials over the last 10 years according to SciFinder
5 Scholar.

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Fig. 2. Schematic diagram showing the development of MOF-derived nanomaterials with various dimensionalities and structures.



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Fig. 3. Schematic illustration of the preparation of MOF-based nanofibers by electrospinning: *Route I* - mixing pre-fabricated MOF nanoparticles with polymer followed by electrospinning; *Route II* - electrospun polymer nanofibers (containing metal ions) followed by in-situ growth of MOF particles.

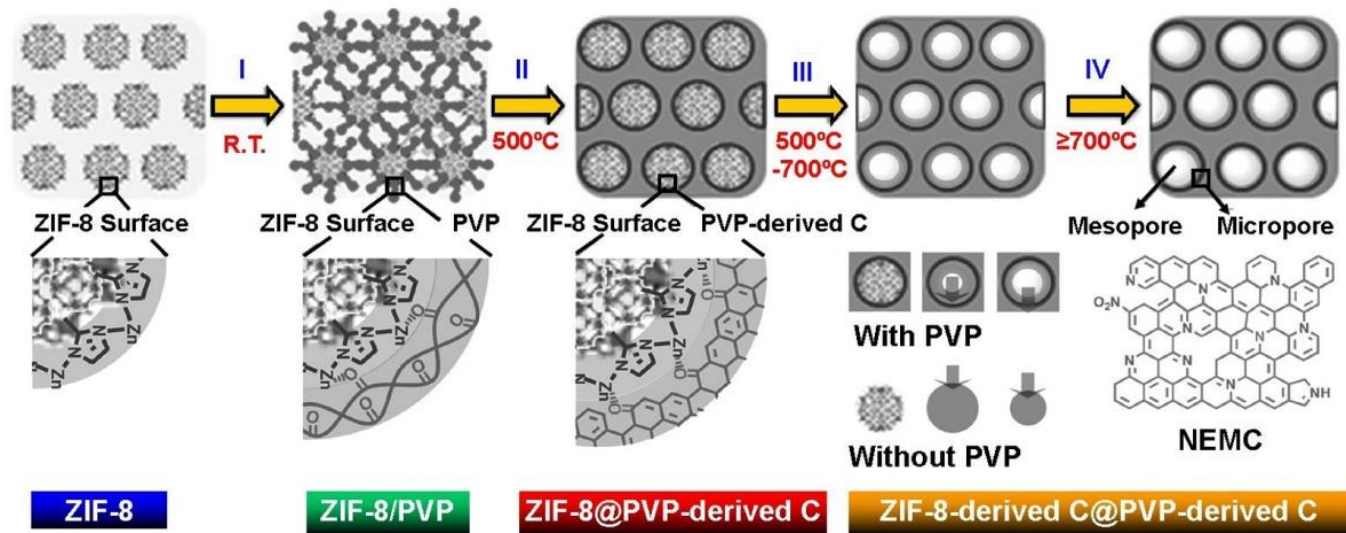
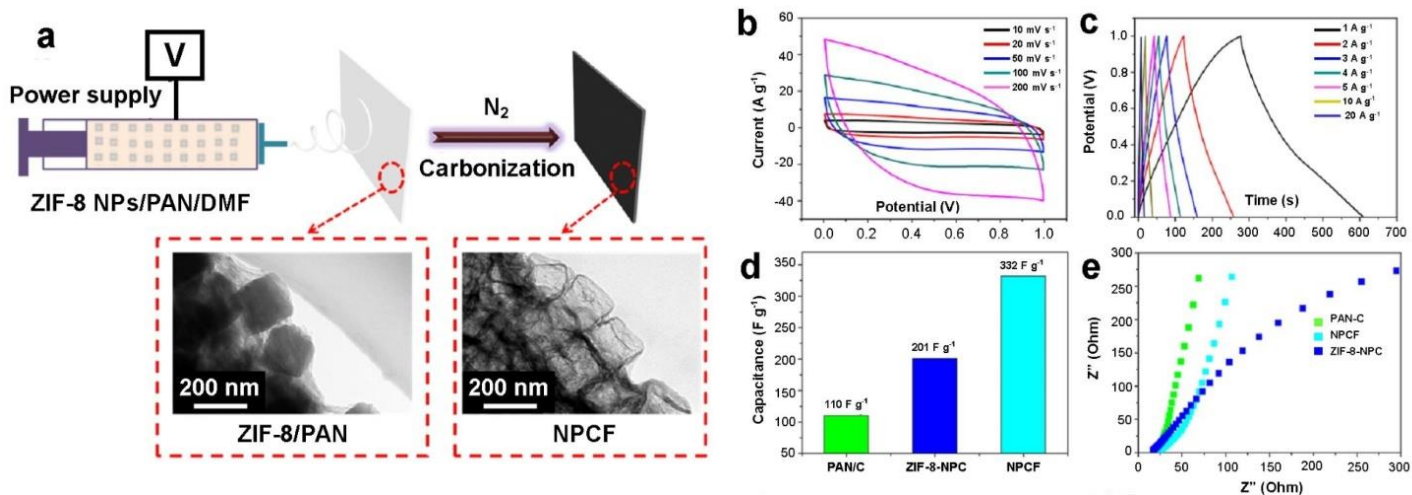


