Integrated and Superaerophilic Ni-O-C Electrode enables Fast and Stable Electrochemical H2O2 Production for Electro-Fenton-like Process

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Abstract

Fast and stable production of hydrogen peroxide ($\text{H}_2\text{O}_2$) through electrochemical pathways is crucial for wastewater treatment applications. With this objective, herein we report an integrated and superaerophilic electrode composed of atomically dispersed Ni-O-C sites enriched carbon nanosheets (IS-NiOC electrode) for electrochemical oxygen reduction to produce $\text{H}_2\text{O}_2$. Both experimental and theoretical results have proven that atomically dispersed Ni-O-C sites enable a low overpotential (260 mV at 0.1 mA cm$^{-2}$) and high selectivity (> 90% at 0.0 ~ 0.5 V vs. RHE) in neutral electrolyte. Compared with a commercial gas-diffusion electrode, the IS-NiOC electrode offers stronger affinity to oxygen bubbles and more robust three-phase contact points, resulting in high current density (~ 106 mA cm$^{-2}$ at 0.25 V vs. RHE) and superior stability (~ 200 h). These merits allow the application of the IS-NiOC electrode in an electro-Fenton-like process, which enables fast degradation of representative organic pollutants in both steady state and flow state.

Introduction

Hydrogen peroxide ($\text{H}_2\text{O}_2$) is an environmentally friendly and highly efficient oxidant with a wide range of applications in fields of wastewater purification, medical treatment and industrial synthesis$^1$–$^4$. At present, over 99% of $\text{H}_2\text{O}_2$ is produced by anthraquinone oxidation process; however, this process requires complex and large-scale infrastructure, generates a substantial volume of waste chemicals, and induces potential hazards in transport or store of the high concentration products$^5$–$^9$. For these reasons, an alternative route for small-scale on-site generation is urged to exploit. Direct electrochemical synthesis of $\text{H}_2\text{O}_2$ by oxygen reduction reaction via a two-electron pathway (2e ORR) has gained a satisfactory solution to address the issues associated with the anthraquinone process due to its low resource consumption, convenience, and green initiative$^8$,$^10$–$^14$. In practice, 2e ORR has already been successfully applied in electro-Fenton(EF)/electro-Fenton-like(EF-like) reactions$^{15}$–$^{17}$, which belong to economical advanced oxidation processes$^{18}$–$^{20}$ for hazardous organic pollutants degradation.

In order to achieve fast and stable $\text{H}_2\text{O}_2$ generation by an electrochemical method, catalytic electrodes with high current density at a low overpotential, high selectivity and excellent stability are desperately needed$^2$,$^{21}$–$^{24}$. Previous studies have already identified various active catalysts, including noble metals and their alloys$^{25}$–$^{28}$, active carbonaceous materials$^{11}$,$^{29}$–$^{32}$ and transition metal-doped carbon materials$^{33}$–$^{36}$. In contrast to tremendous progress in seeking for active catalysts, there are very limited efforts devoted to investigate advanced electrode architectures, which is equally important and nonnegligible as efficient electrode architectures can simultaneous accelerate electron and reactant transport rate.$^{37}$–$^{42}$ The general strategy to solve this problem is employing a Teflon-treated carbon fiber paper (TCFP) as the substrate to increase three-phase contact point (TPCP) by constructing gas-diffusion layer, as shown in Fig. 1a. However, the in-situ generated $\text{H}_2\text{O}_2$ on an electrode surface would easily oxidize the carbon-based catalysts/substrates, and eventually destroy the gas diffusion channels and
operation stability. Direct construction of active materials into micro-/nanostructures on conductive substrates to form integrated electrodes may reinforce the gas diffusion channels to against the destruction.\textsuperscript{38,39,43,44} As schemed in Fig. 1b, we hypothesize that the integrated and superaerophilic electrode establishes a dense and thick gas diffusion layer that can mitigate the H\textsubscript{2}O\textsubscript{2} corrosion process.

Following this line, herein we demonstrated a nickel-incorporated oxidized carbon nanosheets arrays on carbon-fiber paper (CFP) for electrochemical H\textsubscript{2}O\textsubscript{2} production. The nickel-incorporated oxygen-doped carbon catalyst (NiOC) exfoliated from CFP exhibited an outstanding intrinsic activity with an onset potential of \(\sim 0.5\text{V vs. RHE}\) and high selectivity (\(> 82\%)\) over a wide potential range of \(0.0 \sim 0.5\text{V vs. RHE}\) in neutral electrolyte. Proper surface modification of the NiOC electrode surface achieved strong affinity to gas bubbles, resulting in the fabrication of an integrated and superaerophilic NiOC electrode (IS-NiOC electrode) with robust and enriched TPCPs. Combining the advantages of the catalyst and surface property, the IS-NiOC electrode exhibited large current density (\(> 100\text{mA·cm}^{-2}\)) at \(0.25\text{V vs. RHE}\) and remarkable selectivity (\(> 90\%\) at \(0.0 \sim 0.5 \text{V vs. RHE}\)). More importantly, compared with a traditional electrode made by a drop-casting method (D-NiOC electrode, < 10 h stability), the IS-NiOC electrode possessed superior operation stability (\(\sim 200\) h) for electrochemical air reduction to produce H\textsubscript{2}O\textsubscript{2}. We also demonstrated the application of the IS-NiOC electrode in an EF-like process, which realized fast degradation of representative organic pollutants.

