Recent Progress on Carbonaceous Material Engineering for Electrochemical Hydrogen Peroxide Generation

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Received: 29 November 2019 / Revised: 20 January 2020 / Accepted: 12 February 2020 / Published online: 7 March 2020
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Abstract
Electrochemical synthesis of hydrogen peroxide ($\text{H}_2\text{O}_2$) provides a clean and safe technology for large-scale $\text{H}_2\text{O}_2$ production. The core of this project is the development of highly active and highly selective catalysts. Recent studies demonstrate that carbonaceous materials are favorable catalysts because of their low-cost and tunable surface structures. This brief review first summarizes the strategies of carbonaceous material engineering for selective two-electron $\text{O}_2$ reduction reaction and discusses potential mechanisms. In addition, several device designs using carbonaceous materials as catalysts for $\text{H}_2\text{O}_2$ production are introduced. Finally, research directions are proposed for practical application and performance improvement.

Keywords
Hydrogen peroxide · Carbonaceous materials · Heteroatom doping · Device design

Introduction
Hydrogen peroxide ($\text{H}_2\text{O}_2$) is a highly valuable chemical, among the 100 most important chemicals in the world [1]. As an environmentally friendly oxidant, $\text{H}_2\text{O}_2$ can be used not only in various chemical industries and environmental remediation [2] but also as a potential energy carrier for numerous energy conversion processes [3]. At present, the industrial production of $\text{H}_2\text{O}_2$ is an energy-intensive anthraquinone process that requires complex, large-scale installations and expensive catalysts [3]. Furthermore, a large amount of organic waste is produced in this process. More importantly, due to the instability of $\text{H}_2\text{O}_2$, additional security precautions should be taken in transportation and storage [4]. Therefore, having a low-cost and decentralized preparation scheme is necessary to solve the aforementioned problems [5]. Direct synthesis of $\text{H}_2\text{O}_2$ from $\text{H}_2$ and $\text{O}_2$ indicates a new direction; however, the risk of explosion is prominent and needs to be eliminated [6–8]. The direct production of $\text{H}_2\text{O}_2$ by two-electron $\text{O}_2$ reduction reaction (ORR), which offers a fast, clean, and safe $\text{H}_2\text{O}_2$ generation technology, has attracted much attention. The reaction process is expressed as follows:

$$\text{O}_2 + 2\text{H}_2\text{O} + 2e^- \rightarrow \text{H}_2\text{O}_2 + 2\text{OH}^- \quad U^0 = 0.68 \text{ V vs.RHE}$$

(1)

The two-electron ORR process has been practically used since the 1990s, and the key lies on active and selective catalyst screening [9]. Various catalysts have also been employed in electrochemical synthesis of $\text{H}_2\text{O}_2$ from two-electron ORR process, and noble metal-based catalysts show the best performance [10, 11]. However, finite reserves and high costs of precious metals limit their large-scale applications [12]. Alternatively, carbonaceous materials have been identified as efficient catalysts because of their abundance, high stability in electrochemical reactions, and tunable surface and structural properties [12–14].

In early studies in this field, pristine carbon materials were investigated as catalysts for ORR to produce $\text{H}_2\text{O}_2$; the activity and selectivity of such materials were relatively low [15, 16]. Unsatisfactory performance was mainly attributed to the unsuitable electronic structure that cannot offer
an optimized adsorption/desorption energy to the reaction intermediate (*OOH). Thus, numerous strategies, including defect design [13, 17] and heteroatom doping [12, 18–25], have been adopted to optimize the electronic structure to improve the activity and selectivity of carbon materials (Fig. 1). In this brief review, the performance of defects and heteroatom-doped carbonaceous materials are first introduced and their reaction mechanisms are also presented. Then, several devices using carbonaceous materials as catalysts for H$_2$O$_2$ production are introduced. Finally, we propose that topological structure design for high-current density catalysts, long-term run, and other strategies for practical applications should be investigated in the future.