Fig. 4. Formation mechanism of hollow carbon nanofibers (HCNFs) from zeolitic imidazolate framework-8 (ZIF-8)/polyacrylonitrile (PAN) nanofibers.⁸⁴ Copyright 2016, Wiley-VCH.



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 2 **Fig. 5.** (a) Preparation process of nanoporous carbon fibers (NPCFs). (b) CV curves and (d) galvanostatic charge-
 3 discharge curves of NPCFs. (e) Capacitance and (f) Nyquist electrochemical impedance spectra of PAN-derived carbon
 4 (PAN-C), ZIF-8 derived nanoporous carbon (ZIF-8-NPC), and NPCFs.⁷⁹ Copyright 2017, The Royal Society of
 5 Chemistry.

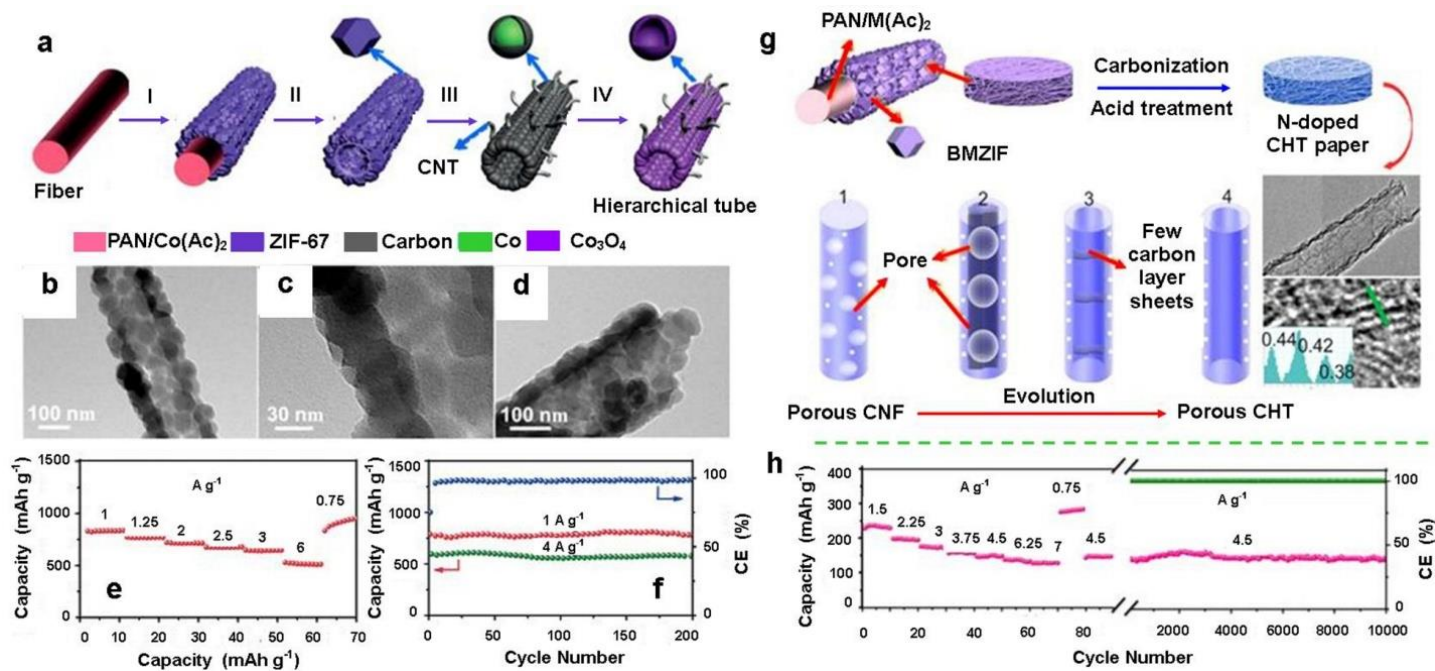


Fig. 6. (a) Schematic illustration of the formation of ZIF-67/PAN-derived hierarchical CNT/Co₃O₄ microtubes. (b-d) TEM images of the synthesized ZIF-67 microtubes. Electrochemical properties of the hierarchical CNT/Co₃O₄ microtubes for lithium-ion batteries (LIBs): (e) rate performance and (f) cycling performance and Coulombic efficiency.⁹⁸ Copyright 2016, Wiley-VCH. (g) Schematic illustration of the synthesis of a series of 1D nanocarbons from porous N-doped carbon fiber networks (CFNs) to carbon hollow tubes (CHTs) with enlarged graphene interlayer spacing. (h) Rate performance and cycling stability of the porous N-doped carbon hollow tubular structures (CHTs) for LIBs.⁹⁹ Copyright 2017, Cell Press.