Results

Synthesis and characterizations of the IS-NiOC electrode. The IS-NiOC electrode was synthesized by a multi-step process, as schemed in Supplementary Fig. 1a. Firstly a conventional solvothermal method was employed to synthesis Ni(OH)\textsubscript{2} nanosheet (average size and thickness of \(\sim 1\mu\text{m}\) and \(\sim 50\text{nm}\), respectively) arrays on CFP, as shown in Supplementary Fig. 1b. Afterwards, the Ni(OH)\textsubscript{2} nanosheet arrays were \textit{in situ} transformed to NiOC catalyst (Fig. 1c, XRD can be seen in Supplementary Fig. 1c) with hierarchical and porous architecture by carbonizing polyoxyethylene (PEO) at a high temperature. Further, the NiOC electrode was modified by a specific concentration of fluorine-based polymers to achieve a superaerophilic surface. To demonstrate the structural advantage, the exfoliated NiOC catalyst was drop-casted on TCPF (D-NiOC electrode) as the contrast electrode. It was revealed that, once an individual air bubble (~ 50 µL) touched the electrode surface, about ~ 3.8s was needed for the bubble to spread out on D-NiOC electrode (Fig. 1g and Supplementary Movie 1) while the required time shrunk to ~ 0.8 s for the IS-NiOC electrode (Fig. 1e and Supplementary Movie 2), clearly demonstrating a much stronger affinity of the IS-NiOC electrode towards air bubbles. This phenomenon was attributed to the superaerophilic layer under neutral electrolyte (1 M Na\textsubscript{2}SO\textsubscript{4}), which could build up a numerous of sturdy air pockets for coalescing with bubbles.

Transmission electron microscopy (TEM, Fig. 2a) revealed that the final product was mainly composed of stubby carbon nanotubes, and high-resolution TEM image (Supplementary Fig. 2) illustrated that Ni nanoparticle (lattice spacing: 0.208 nm) was encapsulated in the multilayered carbon shell (lattice
spacing: 0.367 nm) and located at the tip area of stubby nanotubes. It should be noted that the Ni nanoparticles cannot be washed out even using strong acids. The atomic dispersion of Ni sites on NiOC was confirmed by aberration-corrected high-angle annular dark field scanning transmission electron microscopy (HAADF-STEM, Fig. 2b) where the bright dots corresponding to Ni single-atom sites (marked by white circles, size of the bright spots if ca. 0.2 nm) were homogenously distributed throughout the entire sample. Elemental mappings results further confirmed the homogeneous distribution of Ni, O and C species across the tubular structure (Fig. 2c) despite the presence of inevitable Ni nanoparticles.

Closer inspection on the electronic structure promotes the deep understanding of structure-activity relationship. X-ray photoelectron spectroscopy (XPS) was firstly performed, and the results (Supplementary Fig. 3a) further proved the presence of Ni, O and C elements in NiOC electrode. The existence of fluorine (F) (Supplementary Fig. 3b) in IS-NiOC electrode confirmed the successful modification by polytetrafluoroethylene (PTFE). The deconvoluted spectrum of nickel 2p (Fig. 2d) illustrated the coexistence of metallic and oxidation state nickel\(^{45-47}\), and the deconvoluted spectrum of oxygen 1s (Fig. 2e) revealed that oxygen existed in three forms including C-O-C (533.0 eV)\(^{11,48}\), C = O (531.96 eV)\(^{11,48}\) and Ni-O-C (530.82eV)\(^{48}\). Compared with oxidized carbon material (O-C) without Ni incorporation (Supplementary Fig. 4, synthesis details in SI), the additional Ni-O-C peak indicated that the atomically dispersed Ni sites were mostly coordinated with oxygen. The oxidation state of Ni was also examined by the position of rising edge of Ni K-edge X-ray absorption near edge structure (XANES). As shown in Fig. 2f, the Ni K-edge XANES spectra of NiOC contained different characteristic features to Ni-foil and NiO\(^{49}\). The inset in Fig. 2f highlighted that the rising edge of Ni K-edge XANES spectra was located between those of Ni foil and NiO, clearly showing the dominant unique electronic structure of Ni-O-C.

