Molten-Salt-Assisted Synthesis of Nitrogen-Doped Carbon Nanosheets Derived from Biomass Waste of Gingko Shells as Efficient Catalyst for Oxygen Reduction Reaction

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Abstract: Developing superior efficient and durable oxygen reduction reaction (ORR) catalysts is critical for high-performance fuel cells and metal–air batteries. Herein, we successfully prepared a 3D, high-level nitrogen-doped, metal-free (N–pC) electrocatalyst employing urea as a single nitrogen source, NaCl as a fully sealed nanoreactor and gingko shells, a biomass waste, as carbon precursor. Due to the high content of active nitrogen groups, large surface area (1133.8 m² g⁻¹), and 3D hierarchical porous network structure, the as-prepared N–pC has better ORR electrocatalytic performance than the commercial Pt/C and most metal-free carbon materials in alkaline media. Additionally, when N–pC was used as a catalyst for an air electrode, the Zn–air battery (ZAB) had higher peak power density (223 mW cm⁻²), larger specific-capacity (755 mAh g⁻¹) and better rate-capability than the commercial Pt/C-based one, displaying a good application prospect in metal-air batteries.

Keywords: N-doped carbon nanosheet; gingko shells; oxygen reduction reaction; Zn–air battery

1. Introduction

The oxygen reduction reaction (ORR) plays a key role in fuel cells and metal–air batteries to implement electrochemical energy conversion [1,2]. However, the sluggishness of ORR kinetics at the cathode limits their commercial application [3,4]. Pt and its alloys materials, recognized as the best ORR electrocatalysts, suffer from the resource deficiency, prohibitive cost and inferior durability [5,6]. Consequently, development of earth-abundance, cost-effective and performance-stable catalysts to substitute the precious-based catalysts for ORR is crucial yet challenging [7–9].

Heteroatom-doped carbon materials, such as P-doped carbon xerogel [10], N-doped nanotube [11,12], S-doped carbon [13,14] and B-doped graphene [15], are among the most promising electrocatalysts for ORR due to their advantages of relatively low cost, abundant reserves, good electronic conductivity and environmental friendliness [16,17]. Among them, N-doping has been considered to be one of the most effectual ways to enhance the electrocatalytic activities of carbon materials for ORR [18,19] because the N-doping can improve the electrical conductivity of carbon and change the charge distribution of the carbon atoms close to the dopants and then make them positively charged, which is beneficial to the adsorption and reduction of oxygen at the corresponding sites and improving the ORR activity [7,20,21].

However, the morphology of N-doped carbon materials prepared by traditional methods is hardly controlled, and the content of the N-dopant is relatively low because of the volatilization of the N element at high temperature, which has an undesirable effect on
the electrocatalytic performance of the prepared materials. The employment of nanoreactor during the synthesis process can help relieve these challenges. The assistance of inorganic salts, such as NaCl [22,23], MgO [24], SiO$_2$ [25,26] and Fe$_3$O$_4$ [27], has been considered to be an effective strategy to prepare porous carbon materials. Among them, NaCl is one of the most commonly used because NaCl can not only act as a confining agent to reduce the weight loss of precursors and promote the N-doping during the high-temperature pyrolysis process but also as pore-forming agents to promote the production of macro-pores and a 3D network structure. In addition, NaCl can be conveniently removed by water-washing with no use of any corrosive and harmful chemicals and can be facilely recycled by recrystallization. However, a cost-effective and large-scale synthetic production method is still highly desirable but challenging. During the past decade, various biomass-derived carbon ORR catalysts have aroused ever-growing attention [28–31]. Gingko is mainly used for food and medicine, but the shells are usually thrown away as garbage, which not only causes a huge waste of resources but also leads to environmental pollution.

Herein, we report a molten-salt-assisted strategy to fabricate a 3D high-level nitrogen-doped porous carbon nanosheets network (N–pC) by a facile high-temperature pyrolysis method, employing the carbon-enriched biomass waste of gingko shells and urea as a carbon precursor and NaCl and urea as a fully sealed nanoreactor and a nitrogen source, respectively. The merits, such as large specific surface area, hierarchical pore structure and high active nitrogen (pyridinic and graphitic-N) content, endow the obtained N–pC excellent electrocatalytic ORR performance in alkaline solution, much superior to commercial Pt/C and most of the reported N-doped carbon materials. Inspired by the excellent ORR catalytic performance, the prepared N–pC was further used to assemble Zn–air batteries, which showed high peak power density, large specific-capacity and good rate-capability, making it a competitive alternative to the application in metal–air batteries.