Strategies in Carbonaceous Material Engineering

Defect Engineering in Carbonaceous Materials

**sp$^3$-C and Defects**

Liu et al. [17] proposed that defects and sp$^3$-C in carbonaceous materials were the active sites for reactant adsorption or reactive sites in electrochemical processes, resulting in improvement of oxygen reduction kinetics. Thus, the researchers prepared hierarchically porous carbon (HPC, Fig. 2a) and revealed the positive effect of defects and sp$^3$-C on H$_2$O$_2$ production. When the $I_D/I_G$ value of HPCs increased from 0.84 to 1.17 (Fig. 2b), the H$_2$O$_2$ production rate could be enhanced gradually, indicating a significant contribution of the sp$^3$-C bonds and defects to enhance the electrocatalytic performance. Combined with high surface area and porosity, the material exhibited a high-current density...
efficiency and high H₂O₂ production rate in both acidic and neutral solutions (Fig. 2c).

sp²-C Defects

Chen et al. [13] also investigated the effect of defects in porous carbon catalysts on ORR performance. The mesoporous carbon and microporous carbon prepared with a large number of sp²-C defects (Fig. 2d) exhibited high selectivity and activity for H₂O₂ production. To understand the role of defects in oxygen reduction, the researchers also investigated a wide variety of defect configurations (Fig. 2d) through theoretical calculations. As shown in Fig. 2e, all the defect configurations were located on the right side of the two-electron volcano, which preserved O–O bonds due to weak binding with oxygenated species and contributed to the formation of H₂O₂. Some types of defects had favorable energies of oxygenated species, and their activity may be higher than that of PtHg₄ [10]. This result was consistent with the experimental characterizations that a variety of sp²-C defects improved the performance of porous carbon (Fig. 2f). The important role of defects for two-electron ORR was also investigated and demonstrated in other carbon materials [26–29].

Heteroatom Doping of Carbonaceous Materials

Oxygen-Doped Carbonaceous Materials

In addition to considering the defects of C atomic arrangement, heteroatom doping is another effective strategy to tune the electronic structure of carbonaceous materials. Santos et al. [16] conducted a comparative study on Printex L6 and Vulcan XC-72 R carbon for electrogeneration of H₂O₂ by reducing dissolved oxygen in an alkaline solution. The results showed that the Printex L6 was more effective than Vulcan XC-72 R for H₂O₂ production. The researchers thought the most likely difference between two carbon materials is the presence of more oxygenated acid species in Printex L6, which could provide greater hydrophilicity, thereby increasing the activity for H₂O₂ production [16]. The researchers further studied the materials after pre-treatment with nitric acid or ammonia. The results showed that acid-treated Printex L6 was the most suitable material for the two-electron pathway of the ORR because it contained the largest concentration of oxygenated acid species, which increased the hydrophilic character of the carbon support [30].

To deeply investigate the specific roles of oxygen-containing functional groups in ORR performance improvement, Lu et al. [12] demonstrated a facile and general approach to catalyst development through the surface oxidation of abundant carbon materials (Fig. 3a) to significantly enhance the activity and selectivity (approximately 90%) for H₂O₂ production (Fig. 3c). Combined with theoretical calculations, the results revealed that the carbon atoms adjacent to several oxygen functional groups (–COOH and C–O–C, Fig. 3a, b) were the active sites for oxygen reduction reaction via two-electron pathway. The experimental observations and theoretical calculations provided new insights into catalyst development that may be relevant in the production of industrial chemicals by means of clean, renewable electrical energy [12]. Kim et al. [18] also developed an easily scalable mild thermal reduction in graphene oxide (mrGO, Fig. 3d) to form H₂O₂ from oxygen. Their prepared few-layered mrGO (F-mrGO) exhibited highly selective and stable peroxide formation activity at low overpotentials (< 10 mV) under basic conditions (Fig. 3f), thereby exceeding the performance of current state-of-the-art alkaline catalysts. The performance of H₂O₂ production could be further improved by annealing at 600 °C, as the density of active functional groups increased (Fig. 3e). Their findings revealed that sp²-hybridized carbon near-ring ether defects along sheet edges were the most active sites for peroxide production, providing new insights into the electrocatalytic design of carbon-based materials for effective peroxide production [18].

