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Magnetic cathode stimulates extracellular electron transfer in bioelectrochemical systems

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Abstract: Exploring alternative cathodic catalysts capable of highly catalytic activity is crucial to the expansion of bioelectrochemical systems. Herein, Fe₃O₄@N-mC is developed as a magnetic cathode catalyst for bioelectroreduction of oxygen. The Fe₃O₄@N-mC exhibits better electrocatalytic activity, selectivity (four electron transfer...
pathway), and long-term electrochemical stability in neutral solutions compared to commercial Pt/C catalyst. The microbial fuel cell using Fe₃O₄@N-mC generates a power density of 1141 mWm⁻², which is higher than that of using Pt/C (1022 mWm⁻²). Furthermore, the decline of power density is much lower in reactor with Fe₃O₄@N-mC (4 %) than with Pt/C (8 %). With Fe₃O₄@N-mC, the cell also obtains higher coulombic efficiency (26 %) than that with Pt/C (21.7 %). The outstanding electrocatalytic activity and stability of Fe₃O₄@N-mC show its great potential to be a favorable substitute to Pt/C catalysts in microbial electrochemical energy devices.

Keywords: Encapsulated Fe₃O₄ nanoparticles; Magnetic cathode; Extracellular electron transfer; Oxygen reduction reaction; Bioelectrochemical systems

Introduction

The fast growth of global population and depletion of natural energy resources has caused overwhelming demand on the exploration of environmentally friendly and sustainable energy. Bio-battery such as microbial fuel cells (MFCs) have captured huge notice lately as superior eco-friendly biotechnology for recovering energy from biodegradable wastes.¹,² Exoelectrogenic bacteria in the MFC anode boost the decomposition of biodegradable organic matter and then pass the resulting electrons and protons to the electron receptors on the cathode.³ Oxygen (O₂) is regarded as an ideal terminal electron acceptor due to its high reduction potential (1.229 V), clean product, and prepared accessibility.⁴ There are two pathways for the oxygen reduction reaction (ORR) process. One is to reduce oxygen to H₂O directly via a four-electron pathway, and another one is to reduce oxygen first to H₂O₂ and then further be reduced to H₂O through a two-electron pathway. So, the four-electron pathway is more efficient and desirable in terms of energy recovery.⁵ One of obstacles for the practical
implementations of air-cathode MFCs is the sluggish kinetics of ORR.\textsuperscript{6,7} Traditionally, platinum and its alloys can attain the ORR in four-electron pathway by reducing the overpotential, and thereby resulting in high-efficiency energy recovery and power output. Nevertheless, their high capital cost and susceptibility to time-dependent drift compromise their extensive utilization and commercialization.\textsuperscript{8-10} Thus, great efforts have been done to explore cost-effective materials that possess comparable catalytic performance to platinum for the ORR in cathode.

Non-precious metal catalysts (NPMCs) with distinguish catalytic ORR activity, good stability and popular price have received increasing attention. Among which, nitrogen-doped carbon materials are regarded as favorable catalysts because they can promote the O\textsubscript{2} adsorption, increase active sites and improve the surface hydrophobicity, in addition to its high electrical conductivity and oxidation stability.\textsuperscript{11} As reported previously, nitrogen species can conduct a four-electron transfer process, which can reduce the O–O bond and could function as the active sites in ORR process.\textsuperscript{12} Nevertheless, most of the pure nitrogen-doped catalysts showed less ORR activity than Pt/C catalysts. Thus, it has been further integrated with nonprecious metals (especially, cobalt or iron) to enhance the ORR activity, endurance to poisons, and the durability. Recent studies have shown that the transition metal encapsulated within nitrogen-doped graphitic carbon layer to form a core–shell structured catalyst was a good substitute for Pt/C.\textsuperscript{13,14} In our previous study, we synthesized a core-shell structure Fe-N-C catalyst, which performed higher ORR activity than its intermediates, including N-C.\textsuperscript{15} The outstanding ORR activity of Fe-N-C catalyst compared to N-C can be ascribed to two reasons: (1) Fe-N coordination is generally regarded as the active sites in ORR process, which led a four-electron transfer pathway;\textsuperscript{16} (2) Fe acted as the catalyst in the carbon graphitization process increased the graphitic structures of the hybrid, resulting in
improving electrical conductivity and catalytic activity.\textsuperscript{17} While the iron core of Fe-N-C catalyst in our previous work was too big (around 400 nm) to sufficiently interact with the surrounding N or C, which limits its optimal ORR performance. In view of catalyst design, the introduction of high dispersion Fe\textsubscript{3}O\textsubscript{4} nanoparticles was expected to increase the Fe-N active sites and graphitic carbon to enhance the catalytic activity of Fe-N-C catalyst in ORR process.

