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A highly durable zinc-air battery from a directly integrated $\text{Fe}_x\text{NC@NiFe(OH)}_x$ bifunctional catalyst†

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Rechargeable zinc-air batteries (ZABs) hold great promise for energy storage and conversion due to their high theoretical energy density, cost-effectiveness, and inherent safety. However, progress is constrained by sluggish oxygen electrocatalysis and instability at the air cathode. To address such issues, we resort to a directly integrated pseudo-3D composite electrocatalyst based on carbon cloth, on which Fe/Fe₃C- and N- co-doped carbon nanotubes are directly induced and then used to further intercalate NiFe hydroxide clusters, Fe_xNC@NiFe(OH)_x. This hierarchical electrocatalyst shows enhanced oxygen electrocatalysis (ΔE is 636 mV), rendering high efficiency and durability of ZABs. Such improvement can be attributed to the rationally integrated pseudo-3D structure with high conductivity, high density of active sites, interconnected porosity, and well-bonded components for accelerating electron transfer and ion diffusion while ensuring structural integrity. Moreover, the hierarchical structure increases the electrochemical surface area with superior surface hydrophilicity. As a result, the composite electrocatalyst shows great potential as a binder-free air electrode, as demonstrated in a rechargeable ZAB of a high power density of 85.1 mW cm⁻² and a long period of operation beyond 2000 cycles (350 h) without notable degradation, outperforming noble metal electrodes.

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Introduction

Expedient exploitation of clean energy storage and conversion technologies is imperative to mitigate climate change, enhance energy efficiency, and integrate renewable energy resources. Rechargeable zinc-air batteries (ZABs) hold great promise due to their high theoretical energy density, cost-effectiveness, and inherent safety.^{1–7} Nevertheless, scale-up applications are still hindered by unsatisfactory performance, particularly due to the unstable and sluggish oxygen electrocatalysis at the air cathode.^{8–12} The oxygen evolution and reduction reactions (OER/ORR) proceed by a multi-step proton-coupled electron transfer process, normally necessitating noble metals to expedite the reaction kinetics, such as the benchmarking Pt-based (Pt/C) for ORR and the Ir/Ru-based for OER.^{13–16} However, the exiguous nature and high cost of noble metals also challenge the wide applications of ZABs. Therefore, extensive efforts have been devoted to synthesizing noble metal-free electrodes for highly efficient oxygen electrocatalysis.^{17–25} Cost-

effective bifunctional catalytic electrocatalysts that can accelerate both the ORR and OER have sprung up for high-performance ZABs.^{26–31}

Transition metal (TM) compounds embedded in nitrogen (N)-doped carbon electrocatalysts have been screened for efficient oxygen reduction.^{32–36} Previous experiments and simulations of such composites have demonstrated that the enhancement of Fe/Fe₃C nanoparticles and Fe–N_x coordinated moieties for ORR arises from the interplay between the metal sites and the N-doped carbon lattices, where the well-tuned electronic structure leads to optimized adsorption affinity towards successively generated intermediates.^{37–41} Although Fe/Fe₃C@NC is effective in catalysing ORR, their catalytic performance for OER cannot compete with the state-of-the-art OER catalytic electrodes, nickel-iron hydroxide (NiFe(OH)_x).^{42,43} NiFe(OH)_x is characterized by a layered structure with alternating metal cations and intercalated solvent molecules. Despite high intrinsic activity for OER, it suffers from poor electrical conductivity and insufficiently exposed active sites.^{44–46}

From the above, it is clear that it is challenging for a single component electrode to satisfy the need for oxygen electrocatalysis because of the mechanistic incompatibility of facilitating OER and ORR into reversible reactions.^{47–50} For instance, in an alkaline solution, despite sharing the same reaction intermediates (O*, OH*, OOH*), the additional initial step of the adsorp-