2. Materials and Methods

2.1. Synthesis of the Materials

N–pC was prepared by pyrolyzing the mixture of gingko shells, NaCl and urea. Typically, 0.5 g gingko shells rinsed by deionized water, 1.4 g NaCl and 4 g urea were mixed to refined powder by planetary ball-milling for 4 h. Then, the mixture was initially carbonized under Ar atmosphere at 750 °C for 2 h with a ramp rate of 3 °C min$^{-1}$. Subsequently, the obtained product was refluxed in 3 M HCl (50 mL) for 6 h and washed with deionized water 3 times. After being dried at 80 °C for 24 h, the obtained black powder was further pyrolyzed at 900 °C for 2 h, and then the final product (N–pC) was harvested. For comparison, N–C catalyst was fabricated through the same procedure in the absence of NaCl. C catalyst was also synthesized by direct pyrolysis of the gingko shell powder.

2.2. Characterizations

Rigaku Ultima IV diffractometer (Cu Ka radiation) was employed to obtain the X-ray diffraction (XRD) spectra. Microstructural morphologies were observed employing field emission scanning electron microscopy (SEM, ZEISS SUPRA 55) and transmission electron microscope (TEM, JEM-2100F). To analyze the specific surface areas and pore-size distribution, nitrogen adsorption–desorption isotherms were tested on ASAP 2000 analyzer. X-ray photoelectron spectroscopy (XPS) measurement was implemented employing a Thermo Scientific Escalab 250 Xi.

2.3. Electrochemical Measurements

All the electrochemical tests were implemented on bipotentiostat (CHI760E, Shanghai, China) assembled with a rotation system (Pine Research Instrumentation, Durham, NC, USA) in a three-electrode system. A carbon rod, an Ag/AgCl electrode (in saturated KCl solution) and a glassy carbon electrode (GCE) loaded with various catalyst (N–pC, N–C, C or benchmark 20 wt.% Pt/C) were used as the counter electrode, reference electrode, and working electrode, respectively. To prepare the working electrode, 14 µL the catalyst ink
obtained by dispersing 1 mg catalyst powder into 0.5 mL Nafion solution (0.05 wt.%) via sonication was dropped onto the GCE, keeping the catalyst loading of 141.6 µg cm$^{-2}$. All potentials versus Ag/AgCl electrode ($E_{\text{Ag/AgCl}}$) were converted to the reversible hydrogen electrode ($E_{\text{RHE}}$) based on the formula of $E_{\text{RHE}} = E_{\text{Ag/AgCl}} + E^0_{\text{Ag/AgCl}} + 0.059pH$.

Cyclic voltammetry (CV) and linear sweep voltammetry (LSV) measurements were carried out in 0.1 M KOH electrolyte at a scanning rate of 100 mV s$^{-1}$ and 10 mV s$^{-1}$, respectively. The electron transfer number ($n$) was calculated using the Koutecky–Levich (K-L) equation:

$$\frac{1}{J} = \frac{1}{J_k} + \frac{1}{B\omega^{1/2}}$$  \hspace{1cm} (1)

$$B = 0.62nFC_0D_0^{2/3}v^{1/6}$$  \hspace{1cm} (2)

where $J$ and $J_k$ are the measured disk’s current density and the kinetic current density, respectively, $\omega$ is the angular velocity of rotation, $F$ is the Faraday constant (96,485 C mol$^{-1}$), $C_0$ is the O$_2$ concentration in electrolyte ($1.2 \times 10^{-6}$ mol cm$^{-3}$), $D_0$ is the O$_2$ diffusion coefficient ($1.9 \times 10^{-5}$ cm$^2$ s$^{-1}$), and $v$ represents the kinematic viscosity of the electrolyte ($1.13 \times 10^{-2}$ cm$^2$ s$^{-1}$).

The yield of HO$_2^-$ and the $n$ values for ORR via the RRDE measurements were determined by the equations as below:

$$\text{HO}_2^-\% = 200 \times \frac{I_r/N}{I_d + I_r/N}$$  \hspace{1cm} (3)

$$n = 4 \times \frac{I_d}{I_d + I_r/N}$$  \hspace{1cm} (4)

where $I_d$ and $I_r$ are the current density on disk and ring electrode, respectively, and $N$ is the current collection efficiency of Pt ring, which is equal to 0.37.

