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Published in:

Biomass and Bioenergy

10.1016/j.biombioe.2022.106676

Publication date:

2023

Document Version Publisher's PDF, also known as Version of record

#### Link to publication

Citation for pulished version (HARVARD):

Cao, Y, Sun, Y, Zheng, R, Wang, Q, Li, X, Wei, H, Wang, L, Li, Z, Wang, F & Han, N 2023, 'Biomass-derived carbon material as efficient electrocatalysts for the oxygen reduction reaction', Biomass and Bioenergy, vol. 168, 106676. https://doi.org/10.1016/j.biombioe.2022.106676

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## Biomass and Bioenergy

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# Biomass-derived carbon material as efficient electrocatalysts for the oxygen reduction reaction

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#### ARTICLE INFO

#### Keywords: Oxygen reduction Biomass Carbon Catalyst

#### ABSTRACT

Despite the abundance of carbon in nature, a significant portion of the existing biomass carbon materials in livestock, agriculture, and marine fishery industry are currently being wasted. Utilizing sustainable carbon materials as an alternative to noble Pt-based catalysts is crucial step to convert widely available and low-cost biomass resources into clean energy systems. Therefore, the rational synthesis of carbon-based catalysts for oxygen reduction reaction (ORR) has become a hot research focus in the field of electrochemistry. In this study, the recent progress in the synthesis of ORR electrocatalysts using sustainable biomass resources was reviewed; the activation and synthesis strategies of various biomass resources, as well as the microstructure and oxygen reduction performance of the prepared carbon-based catalysts were investigated. It is hoped that this review article will promote the understanding of various parameters from biomass as precursors for catalyst preparation and make contribute to the transition of biomass resources from the wasted carbon materials to the main catalysts in future energy devices.

#### 1. Introduction

In view of the growing energy demand and the increasing concerns about environmental issues, significant importance has been attached to the renewable and clean energy [1-5]. Among the sustainable energy resources, wind energy and solar energy are of most abundance and easy availability [6,7]. However, such energy resources are highly dependent on climate, leading to great challenges for the efficient utilization of them [8-10]. Hence, in order to realize the efficient utilization of clean energy, it is urgent to develop fuel cell, battery, supercapacitor and other novel energy equipment [11-14], among which fuel cells have attracted great attention because of their wide applications [15-18]. ORR, as the core of fuel cell devices, determines the efficiency of energy conversion and storage [19-24]. However, ORR is slow in operation, and it requires precious metal catalysts (e.g., platinum) [25-29]. Thus, the scarcity,

poor stability and high cost of Pt-based ORR catalysts have hindered the commercial application of fuel cells [30–34]. Therefore, it is crucial to find alternative ORR catalysts which are not only earth-abundant and low cost, but also of high catalytic activity [35–39]. Porous carbon materials have been studied widely because of their excellent electrical conductivity [40,41], tunable morphology [42], high surface area [43], abundant precursors [8], and chemical and structural stability [44]. Despite the promising catalytic activities of carbon nanotubes, graphene, and other carbon materials, their large-scale commercialization is limited due to the excessive cost and synthesis processes [45–47]. Therefore, the advanced carbon-based catalysts derived from biomass resources have received extensive attention (see Table 1).

Carbon can combine with other elements such as nitrogen, oxygen, hydrogen and sulfur to provide energy and structural tissue for living organisms in the biosphere [48,49]. Biomass resources are classified into

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**Table 1**ORR electrocatalysts obtained from biomass.

Biomass source	Other precursors	Synthesis strategy	Surface area (m <sup>2</sup> g <sup>-1</sup> )	Pore volume (cm <sup>3</sup> g <sup>-1</sup> )	$E_{ m Onset}$	$E_{1/2}$	Electrolyte	Reference
Shrimp shells	Pyrrole and FeCl <sub>3</sub>	HTC, Pyrolysis	806.7	0.205	-0.017 V νs. Ag/AgCl	−0.15 V vs. Ag/AgCl	0.1 M KOH	[80]
Soybean	ZnCl <sub>2</sub>	Activation, pyrolysis and graphitization	949	_	-0.02 V vs. Ag/AgCl	-0.12 V vs. Ag/AgCl	0.1 M KOH	[93]
Shrimp shells	-	HTC, Silica templating and pyrolysis	360.2	-	-0.06 V vs. Ag/AgCl	-0.21 V vs. Ag/AgCl	0.1 M KOH	[101]
Duck blood	FeCl₃·6H₂O	Pyrolysis, ball milling and pyrolysis	-	-	0.002 V vs. Ag/AgCl	0.137 V vs. Ag/AgCl	0.1 M KOH	[103]
Pig blood (PB)	$\rm FeCl_3{\cdot}6H_2O$	Carbonization, ball milling, acid leaching and pyrolysis	641	_	-0.04 V vs. Ag/AgCl	-0.16 V vs. Ag/AgCl	0.1 M KOH	[51]
Blood protein	Vulcan XC-72R	Pyrolysis, ball milling, pyrolysis, acid leaching and pyrolysis again	93	0.18	0.90 V vs. RHE	0.78 V vs. RHE	0.1 M KOH	[107]
Cattle bone	Co <sub>3</sub> O <sub>4</sub>	Precarbonized, mixed with Co (OAc) <sub>2</sub> , centrifuging, washing and drying	1070	-	0.960 V vs. RHE	0.853 V <i>vs</i> . RHE	0.1 M KOH	[108]
Eggs	-	Stirring with TEOs, precarbonized and anneal	970	-	0.84 V vs. RHE	0.69 V vs. RHE	0.1 M KOH	[112]
Eggs	FeCl <sub>3</sub> ·6H <sub>2</sub> O, multi-walled carbon nanotubes (MWNT)	Mixed with FeCl <sub>3</sub> ·6H <sub>2</sub> O and MWNT, pyrolysis and pyrolysis again	680	-	-0.007 V vs. Ag/AgCl	-0.133 V vs. Ag/AgCl	0.1 M KOH	[113]
Egg-yolk	$g$ - $C_3N_4$	Grind and pyrolysis	447	-	0.89 V vs. RHE	-	0.1 M KOH	[114]
Human hair	NaOH	Precarbonized, activation and pyrolysis	1814	-	-0.016 V vs. Ag/AgCl	-	0.1 M KOH	[116]
Human urine	-	Drying and pyrolysis	811	_	-0.03 V vs. Ag/AgCl	_	0.1 M KOH	[117]
Biomass source	Other precursors	Synthesis strategy	Surface area (m <sup>2</sup> g <sup>-1</sup> )	Pore volume (cm <sup>3</sup> g <sup>-1</sup> )	E <sub>Onset</sub>	$E_{1/2}$	Electrolyte	Reference
Porphyra	$\rm FeCl_3{\cdot}6H_2O$	Grind and pyrolysis	1533.7	0.39	0.96 V vs. RHE	0.84 V <i>vs.</i> RHE	0.1 M KOH	[118]
Porphyra	Hemin	Activation and carbonization and pyrolysis	1301.5	197.6	-	0.87 V vs. RHE	0.1 M KOH	[119]
Catkins	FeCl <sub>3</sub> , KOH and melamine	Stir, pyrolysis and acid leaching	461.5	-	−0.098 V vs. Ag/AgCl	−0.194 V vs. Ag/AgCl	0.1 M KOH	[120]
Гоfu	Na <sub>2</sub> PdCl <sub>4</sub> , HAuCl <sub>4</sub> and H <sub>2</sub> PtCl <sub>6</sub> ·6H <sub>2</sub> O	Activation, carbonization and pyrolysis	3814	2.08	1.04 V <i>vs</i> . RHE	0.91 V vs. RHE	0.1 M HClO <sub>4</sub>	[106]
Гоfu	Urea and NaCl	Pyrolysis	988.2	0.689	0.96 V <i>vs</i> . RHE	0.81 V vs. RHE	0.1 M KOH	[123]
Pomelo peel	Co(NO <sub>3</sub> ) <sub>2</sub> ·6H <sub>2</sub> O and KOH	Activation and pyrolysis	-	1.28	0.87 V <i>vs</i> . RHE	0.78 V <i>vs</i> . RHE	0.1 M KOH	[125]
Peanut shells	Co(OH) <sub>2</sub> and NaH <sub>2</sub> PO <sub>2</sub>	Precarbonized, CO <sub>2</sub> reduction	671.4	_	0.91 V vs. RHE	0.81 V <i>vs</i> . RHE	0.1 M KOH	[74]
Coconut shell	NH <sub>4</sub> F and dicyandiamide	Carbonization, activation and pyrolysis	1028.3	0.83	0.985 V vs. RHE	0.834 V vs. RHE	0.1 M KOH	[131]
Corn silk	FeCl <sub>3</sub> and KOH	HTC, precarbonized and pyrolysis	1038.9	0.83	0.957 V vs. RHE	0.852 V vs. RHE	0.1 M KOH	[133]
o-Glucose and Cellulose	Soya bean flour	HTC, freeze drying and pyrolysis	449	0.25	−0.06 V <i>vs.</i> RHE	-	0.1 M KOH	[138]
Glucose	NH <sub>3</sub> , and thiophene	Silica templating, pyrolysis, heteroatoms doping, ammoxidation	1023	0.973	0.92 V vs. RHE	-	0.1 M KOH	[139]
Glucose	Dicyandiamide and FeSO <sub>4</sub> ·7H <sub>2</sub> O	Silica templating and pyrolysis	906.3	2.89	0 V vs. Ag/ AgCl	−0.15 V vs. Ag/AgCl	0.1 M KOH	[140]
Glucose	Melamine and FeSO <sub>4</sub> ·7H <sub>2</sub> O	Pyrolysis	393	0.99	0.874 V vs. RHE	-	0.1 M KOH	[141]
Glucose	Prussian blue (PB)	Grind and pyrolysis	418	-	−0.055 V vs. Ag/AgCl	−0.152 V vs. Ag/AgCl	0.1 M KOH	[142]
Corn starch	Urea	Pyrolysis	1568.85	-	−0.03 V vs. Ag/AgCl	-0.15 V vs. Ag/AgCl	0.1 M KOH	[143]
Starch	Flake graphite, FeCl <sub>3</sub> and KSCN	HTC and pyrolysis	749.5		0.95 V vs. RHE	0.83 V vs. RHE	0.1 M KOH	[63]
Biomass source	Other precursors	Synthesis strategy	Surface area (m <sup>2</sup> g <sup>-1</sup> )	Pore volume (cm <sup>3</sup> g <sup>-1</sup> )	E <sub>Onset</sub>	E <sub>1/2</sub>	Electrolyte	Referenc
Chitin	-	Pyrolysis	526	-	0.054 V vs. Ag/AgCl	-	0.1 M KOH	[145]
Chitin	$ZnCl_2$	HTC, pyrolysis, activation and pyrolysis	300.7	0.31	-0.15 V vs. SCE	-	0.1 M KOH	[148]
Chitosan	Fe(NO <sub>3</sub> ) <sub>3</sub> ·9H <sub>2</sub> O, TiO <sub>2</sub> and NaOH	Stirring, pyrolysis and wash	1190	1.10	-0.01 V vs. Ag/AgCl	-	0.1 M KOH	[149]
Chitosan	FeCl <sub>3</sub>	Pyrolysis	543	-	-0.08 V vs. Ag/AgCl	-	0.1 M KOH	[150]