Recent studies proved that magnetic anode and pulse electromagnetic fields enhanced the extracellular electron transfer (EET) of bioelectrochemical systems (BESs).\textsuperscript{18,19} We hypothesize that magnetic materials at the cathode might impact EET of BESs. Herein, we developed a magnetic core-shell structure catalyst (Fe\textsubscript{3}O\textsubscript{4}@N-mC), in which Fe\textsubscript{3}O\textsubscript{4} nanoparticles of around 8 nm were embedded in mesoporous nitrogen-doped carbon. A comprehensive comparison between the Fe\textsubscript{3}O\textsubscript{4}@N-mC and the Pt/C catalysts was conducted in catalytic activity, power production and long-term electrochemical stability. The Fe\textsubscript{3}O\textsubscript{4}@N-mC exhibited superior catalytic activity and stability to a commercial Pt/C catalyst, thus creating a new cost-effective catalyst for BESs.

Experimental Section

Preparation and characterizations of Fe\textsubscript{3}O\textsubscript{4}@N-mC

Fe(acac)\textsubscript{3} (0.7 g) was dissolved in triethylene glycol (TREG, 20mL), and then heated at 200 °C for 8 h to get Fe\textsubscript{3}O\textsubscript{4} nanoparticles as previously reported.\textsuperscript{20} Fe\textsubscript{3}O\textsubscript{4} (100 mg) was added to 1 mol/L of hydrochloric acid (100 mL) which dissolved sodium dodecyl sulfate (SDS, 0.288g) and poly (ethylene oxide)\textsubscript{20}-block-poly (propylene oxide)\textsubscript{70}-block-poly (ethylene oxide)\textsubscript{20} (P123, 0.15g) in advance. After 30 minutes ultrasonically dispersed, the mixture was added with aniline (0.36 g) and ammonium persulfate (0.9
g) in succession at 4 °C and vigorously stirred for 6 h to get Fe₃O₄@mPANI. Lastly, the obtained product was heated at 700 °C for 6 h in N₂ atmosphere to obtain Fe₃O₄@N-mC. Characterizations of Fe₃O₄@N-mC were tested as previously described.¹⁵ Details were shown in Supplementary Material.

**Electrochemical characterization of Fe₃O₄@N-mC**

The electrochemical property of catalysts were performed in a three electrode electrochemical cell using phosphate buffer saline as electrolyte (PBS, 50 mM), and furnished with a platinum sheet as counter electrode (2.25 cm²) and an Ag/AgCl electrode as reference electrode. The current density was recorded at a scan rate of 10 mV s⁻¹ with varied rotation rate from 100 to 1600 rpm. The ORR property of the catalysts was analyzed on RRDE-3A apparatus (BAS Inc., Japan) with a scan rate of 10mV s⁻¹, and the rotating speed was varied from 400 to 3600 rpm progressively. The glassy carbon (GC) working electrode was uniformly covered with a catalyst ink (0.1 mg cm⁻²), which was prepared by dispersing 5 mg electrocatalysts in 5 mL deionized (DI) water, then it was ultrasonicated for 30 min. The electrode was fixed by adding 5 μL Nafion solution (5 wt%). The rotating ring-disk electrode (RRDE) tests were conducted at a constant ring potential of 1.0 V (vs Ag/AgCl) with a rotating speed of 1600 rpm. The electron transfer number (n), kinetic current (𝑖_𝐾), and peroxide yield (H₂O₂) during ORR was calculated as previously reported.¹⁵