The durability and the possible methanol crossover effect of N–pC and benchmark Pt/C catalyst were evaluated by chronoamperometric measurement at a rotation of 1600 rpm and a fixed potential of 0.7 V (versus RHE) in the O$_2$-saturated 0.1 M KOH solution.

The homemade Zn–air battery (ZAB) was equipped with a polished pure Zn plate (0.5 mm in thickness) as the anode, carbon paper (2 cm $\times$ 2 cm) uniformly coated with the catalyst (N–pC or benchmark 20 wt.% Pt/C, mass loading: 1 mg cm$^{-2}$) as the air cathode and 6 M KOH solution as the electrolyte.

3. Results and Discussion

The synthetic procedures of N–pC are illustrated in Scheme 1. Gingko shells, NaCl and urea were first mixed to a refined powder by ball-milling. Then, the powder was initially carbonized at 750 °C (slightly lower than the melting point of NaCl to avoid the structural collapse of newly formed carbon materials). After the removal of NaCl by water washing and further pyrolysis at 900 °C to increase the graphitization degree and conductivity, the final product (N–pC) was obtained. The compositions of the C, N–C and N–pC were characterized by XRD. As shown in Figure 1, the diffraction peaks of C, N–C and N–pC at 26° and 44° correspond to the (002) and (101) planes of graphite carbon, respectively, indicating that gingko shells have been successfully transformed into carbon materials under high-temperature pyrolysis. The characteristic peaks of NaCl cannot be observed in N–pC, indicating that the NaCl template can be removed by simple washing. In addition, we notice that the peak intensity weakens in the order of C, N–C and N–pC, suggesting that N-doping can increase the number of defect sites and the addition of NaCl can further induce the formation of more defect sites.
The morphologies of the synthesized materials were observed by SEM and TEM. As shown in Figure 2a, the C catalyst obtained from the direct pyrolysis of gingko shells without the addition of urea and NaCl presents a compact and irregular block, whereas N–C has a significantly much looser and more porous morphology (Figure 2b), which mostly results from the large amount of pyrolysis gas of urea. With the further addition of NaCl, acted as sealed nanoreactor, the N–pC shows a 3D porous networks structure composed of graphene-like nanosheets (Figure 2c), which is beneficial to the increase of the specific surface area, the exposure of more active sites, and the improvement of transport rate of various substances involved in the reaction. TEM results are consistent with SEM. As shown in Figure 2d,e, the N–pC consists of a large number of graphene-like thin nanosheets, which interleave with each other, resulting in the formation of hierarchical pore structure and somewhat distorted of the carbon crystal faces. According to the elemental mapping (Figure 2f), the elements C, N and O are homogeneously distributed in the nanosheet, indicating the successful doping of N in N–pC catalyst.

The effect of urea and NaCl was further investigated by nitrogen adsorption/desorption measurements. As shown in Figure 3a, a distinct nitrogen uptake occurs at relative low pressure and a hysteresis loop appears at medium pressure in the isotherm curves of C, N–C and N–pC, indicating the existence of micro- and meso-pores in the three samples, which may be ascribed to the gas released from the decomposition of urea and gingko shells in the carbonization process. In addition, the isothermal adsorption curve of N–pC shows a sharp adsorption in the relative high-pressure region, demonstrating the existence of macropores on the material as the result of the occupying space of NaCl. The corresponding pore size distribution of the three materials (Figure 3b) further confirms their porous structures. Due to the different pore structure, the specific surface area and pore volume of C (641.9 m$^2$ g$^{-1}$, 0.365 cm$^3$ g$^{-1}$), N–C (1102.6 m$^2$ g$^{-1}$, 1.026 cm$^3$ g$^{-1}$) and N–pC (1133.8 m$^2$ g$^{-1}$, 1.22 cm$^3$ g$^{-1}$) are different. The large specific surface area and pore volume of N–pC means it can fully expose the active sites and most effectively accelerate mass transport, thus improving the electrochemical catalytic performance.
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Figure 2. SEM images of (a) C, (b) N–C and (e) N–pC; (d) TEM; (e) HRTEM; and (f) element mapping images of N–pC.