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tion of OH⁻ ions for OER or the activation of O₂ molecules for ORR is crucial for accelerating the whole catalytic process, which challenges the single component to realize bifunctional specificity.^{51–54} In comparison, two-component heterojunction catalysts can integrate the merits of both and realize high catalytic activities for both OER and ORR.⁵⁵ For example, atomic Co–N–C and Co₃O₄ nanoparticles were selected as ORR and OER electrocatalysts, respectively, yielding a robust bifunctional electrode for rechargeable ZABs,⁵⁶ with a relatively long lifespan of 2250 cycles at 5 mA cm⁻² and 365 cycles at 10 mA cm⁻². Gao *et al.* proposed a novel composite electrocatalyst constructed by bimetallic phosphide CoFeP nanoparticles embedded in N,P co-doped carbon matrix, where the former provides ample OER active sites and the latter promises high ORR activity.⁵⁷ The corresponding ZAB attains a large power density of 143.5 mW cm⁻² and long operation life of 1200 cycles (200 h). Recently, Lin and co-workers innovatively integrated wrinkled MoS₂ nanospheres with a layer of single Fe–N–C shell as OER and ORR active components, respectively.⁵⁸ The Fe–N–C shell protects the MoS₂ core from oxidation, enhancing the cycling stability at a power density of 78 mW cm⁻² for the corresponding wearable ZABs. However, most studies focus more on synthesizing oxygen catalysts, ignoring the drawback of these planar electrodes where additional binder buries active sites, and the electron transfer/ion diffusion is blocked. Moreover, the porous electrodes are crucial components of high-performance electrochemical batteries, which enlarge the electrodes/electrolyte interface to three-dimension. While the rate depends on both the specific surface area of oxygen catalysts and the efficacy of electron transfer and ion diffusion. Additionally, oxygen electrocatalysis occurs on the air–liquid–solid triple phase junction, requiring high mobility of the reactive species, namely, O₂ in air, ions in liquid (electrolyte) and electrons in solid (electrode), readily accessible active sites, and sustained structural durability.^{59–61} There is a lack of consideration of such heuristic design principles for the fabrication of bifunctional composite electrodes regarding the selection of active components and electrode structures.^{62–65}

Here, a long-lifespan rechargeable zinc-air battery is achieved by constructing a pseudo-3D composite electrocatalyst that combines “directly-grown-on-support” Fe/Fe₃C- and N-co-doped carbon nanotubes with NiFe(OH)_x (Fe_xNC@NiFe(OH)_x). In this way, OER active sites provided by NiFe(OH)_x clusters can be connected to the electron transport pathway due to the high conductive Fe_xNC network. Correspondingly, the hydrophilicity of NiFe(OH)_x also promises intimate contact between Fe_xNC with oxygen reactant. Such a unique configuration promises effective electron/ion transfer, abundant active sites, interfacial hydrophilicity for oxygen diffusion, and structural integrity. Consequently, the composite electrode shows remarkable ORR and OER performance with a voltage gap of 636 mV, far better than their respective individual components. The ZAB with the binder-free air composite cathode (Fe_xNC@NiFe(OH)_x) shows higher power density and longer operation life than the noble electrode (Pt/C//RuO₂).

Experimental section

Chemicals

Potassium ferricyanide, ferric chloride, dicyandiamide, concentrated HNO₃, concentrated HCl, iron(II) sulfate heptahydrate, nickel(II) nitrate hexahydrate, Pt/C (20%), and RuO₂ were all used without further purification.

Material synthesis

Fabrication of Fe_xNC. Typically, commercial carbon cloth was treated in concentrated HNO₃ under ultrasonication for 1 h and rinsed with deionized water. 0.496 g K₃Fe(CN)₆ was dissolved in 15 ml deionized water and then poured into FeCl₃ solution (25 ml, 0.41 g), followed by stirring at room temperature for 5 min to form a homogeneous solution. Then, a piece of pre-treated carbon cloth (2 cm × 3 cm) was added to the solution and heated at 60 °C for 6 h. After cooling to room temperature, the carbon cloth was taken out and washed thoroughly with deionized water and dried in the vacuum oven at 80 °C overnight.⁶⁶ The above carbon cloth and 1.0 g dicyandiamide were put into the same porcelain boat with a cover. Then, the sample was calcined at 400 °C for 2 h in N₂ atmosphere and then annealed at 850 °C for another 2 h. The heating rate was kept at 3 °C min⁻¹. To remove unstable iron species, the annealed electrode was immersed in 2 M HCl for 12 h at 65 °C. Afterward, the electrode was washed with water and dried at 80 °C for 6 h. The loading amount of active material on carbon cloth was ~0.8 mg cm⁻².