**Electrode Preparation and Operation Condition for MFCs**

The anode of MFC was carbon fiber brush. The gas diffusion layer (GDL) of MFC was made by rolling the mixture of carbon black and 60 wt% PTFE (mass ratio of 7:3) on a
stainless steel mesh, then sintered at 340 °C for 30 min. The catalyst layer (CL) was made by rolling the mixture of the catalysts and PTFE (mass ratio of 6:1) onto the opposite side of stainless steel according as previously reported. Single-chamber MFCs were constructed with an effective volume of 28 mL. The cathode had a projected area of 7 cm$^2$, which was connected with anode through a 1000 Ω external resistance by external wire. Per liter of the culture solution contained sodium acetate (2g), phosphate buffer solution (PBS, 50 mM) mineral solution (10 mL), and vitamin solution (10 mL). Activated sludge in secondary clarifier of wastewater treatment plant (Harbin, China) serves as inoculum. The Fe$_3$O$_4$@N-mC composites were tested as the cathodic catalysts for MFCs and commercial Pt/C (20 wt%) was used (5 mg cm$^{-2}$) for comparison. All MFC reactors were worked in fed-batch mode at the temperature of 30 ± 2 °C.

**Analytical and electrochemical techniques of MFCs**

The cell voltages were recorded automatically every 10 min over an external resistance (1000 Ω) by using a data acquisition system (Model 2700, Keithley Instruments Inc., USA) as previously reported. The electrochemical performances of the air cathode were tested by a potentiostat Autolab Potentiostat/Galvanostat (Autolab PGSTAT 128N, MetrohmAutolab Inc., Netherlands). LSV was performed from open circuit potential (OCP) to −0.5 V with a scan rate of 1 mV s$^{-1}$ while 50 mM PBS solution was used as the electrolyte in the tested MFCs. LSV was tested in a three-electrode configuration, in which the air cathode was used as working electrode, an Ag/AgCl electrode was used as reference electrode, and the anode was used as counter electrode.
Changing the resistance of the external circuit from 3000 to 50 Ω to get the polarization curve. The power density was calculated by using the equation of \( P = UI/A \), in which \( U \) stands for voltage, \( I \) stands for current, and \( A \) stands for the surface area of the cathode. Electrode (anode and cathode) potentials were recorded during polarization test by using Ag/AgCl reference electrode. The APHA standard method was used to measure the chemical oxygen demand (COD) of reactors. Coulombic efficiency (CE) was the percentage of practical electron charge produced in one cycle versus the available electrons from substrate metabolism.\(^{24}\)

**Results and discussion**

**Structure characteristics**

The TEM image shows \( \text{Fe}_3\text{O}_4 \) nanoparticles obtained in the first step are about 8 nm in diameter, and with good dispersing property (Fig. 1a). The observed morphology of \( \text{Fe}_3\text{O}_4@\text{N-mC} \) indicates that the catalyst was uniformly dispersed as expected (Fig. 1b). The flocculates with lamellar structure can be seen as the graphite layers embedding many small dark particles belonged to \( \text{Fe}_3\text{O}_4 \). SEM images revealed the catalyst had mesoporous spherical morphologies with nano-sized pores (Fig. 1c). Thus, the product might be consisted of a carbon shell and \( \text{Fe}_3\text{O}_4 \) core. The element mapping images demonstrate the existence of C (Fig. 1d), Fe (Fig. 1e) and N (Fig. 1f) elements. The surface area and porous structure of \( \text{Fe}_3\text{O}_4@\text{N-mC} \) were measured by nitrogen adsorption–desorption isotherms. A typical type-IV isotherm was attributed to its mesoporous structure. The BET surface area was 154.3 m\(^2\) g\(^{-1}\) and a uniform mesoporous size distribution with an average size of 4.2 nm was obtained from analysis of the desorption curve (Fig. S1). The mesoporous structure was beneficial to the
accessibility of active sites and facilitates mass transfer kinetics.