2e ORR performance of IS-NiOC electrode. The electrochemical \(\text{H}_2\text{O}_2\) production in neutral media is of practical significance as it can accommodate high concentration of \(\text{H}_2\text{O}_2\) with negligible decomposition\(^2,50\). The intrinsic 2e ORR performance of the as-grown NiOC (exfoliated from the NiOC electrode) was firstly evaluated by casting the ink onto the rotating ring disk electrode (RRDE, collection efficiency is ~ 0.35, Supplementary Fig. 5) in 0.1 M phosphate buffered saline (PBS). The oxygen reduction current was measured on a disk electrode (black solid line in Fig. 3a), and the amount of formed \(\text{H}_2\text{O}_2\) was simultaneously quantified by the Pt ring electrode (black dashed line in Fig. 3a). The corresponding \(\text{H}_2\text{O}_2\) selectivity was plotted (black line) in Fig. 3b as a function of potential. For comparison purposes, oxidized carbon (O-C) catalyst without Ni incorporation was synthesized and tested under the same condition. Notably, the NiOC catalyst possessed an onset potential of ~ 0.50 V (vs. RHE) at 0.01mA·cm\(^{-2}\) and achieved selectivity of > 82.8% at a wide potential range of 0-0.50V (vs. RHE) in neutral media. Analogously, the O-C catalyst exhibited a similar selectivity at a narrower potential range (0 ~ 0.33 V vs. RHE), but a strong distinction occurred on the overpotential (\(\Delta = 0.17\) V). In addition, a faster ORR kinetics was found on the NiOC catalyst than that of O-C catalyst, as reflected in Tafel slopes (118.2 mV dec\(^{-1}\) for NiOC and 161.0 mV dec\(^{-1}\) for O-C, Supplementary Fig. 6). In view of the former XPS and XANES analysis, we thus drew a conclusion that such a huge ORR performance disparity was
attributed to the presence of atomically-dispersed nickel in catalyst. Moreover, the NiOC catalyst was superior to those of reported carbon-based catalysts (O-CNTs\textsuperscript{11}, Fe-CNT\textsuperscript{33}, g-N-CNTs\textsuperscript{51}, summarized in Table S1) and represented an outstanding activity and selectivity of the catalyst for 2e ORR in neutral media.

To understand the intrinsic high activity of NiOC system, DFT calculations were performed. For the 2e ORR, there are two reaction steps\textsuperscript{52}:

\begin{align}
*{O}_2 + H_2O + e^- & \rightarrow *{OOH} + O^- \quad (1) \\
*{OOH} + e^- & \rightarrow H_2O_2 + * \quad (2)
\end{align}

where the asterisk (*) represents the active site. As shown in Fig. 3c, the energy of *OOH species on C-O surface was related to a thermodynamical barrier of +0.77 eV under external potential of 0.70 eV. Once the Ni single atom coordinates to oxygen forming the Ni-O-C bond, the binding between hydroperoxide group and carbon is significantly strengthened. Generation of $H_2O_2$ possesses a smaller barrier of -0.53 eV. Herein the Ni-O-C system should exhibit an improved performance than pristine C-O system. Furthermore, Bader charge analysis shows that the atomic charge of the potential active carbon site changes from positive (0.77 |e|) to negative (-0.53 |e|) after the Ni coordination with O, indicating that electron transfers from nickel to carbon substrate. As a result, the accumulation of electron facilitates the reduction of oxygen to generate $H_2O_2$.\textsuperscript{53}