Figure 3. (a) N2 adsorption/desorption isotherms (inset shows the enlarged plots) and (b) the related pore size distribution curves of C, N–C and NpC.

The surface composition of the C, N–C and N–pC samples were investigated by XPS measurements. Figure 4a shows the full spectrum of the three materials. Compared with C, both N–C and N–pC contain N, indicating that the addition of urea can successfully realize N-doping. However, the N content of N–pC (5.18%) is higher than that of N–C (4.65%) due to the existence of NaCl’s confining effect acting as a fully sealed nanoreactor, which makes the volatilization and escape of N-containing gas generated by urea pyrolysis reduced. As can be observed from the high-resolution C 1s spectrum of N–pC (Figure 4b), there are four peaks located at 284.67, 285.72, 286.98 and 289.50 eV assigned to the C = C, C = N/C–O, C–N/C = O and O–C = O, respectively [32]. The presence of C = N and C–N bonds suggests that N atoms have been successfully incorporated into the carbon skeleton. Additionally, the N 1s spectra (Figure 4c) can be well separated into four species, corresponding to pyridine N at 398.07 eV, pyrrole N at 399.1 eV, graphitic N at 400.72 eV and oxidized N at 403.4 eV [33]. It has been reported that pyridine N and graphitic N can improve the electrocatalytic ORR performance of the materials, in which pyridine N can promote O2 adsorption on the neighboring carbon and graphitic N can enhance the electrical conductivity [34–36]. Combining Figure 4a,d, it is easy to find that N–pC possesses a much higher content of pyridine N and graphitic N than N–C. Therefore, the excellent ORR electrocatalytic performance of N–pC will be expected.
Cyclic voltammetry (CV) was conducted to evaluate the electrocatalytic ORR performance of C, N–C and N–pC in O₂-saturated 0.1 M KOH electrolyte. As it can be observed from Figure 5a, the CV curves of C, N–C and N–pC show an obvious cathode reduction peak, among which N–pC gives the most positive oxygen reduction peak and the highest peak current density, indicating that N–pC exhibits the highest electrocatalytic activity among the three catalysts. The linear sweep voltammetry (LSV) curves were also measured to further evaluate the ORR activity. As shown in Figure 5b, the C, N–C and commercial Pt/C catalysts have the onset potential (E_{onset}) of 0.756 V, 0.913 V and 1.012 V and the half-wave potential (E_{1/2}) of 0.585 V, 0.773 V and 0.836 V, respectively. Compared with those catalysts, the N–pC has the more positive onset potential (1.020 V) and half-wave potential (0.863 V) and higher limiting current density. In addition, as presented in Figure 5c, the Tafel slope of N–pC (59 mV dec⁻¹) is slightly smaller than that of commercial Pt/C (62 mV dec⁻¹), suggesting that N–pC has a comparable ORR kinetic process to that of commercial Pt/C. All these results suggest that N–pC has better electrocatalytic ORR activity than C, N–C, Pt/C and most of the reported metal-free catalysts (Table S1) in alkaline media. The superior catalytic performance of N–pC can be attributed to the incorporation of highly active N and the 3D hierarchical porous network structure.

To further evaluate the reaction kinetics of the ORR, the polarization curves at different rotation rates of N–pC were acquired (Figure 5d). Based on the corresponding Koutecky–Levich (K–L) plots, the transferred electron number (n) of the C catalyst are calculated to be about 2–2.5, suggesting that the ORR on the C electrode follows a two-electron and two-step mechanism. For the N–C catalyst, the n values are about 3.5, meaning that the ORR current efficiency is widely increased after the N-doping. On the contrary, the values of n for N–pC are close to 4 in a wide potential range, implying that a 4e⁻¹ process occurs on the N–pC electrode (Figure 5e). The RRDE experiments of the C, N–C and N–pC catalysts were carried out to further quantify the n and HO₂⁻ yield. As shown in Figure 5f, the n values for the C, N–C and N–pC are about 2.4, 3.5 and 4, respectively, entirely consistent with the RDE measurement results. Meanwhile, the HO₂⁻ yield at the N–pC is approximately equal to 0 in the investigated potential range, which is much lower than that at N–C and C.
In addition, the tolerance for methanol and durability are also important factors in fuel cells. The chronoamperometric response of N–pC and Pt/C catalysts to methanol crossover was tested in the O₂-saturated 0.1 M KOH (Figure 6a). After the injection of 1 M methanol, an obvious current attenuation is observed on Pt/C due to methanol oxidation, while there was a negligible change in N–pC, indicating that the N–pC catalyst shows much better tolerance for methanol than commercial Pt/C catalyst. Furthermore, the durability was also evaluated (Figure 6b). After a continuous operation for 16,000 s, the current density on N–pC remains at about 90%, whereas it only maintains 79% on Pt/C under the same condition, confirming that the N–pC catalyst has higher durability than the Pt/C catalyst.