![TEM images and element mapping](image)

**Fig. 1** TEM images of Fe$_3$O$_4$ (a) and Fe$_3$O$_4$@N-mC (b); SEM(c) image of Fe$_3$O$_4$@N-mC; the element mapping images for C (d), Fe (e) and N (f) of Fe$_3$O$_4$@N-mC.

![XRD analysis](image)

**Fig. 2** XRD analyses of the main crystalline phases in Fe$_3$O$_4$ and Fe$_3$O$_4$@N-mC composite.
The microstructure of Fe₃O₄@N-mC was further verified by the XRD (Fig. 2). There were seven distinct diffraction peaks corresponds to the (220), (311), (400), (422), (511), (440), and (533) of Fe₃O₄ (JCPDS No. 65-3107), which confirmed the existence of Fe₃O₄ in the hybrid. The peak observed at around 25.5° belonged to the (002) planes of graphitic carbon and it should be ascribed to the iron nanoparticles playing as the graphitization catalyst to improve the graphitization degree of carbon layer during pyrolysis.¹⁷,²⁵ The existence of the graphitic carbon was also verified by Raman spectroscopy (Fig. S2). The signal peaks showed a G band and a weaker D band at around 1595 cm⁻¹ and 1343 cm⁻¹, respectively. The G-band was usually ascribed to the stretching of sp²-bonded on carbocycle or long-chain of ordered graphite, whereas, the D-band is related to defects in the graphite.²⁶ Therefore, the G/D intensity ratio was regarded as a measurement for order quantity in graphitic materials. The I_G/I_D ratio of Fe₃O₄@N-mC was 1.19, which indicated more order crystalline graphite and could be beneficial for ORR since it increases the electricity conductivity of the derived carbon.⁸

The XPS analysis revealed the content and chemical state of elements on Fe₃O₄@N-mC (Fig. 3). The survey scan of Fe₃O₄@N-mC showed that there were four kinds of elements, carbon (81.28%), oxygen (10.08%), nitrogen (5.01%), and iron (3.63%) (Fig. 3a). The XPS survey of C1s declared four kinds of C function group ascribed to C=C (284.8 eV), C-O (286 eV), C=O or C=N (287.5 eV), and C-N (289.9 eV) (Fig. 3b). It is generally believed that peak ascribed to C=O and C=N indicated the nitrogen is successfully doped into the carbon matrix.²⁷ The high-resolution N1s spectra were fitted with three peaks at 398.4 eV, 399.5 eV, and 401 eV, belonging to pyridinic N, pyrrolic N, and graphitic N, respectively (Fig. 3c).²⁸ In addition, the difference of binding energies between Fe-N and pyridinic N is slight,
Fig. 3 XPS survey spectrum (a) and high-resolution C1s (b), N1s (c) and Fe 2p (d) spectra of Fe₃O₄@N-mC catalyst.

so the peak at 398.4 eV should also be affiliated with Fe-N. Different nitrogen functionalities doped into graphitic carbon matrix is beneficial for ORR catalysis. Previous studies have reported that pyrrolic N play a role in reducing O₂ to H₂O₂, while pyridinic N can improve the onset potential and play a role in reducing H₂O₂ to H₂O and graphitic N contributes to the improvement of limiting diffusion current. Meanwhile, the pyridinic N is regarded as the main site for binding the transition metal ions, and the Fe-N dominates a four-electron transfer pathway for ORR. The Fe 2p profile can be reclassified into two components: Fe 2p₁/₂ and Fe 2p₃/₂, corresponding to 723.6 eV and 710 eV, respectively (Fig. 3d). No peaks detected between 710 eV to 724 eV proved that the purity of the Fe₃O₄ sample without γ-Fe₂O₃. Fe 2p peak at 711 eV was contributed to coordinating to nitrogen according to previous report.
Considering above, the distinctive structure and composition of Fe$_3$O$_4$@N-mC catalyst, which in detail is nitrogen-doped mesoporous graphitic carbon matrix shell embedded with Fe$_3$O$_4$ nanoparticles core, could to a large extent improve ORR activity. Fe$_3$O$_4$ nanoparticles core could also activate graphitic carbon layers and coordinate with nitrogen to form well-supplied available active sites and fast the transfer of ORR-relevant species, and afterward enhance the ORR electrocatalysis.