The superaerophilic property of an electrode has already been demonstrated effective in accelerating the gas diffusion process, thereby greatly enhancing the current increasing rate in diffusion region\textsuperscript{39}. In this case, a fast-current increase rate as well as a large current density was observed in the IS-NiOC electrode by virtue of the advantages of catalyst and architecture. To evaluate the performance under high current density, a steady state ORR test was performed in with continuous oxygen bubbling by using H-Cell (as shown in Supplementary Fig. 7). The D-NiOC electrode with weaker affinity to gas bubbles was also tested for comparison. The typical ORR polarization curves with IR-correction of D-NiOC electrode and IS-NiOC electrode in neutral media were shown in Fig. 3d. The IS-NiOC electrode displayed an ultrahigh ORR performance with a rapid and stable current increase ($\sim$75 mA·cm\textsuperscript{-2} per 100 mV) and a large current density ($\sim$122 mA·cm\textsuperscript{-2} at 0.2 V vs. RHE, with IR correction). On the contrary, the D-NiOC electrode exhibited a limited current density (37.9 mA·cm\textsuperscript{-2} at 0.2 V vs. RHE, with IR correction) and slower current increasing rate ($\sim$17.6 mA·cm\textsuperscript{-2} per 100 mV). This phenomenon was attributed to lack of hierarchical architecture on D-NiOC electrode (as shown in Supplementary Fig. 8a), which weakened the interaction with gas bubbles. Moreover, benefiting from the superaerophilic nanostructure, the unique surface of IS-NiOC electrode could guarantee an unobstructed channel for gas reactants by constructing the robust TPCP and thick gas transfer channel.
Tafel analysis (Fig. 3e) illustrated that the IS-NiOC electrode provided a wider potential and current range ($\Delta = 0.1347 \text{ V}, 0-15.46 \text{ mA cm}^{-2}$) than that of D-NiOC electrode ($\Delta = 0.071 \text{ V}, 0-8.55 \text{ mA cm}^{-2}$) for linear portion of the curve, indicating that the gas diffusion process on IS-NiOC electrode was much faster than that of D-NiOC electrode. For the IS-NiOC electrode, the corresponding selectivity and production rate of $\text{H}_2\text{O}_2$ under different potential (without IR-correction) in neutral media were shown in Fig. 3f, where a high $\text{H}_2\text{O}_2$ selectivity of 90.4% and $\text{H}_2\text{O}_2$ production rate of 59.3 mg·cm$^{-2}$·h$^{-1}$ at 0 V vs RHE were observed. This performance enhancement was also applicable in alkaline electrolyte (1 M KOH), where the IS-NiOC electrode exhibited a large current density ($\sim 250 \text{ mA·cm}^{-2}$ at 0.3 V vs. RHE, without IR correction) and splendid $\text{H}_2\text{O}_2$ selectivity ($\sim 95$%), as shown in Supplementary Fig. 9. The IS-NiOC electrode can also reduce the oxygen in air with relatively lower concentration to form $\text{H}_2\text{O}_2$ in a neutral system, where the working electrode was floated horizontally on the surface of the electrolyte (as shown in Supplementary Fig. 10). The up-side surface can absorb air as the reactants and simultaneously release the liquid products from the down-side surface. For the IS-NiOC electrode, the 2e ORR current density in air atmosphere decreased to $\sim 20\%$ of that in oxygen atmosphere (Fig. 3g), which was consistent with the changing of oxygen concentration. In addition, the performance of IS-NiOC electrode and D-NiOC electrode in air got a similar tendency with those in oxygen atmosphere, as shown in Fig. 3h and 3i. The Tafel plots (Supplementary Fig. 11) indicated that, compared with the ORR potential of IS-NiOC electrode in pure oxygen, a negative potential shift ($55\text{mV}$) was observed for the same electrode in air. This potential shift was very close to the theoretical value of 46 mV (based on Nernst Equation, more details can be found in Supporting Information). The Tafel slopes of IS-NiOC electrode were $93.4 \text{ mV·dec}^{-1}$ and $92.3 \text{ mV·dec}^{-1}$ in air and pure oxygen, respectively, indicating that the decrease of oxygen concentration did not affect the ORR kinetics significantly.

It was found that the ORR performance of the IS-NiOC electrode was highly dependent on the PTFE-modification parameters, as shown in Fig. 3j. A very limited current density (1 mA cm$^{-2}$ at 0V) was observed on the electrode without PTFE-modification, mainly because of the weak oxygen bubble adhesion capability (Supplementary Fig.12). The IS-NiOC electrode soaked in a diluted PTFE concentration (0.05 wt%) cannot afford a strong oxygen bubble adhesion, whereas increasing the concentration would impede the electron transport. These two metrics should be simultaneous optimized for achieving a superior current density. After intensive experimental efforts, an optimal PTFE concentration came to 0.1 wt%. More noteworthy, a significantly prolonged 2e ORR stability was observed on IS-NiOC electrode when performing air reduction in neutral system. Under the same test condition, the IS-NiOC electrode (modified by 0.1 wt% PTFE) could continuously produce $\text{H}_2\text{O}_2$ ($\sim 25 \text{ mA·cm}^{-2}$) for over 200 h (navy plots in Fig. 3k), while the promising initial current density was only maintained for a few hours (<10 h) for D-NiOC electrode with the same mass loading (dark red plots in Fig. 3k). It is worth mentioning that, the $\text{H}_2\text{O}_2$ selectivity was kept at a stable value ($\sim 90\%$) during all the processes for both IS-NiOC and D-NiOC electrodes, indicating that the decay of the D-NiOC electrode was not caused by catalysts degradation. In a deeper analysis, it is revealed that both the IS-NiOC electrode and D-NiOC electrode suffered from the surface corrosion from the $\text{H}_2\text{O}_2$ production since the surface liquid contact
angles (LCAs, Supplementary Fig. 13) got a variable decrease after a period of running. The significant difference is attributed to the robust TPCP of IS-NiOC electrode which could alleviate the corrosion to a great extent. To validate this assumption, an IS-NiOC electrode with a thinner gas film (soaked in 0.1 wt% PTFE for shorter time, ~8 min) was subject to a stability running. As expected, the latter one showed a slightly worse stability (red plots in Supplementary Fig. 14). Based on the above information, we believe that the robust gas diffusion layer plays a vital role in stability enhancement. The surface properties of IS-NiOC electrode and D-NiOC electrode before and after long-term reaction were thus characterized in detail. The air diffusion side for both IS-NiOC and D-NiOC electrodes is still intact after long-term testing, indicating that oxygen can be continuously diffused into the electrode. However, as for the water diffusion side, the IS-NiOC and D-NiOC electrodes show a considerable difference in the affinity to water. As shown in Supplementary Fig. 13, a significant decrease on LCA (from 150.8° to 53.9°) was found for the D-NiOC electrode after long-term testing, while a slight decrease (from 154.2° to 119.5°) was observed in the LCAs of the IS-NiOC electrode. Considering the ORR reaction only occurred at TPCP, the fast-current density degradation was directly caused by TPCP vanishment.