(continued on next page)

animal biomass, plant (lignocellulose) biomass, as well as carbohydrates and polysaccharides [50]. Animal biomass includes animal waste, such as blood [51], excrement, feathers [52,53], and human feces, hair [54], and so on. Lignocellulose biomass, including soybean shell [55], eggplant [56], seaweed [57], leaves and grasses [58] and so on, is mainly generated through photosynthesis [59]. Carbohydrates and polysaccharides mainly include fructose [60,61], glucose [62] and its derivatives, starch [63], chitin [64], etc. Large amounts of biomass from human activities, agriculture, and marine fishing are wasted, which causes certain ecological impacts, greenhouse gas emissions and an increasing waste-management burden [65-67]. Therefore, it makes sense to convert sustainable biomass into advanced carbon-based materials. In addition, such advanced carbon-based materials play a significant role as oxygen reduction catalysts in new energy technologies [33,68-72]. The natural structure and rich elemental composition of biomass materials provide the carbon-based catalysts with porous structures [73–75] as well as abundant heteroatom doping [8], and also they can fully expose the active sites [76], which can enhance the catalytic activity. Therefore, numerous studies have focused on porous carbon-based materials and their applications in ORR [156–158]. Herein, we aspire to promote the selection of targeted biomass resources and the optimization of carbon-based material structures, so as to accelerate the development of highly active oxygen reduction catalysts.

Here, we outline the recent progress in oxygen reduction catalysts using biomass resources as precursors. Sections 2 and 3 describe in detail the synthetic strategies of matter-derived carbon nanomaterials including activated pore-making strategies, the physicochemical structures and oxygen reduction activity from plants, animals and Carbohydrates and polysaccharides. Finally, we offered personal views and recommendations on the synthesis of biomass carbon.

#### 2. Synthetic method

#### 2.1. Hydrothermal carbonization

Hydrothermal carbonization (HTC) is an environmentally friendly preparation method that converts biomass to practical carbon materials through dehydration, condensation, polymerization and other processes [50,77–79]. HTC is carried out in a mild aqueous medium below 200 °C, and the prepared carbon materials need to be further activated and pyrolysis to increase their pore volume and specific surface area (SSA).

Zhang et al. prepared N-doped carbon nanodots (N-CNs) using shrimp shells as precursors by a simple hydrothermal carbonization process, and then polymerized pyrrole in FeCl $_3$  solution and to obtain carbonized at high temperature to obtain Fe–N-PGC-800 (Fig. 1A) [80]. It can be clearly seen that pyrolysis in high temperature results in the conversion of pyrrole nitrogen to pyridine nitrogen. The prepared Fe–N-PGC-800 material exhibited a large SSA of 806.7 m $^2$  g $^{-1}$  and presented a mesoporous structure, which contributed to the improvement of ORR performance. Gao et al. used willow leaf as carbon source to obtain bifunctional N-doped carbon quantum dots (N-CDs) by a green and environmentally friendly hydrothermal carbonization process (Fig. 1B) [81]. According to TEM images, N-CDs was very uniform with a diameter of about 2–4 nm. The supernatant (SP1) showed intense fluorescence under ultraviolet irradiation, which could be used as fluorescent ink. The catalyst (SP2) by pyrolysis had outstanding ORR

activity and good durability.

#### 2.2. Chemical and physical activation

Chemical activation usually involves impregnating biomass precursors with  $\rm K_2CO_3$ , KOH, NaOH and ZnCl<sub>2</sub> solutions, etc., followed by pyrolysis and final pickling to remove chemical reagents [82–86]. The synthesis of porous carbon materials using the physical activation method is divided into two steps. First, initial carbonization is performed in an inert environment such as  $\rm N_2$  and Ar, and then activated at a high temperature of 600–1000 °C using  $\rm N_2$ , CO<sub>2</sub>, etc [73,87–90]. The activation method can be very successful in increasing the surface area and pore volume of biomass carbon materials, and their porosity can be well controlled [91].

Quan et al. prepared porous carbon catalysts by HTC and chemical activation method using osmanthus fragrans and sterchilia scaphigera as carbon sources, labeled as OPC and SPC, respectively [92]. OPC had a Tremella-like structure, dominated by micropores; while SPC had a honeycomb-like structure, dominated by mesopores (Fig. 1E and F). Besides, high-performance carbon-based catalysts were obtained by carbonizing soybean at high temperature with ZnCl<sub>2</sub> as an activated pore-forming agent [93]. It was shown that the BET SSA (949  $\rm m^2~g^{-1})$  of the catalyst can be significantly increased due to the activation of ZnCl<sub>2</sub>, so that it had a higher active center density, which helped to improve the catalytic activity.

#### 2.3. Hard and soft templating

Ordered inorganic solids such as mesoporous silica [94–98] are used as sacrificial hard template to guide the formation of pores during high temperature pyrolysis. However, the hard templating method has the following disadvantages. First, template must be removed using NaOH, HF and other toxic chemicals [99,100]. In addition, control over the pore diameter of porous material is also limited to the pores row of carbons that adopt the inverse structure of template pores. To avoid the use of toxic chemicals, soft templates is a good choice for preparing porous carbon materials. Soft template methods involve co-assembly between carbon precursors and surfactants in solution.

Liu et al. successfully reported a three-dimensional (3D) N-doped porous carbon (NPC) derived from low cost shrimp shells through simple SiO<sub>2</sub> spheres assisted, high temperature pyrolysis and HF acid leaching (Fig. 1G) [101]. It was found that the average pore size of three (3D) NPC was about 200 nm due to the removal of silica, and the carbon wall of 3D NPC exhibited an unusually distortional graphitic carbon lattice structure (Fig. 1H and I), indicating that more defects were formed in 3D NPC, thereby contributing to the enhanced electrocatalytic activity. Liu et al. developed N-doped porous carbon ORR catalyst (N-HPCMs) based on chelation-assisted multicomponent co-assembly strategy using banana peels as precursors and aluminum matrix composites and Pluronic F127 as co-templates (Fig. 1J) [102]. The N-HPCMs had many obvious three-dimensional mesoporous and microporous structures after HF treatment. It was found that this chelation-assisted multicomponent co-assembly strategy can tune the SSA and pore volume of N-HPCMs by simply changing the doses of Al<sup>3+</sup> and F127.