Sa et al. [31] also reported high activity for H₂O₂ production on carbon nanomaterials with enriched oxygenated carbon edges. Zhou et al. [32] studied activated carbon (AC) and revealed that the oxygen-containing groups on the AC function as oxygen reduction reaction sites for H₂O₂ electrogeneration. Furthermore, the oxygen functional groups on carbon materials were increased by the treatments of acid, alkali, electrochemical oxidation, and glow discharge plasma. All of these materials showed excellent properties of two-electron ORR [33–38]. These reports have proven that oxygen functional groups on carbon materials are effective for selective O₂ reduction to produce H₂O₂.

Nitrogen-Doped Carbonaceous Materials

Nitrogen doping in carbonaceous materials has been used in oxygen reduction reaction. The doped nitrogen species play a crucial role in reaction selectivity. Indeed, the nitrogen can be doped in the forms of pyridinic-N, pyrrolic-N, graphitic-N, quaternary-N, and N-oxide, as presented in Fig. 4a and characterized in Fig. 4b [21]. Lee et al. [39] found that N-doped carbon exhibited high electrocatalytic activity toward the two-electron ORR. Therefore, they fabricated N-doped porous carbon for electrocatalytic production of H₂O₂ and the selectivity reached as high as 93%. Although several researchers have proposed that pyridinic, pyrrolic, and graphitic nitrogen in porous carbon might contribute to the two-electron ORR, the specific active sites in N-doped carbon have not been clearly revealed [40–43].
The specific action of nitrogen doping in carbon materials was demonstrated by subsequent reports. Perazzolo et al. [19] investigated nitrogen-doped mesoporous carbon as metal-free catalysts for $\text{H}_2\text{O}_2$ production, and a linear dependence of the nitrogen content on two-electron ORR performance was presented. They believe that the activity of nitrogen-doped carbon was related to the protonation/deprotonation of pyridinic-N, which led to increased activity at acid pH due to the protonation of nitrogen. Iglesias et al. also revealed that the pyridinic-N encounters difficulty in weakening the O–O bond in acid solutions due to its protonation [44]; thus, it was more conducive to the generation of $\text{H}_2\text{O}_2$, which was consistent with previously reported results. By conducting comparison with other nitrogen-doped carbonaceous materials, the researchers also found that the high $\text{N}_{\text{pyridinic}}/\text{N}_{\text{pyrrole}}$ ratio in their catalysts would contribute to favorable selectivity [20].

Strasser et al. [21] deeply investigated the role of nitrogen-doped species in two-electron ORR based on porous nitrogen-doped carbon materials. They found different activities and selectivity in various electrolytes (Fig. 4c). It was reported that the contents of pyridinic-N decreased in acid conditions and the graphitic-N decreased in neutral and alkaline conditions after electrochemical measurement. Meanwhile, the pyrrolic-N increased in all measurement environments. When C atoms neighboring pyridinic-N and graphitic-N are attached to the OH group, the N1s binding energy becomes extremely close to the pyrrolic-N, which is consistent with the content variety of nitrogen. Thus, the researchers believed that the pyridinic-N played a significant role in acid solutions, whereas the graphitic-N boosted the process of two-electron ORR in neutral and alkaline conditions, which is consistent with a subsequent report [45]. The C atoms neighboring N-doped carbons were revealed as the active sites, and the role of pyridinic-N for two-electron ORR in acid conditions was affirmed again. As previously reported, pyridinic-N facilitates the charge transfer from the $\pi$ orbital to the antibonding orbital in $\text{O}_2$ to weaken the O–O bond.
bond because of its delocalized lone pair electrons [46]. The pyridinic-N could prevent this effect by occupying the lone pair electrons to form an N–H bond; thus, two-electron pathway occurred in acidic conditions, which was consistent with Refs. [19, 20, 47]. Although the pyridinic-N exhibited excellent selectivity for two-electron ORR, Strasser et al. [45] found that the selectivity would be decreased when the content of nitrogen-doped sites was extremely high. Therefore, having suitable nitrogen content was necessary for hydrogen peroxide production.