**Electrocatalytic ORR activity of Fe$_3$O$_4$@N-mC**

The electrocatalytic activity of the Fe$_3$O$_4$@N-mC toward ORR was assessed via rotating disk electrode (RDE) experiments compared with commercial catalyst (Pt/C, 20 wt%) (Fig. 4 and Fig. S3). When the rotating speed is 1600 rpm, polarization curves of Fe$_3$O$_4$@N-mC showed a slightly lower onset potential (0.302 V vs. Ag/AgCl) but higher limiting current density (3.54 mA cm$^{-2}$) than those observed in Pt/C (0.34 V and 2.99 mA cm$^{-2}$) (Fig. 4b). The ORR kinetics calculated based on Koutecky-Levich (K-L) equation demonstrated the rotation-dependent currents at different potential values. The linear K-L plots of Fe$_3$O$_4$@N-mC electrodes with good parallelism implied first-order reaction kinetics of oxygen concentration. The electron transfer number of Fe$_3$O$_4$@N-mC (n=3.86) was higher than that of Pt/C catalyst (n=3.77) (Fig. 4c and Fig. S4). The results indicate that the ORR in Fe$_3$O$_4$@N-mC dominantly followed the four-electron pathway reaction.
Fig. 4 Polarization curves of Fe$_3$O$_4$@N-mC recorded by RDE at various rotating speed (a), Linear scan voltammetry of Fe$_3$O$_4$@N-mC and Pt/C with a rotating speed of 1600 rpm (b), the Koutecky–Levich plots of Fe$_3$O$_4$@N-mC at different potential (c), polarization curves of Fe$_3$O$_4$@N-mC recorded by RRDE with a rotating speed of 1600 rpm (d), electron transfer number (n) of Fe$_3$O$_4$@N-mC and Pt/C (e), and the stability of the Fe$_3$O$_4$@N-mC and Pt/C with a constant potential of −0.4 V. The RDE and RRDE tests were both conducted in PBS (50 mM) saturated with oxygen at room temperature. For further insight into the selectivity of ORR pathway for Fe$_3$O$_4$@N-mC, the RRDE technique was used to monitor the electron transfer number by detecting the disk and
ring current of Fe₃O₄@N-mC at a constant rotating speed of 1600 rpm (Fig. 4d). The measured H₂O₂ yield of Fe₃O₄@N-mC varied strongly (3.4-16.8%) under different measured potentials, which was lower than that of Pt/C at the potential range of -0.2 to -0.7 V (Fig. S5). Based on the ring and disk currents results, the calculated electron transfer numbers for Fe₃O₄@N-mC (3.67-3.81) were consistent with the values calculated by K-L equation, which were higher than that of Pt/C cathode (3.62-3.72) (Fig. 4e). These results indicated that four-electron pathway is dominant in the ORR procedure of Fe₃O₄@N-mC. The superior ORR might be strongly associated with its mesoporous structures, and uniformly dispersion of nitrogen and metal coordinated metal in graphite carbon matrix. Moreover, as revealed by the RRDE experiments, Fe₃O₄@N-mC shows superior ORR catalytic activity to that of Pt/C, implying it is a promising alternative non-precious metal catalyst for ORR.

The durability of the obtained Fe₃O₄@N-mC catalyst was compared to commercial Pt/C catalyst via chronoamperometric tests, which was performed in 50mM PBS saturated with oxygen and applied with the constant potential of -0.4 V (Fig. 4f). After 8000 s testing, the current−time (i−t) chronoamperometric response showed that Fe₃O₄@N-mC revealed a slow attenuation with relative higher current retention (70 %) than Pt/C (58 %), which demonstrated higher long-term stability of Fe₃O₄@N-mC in neutral electrolyte. Numerous active sites dispersed in the mesoporous carbon matrix of Fe₃O₄@N-mC may prevent the iron and iron related active sites migration and aggregation, and thus, might be contribute to the electrochemical durability. Additionally, even if the outer active sites of Fe₃O₄@N-mC catalysts were destroyed
over time in the durability tests, the inner active sites can also expose and access to the electrochemical reaction interface through mesoporous channels to support the ORR durability.