Table 1 (continued)

Biomass source	Other precursors	Synthesis strategy	Surface area (m <sup>2</sup> g <sup>-1</sup> )	Pore volume (cm <sup>3</sup> g <sup>-1</sup> )	$E_{ m Onset}$	$E_{1/2}$	Electrolyte	Reference
Chitosan	Triphenyl phosphine and FeCl <sub>3</sub>	HTC and pyrolysis	845	0.66	−0.014 V vs. AgCl	-	0.1 M KOH	[151]
Chitosan	Co(Ac) <sub>2</sub> and urea	Stirring, pyrolysis and wash	337	-	-	0.839 V vs. RHE	0.1 M KOH	[154]
Chitosan	$C_4H_6CoO_4$ • $4H_2O$ and Mg $(OH)_2$	Freeze drying, pyrolysis and pyrolysis again	1716	1.55	$-0.075~\mathrm{V}~vs.$ AgCl	−0.151 V vs. AgCl	0.1 M KOH	[155]

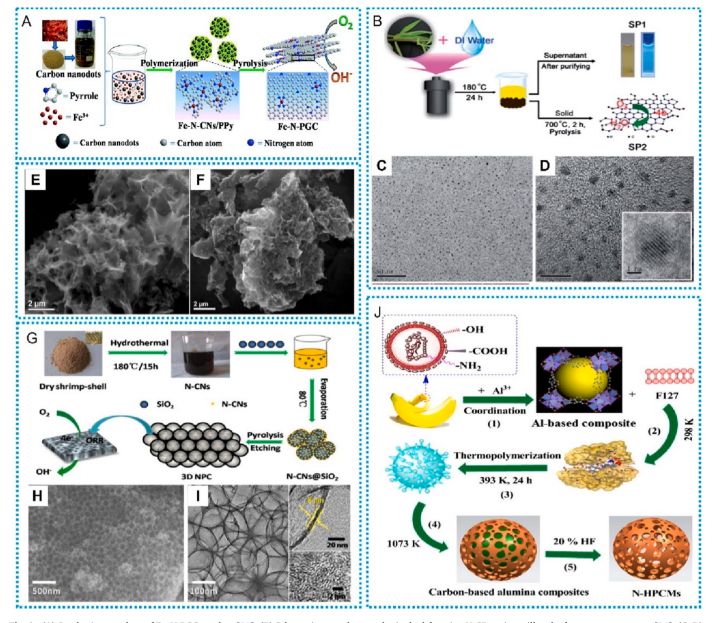


Fig. 1. (A) Synthesis procedure of Fe-N-PGC catalyst [80]. (B) Schematic procedure to obtain dual-function N-CDs using willow leaf as precursor system [81]. (C, D) TEM of the N-CDs material. (E, F) SEM of OPC and SPC materials [92]. (G) Synthesis procedure of three-dimensional NPC using low cost shrimp shells as precursor [101]. (H) SEM and (I) TEM of 3D NPC. (J) Schematic procedure to obtain N-HPCMs [102].

#### 3. Biomass-derived carbon material as ORR electrocatalysts

#### 3.1. Animal-based ORR electrocatalysts

Animals, including shrimp, fish, pigs and their waste (horns, blood, excrement, skins, bones, skins, cocoon silk) and human waste (hair and urine), etc., can be used to prepare carbon-based catalysts. These huge amounts kinds of biomass generated everyday which put enormous pressure on the environment. Rational use of this renewable energy is essential for ecological protection, waste management, and economic considerations. Considering that the ORR performance of catalysts can be enhanced by introducing dopants during post-processing, strategies for synthesizing oxygen reduction catalysts from various parts in animals, as well as their carbon structure and ORR performance, were described in detail in this chapter.

Blood proteins that are usually discarded as waste include abundant hemoglobin.

Duck blood is a waste produced from the process of slaughtering

ducks to prepare the Fe-doped carbon materials (CDB-Fe) for ORR [103]. Hemoglobin itself contains catalytically Fe active centers. CDB-Fe was composed of many irregular particles with diameters between 200 nm and 500 nm, shown in Fig. 2A and B. With the presence of Fe, the increased nitrogen and pyridinic N content of CDB-Fe suggested that the successful doping of Fe contributed to the increase of nitrogen content and the change of coordination ratios of various C–N bonds. The onset potential and half-wave potential of ORR for CDB-Fe were 2 mV and 137 mV, respectively, which were comparable to the performance of Pt/C catalyst (Fig. 2C and D).

Pig blood can be used to synthesize porous carbon materials and serve as an electrocatalyst. Pig blood is a sustainable animal resource which is rich in nitrogen due to its high protein content. Maruyama et al. obtained an catalyst using animal hemoglobin of oxygen reduction for a proton exchange membrane (PEM) fuel cell [104]. Catalytic performance was significantly enhanced by a two-step carbonization pyrolysis. In addition, the catalytic activity was much higher in the carbonization atmosphere of  $\rm CO_2/Ar$  (25:75) mixed gas than that in pure air. First, this

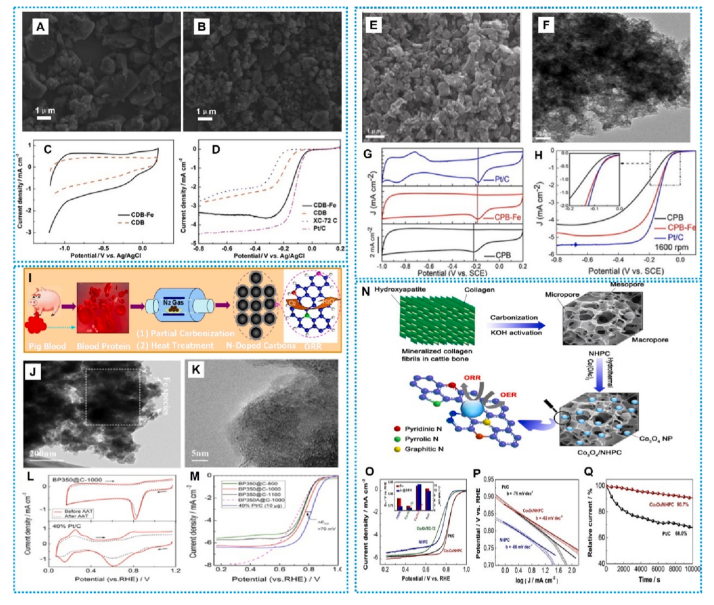


Fig. 2. (A, B) SEM of carbon materials from duck blood before and after iron treatment, respectively [103]. (C) CV curves and (D) LSV curves of CDB-Fe catalyst in 0.1 mol L $^{-1}$  KOH solution. (E) SEM and (F) TEM of pig blood-derived carbon material [51]. (G) CV curves and (H) LSV curves of CPB-Fe and Pt/C catalysts. (I) Schematic diagram of the synthesis of N–CNS using blood proteins as precursors [107]. (J) TEM and (K) HR-TEM images of BP350@C-1000. (L) CV curves and (M) LSV curves of these catalyst. (N) Schematic procedure to obtain the  $Co_3O_4/NHPC$  ORR catalyst [108]. (O) LSV curves of  $Co_3O_4/NHPC$  and Pt/C catalyst in  $O_2$ -saturated 0.1 M KOH solution. (P) Tafel plots of these electrocatalysts. (Q) The i-t chronoamperometry curves of  $Co_3O_4/NHPC$  and Pt/C catalyst.

enhanced the SSA and pore volume of the carbonized material, making it easier to expose catalytically active sites, and second, it could converted  ${\rm Fe}^{3+}$  to  ${\rm Fe}^{2+}$ , which acted as oxygen bonding sites in the ORR.

Some biomass itself contains catalytic active centers, which can be used not only as carriers but also directly as catalysts. The presence of Fe in the carbonization process profoundly affected the degree of graphitization, formation of pyridinic N, and ORR catalytic activity of the prepared electrocatalyst. Later Zhang et al. successfully prepared a N self-doped carbon electrocatalyst with the aid of metal iron based on pig blood (Fig. 2E and F) [51]. The contents of pyridine N and graphite N of Fe-free catalyst (CPB) were 30.1% and 39.4%, respectively, while those of Fe-containing catalyst (CPB–Fe) were 35.1% and 39.2%, respectively. The limited current density ( $J_{\rm L}$ ) can be improved with graphitic N of ORR catalyst, while the onset potential ( $E_{\rm onset}$ ) can be enhanced with pyridinic N [105,106]. CPB exhibited an obvious ORR peak for ORR at -0.22 V  $\nu_{\rm S}$ . Ag/AgCl. After iron was added to PB, the ORR peak of CPB-Fe moved to a higher positive potential at -0.19 V. Besides, CPB-Fe

showed a positive  $E_{\rm onset}$  (-0.04 V vs. Ag/AgCl) and half-wave potential ( $E_{1/2}$ ) of -0.16 V vs. Ag/AgCl, which were comparable to those of commercial Pt/C catalyst (Fig. 2G and H).