**EF-like performance of IS-NiOC electrode.** Electrochemical advanced oxidation processes (EAOPs) have attracted substantial attention owing to the environmental versatility, high efficiency and safety\(^\text{54}\). Among all the EAOPs, the EF process is most popular, in which the oxidant (H\(_2\)O\(_2\)) is formed *in-situ* from oxygen at the cathode surface in acidic solutions. However, generation of large values of iron-containing sludge and limited operating pH range (an optimal run is performed at pH ~3) of EF process urges the development of EF-like process which can be operated in a completely heterogeneous and neutral system.\(^\text{55,56}\) Besides stable and fast H\(_2\)O\(_2\) generation, a desirable electrode should also possess the capability to activate H\(_2\)O\(_2\) to degrade organic pollutants (i.e. Fenton-like process)\(^\text{57-59}\). It is reported that H\(_2\)O\(_2\) can be activated to reactive oxygen species (ROS) along with the valence state change of Ni in Ni-based catalysts\(^\text{60}\). In this case, we thus evaluated the Fenton-like performance of NiOC catalyst in neutral system and observed that NiOC catalyst exhibited a more effective degradability compared with that of O-C catalyst under the same experiment conditions (Supplementary Fig. 15). Thus, combining the advantages of rapid H\(_2\)O\(_2\) production and Fenton-like activity in neutral system, the IS-NiOC electrode is supposed to show a high EF-like performance.

To evaluate the EF-like performance, a steady-state three-electrode system enabling on-site H\(_2\)O\(_2\) production and *in-situ* organic (Indigo) degradation was constructed, as shown in Fig. 4a. A Nafion membrane was adapted to separate the cathode and anode reaction, which could avoid the oxidation of generated H\(_2\)O\(_2\) on the anode surface. The IS-NiOC electrode was employed as the cathode to reduce oxygen to H\(_2\)O\(_2\), which was further converted to ROS assisted with NiOC catalyst. As shown in Fig. 4b and 4c, the IS-NiOC electrode took 60, 40 and 24 minutes to complete degrade Indigo (50 ppm in 15 ml) under the current densities of 10, 20, and 50 mA cm\(^{-2}\), respectively. In order to simulate the industrial pollutants degradation process, a two-electrode device (without any membrane) with an electrolyte circulating system was constructed, as schemed in Figure 4d and Supplementary Fig. 16. The IS-NiOC electrode was
employed as the cathode and the dimensionally-stable boron-doped diamond (BDD) electrode was used as the anode. The employment of BDD electrode enables accelerated organics degradation process as it is well-known to produce strong oxidant (hydroxyl radical, \( \cdot \text{OH} \)) through the discharge of water at high oxidation potentials. The oxygen gas flowed over the anode surface to provide reactant gas, and 50 mL electrolyte containing representative organic pollutant (50 ppm bisphenol A in 0.05 M \( \text{Na}_2\text{SO}_4 \)) flowed through the channel. As shown in Fig. 4e, both the BPA and total organic carbon (TOC) removal rate increased under an operation current of 40 mA (the operation voltage is almost constant at \( \sim 3.3 \) V). After 8 hours reaction, the BPA and TOC removal rate reached \( \sim 95\% \) and \( \sim 62\% \), respectively. Overall, the results demonstrated the great application potential of IS-NiOC electrode in EF-like process for wastewater treatment.