![Diagram](image.png)

**Fig. 5** Voltage production of MFCs with the cathode of Fe₃O₄@N-mC or Pt/C at external resistance of 1000 Ω.

**Performance of MFC supplied with Fe₃O₄@N-mC cathode**

In order to test the feasibility of Fe₃O₄@N-mC as the cathode catalyst of MFC, air-cathode single-chamber MFCs were set up. The performance of MFCs with Fe₃O₄@N-mC as the cathode catalyst (Fe₃O₄@N-mC-MFC) was compared with MFCs with Pt/C as the cathode catalyst (Pt/C-MFC). The LSV scan test for cathodes showed that Fe₃O₄@N-mC cathodes had higher current density (10.94 A m⁻²) than that of Pt/C cathode (10.07 A m⁻²), suggesting the distinguished ORR catalytic activity of Fe₃O₄@N-mC. The results were in good agreement with that obtained by RDE (Fig. S6).

The reactors were operated for 21 cycles in 72 days. The output voltages of Fe₃O₄@N-
mC-MFC were similar with that of Pt/C-MFC in the first 6 cycles possessing the same highest output voltage of 560 mV, however, it turned to be higher than that of Pt/C-MFC at the later cycles (Fig. 5). Significantly, the voltage value decreased slowly to 496 mV for Fe₃O₄@N-mC-MFC, whereas it was decreasing to 420 mV for Pt/C-MFC. Based on the above, we can draw the conclusion that MFCs using Fe₃O₄@N-mC as cathodic catalysts performed better and more stable than Pt/C-based MFC.

**Fig. 6** Power, voltage, and the corresponding anode and cathode potentials of MFCs with the cathode catalysts of Fe₃O₄@N-mC and Pt/C at the 5th cycle, 14th cycle, and 21th cycle.

The power output and polarization curves were both performed after operating for 5 cycles, 14 cycles, and 21 cycles (Fig. 6). The power density of MFCs with Fe₃O₄@N-mC were 989 mW m⁻² after 5 cycle operation, which were slightly lower than that with
Pt/C cathodes (1187 mW m\(^{-2}\)). More importantly, after 14 cycles operation, the power density of Fe\(_3\)O\(_4\)@N-mC-MFC increased by 15.4 % to 1141 mV m\(^{-2}\), which was higher than that detected in Pt/C-MFC (1022 mV m\(^{-2}\)). Furthermore, after 21 cycles operation, the power density of all MFCs showed downtrend with 1095 mW m\(^{-2}\) for Fe\(_3\)O\(_4\)@N-mC-MFC and 940 mW m\(^{-2}\) for Pt/C-MFC. According to polarization curves, all MFCs showed low potential at high current density region in initial operation stage, implying their high diffusion resistance. In the case of Fe\(_3\)O\(_4\)@N-mC-MFCs, the potential at high current density region increased in fourteenth cycle, which led to the increase in power density. Thereafter, it had hardly any change at all after 21 cycle operation compared to Pt/C-MFC. It is obvious that Pt/C-MFC had much lower potential at medium current density region after long time operation, which illustrated that the activation resistance of Pt/C-MFC increased with the running time. The increase of activation resistance might cause by the biofouling on catalysts surface and gradually decreased ORR activity, which also could lead to the lower power density of Pt/C-MFC. Furthermore, the anode and cathode potentials versus current density also showed the unanimous anode potential and diverse cathode potential. The cathode potential decreased followed the same trend of polarization curves, which demonstrated that the differences in power generation were ascribed to the different ORR properties of the cathodes. Moreover, the coulombic efficiency was also detected to analyze water-processing ability of MFCs. With the similar COD removal rate, the Fe\(_3\)O\(_4\)@N-mC-MFCs reached to higher coulombic efficiency (26 %) compared to that of Pt/C-MFCs (21.7 %) (Fig. S7).
These results demonstrate that Fe₃O₄@N-mC can be a favorable alternative for commercial Pt/C catalyst in MFC. The power density obtained in Fe₃O₄@N-mC- MFCs (1141 mWm⁻² or 36.7 W/m³) was also higher than that of works previously reported (Table S1). For instance, the power density was 730 mW m⁻² for Fe₃O₄@N-mC with non-nanometer Fe₃O₄ core, which we reported previously,¹⁵ 12.54 W/m³ for polyaniline and iron-based catalysts (PANI-Fe₉₀₀),³⁴ 399 mWm⁻² for Co-N-doped carbon (Co-N-C),³⁵ and 745 mWm⁻² for N/Fe co-doped carbon (N/Fe–C).³⁶ The outstanding ORR activity and better durability of Fe₃O₄@N-mC as MFC cathode was inevitably ascribed to the its well-defined structure features: (1) the ordered mesoporous structure results in more accessible active sites to promote proton and oxygen transportation and adsorption, suppress the agglomeration of the Fe₃O₄ nanoparticles, and thus, speed up the mass exchange of ORR;¹⁴ (2) numerous nitrogen functional groups distributed in carbon matrix have considered to facilitate oxygen reduction, and on the basis of that the coordination of N and Fe can offer more active sites favorable to ORR;⁹,¹⁶ (3) the introduction of Fe₃O₄ catalyze the carbonization of PANI to provide more order graphite carbon, which with high conductivity to enhance the electron transfer;¹⁷, ³⁷ (4) the core-shell structure of Fe₃O₄@N-mC can prevent acid-leaching, oxidation and aggregation of Fe₃O₄ to achieve the high durability of catalysts.³⁸