Chen successfully prepared an ORR electrocatalyst by using blood protein and carbon nanospheres at different pyrolysis temperatures (Fig. 2I) [107]. As shown in Fig. 2J and K), many Fe-containing agglomerations and an amorphous structure were shown in BP350@C-1000, indicating more structural defects and more exposed edges can be formed, which was conducive to the improvement of oxygen reduction activity. In the pyrolysis process, the metallic Fe of raw blood protein can enhance the conversion of oxidative N to planar N, as well as the co-incorporation of pyridine and pyrrolic N groups into the carbon material, which further generated more defects in the carbon structure. Pyridine and pyrrolic N may be the most influential N functional groups on ORR activity and can serve as the ORR active sites of catalysts. In conclusion, the catalyst had outstanding catalytic activity for ORR (Fig. 2L and M).

As a kind of biomass waste, animal bones such as fish scales [109, 110], pig bones and cow bones [108,111], are widely used as low-cost carbon sources to obtain carbon materials. Collagen in animal bones can provide C and N resources to synthesize heteroatom-doped carbon materials, while hydroxyapatite crystals can serve as hard templates to generate pore structures, therefore bones can be pyrolyzed into carbon precursors for ORR. Carbon materials with high content of N doping and typical 3D hierarchical porous structures can be produced. Co<sub>3</sub>O<sub>4</sub> nanoparticles anchored on N-doped 3D hierarchically porous carbon material with (Co<sub>3</sub>O<sub>4</sub>/NHPC) as an high-efficiency ORR catalyst derived from cattle bone can be obtained through a simple hydrothermal synthesis process (Fig. 2N) [108]. The high SSA of Co<sub>3</sub>O<sub>4</sub>/NHPC as high as 1070 m<sup>2</sup> g<sup>-1</sup>, hierarchical porous structure and high content of N doping (4.93 wt %) facilitated the uniform anchoring of Co<sub>3</sub>O<sub>4</sub> nanoparticles on the carbon support and enhanced the mass transfer capacity. Notably, the  $Co_3O_4/NHPC$  catalyst exhibited outstanding ORR activity. The  $E_{1/2}$ was 0.853 V vs. RHE, the tafel slope was  $63 \text{ mV} \text{ dec}^{-1}$ , and after a test lasting for 10 000 s, the relative current loss was only 1.74% (Fig. 2Q).

Eggs are rich in protein, cholesterol and lecithin, etc., which can be used as precursors of carbon and nitrogen. Wu et al. synthesized nitrogen, phosphorus-doped carbon microspheres with mesoporous structure (egg-CMS) by a hard templating method, using eggs as carbon precursors through a spray-drying process, pyrolysis, and template removal (Fig. 3AC) [112]. The prepared carbon microspheres had regular mesoporous structure and large SSA (970 m<sup>2</sup> g<sup>-1</sup>). When prepared as an oxygen reduction catalyst, the egg-CMS catalyst had a large limiting current (4.36 mA cm<sup>-2</sup>), a positive  $E_{1/2}$  (0.84 V vs. RHE), shown in Fig. 3D. In terms of stability, 92% of the initial current can be maintained after 20 h of testing (Fig. 3E). It is demonstrated that the egg-CMS catalyst can be used in fuel cells.

Zhang et al. synthesized Fe and N doped carbon materials (CE-Fe-MWNT) used eggs as the N source and carbon source, transition metal iron as the iron source, combined with multi-walled carbon nanotubes (MWNT), by cooking denaturation and secondary heat treatment [113]. TEM showed that the carbon material obtained by high-temperature pyrolysis can be well combined with MWNTs and had a porous

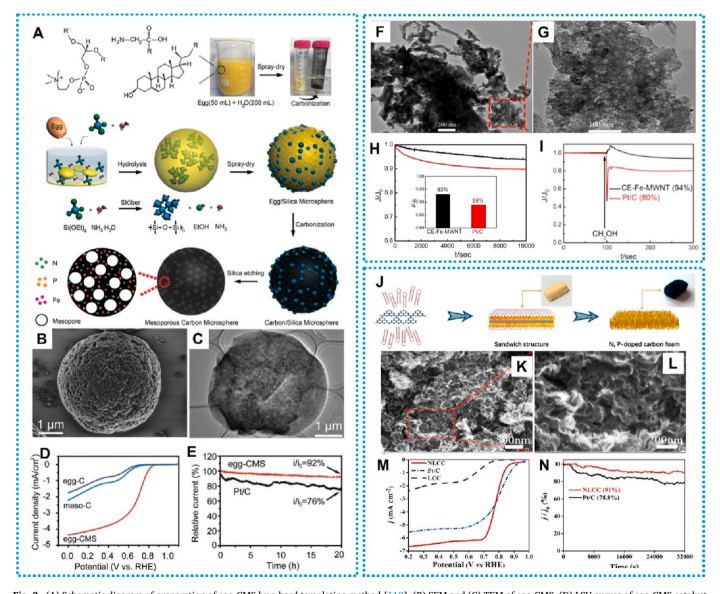


Fig. 3. (A) Schematic diagram of preparation of egg-CMS by a hard templating method [112]. (B) SEM and (C) TEM of egg-CMS. (D) LSV curves of egg-CMS catalyst in  $O_2$ -saturated 0.1 M KOH solution. (E) The chronoamperometry curves of egg-CMS and Pt/C catalyst. (F, G) TEM images of the CE-Fe-MWNT material [113]. (H) The methanol tolerance curves of CE-Fe-MWNT and Pt/C catalyst. (I) Chronoamperometric responses of CE-Fe-MWNT and Pt/C catalyst after methanol introduction. (J) Schematic procedure to obtain NLCC by in-situ foaming technology [114]. (K, L) SEM image of NLCC. (M) LSV curves of NLCC catalyst. (N) The i-t chronoamperometry curves of NLCC and Pt/C catalyst.

structure (Fig. 3F and G), which was beneficial to promote ORR mass transfer and thus enhance ORR activity. The MWNTs with high graphitization acted as a stable matrix to immobilize ORR active sites because of its perfect electrical conductivity and stability. Electrochemical tests showed that the CE-Fe-MWNT catalyst had excellent ORR activity in alkaline electrolyte, outstanding stability (Fig. 3H) and methanol resistance (Fig. 3I), making it a viable alternative to expensive Pt-based catalysts. Xian et al. used crude lecithin extracted from eggs as a phosphorus source and C<sub>3</sub>N<sub>4</sub> as a self-sacrificing template to obtain nitrogen and phosphorus-doped carbon materials (NLCC) by in-situ foaming technology (Fig. 3J) [114]. The unique structure of the lecithin molecule can encapsulate g-C<sub>3</sub>N<sub>4</sub>. During the pyrolysis, g-C<sub>3</sub>N<sub>4</sub> could be decomposed, and the mixture was gradually expanded, and finally a special carbon foam composed of many multi-folded morphology nanosheets was formed (Fig. 3K and L). In alkaline electrolyte, the NLCC catalyst showed a significant  $J_{\rm L}$  of 6.7 mA cm<sup>-2</sup> (Fig. 3M), which was significantly greater than that of Pt/C catalyst (5.5 mA cm<sup>-2</sup>). In terms of stability, after about 9 h of chronoamperometry, the NLCC catalyst only lost 9% of the current density, proving its excellent long-term durability (Fig. 3N).

Human hair is composed of  $\alpha$ -keratin, containing 15–16% N and 4.5–5.5% S, which can be used to synthesize N, S doped carbon catalysts for ORR [115]. Kalaiselvi et al. activated pre-carbonized hair with NaOH and then pyrolyzed at high temperature to obtain hair-derived carbon materials (HCC) for anode materials of sodium-ion battery and cathode ORR catalysts [54]. Due to its unique structural characteristics and large number of heteroatoms, the HHC exhibited excellent rate performance and long-term cycling performance and when used in ORR, HCC catalyst

showed good ORR performance and long-term durability (Fig. 4A and B).

Chaudhari et al. prepared N, S-doped carbon (HC-X, X represents the carbonization temperature) by pre-carbonization, NaOH activation, and high-temperature carbonization by using hair as nitrogen and sulfur sources (Fig. 4C) [116]. The unique preparation strategy including mild NaOH activation and further high-temperature graphitization enhanced the electrical conductivity of carbon materials and preserving valuable N, S heteroatoms. HC-900 as a catalyst exhibited excellent ORR performance, four-electron selectivity (electron transfer was between 3.80 and 3.90), and long-term stability (HC-900 lost 14% of current density after 50000 s), suggesting that hair-derived carbon materials can be promising alternatives to Pt/C catalysts (Fig. 4D-G). The author further proposed a "proof of concept" that used urine to produce a new type of carbon called "urine carbon" (URC) (Fig. 4H and I) [117]. High porosity can be created by removing the inherent salt particles in the URC materials, and multiple heteroatoms can be doped into URC materials via high-temperature carbonization without the use of complex and expensive templates and the addition of additional precursors of heteroatoms. Electrochemical tests showed that the mass activities of the URC-900 and URC-1000 were increased by 22% and 24%, respectively, compared with other catalysts. Surprisingly, URC-1000 had low BET surface area and less content of nitrogen, but still exhibited the outstanding ORR activity. This is due to lower resistivity, presence of numerous heteroatoms, high electrical conductivity and a large number of pore structures (Fig. 4J and K).