**Discussion**

In conclusion, we have successfully fabricated an integrated and superaerophilic electrode with a hierarchical architecture of NiOC catalyst for fast and stable \( \text{H}_2\text{O}_2 \) electrochemical generation in neutral media. The NiOC catalyst was demonstrated effective in selective production of \( \text{H}_2\text{O}_2 \) from electrochemical oxygen reduction, and the superaerophilic surface property further accelerated both oxygen diffusion and electron transport rates, thereby greatly improving the 2e ORR current density and stability. We also demonstrated the application of IS-NiOC electrode in EF-like process, which realized effective degradation of representative organic pollutants. The concept and principles in this work should be applicable for designing electrochemical \( \text{H}_2\text{O}_2 \) production electrodes in the future.

**Method**

**Synthesis of IS-NiOC electrode:** The IS-NiOC was fabricated by a three-step method. Firstly, \( \text{Ni(OH)}_2 \) nanosheets were constructed on microporous carbon fiber paper (CFP, fuel cell store) by solvothermal method. In a typical procedure, \( \text{Ni(NO}_3\text{)}_2 \cdot 6\text{H}_2\text{O} \) (1 mmol) and \( \text{CO(NH}_2\text{)}_2 \) (10 mmol) were dissolved in 36 mL of methanol to form a clear solution by continuous stirring. CFP which was pre-treated by oxygen plasma and the as-obtained solution were transferred to a 40 mL Teflon-lined stainless-steel autoclave. Then, the autoclave was kept at 120°C for 12 hours, and then allowed to cool down to room temperature naturally. Thus, the \( \text{Ni(OH)}_2 \) nanosheet arrays on CFP were obtained and subsequently rinsed with distilled water and ethanol each for 5 min with the assistance of ultrasonication, and dried at 80°C for 2 hours. Secondly, the \( \text{Ni(OH)}_2 \@\text{CFP} \) nanosheet arrays was propped on a porcelain boat which was filled with polyoxyethylene (PEO, Mv~10,000) powder and the porcelain boat was put into the tube furnace with an Ar gas flow (40 sccm). The tube furnace was heated up to 900 °C in 90 min and was held at that temperature for 3 hours to carbonize the PEO and grow carbon materials in-situ on the CFP. The distance between the CFP and the PEO powder was kept between 0.5 and 1 cm, and the PEO amount was kept between 0.5 and 2 g. Lastly, the as-prepared NiOC electrode was soaked in PTFE (0.1 wt%) for 10 min, and then heated at 350 °C for 30 min in air atmosphere. The mass loading of the catalyst was measured as \( \sim 3\text{mg cm}^{-2} \).
**Synthesis of D-NiOC electrode:** The same mass loading (~3mg·cm\(^{-2}\)) of the NiOC catalyst was fabricated by the drop-casting method. Specifically, the NiOC catalyst was dispersed in ethanol solution with a concentration of 1 mg mL\(^{-1}\). Then the solution with NiOC was loaded on PTFE modified CFP by drop-casting.

**Synthesis of O-C catalysts:** 1g active carbon and 1g PEO was put into the tube furnace after mixing up, the tube furnace was heated up to 900°C in 90 min and held for 3 hours under Ar atmosphere.

**Characterizations:** The structural information of sample was characterized using field-emission SEM (Zeiss SUPRA 55) operating at 20 kV and high-resolution TEM system (JEOL 2100) operating at 200 kV. XPS spectrum was carried out by using a model of ESCALAB 250. X-ray powder diffraction patterns were recorded on an XRD (Rigaku D/max 2500) at scan rate of 10° min\(^{-1}\). The aberration-corrected HAADF-STEM measurements were taken on a JEM-ARM200F instruments at 200 keV. X-ray absorption spectroscopy (XAS) was conducted at the Shanghai Synchrotron Radiation Facility (SSRF).

**Surface Characterizations to Oxygen Bubbles:** This study characterized the wetting ability of the electrodes by measuring the contact angles of a Na\(_2\)SO\(_4\) solution using Optical contact angle and Surface/interfacial tension measuring system (OSA 60G, LAUDA Scientific, Germany). For these experiments, 50 μL of the electrolyte were dropped on the electrode surface, and the LCA was measured in ambient air at room temperature. The air-bubble CA with the volume of ~50 μL was measured by the captive-bubble method.

**Electrochemical RRDE Characterizations:** For the preparation of the catalytic electrodes, the NiOC catalysts were dispersed in ethanol to achieve a catalyst concentration of ~5 mg·mL\(^{-1}\) with 5 wt% Nafion solution. After sonication for 60 min, 10 μL of the catalyst ink was drop-dried onto a glassy carbon disc (area: 0.247 cm\(^2\), PINE). The electrochemical tests were performed in a computer-controlled CHI working station (Shanghai CHENHUA) with a three-electrode cell at room temperature. The glass carbon electrode loaded with catalyst was used as the working electrode. A graphite rod and a Ag/AgCl were used as the counter and reference electrode, respectively. 0.1 M phosphate buffered saline was chosen as the electrolyte. The ORR activity and selectivity were investigated by polarization curves and rotating ring-disk electrode measured in oxygen-saturated electrolyte at a scan rate of 10 mV·s\(^{-1}\).