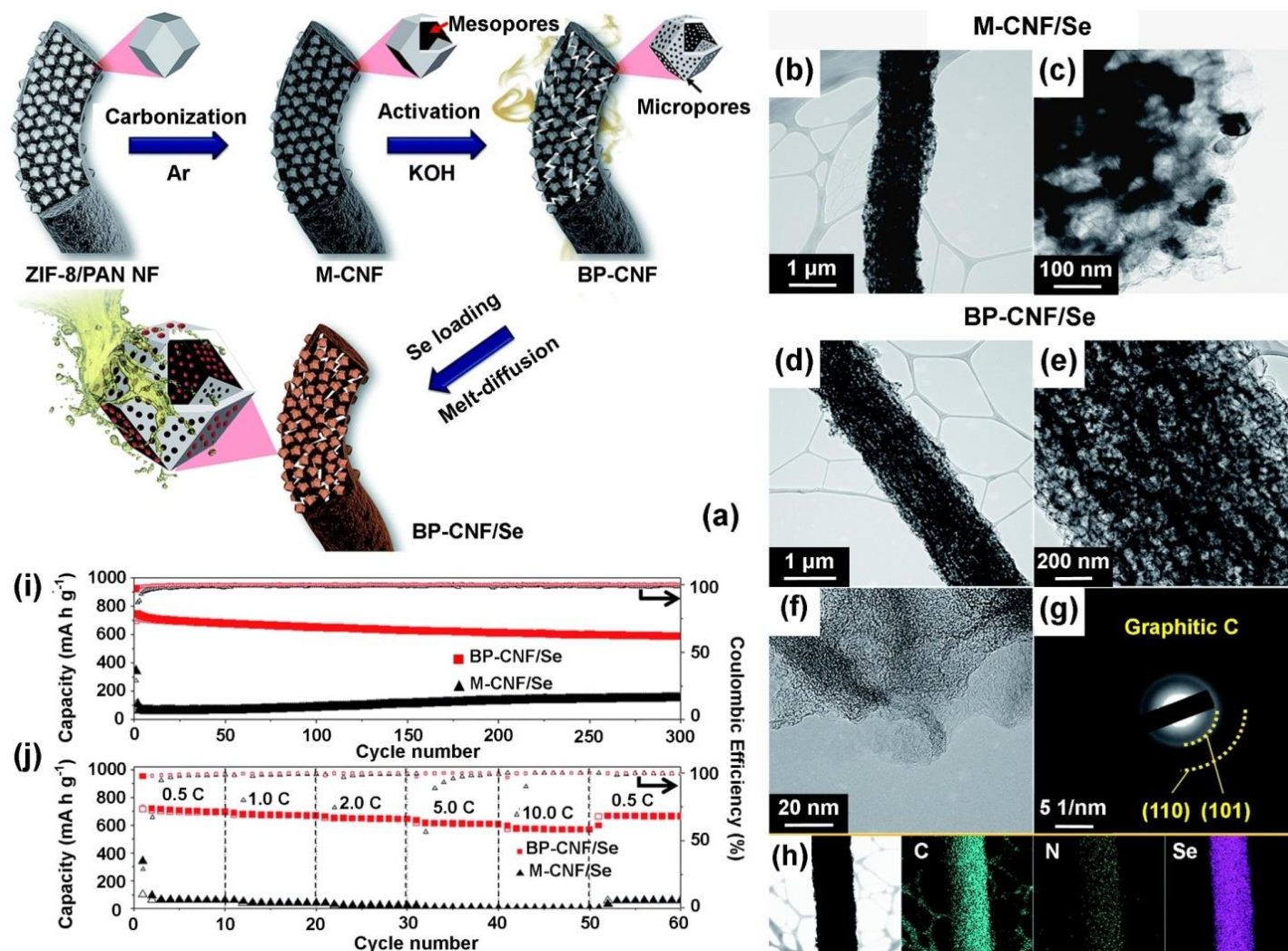


Fig. 7. (a) Formation mechanism of bimodal porous nitrogen-doped carbon nanofiber (BP-CNF) homogeneously filled with chain-like Se. (b, c) TEM images of the mesoporous carbon nanofibers (M-CNF). (d, e) TEM images, (f) HRTEM image, (g) SAED pattern, and (h) elemental mapping images of Se-loaded BP-CNF (BP-CNF/Se). Electrochemical properties of BP-CNF/Se and Se-loaded M-CNF (M-CNF/Se) for lithium-selenium (Li-Se) batteries: (i) cycling performance at a current density of 0.5 C and (j) rate performances with the Coulombic efficiencies¹⁰⁵ Copyright 2017, The Royal Society of Chemistry.