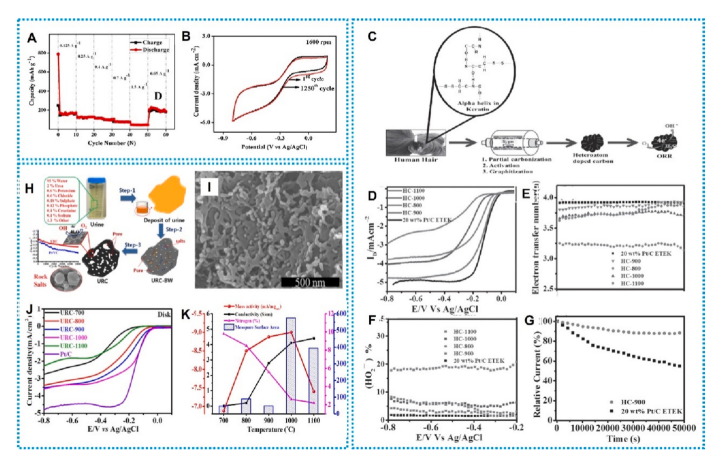


Fig. 4. (A) Rate performance behavior of HCC anodes [54]. (B) Durability test of HCC catalyst. (C) Schematic procedure to obtain N, S-doped carbon by using hair as nitrogen and sulfur sources [116]. (D) LSV curves of HC-X catalyst in 0.1 M KOH. (E) The electron number of HC catalyst at different pyrolysis temperatures. (F) Peroxide yield of HC catalyst at different pyrolysis temperatures. (G) The i-t chronoamperometry curves of HC-900 and Pt/C catalyst. (H) Schematic diagram of preparation of urine carbon (URC) [117]. (I) SEM image of URC-1000. (J) LSV curves of URC catalyst at different pyrolysis temperatures and Pt/C. (K) Comparative relationship between conductivity, nitrogen content and mesoporous surface area and mass activity of URC materials.

#### 3.2. Plant-based ORR electrocatalysts

Different from animal biomass, plant biomass usually selects specific types of plant biomass with specific structures and enhances electrocatalyst performance with appropriate synthetic strategies. The researchers had done many attempts to convert abundant and inexpensive plant (leaves, fruit, roots or flowers) biomass into ORR efficient ORR catalyst. In this section, we had summarized unique characteristics of biomass, which was important for ORR performance.

Porphyra itself contains proteins and taurine, which can be used as nitrogen, sulfur precursor. On this basis, Zhang et al. obtained nitrogen and sulfur heteroatom doped Fe–N–C material (Fe-NSDC) by mixing and grinding FeCl<sub>3</sub>·6H<sub>2</sub>O with porphyra as precursor and pyrolysis at high

temperature (Fig. 5A) [118]. Interestingly, the researcher used ultrasound-assisted strategy to remove taurine contained in porphyra and prepared iron-doped carbon catalyst (Fe-NDC) without sulfur, so as to compare the effect of sulfur and Fe-N $_{\rm x}$  on improving oxygen reduction activity. Because of the catalysis of Fe, the Fe-NSDC catalyst exhibited a three-dimensional layered sheet-like structure. And the thin-layer sheet-like structures with porous and wavy corrugated structures of Fe-NSDC can provide additional active site and facilitated electron transfer. Fe-N $_{\rm 3}$ |S can be found as Fe-NSDC catalytic active site by XAS analysis. The Fe-NSDC catalyst exhibited ultra-high performance with  $E_{1/2}$  up to 0.84 V vs. RHE, which was superior to Pt/C catalyst performance (Fig. 5B and C). It was found that the addition of S can adjust the charge distribution of Fe-NSDC catalyst, resulting in excellent ORR

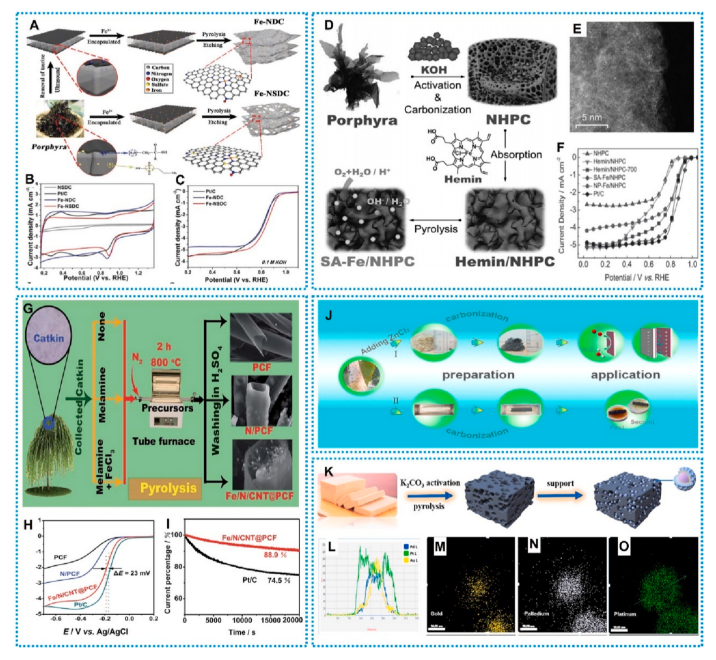


Fig. 5. (A) Schematic procedure to obtain Fe-NSDC with porphyra as precursor [118]. (B) CV curves and (C) LSV curves of Fe-NDC, Fe-NSDC and Pt/C catalyst. (D) Synthesis procedure of SA-Fe/NHPC catalyst [119]. (E) HAADF-STEM of SA-Fe/NHPC catalyst. (F) LSV curves of SA-Fe/NHPC catalyst. (G) Schematic diagram of preparation of Fe/N/CNT@PCFs by direct carbonizing of catkins [120]. (H) LSV curves of Fe/N/CNT@PCF and Pt/C catalyst. (I) The durability tests of Fe/N/CNT@PCF and Pt/C catalyst. (J) The formation process of carbon with high porosity by using catkins as precursor [121]. (K) Synthesis procedure of Au@Pd@Pt-NSC-900 composite [106]. (L) Line-scanning of Au@Pd@Pt core-shell material. (M ~ O) Elemental mapping of Au, Pd and Pt.

activity.

On the basis of the confinement effect of porous carbonaceous material and Fe element, Zhang et al. obtained nitrogen-doped hierarchical porous carbon-supported single-atom Fe (SA-Fe/NHPC) catalyst by rapid pyrolysis of hemin-adsorbed bio-porphyrin carbon (Fig. 5D) [119]. A large number of micropores were embedded in the carbon layer of SA-Fe/NHPC. At the same time, the Fe atoms in micropores are evenly distributed, and there is no obvious aggregation of metal atoms (Fig. 5E). The as-prepared SA-Fe/NHPC catalyst exhibited great ORR activity ( $E_{1/2}=0.87~Vvs.$  RHE) in alkaline media (Fig. 5F), ORR stability ( $E_{1/2}$  negative shift by 1 mV after ADT) and methanol tolerance, which were greater than those of Pt/C catalysts.

Catkins contain C, N and other element, which can be used as natural, inexpensive and sustainable precursor and template materials to prepare heteratomic self-doping carbon catalysts. Li et al. synthesized Fe and N co-doped carbon nanotubes@hollow carbon fiber catalysts (Fe/N/CNT@PCFs) by direct carbonization of catkins, melamine, and  $FeCl_3\cdot 6H_2O$  (Fig. 5G) [120]. The addition of melamine to the carbon material as a nitrogen source greatly increased the nitrogen content. Through characterization, it could be found that the presence of Fe

element enables the catalytic growth of many carbon nanotubes on the walls of Fe/N/CNT@PCFs. The special structure of Fe/N/CNT@PCFs endowed it with large SSA. In addition, the presence of Fe also promoted the formation of pyridine nitrogen species, which further enhanced the ORR performance. After testing, Fe/N/CNT@PCFs displayed the most positive  $E_{\rm onset}$  of 98 mV and  $E_{1/2}$  of 194 mV (Fig. 5H). And after 20000 s test, the relative current of Fe/N/CNT@PCF was as high as 88.9% (Fig. 5I). Gao et al. took catkins as precursor and synthesized carbon with high pore volume (1.31 cm³ g⁻¹) and surface area (1462.5 m⁻² g⁻¹) by pyrolysis of ZnCl₂ (Fig. 5J) [121]. The catalyst was better than the Pt/C catalyst in ORR activity, long-term durability and methanol resistance.