**Other Doped Carbonaceous Materials**

Similar to N-doped carbon catalysts that activated π electrons and induced charge redistribution, fluorine [22, 48], multiple elements [23, 25], doped carbon, and metal-nonmetal co-doped carbon [24, 49] also exhibited excellent performance for two-electron ORR. Zhao et al. [22] discussed fluorine-doped HPC for two-electron reduction. The experimental results and density functional theory (DFT) calculations revealed that incorporating CF2,3 into carbon plane promoted the activation of O2 and facilitated desorption of intermediate OOH, which was crucial to H2O2 synthesis. The catalyst was also used in an electro-Fenton system, which could effectively remove single organic pollutants and even real refinery wastewater [48].

Two heteroatom co-doped carbonaceous materials have also been investigated for H2O2 production. For example, hexagonal boron nitride (h-BN) domains in boron–carbon–nitrogen materials showed better performance toward the two-electron ORR than the structures with single B or N doping. As determined by DFT calculations, the interface between h-BN domains and graphene exhibited unique catalytic behavior that can preferentially drive the production of H2O2 [23]. Considering the promising activity and selectivity of oxygen functional groups and the nitrogen-doping modification in decreasing the overpotential, Zhao et al. [25] examined the synergistic effect of nitrogen dopants and –COOH groups on carbonaceous materials. Through theoretical calculations, the researchers found that the theoretical overpotential for two-electron ORR would be lower when the two groups combined in one carbon. To amplify the synergistic effect, they also designed metal-free carbon aerogel integrated with nitrogen dopants and surface –COOH groups for H2O2 production, which exhibited complete 100% selectivity for H2O2 with high yields and stability [25]. Furthermore, Cardoso et al. [50] discussed the performance varieties before and after electrochemically stabilized measurements on the N and O group-modified graphene and graphene oxide nanoribbons. The varieties of activity could be attributed to the changes of active functional groups in the process of long-term test. The results provided directions for the selection of stable catalysts in long-term preparation of H2O2.

Besides nonmetal modification, metal and nonmetal co-doped carbon materials are also favorable for oxygen reduction to H2O2 production. Oxygen could be effectively reduced on the Co–Nx–C site, but the main product is water, whereas the oxygen functional groups on carbon materials exhibited high selectivity for two-electron ORR with limited kinetics. Li et al. [49] prepared a Co–Nx–C site and oxygen functional group co-modified carbon catalyst combined...
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with all advantages of aforementioned two materials, which improved activity and selectivity. Similarly, the transition metal single-atom coordination Fe–C–O as an efficient \( \text{H}_2\text{O}_2 \) catalyst exhibited excellent performance in alkaline and neutral pH [24]. Among the metal (Fe, Co, and Mn) atom coordinations (see Fig. 5a), Fe coordination exhibited the best activity and selectivity, which was in agreement with their theoretical conclusions (Fig. 5b, c). This result provided many choices of coordination combinations for improving the performance of two-electron ORR. All of these materials present new strategies of catalyst design for oxygen reduction to \( \text{H}_2\text{O}_2 \).