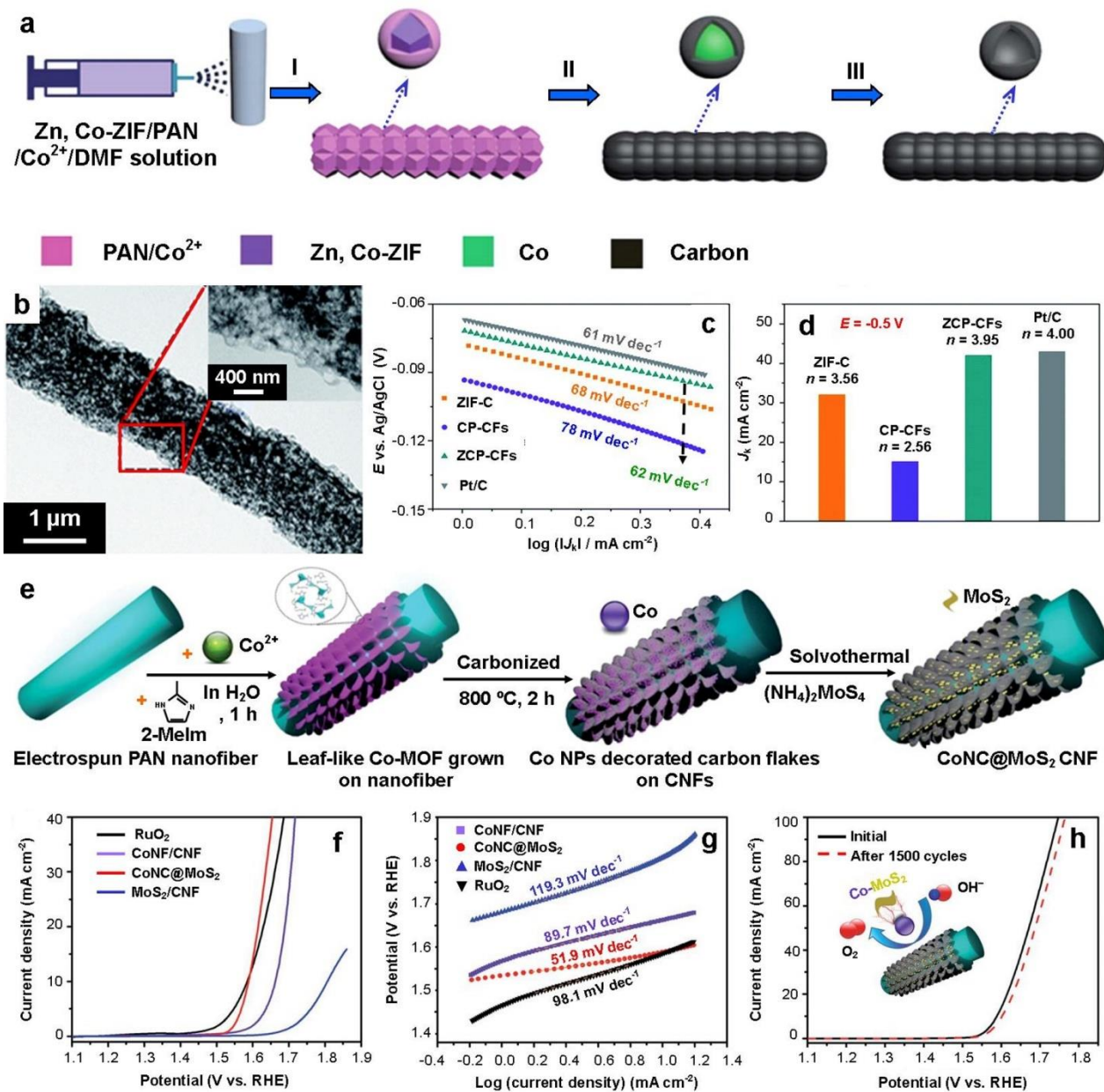


Fig. 8. (a) Schematic illustration of the preparation of the Zn, Co-ZIF-based hierarchical carbon fibers (ZCP-CFs) and (b) the corresponding TEM image of ZCP-CFs. (c) Tafel plots of ZIF-C, CP-CFs, ZCP-CFs and Pt/C. (d) Kinetic-limiting current density of ZIF-C, CP-CFs, ZCP-CFs-9 and Pt/C.⁸⁰ Copyright 2017, The Royal Society of Chemistry. (e) Schematic illustration of the fabrication of free-standing MoS₂ nanosheets grafted Co-N-doped carbon flakes grown on electrospun carbon nanofibers (CoNC@MoS₂/CNF). (f) Polarization curves of CoNC@MoS₂/CNF, MoS₂/CNF, CoNC/CNF, and RuO₂ in 1 M KOH in the OER. (g) OER Tafel plots of CoNC@MoS₂/CNF, MoS₂/CNF, CoNC/CNF, and RuO₂ in 1 M KOH. (h) OER cycling stability of CoNC@MoS₂/CNFs in 1 M KOH.¹¹⁷ Copyright 2017, The Royal Society of Chemistry.

1 **TABLE OF CONTENT DESCRIPTION**

2 This review summarizes the advances in the derivation of one-dimensional porous and hollow carbon nanofibers from
3 metal-organic frameworks for energy storage and conversion.

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