As a common soy product, beancurd has protein and carbohydrates [122], making itself as a cheap precursor. It can be converted into nitrogen-doped carbon by pyrolysis and other processes. However, the pure carbon materials prepared from direct carbonizing of beancurd have some disadvantages such as little SSA and poor catalytic activity. On this basis, Sun et al. reported a tofu-derived porous carbon (NSC-900) by activation of  $K_2CO_3$  and pyrolysis and then loaded the Au@Pd@Pt core-shell particles onto the NSC-900 to obtain composite

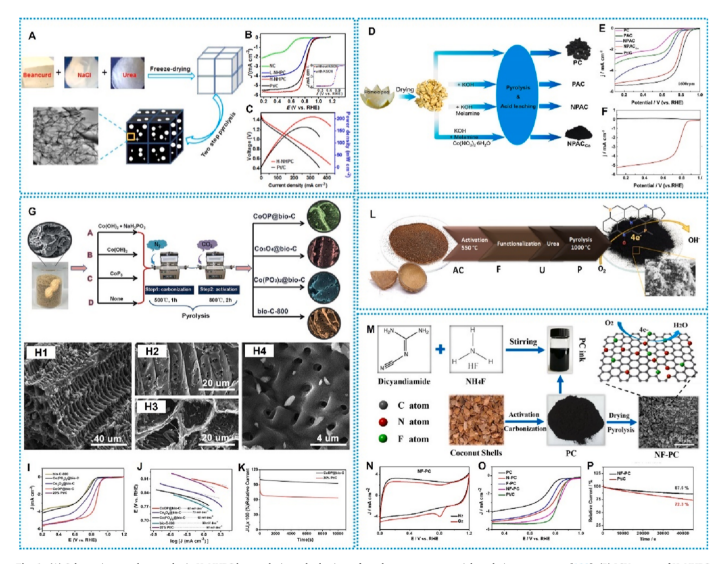


Fig. 6. (A) Schematic procedure to obtain H–NHPC by pyrolysis method using tofu and urea as raw materials and nitrogen sources [123]. (B) LSV curses of H–NHPC and other catalysts. (C) The discharge curves of the H–NHPC-based and Pt/C-based Zn– $O_2$  battery and corresponding power density. (D) Synthesis procedure of NPAC<sub>Co</sub> catalyst [125]. (E) LSV curses of NPAC<sub>Co</sub> and Pt/C catalyst. (F) RDE voltammograms of NPAC<sub>Co</sub>. (G) Synthesis procedure of CoOP@bio-C using peanut shells as precursors [74]. (H1 $\sim$ H4) SEM of CoOP@bio-C. (I) LSV curses of CoOP@bio-C and other catalysts. (J) Tafel curses of CoOP@bio-C and other catalysts. (K) The i-t chronoamperometry curves of CoOP@bio-C and other catalysts. (L) Schematic diagram of preparation of AC-F-U-P [130]. (M) The formation process of NF-PC [131]. (N) CVs of NF-PC catalyst. (O) LSVs of NF-PC and other catalysts. (P) The i-t chronoamperometry curves of NF-PC and Pt/C catalyst.

material (Fig. 5K–O) [106]. Due to the activation of  $K_2CO_3$ , the as-synthesized composite had an ultrahigh SSA (3814 m<sup>2</sup> g<sup>-1</sup>) and porous structure (2.08 cm<sup>3</sup> g<sup>-1</sup>). It was noteworthy that the higher SSA promoted mass transfer, which made the Au@Pd@Pt-NSC-900 have higher ORR performance compared to Pt/C. The specific activities and mass activities of the as-synthesized catalyst were 5 and 13 times as high as those of Pt/C catalyst.

A carbon material (H–NHPC) with porous structure (0.689 cm³ g $^{-1}$ , 988.2 m² g $^{-1}$ ) was prepared by pyrolysis method using tofu and urea as raw materials and nitrogen sources (Fig. 6A) [123]. Under the action of NaCl, H–NHPC consisted of graphene-like carbon nanosheets that were interconnected to present a 3D hierarchical porous structure. The H–NHPC catalyst showed extreme catalytic activity with a  $E_{1/2}=0.81~\rm V$  vs. RHE and four-electron selectivity (Fig. 6B). In addition, the application of H–NHPC to zinc-air batteries with high power density and large specific capacity (Fig. 6C). Zhai et al. prepared a novel Co–N–C catalyst by one-step pyrolysis with soymilk as raw material and the addition of cobalt nitrate [124]. Due to the existence of the Co–N–C structure, the catalyst showed ed outstanding ORR activity in alkaline electrolyte.

Nitrogen, cobalt activated biochar (NPAC<sub>Co</sub>) was obtained by hightemperature carbonization method using cobalt nitrate as cobalt source, potassium hydroxide as chemical activator and pomelo peel as precursor (Fig. 6D) [125]. The successful doping of cobalt enhanced the graphitization of carbon and also increased the doping amount of nitrogen. In addition, pyridine nitrogen content was also increased in NPAC<sub>Co</sub>, which was beneficial to improve the ORR activity. For ORR, NPAC<sub>Co</sub> catalyst exhibited high  $E_{\text{onset}}$  of 0.87 V vs. RHE, high  $E_{1/2}$  of 0.78 V vs. RHE and low H<sub>2</sub>O<sub>2</sub> yield (Fig. 6E and F), comparable to those of Pt/C. The Fe<sub>3</sub>C/WC/GC nanocomposites were prepared by a facile carbothermic reduction method derived from pomelo peel [126]. The incorporation of iron can change the amorphous carbon into Fe<sub>3</sub>C and contributed to the graphitization of carbon. In addition, during the pyrolysis process, the interaction between iron and tungsten was very close, which could be well dispersed and embedded into the carbonized matrix. After testing, Fe<sub>3</sub>C/WC/GC nanocomposites catalyst had high ORR activity and durability in a neutral pH environment. According to DFT calculations, Fe<sub>3</sub>C can provide an active site for ORR, and its (001) surface can dissociate the O-O bond to overcome the ORR barrier.

Waste peanut shells can be used as carbon precursors to synthesize porous carbon materials by pyrolysis and other methods. Wu et al. obtained a Co, S co-doped carbon material (CoOP@bio-C) with microporous structure by CO<sub>2</sub> reduction and high-temperature pyrolysis strategy using peanut shells as carbon precursors (Fig. 6G, H1~H4) [74]. Due to the unique layered 3D open porous structure of CoOP@bio-C materials, the active sites of S and external doping Co<sub>3</sub>O<sub>4</sub> and Co(PO<sub>3</sub>)<sub>2</sub> were increased, which made it have remarkable ORR catalytic activity. The  $E_{\rm onset}$  and  $E_{\rm 1/2}$  of CoOP@bio-C catalyst in 0.1 M KOH solution were 0.91 V vs. RHE and 0.81 V vs. RHE, respectively, the tafel slope was 57 mV  ${\rm dec}^{-1}$  and the current density lost only 20% after 10 000 s (Fig. 6I–K).

As a natural lignocellulosic material, coconut shell is also suitable for preparing carbon materials with porous structure because of its abundant carbon content, excellent structure and low ash level [87]. Porous carbon materials with SSA of about  $800-1500~{\rm m}^2~{\rm g}^{-1}$  were successfully fabricated using raw coconut shell as precursors [127–129]. Borghei et al. used coconut shell residue as carbon precursor, phosphoric acid as activator and urea as nitrogen source, to prepare N, P-doped porous carbon (AC-F-U-P) by chemical activation method (Fig. 6L) [130]. Phosphate can form connections with lignocellulose, thus preventing the shrinkage and collapse of the structure for AC-F-U-P material during high-temperature carbonization. The obtained AC-F-U-P had a large SSA (1216  ${\rm m}^2~{\rm g}^{-1}$ ) and plenty of mesopores. And the prepared biomass carbon catalyst exhibited excellent long-term stability.

Wang et al. cleverly manufactured N, F co-doped porous carbon (NF-PC) by a economical and environmentally friendly strategy derived from green coconut shell (Fig. 6M) [131]. The microporous structure and the

cooperation between N and F element, enabled the NF-PC to exhibit outstanding ORR activity ( $E_{\rm onset} = 0.985$  V  $\nu$ s. RHE,  $E_{1/2} = 0.834$  V  $\nu$ s. RHE) and long-term stability (after 45 000 s, the current density of NF-PC catalyst remains 87.5% of the initial current density), even surpassing that of Pt/C catalyst (Fig. 6NQ).

N-doped partly-graphitized carbon (NPGC) was prepared as a cathode ORR catalyst using cornstalks as precursor, which had outstanding ORR performance and high power density of 1122 mW m $^{-2}$  and long-time running durability [132]. Nitrogen, phosphorus, and iron trimeric nanoporous carbon also prepared from corn silk as precursors provided a good foundation for the development of economical, environmentally friendly and high-yield catalysts [133].