**Device Development for \( \text{H}_2\text{O}_2 \) Production**

Device design is also a vital segment for the practical application of oxygen reduction to \( \text{H}_2\text{O}_2 \). Yamanaka et al. [15] used a previously reported solid polymer electrolyte (SPE) [51] to prepare \( \text{H}_2\text{O}_2 \) from \( \text{O}_2 \) and water. Neutral \( \text{H}_2\text{O}_2 \) solutions could be continuously produced without any toxic by-products through the method of complete exposure solid polymer electrolyte (Exp-SPE). As the electrosynthesis process should proceed at the three-phase boundary, the researchers improved the production rate by modifying the cathode with Nafion. The cathode side was coated with Nafion to maximize the active surface exposure, which resulted in significant enhancement of the electrocatalytic performance compared with the unmodified electrode [15]. Thereafter, the researchers designed a new apparatus by modifying the SPE and proposed an electrolysis method for direct production of neutral \( \text{H}_2\text{O}_2 \) solution from \( \text{O}_2 \) and water. A detailed diagram of their installation is presented in Fig. 6a. The core components included cathode plate, Nafion-117 film, anode plate, and current collector, which are closely connected through flanges, bolts, nuts, and Teflon gaskets [52]. The apparatus provided a three-phase boundary for the electrosynthesis of \( \text{H}_2\text{O}_2 \), and the effect of boundary in the system was observed to have outstanding improvement performance through exposure cathode. Then, the researchers modified the cathode chamber with previous Nafion modification strategy [15]. Figure 6b shows the effect of Nafion mass loading on the productivity of \( \text{H}_2\text{O}_2 \). The formation rate and concentration of \( \text{H}_2\text{O}_2 \) increased with the increase in Nafion mass loading and reached a plateau above 10 mg loading.

Chen et al. [26] also designed a low-cost, membrane-free reactor for oxygen reduction to \( \text{H}_2\text{O}_2 \) production. Figure 6c shows the schematic of the reactor consisting of three core segments: working electrode in chamber 1, production storage in chamber 2, and chamber 3 for counterelectrode. The membrane-free system was achieved by the hydrophobic polymer-coated gas diffusion layer, which prevents water seepage but allows gas diffusion. The device could obtain 0.3 wt% \( \text{H}_2\text{O}_2 \) through a two-hour operation without any chemical stabilizer additives. Moreover, a higher product concentration could also be achieved with a chemical stabilizer (Fig. 6d). The researchers also proposed four design criteria for \( \text{H}_2\text{O}_2 \) production through \( \text{O}_2 \) reduction: (1) Decomposition of \( \text{H}_2\text{O}_2 \) should be minimized, (2) large production rate with high-current efficiency can be achieved at sufficiently high-current densities and low applied cell potentials, (3) product crossover should be prevented, and (4) the operation should be at ambient conditions [26].

**Outlook**

With their outstanding advantages in costs, reserves, stability, activity regulation, and selectivity for two-electron ORR, carbonaceous materials are high-potential catalysts.

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Fig. 5  a All types of Fe atoms coordinated in 2D carbon with and without O species (Fe, O or N, and C atoms indicated by green, red, and gray markers). b Volcano plot for 2\( e^- \) ORR process to \( \text{H}_2\text{O}_2 \) (*OOH adsorption at C and Fe indicated by red and blue markers, respectively). c Electrochemical measurement of CNT and Fe-, Pd-, Co-, and Mn-CNT catalysts in 0.1 mol/L KOH at 1600 r/min. Copyright 2019, Springer Nature
for industrial electrochemical production of H$_2$O$_2$. However, the structure–activity relationship (especially the performance of single functional groups) remains unclear and needs to be further investigated by controllable experiments and precise DFT calculations. Other suitable catalysts can be found through DFT computations [53]. In addition, further attention should be given to topological structure design for active catalysts (actual determination of selectivity should be noted [54]) with high-current density, long-term run, and depressing the decomposition of H$_2$O$_2$ at high overpotentials. The catalysts may be low stability at high-current density, and reasonable experiments and characterization techniques should be applied to find the factors of material degradation [14] and provide targeted solutions. Combined with rational device design, the two-electron ORR process is expected to have other commercial applications in the future.

Acknowledgements This study was supported by Tianjin Science and Technology Project (No. 19YFSLQY00070), Foundation for State Key Laboratory of Organic–Inorganic Composites, Beijing University of Chemical Technology (No. oic-201901004), and Hundred Talent Program of the Chinese Academy of Sciences.

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Fig. 6  a Cell structure illustration of Exp-SPE electrolysis (including cell assembly and details of Teflon plate and current). b The effect of Nafion mass loading at the modified AC + VGCF cathode on Exp-SPE synthesis of H$_2$O$_2$. Copyright 2011, American Chemical Society.

c Design diagram of membrane-free device. d Electrochemical measurement of membrane-free device. Copyright 2017, The Royal Society of Chemistry.
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