H\(_2\)O\(_2\) selectivity of the NiOC catalyst on the rotating ring-disk electrode was calculated based on the current of both disc and ring electrodes (equation (1)). A potential of 1.2 V (versus RHE) was applied on the ring of the working electrode at a speed of 1600 r.p.m. during the entire testing process.

\[
H_2O_2 \text{ yield: } H_2O_2 \text{ (%) } = 200 \times \frac{I_R/N}{I_D + I_R/N}
\]

(1)
where $I_R$ is the ring current, $I_D$ is the disk current, $N$ is the collection efficiency (experimental calibration (~0.35) in Supporting Information (Supplementary Fig. 5)).

**Practical Electrochemical Characterizations:** To avoid the negative effect of Ohm resistance, an open system was employed to measure the current density (as shown in Supplementary Fig. 7). Practical ORR selectivity was performed in a two-compartment cell with Nafion 117 membrane as separator. Both the cathode compartment and the anode compartment were filled with the same electrolyte (20 mL of 1 M Na$_2$SO$_4$ or 1 M KOH). The electrolyte in anode compartment was saturated by oxygen gas, and 1 cm$^2$ electrode (IS-NiOC electrode, or D-NiOC electrode) was soaked in electrolyte as the working electrode. A Ag/AgCl or Hg/HgO electrode was employed as the reference electrode for neutral and alkaline systems, respectively. A graphite rod was placed in the cathode compartment as the counter electrode. To evaluate the H$_2$O$_2$ selectivity, the electrode was operated for several minutes under a constant potential, then the electrolyte was collected to quantify the H$_2$O$_2$ concentration. Theoretically, 1 C of electric quantity will produce 8.82 ppm H$_2$O$_2$ (equation (2)) under this condition (20 mL electrolyte).

$$\frac{1C \times 6.25 \times 10^{18} e/C}{6.02 \times 10^{23} e/mol \times 2} \times \frac{34 \ g}{mol \ 20 \ mL} = 8.82 \times 10^{-6} \ g \ mL^{-1} = 8.82 \ ppm \quad (2)$$

**Electrochemical Air Reduction Characterizations:** The electrochemical air reduction was performed by using the home-made device (as shown in Supplementary Fig. 10). A graphite rod and a Ag/AgCl were used as the counter and reference electrode, respectively. A flowing 1 M Na$_2$SO$_4$ solution was employed as the electrolyte. The selectivity was performed by calibrating the collected electrolyte.

**Fenton-like degradation:** A 50 mL of 10 mg/L RhB solution and 10 mg catalyst were transferred into the conical flask, as shown in Supplementary Fig. 15. The conical flask was kept under sonication until the catalyst homogeneous dispersion. Then, the conical flask was fixed on shaker and keep shaking for 12 hours to complete adsorption. 3 mL of 30 wt% H$_2$O$_2$ was put into the solution. To analysis the concentration of the organic, 1 mL degradation solution was extraction, and 1 mL of 0.5 M Na$_2$SO$_3$ was injected to stop the oxidizing of the remaining H$_2$O$_2$. After the centrifugation, the concentration was quantified by the UV-vis.

**Steady State of electro Fenton-like degradation:** An H-Cell was employed as the reaction device, and Nafion 117 membrane as the separator. Both the cathode compartment and the anode compartment were filled with the same electrolyte (20 mL of 0.05 M Na$_2$SO$_4$ and 50 ppm Indigo). The electrolyte in anode compartment was saturated by oxygen gas, and 1 cm$^2$ of IS-NiOC electrode was soaked in electrolyte as the working electrode. A Ag/AgCl was employed as the reference electrode and a graphite rod was placed in the cathode compartment as the counter electrode. The reaction current was set as 10, 20 and 50 mA, to collect the degrade tendency, 0.5 mL of the reaction solution was extracted and mixed with 0.5 mL Na$_2$SO$_3$. The concentration of Indigo was ensured by the UV-vis.
Flow State of electro Fenton-like degradation: The two-electrode and flow-state device was home-made, see details in Supplementary Fig. 16. The IS-NiOC electrode was employed as the cathode and the BDD electrode was employed as the anode (reaction on BDD electrode was as eq (3)). The working area for anode was 2*2 cm$^2$, and the distance between cathode and anode was 2 mm. The oxygen gas flowed in and out over the surface of the anode, and the electrolyte was pumped through the channel from the cathode by the peristaltic pump.