#### 3.3. Glucose derivatives

Glucose, a component of cellulose, was used as a precursor for carbon supports of platinum nanoparticle. Yan et al. prepared mesoporous bowl-like carbons (BLCs) derived from glucose and silica spheres as hard template (Fig. 7A) [134]. BLCs had high SSA (1108.3  $\rm m^{-2}~g^{-1})$  and large porosity volume, which were conducive to the uniform dispersion and loading of Pt particles, and significantly enhanced mass transfer process of ORR. CV exhibited that ORR mass current density of the prepared catalyst was 1.6 times as high as that of the Pt/C.

To get rid of the dependence on Pt, Liu et al. fabricated Ag nanoclusters anchored on carbon nanodots (Ag NC/CNs) by a green method using glucose (Fig. 7B). The as-prepared Ag nanoclusters exhibited excellent catalytic performance for ORR, as well as a four-electron transfer process [135]. Wassne et al. obtained carbon spheres by hydrothermal carbonizing glucose and then prepared spherical core-shell composite particles (TaON@CN) by sol-gel deposition of tantalum oxide (Fig. 7C) [136]. The core-shell composite particles consist of a spherical carbon nitride core and a tantalum (oxy)nitride shell. Electrochemical tests showed that the ORR activity of core-shell composite was significantly higher than that of pure tantalum (oxy)nitride.

Titirici et al. used a novel method to fabricate N-doped carbonaceous aerogels based on the hydrothermal carbonization of glucose derivatives [137]. The as prepared aerogel was obtained from carbohydrate-based derivatives and phenolic compounds, with a N doping level of 3-5 wt % and a SSA of 450 m<sup>2</sup> g<sup>-1</sup>. Later, Titirici and co-workers futher synthesized N-doped porous carbon aerogels using glucose and cellulose as carbon sources, soybean protein as nitrogen sources and pore-forming guide agent [138]. The porosity of the carbon aerogels was enhanced by heat treatment, and the catalysts exhibited good catalytic activity for ORR. The addition of S had a positive effect on ORR performance except N doping after testing. Qiu et al. prepared regular mesoporous and macroporous N-S-CMK-3 catalyst by hard template method using glucose and thiophene as precursors (Fig. 7D) [139]. And the effects of N and S on the ORR performance of N-S-CMK-3 catalyst were studied. The results showed that N-S-CMK-3 catalyst had excellent ORR performance such as 0.92 V vs. RHE for  $E_{\text{onset}}$  (Figs. 7E), 68 mV dec<sup>-1</sup> for tafel slope, because of the synergy influence of graphite N and thiophene S structure.

Recently, it has been shown that catalytic performance can be enhanced by introducing transition metals into carbon materials rich in nitrogen. Li et al. fabricated Fe, N doped carbon catalysts (3DHP/Fe–N–C-T, T stands for pyrolysis temperature) with high SSA (906.3 m<sup>2</sup> g<sup>-1</sup>), mesoporous and macroporous structure by high temperature pyrolysis using SiO<sub>2</sub> spheres as sacrificial template, glucose as precursor and dicyandiamide as nitrogen source, respectively (Fig. 7F and G) [140]. The  $E_{\rm onset}$  (about 0 V vs. Ag/AgCl) and  $E_{\rm 1/2}$  (-0.15 V vs. Ag/AgCl) of 3DHP/Fe–N–C-900 in alkaline medium were comparable to Pt/C catalysts due to cooperation of Fe<sub>3</sub>O<sub>4</sub>, and N group (Fig. 7H).

Lai et al. fabricated Fe–N/G catalysts by an in-situ self-sacrificial template method to using melamine and glucose as precursors (Fig. 7I) [141]. Under high temperature pyrolysis, Fe/Fe<sub>3</sub>C was encapsulated in carbon nanocapsules and then evenly formed on the surface of NG

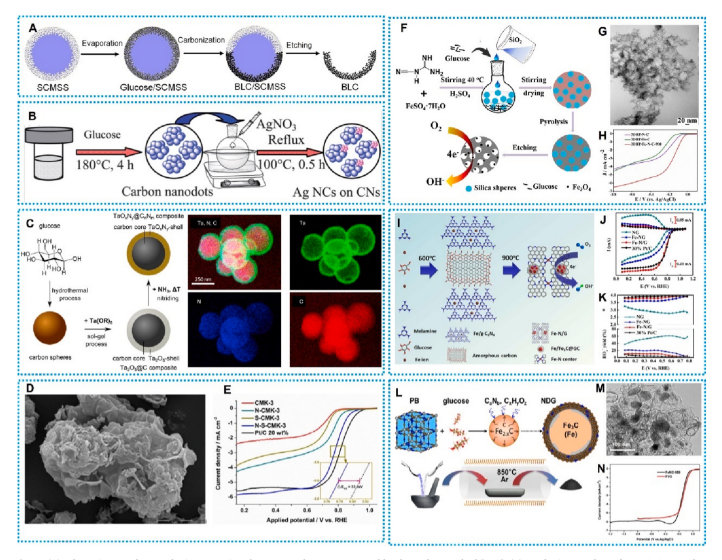


Fig. 7. (A) Schematic procedure to obtain BLCs using glucose as carbon precursor and hard template method [134]. (B) Synthesis procedure of Ag NC/CNs catalyst [135]. (C) The formation process of core-shell TaON@CN and EDS mapping images of TaON@CN [136]. (D) SEM image of N-S-CMK-3 catalyst. (E) LSV curves of N-S-CMK-3 and Pt/C catalyst [139]. (F) Synthesis procedure of 3DHP/Fe-N-C-T [140]. (G) TEM images of 3DHP/Fe-N-C-900. (H) LSV curves of 3DHP/Fe-N-C-900 and other catalysts. (I) Schematic procedure to obtain Fe-N/G [141]. (J) RRDE of Fe-N/G and other catalysts. (K) HO<sub>2</sub> yield and electron transfer number of the corresponding catalyst. (L) Synthesis procedure of FeNC-850 [142]. (M) TEM of FeNC-850. (N) LSV curves of FeNC-850 and Pt/C.

material with a highly open mesoporous structure. The Fe–N/G catalyst had excellent catalytic activity and four-electron selectivity comparable to those of Pt/C because of its highly active Fe-N $_{\rm x}$  sites (Fig. 7J and K). Likewise, metal particles can be encapsulated in graphite carbon layers. Yang et al. obtained an Fe–N co-doped catalyst (FeNC-850) through pyrolysis process using glucose and Prussian blue (PB) as precursors (Fig. 7L and M) [142]. During high temperature carbonization, Prussian blue can be decomposed into iron species and iron species catalyzed the growth of graphite layers to form Fe/Fe<sub>3</sub>C encapsulation structures. After electrochemical testing, it was found that the prepared FeNC-850 catalyst had excellent ORR performance (Fig. 7N). In addition, when the timing current was carried out in alkaline medium, the current loss was 17.85% after 10000 s, which was less than that of Pt/C catalyst (33.53%), confirming that FeNC-850 catalyst had good long-term stability.

### 3.4. Starch

Zhao et al. prepared highly active metal-free N-containing carbon nanosheets (NCNs) by one-step pyrolysis method derived from cornstarch and urea [143]. The RDE tests revealed that NCN possessed a

four-electron transfer process and exhibited excellent ORR performance and durability, making it an alternative to Pt-based catalysts. Cheng et al. used soluble starch (SS) to encapsulate noble nanoparticles, followed by pyrolysis at low temperature to synthesize ultra-thin carbon layers (UTCL) stabilized Pt catalyst (Pt-UTCL/C) with open frameworks [144]. The special structure of UTCL reduced the loss of ECSA and effectively promoted the ORR mass transfer process. UTCL can tightly fix Pt nanoparticles on the carbon matrix, inhibited the agglomeration and separation of Pt nanoparticles, and slowed down their dissolution growth process. Electrochemical tests indicated that the Pt-UTCL/C catalyst had high stability, which was comparable to Pt/C catalysts. Fe, S-modified N-doped graphene/carbon nanosheets (Fe, S/NGC) were obtained by hydrothermal reaction and high-temperature carbonization with graphene oxide (GO), starch, FeCl<sub>3</sub> and KSCN [63]. It was precisely shown that Fe, S/NGC had a highly interpenetrated three-dimensional porous structure which was composed of many interconnected nanoflakes with a thickness of 14 nm. Due to the three-dimensional porosity of Fe, S/NGC-900 and synergistic effect of N, S and Fe elements, the obtained Fe, S/NGC-900 catalysts exhibited positive  $E_{onset}$  of 0.95 V  $\nu s$ . RHE, high  $E_{1/2}$  of 0.83 V vs. RHE and good kinetic such as low Tafel slopes of 67 mV dec<sup>-1</sup>.