Figure 6. Comparison of (a) methanol tolerance and (b) stability for N–pC and 20% Pt/C catalysts measured at 0.7 V versus RHE in O₂-saturated 0.1 M KOH at a rotation speed of 1600 rpm.

To investigate the potential use of the N–pC catalyst in real batteries, the home-made Zn–air battery (ZAB) was constructed using 6 M KOH solution as the electrolyte, polished pure Zn plate as the anode and N–pC or Pt/C as the air cathode, respectively. The open circuit voltage provided by ZAB using the N–pC catalyst is 1.50 V (Figure 7a), higher than that of 1.43 V for the commercial Pt/C-based one. The polarization and power density curves of the ZAB were given in Figure 7b. The peak power density of the battery equipped with N–pC catalyst achieves 223 mW cm⁻² at a high current density of 347.8 mA cm⁻², higher than the Pt/C catalyst (182.6 mW cm⁻² at 304.9 mA cm⁻²), suggesting the N–pC...
can be employed as a potential cathode electrocatalyst for ZAB. The discharge curves of N–pC-based ZAB at different current densities were measured by typical constant current discharge test. As shown in Figure 7c, the voltage can quickly reach equilibrium when the current changes, confirming that the structure of N–pC is beneficial to the mass-transfer. When the discharge current density recovers to the initial value (10 mA cm\(^{-2}\)) after the high current discharge (50 mA cm\(^{-2}\)), the discharge voltage can be well restored to the initial value, suggesting the good stability of the N–pC. Finally, the ZABs were galvanostatically discharged at 50 mA cm\(^{-2}\) to investigate their stability and specific capacity. As shown in Figure 7d, the N–pC-based ZAB has higher and more stable voltage than that of the Pt/C-based one during the discharge process. When normalized to the mass of consumed Zn, the specific capacity of the ZAB prepared with N–pC exhibits 755 mAh g\(^{-1}\), higher than that of 709 mAh g\(^{-1}\) for Pt/C and similar or even better than recently reported catalysts (Table S2). All these results suggest that the N–pC catalyst has a good application prospect in ZAB.

![Graphs showing discharge performance of N–pC and Pt/C-based batteries](image)

**Figure 7.** (a) Open circuit voltages and (b) discharge polarization curves of N–pC and Pt/C-based batteries; (c) The rate capability behavior of N–pC-based battery; (d) The long-term galvanostatic curves of the Zn–air batteries at 50 mA cm\(^{-2}\) using N–pC and Pt/C as ORR catalysts.

### 4. Conclusions

In summary, we successfully prepared a 3D, high-level nitrogen-doped, metal-free (N–pC) electrocatalyst employing urea as a single nitrogen source, NaCl as a fully sealed nanoreactor and gingko shells, constantly renewable and naturally available, as a carbon precursor. Due to the high content of active nitrogen, large surface area (1133.8 m\(^2\) g\(^{-1}\)) and 3D hierarchical porous network structure, the N–pC showed a high onset potential of 1.020 V and a half-wave potential of 0.863 V, and superior tolerance for methanol and stability in alkaline media. Additionally, in ZAB, N–pC showed higher peak power density (223 mW cm\(^{-2}\)), larger specific-capacity (755 mAh g\(^{-1}\)) and better rate-capability than the commercial Pt/C-based ZAB, displaying a good application prospect in metal–air batteries.

**Supplementary Materials:** The following are available online at [https://www.mdpi.com/article/10.3390/pr9122124/s1](https://www.mdpi.com/article/10.3390/pr9122124/s1) Table S1: Comparison of electrocatalytic ORR performance between N–pC and state-of-the-art metal-free catalysts reported in the literatures in alkaline electrolyte; Table
S2: Comparison of the performance of primary Zn-air batteries assembled with various cathodic electrocatalysts.

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