\[ BDD + H_2O \rightarrow BDD(\cdot OH) + H^+ + e^- \quad (3) \]

$H_2O_2$ concentration measurement: The $H_2O_2$ concentration was measured by a traditional titanium sulfonate Ti(SO$_4$)$_2$ titration method based on the mechanism that a yellow solution of H$_2$TiO$_4$ was produced by $H_2O_2$ (equation (4)). Thus, the concentration of H$_2$TiO$_4$ can be measured by ultraviolet-visible spectroscopy. The wavelength used for the measurement was 408 nm.

\[ Ti^{4+} + H_2O_2 + 2H_2O = H_2TiO_4 + 4H^+ \quad (4) \]

Therefore, the concentration of $H_2O_2$ (M) can be determined as M(H$_2$TiO$_4$).

The titanium sulfonate Ti(SO$_4$)$_2$ solution (2 mM) was prepared by dissolving 2 mmol Ti(SO$_4$)$_2$ in 1 L 1 M H$_2$SO$_4$ solution. To quantify $H_2O_2$ concentration, 1 mL or 2 mL Ti(SO$_4$)$_2$ solution was added into 1 mL neutral or alkaline media, respectively. To obtain the calibration curve, $H_2O_2$ with known concentration was added to Ti(SO$_4$)$_2$ solution and measured by UV-vis (calibration curves in neutral and alkaline media are shown in Supplementary Fig. 17). Based on the linear relationship between the signal intensity and H$_2$TiO$_4$ concentration, the $H_2O_2$ concentration of the samples could be obtained.

**Declarations**

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**Author contributions**
W.X., Z.L (Zheng Liang), and Z.L (Zhiyi Lu). designed research; W.X., B.Z., and X.C. performed research; W.X., X.S., and Z.L. (Zhiyi Lu) contributed new reagents/analytic tools; F.S and H.W did the GC and TOC tests. S. G and Z.T did the DFT calculations; W.X., B.Z., X.C., X.S., and Z.L. (Zhiyi Lu) analyzed data; W.X., Z.L (Zheng Liang), N.H, Tanja Kallio, L.C and Z.L. (Zhiyi Lu) wrote the paper.

Additional information

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**Figures**
Figure 1

(a-b) Schematic illustration of surface properties of conventional gas-diffusion electrode and superaerophilic electrode before and after long-term electrochemical oxygen reduction to H2O2; (c) typical SEM image of NiOC electrode (scale bar: 5μm); (d-e) air bubble adhesion behaviors on D-NiOC (d) and IS-NiOC (e) electrode under neutral electrolyte.

Figure 2

(a-b) High-resolution TEM (scale bar: 100 nm) and aberration-corrected HAADF-STEM (scale bar: 2 nm) images of NiOC catalyst; (c) STEM-EDX mapping results of NiOC catalyst (scale bar: 100 nm); (d-e) high-resolution XPS spectra of Ni 2p and O 1s in NiOC electrode; (f) Ni K-edge XANES spectra in NiOC electrode.
Figure 3

(a) Electrochemical oxygen reduction polarization curves at 1,600 r.p.m. (solid lines) and simultaneous H2O2 detection currents at the ring electrode (dashed lines) for the NiOC catalyst and O-C catalyst in 0.1 M PBS solution; (b) relevant calculated H2O2 selectivity; (c) free energy diagram of 2e ORR on NiOC and O-C at equilibrium potential of the reaction; (d) LSV curves (with IR-correction) of IS-NiOC electrode and D-NiOC electrode recorded in 1 M Na2SO4 solution with oxygen-bubbling; (e) corresponding Tafel slopes of...
IS-NiOC and D-NiOC electrodes; (f) corresponding selectivity and production rate of IS-NiOC electrode; (g-i) relevant performance of IS-NiOC and D-NiOC electrodes under air reduction; (j) current density in oxygen atmosphere varieties with the PTFE concentration; (k) long-term stability and selectivity of IS-NiOC and D-NiOC electrodes for H2O2 production.

Figure 4

(a) Schematic illustration of 3-electrode steady-state device for organic degradation; (b-c) operation voltages and degradation rates under different currents (conditions: 0.05 M Na2SO4, 50 mg/L Indigo, oxygen gas flow rate 80 sccm); (d) schematic illustration of flow state device, where the IS-NiOC and BDD electrode were employed as the cathode and anode, respectively; (e) BPA and TOC removal rates of the device constructed by IS-NiOC and BDD electrode electrodes (conditions: 0.05 M Na2SO4, 50 mg/L BPA, oxygen gas flow rate was 80 sccm, electrolyte recycle flow rate was 2 mL min-1, the volume of the circulating solution was 50 mL).

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