#### 3.5. Chitin and chitosan

Chitin can serve as both carbon and nitrogen sources at the same time to obtain self-nitrogen-doped carbon materials due to its rich nitrogen content [145]. Li et al. employed a facile solution-chemical method to extract chitin aerogels with micropores and mesopores from shrimp shells, followed by carbonization to obtain N-doped carbon materials (Chitin-900) (Fig. 8A). The prepared Chitin-900 catalyst had a typical four-electron oxygen reduction path (Fig. 8B). Wu et al. obtained N-doped carbon/graphite C<sub>3</sub>N<sub>4</sub> (C-Chitin/g-C<sub>3</sub>N<sub>4</sub>) catalyst by dissolving and pyrolysis of g-C<sub>3</sub>N<sub>4</sub> and chitin at high temperature (Fig. 8C) [146]. The C-chitin/g- $C_3N_4$  composite exhibited a unique three-dimensional (3D) loofah-like network structure, which enhanced the contact opportunity between the cathode ORR catalyst and the electrolyte, promoted mass transfer, and was beneficial to expose the ORR catalytic active sites. Benefiting from the well-designed porous structure, the C-Chitin/g-C<sub>3</sub>N<sub>4</sub> catalyst exhibited obvious ORR activity (Fig. 8D) and excellent methanol resistance, etc.

Titirici et al. obtained nitrogen-doped carbon by a one-pot pyrolysis method using lobster shells and artificial lobster shells which was composed of chitin and CaCO<sub>3</sub> particles as precursors, respectively

[147]. Due to the relatively uniform distribution of CaCO<sub>3</sub> particles in the artificial shrimp shell, chitin-derived carbon had a higher SSA. The chitin-derived catalyst had higher  $E_{\text{onset}}$  and lower  $H_2O_2$  yield after tests. Activated carbon sheets (ACS) were synthsized by hydrothermal carbonization using chitin as precursor and zinc chloride as activator (Fig. 8E) [148]. Chitin-based carbons had sheet-like morphology and are composed of porous structures. The ACS catalyst exhibited obvious catalytic activity, excellent long-term durability and methanol resistancein alkaline electrolyte, comparable to Pt/C catalysts. (Fig. 8F), Chitosan is a sustainable and inexpensive biomass precursor capable of forming N-active sites in carbon materials. Zhang et al. fabricated metal-free mesoporous N-doped carbon materials by adding different kinds of pore-forming agents and using chitosan as precursor (Fig. 8G and H) [149]. Interestingly, the SSA (1190 m<sup>2</sup>/g<sup>-1</sup>) of N-doped mesoporous biomass carbon (NC-3) increased significantly after adding ferric nitrate as a pore former. The synthesized NC-3 catalyst contained a large amount of pyridine N, indicating that Fe doping can regulate the generation of pyridine N, thereby promoting the four-electron reaction and improving the ORR activity (Fig. 8I and J). Highly graphitized graphene-like N-doped mesoporous carbon (STST, T stands for temperature) by high temperature pyrolysis using FeCl<sub>3</sub> as a soft template

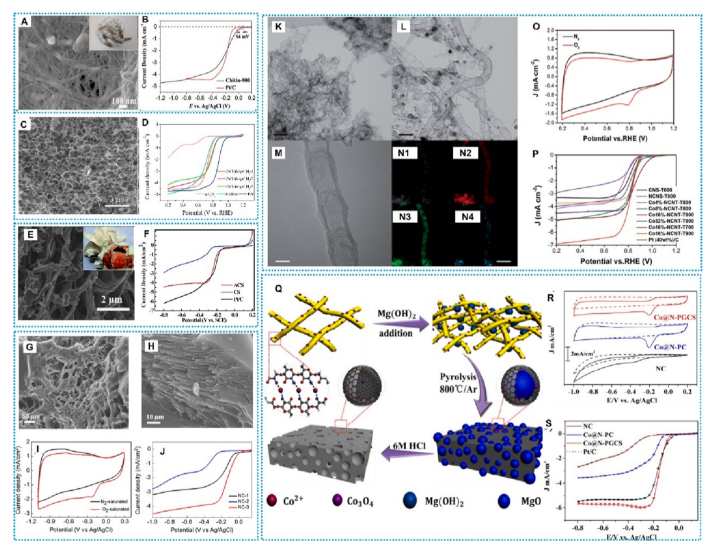


Fig. 8. (A) SEM of Chitin-900 [145]. (B) LSV of Chitin-900 and Pt/C catalyst. (C) SEM of C-chitin/g-C<sub>3</sub>N<sub>4</sub> [146]. (D) LSV curves of C-chitin/g-C<sub>3</sub>N<sub>4</sub> and other catalysts. (E) SEM image of ACS [148]. (F) LSV curves of ACS and Pt/C catalyst. (G, H) SEM image of NC-3 [149]. (I) CV curves of NC-3 catalyst. (J) LSV curves of NC-3 and Pt/C catalyst. TEM images of (K) nitrogen doped carbon and (L) Concnt-1800 [154]. (M) HRTEM image of Co-NCnt-1800. (M) STEM images and (N1~N4) EDS mappings of Co-NCnt-1800. (O) CV curves of Co-NCnt-1800 catalyst. (P) LSV curves of Co-NCnt-1800 and Pt/C catalyst. (Q) Synthesis procedure of Co@N-PGCS [155]. (R) CV and (S) LSV curves of Co@N-PGCS and corresponding catalysts.

[150]. The carbon material structure and N species can be regulated by tuning the carbonization temperature. STS800 pyrolyzed at 800 °C showed excellent ORR performance due to high graphitization and abundant pyridinic N and graphitic N species. Later, Tong et al. used chitosan as the precursor, triphenylphosphine as the phosphorus source, and ZnCl<sub>2</sub> as the activated pore-forming agent to prepare N, P double-doped porous graphite carbon materials by hydrothermal reaction and carbonization [151]. P doping can significantly increase the BET SSA and pore volume of carbon materials and enhance charge delocalization to obtain carbon structures with more defects and edge sites. Compared with the Pt/C, the as-prepared catalyst showed comparable ORR activity ( $E_{\rm onset} = -0.014$  V vs. AgCl) and better methanol resistance.

Transition metal/carbon ORR catalysts with strong M-N-C bonds exhibit synergistic effects in further improving their ORR performance [152,153]. Zhang et al. used chitosan to obtain cobalt-doped nitrogen-containing carbon nanotube (Co-NCNTs) ORR catalysts through high temperature carbonization pyrolysis and catalyzed by transition metal cobalt (Fig. 8K-N4) [154]. It was found that the doping of cobalt changed the shape of the carbon matrix from nanosheet to nanotube (Fig. 8K and L). The successful synthesis of Co-N-C active sites was crucial for improving the ORR performance of the Co-NCNTs. The  $E_{1/2}$  of the Co-NCNT catalyst was 0.839 V vs. RHE, which was comparable to that of Pt/C (Fig. 8O and P). Co, N co-doped carbon nanosheets (Co@N-PGCS) were obtained by a facile Mg(OH)2 nanocasting strategy using chitosan as carbon source (Fig. 8Q) [155]. The structure of Co@N-PGCS was mainly mesoporous and had a large BET SSA (1716 m<sup>2</sup> g<sup>-1</sup>) and pore volume (1.55 cm<sup>3</sup> g<sup>-1</sup>), thus promoting the diffusion of reactive species. Compared with Pt/C, Co@N-PGCS exhibited excellent activity with -0.075 V vs. AgCl for  $E_{\text{onset}}$  and -0.151 V vs. AgCl for  $E_{1/2}$ in 0.1 M KOH (Fig. 8R and S).

#### 4. Conclusion and outlook

ORR plays an essential role in the performance of energy conversion storage devices. Therefore, technological progress towards economical, efficient, and sustainable materials is very necessity. The work above showed that biomass from plants, animals, and carbohydrates holded out great prospects as precursors for superior ORR electrocatalysts. Although biomass carbon is versatile, the advantages and disadvantages of each biomass carbon are obvious. Animal biomass carbon can contain elements that are unique to the catalytic activity center, such as iron in pig blood biomass carbon, but it is difficult to have fixed morphological characteristics in animal biomass carbon. Plant biomass carbon can be utilized with the original morphological characteristics of the plant, for example, stem biomass can retain the original tube packing structure, but basically all biomass carbon pore structures are difficult to regulate, and the pore structure is very complex. Glucose derivatives and chitosan-like substances can be made into nanospheres, as well as can be combined with drinking multi-catalytic active centers, but the relationship between their catalytic active centers and heteroatoms in biomass is not clear.

However, the differences between the same organisms growing in different areas make reproducibility difficult. Hence, in order to develop superior catalysts, this issue should be addressed. First, more attention should be paid to the properties of various biomass, and a general reproducible preparation route that is more suitable for different organisms should be designed. Second, standard experimental parameters which are essential to the reproducibility should be well controlled, including the manufacturer and origin. Finally, the general applicability of similar fractions across various biomass should be considered so as to further develop reproducibility. It is believed that a mass of rational preparation process will emerge in the next few years for the manufacture of new sustainable carbon-based materials from biomass resources, and it is hoped that this research will contribute to a wider application of sustainable carbon-based materials.

#### Data availability

No data was used for the research described in the article.

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