In conclusion, a mesoporous Fe₃O₄@N-mC catalyst was developed by employing SDS and P123 soft template agents. The Fe₃O₄ @N-mC can considerably lessen the expense, while exhibited higher ORR activity (limiting current density, 3.54 mA cm⁻²), higher selectivity (electron transfer number, 3.66-3.94), and better long-term electrochemical...
stability in neutral solutions in comparison with the commercial Pt/C catalysts. The superior activity and durability toward the ORR of Fe$_3$O$_4$@N-mC was attributed to its well-defined mesoporous core-shell structure and numerous nitrogen functional groups dispersed in the carbon matrix. When it was applied in MFCs as cathodic catalysts, the Fe$_3$O$_4$@N-mC-MFC exceeded the Pt/C-MFC in durability performance, power production, and coulombic efficiency. Results demonstrate that the nonprecious metal catalysts had great potential efficient catalyst to replace commercial precious catalysts such as Pt.

**ASSOCIATED CONTENT**

**Supporting Information**

The supporting information is available free of charge on ACS publication website at [http://pubs.acs.org](http://pubs.acs.org).

Characterization methods; Comparison of power density between MFCs with different nonprecious metal catalysts (NPMCs); Nitrogen adsorption–desorption isotherms and Raman spectras of Fe$_3$O$_4$@N-mC catalyst; RDE polarization curves and Koutecky–Levich plots of Pt/C; Peroxide species detected on ring electrode of Fe$_3$O$_4$@N-mC and Pt/C; The LSV scan, COD removal rate, and Coulombic efficiency of MFC cathodes with Fe$_3$O$_4$@N-mC and Pt/C catalysts;

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Declarations of interest

None

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N-doped carbon electrocatalysts for high-performance zinc-air batteries and MFCs.


Synopsis: This work focuses on exploring the cost-effective cathode catalyst for microbial fuel cells, which is an eco-friendly biotechnology for recovering energy from biodegradable wastes.
Fig. 1 SEM TEM

236x147mm (150 x 150 DPI)
Fig. 2 XRD

88x87mm (300 x 300 DPI)
Fig. 3 XPS

74x78mm (300 x 300 DPI)
Fig. 4 ORR

61x87mm (300 x 300 DPI)
Fig. 5 Output voltage

99x95mm (300 x 300 DPI)
Fig. 6 Power density

146x123mm (300 x 300 DPI)