Fabrication of Fe_xNC@NiFe(OH)_x. Electrodeposition of nickel–iron hydroxide was carried out in a standard three-electrode electrochemical cell, employing the prepared Fe_xNC as the working electrode, carbon rod as the counter electrode, and saturated calomel electrode (SCE) as the reference electrode.⁵³ The aqueous electrochemical bath consists of 0.436 g of Ni(NO₃)₂·6H₂O and 0.417 g of FeSO₄·7H₂O. After maintaining a fixed electrodeposition current density of 20 mA cm⁻² for 5 min, the Fe_xNC@NiFe(OH)_x electrode was obtained with a loading amount of ~2.8 mg cm⁻² NiFe(OH)_x. Different deposition times of 0.3 min and 10 min were also investigated to optimize the performance.

Fabrication of Pt and RuO₂ on carbon cloth. A sample of 4 mg Pt/C (20%) was dispersed in a solvent mixture with 500 μl ethanol, 470 μl water, and 30 μl Nafion (0.5 wt%), followed by mixing with ultrasonication for 3 h to obtain a homogeneous ink. 200 μl of the catalyst ink was pipetted onto a piece of carbon cloth as the electrode (1 cm × 1 cm). The loading mass of 20% Pt/C was ~0.8 mg cm⁻². Commercial RuO₂ was also used for comparison of OER, where 10 mg RuO₂ was dispersed in the above solution and sonicated for 3 h. The loading mass of RuO₂ suspensions on carbon cloth was ~2.8 mg cm⁻².

Material characterization

The morphology and microstructure of the samples were characterized by scanning electron microscopy (SEM MAIA3) and transmission electron microscopy (JEM-2010, JEOL).

Powder X-ray diffraction (Bruker D8, PXRD) was used to identify the phase structures of the electrode components. The surface chemistry of the electrode was analyzed by X-ray photoelectron spectroscopy (XPS, Kratos Axis Ultra). The contact angle measurement instrument was used to examine the hydrophilicity of the electrode with the electrolyte (Dataphysics, OCA 20). Nitrogen adsorption-desorption was performed on the adsorption apparatus (Micromeritics, ASAP2460). As-prepared electrodes were cut into small fragments using scissors for tests. The surface area and pore size distribution were fitted by the Brunauer-Emmett-Teller (BET) method and the Barrett-Joyner-Halenda (BJH) method, respectively.

Electrochemical test

Electrochemical tests were carried out on an electrochemical station (CHI 760E) using a standard three-electrode system. A saturated calomel electrode (SCE) and carbon rod worked as the reference electrode and counter electrode, respectively. The electrolyte was 0.1 M KOH for OER and ORR. Catalysts supported on carbon cloth were directly used as the working electrode. All potentials in this work were calibrated to the reversible hydrogen electrode (RHE) by the following equation:

$$E_{\text{RHE}} = E_{\text{SCE}} + 0.241 + 0.059 \text{ pH}$$

OER and ORR performances were accessed *via* linear sweep voltammetry (LSV) with 95% *iR* compensation. OER was evaluated with a scan rate of 2 mV s⁻¹ from 0.2 to 1.0 V *vs.* SCE, and ORR was measured at a scan rate of 10 mV s⁻¹ from 0 to -0.6 V *vs.* SCE. The electrochemical active surface area (ECSA) of the electrodes was estimated by comparing double-layer capacitances (*C_{dl}*), which were obtained by scanning cyclic voltammetry (CV) at different rates of 10, 30, 50, 70, and 90 mV s⁻¹ in the non-faradaic potential region (0–0.1 V *vs.* SCE). From the linear fitting of the line of the current density difference at 0.050 V *vs.* SCE against the above scan rates, the slope is *C_{dl}*. ZAB test was performed under an atmospheric environment using a homemade cell. A polished zinc plate (500 μm in thickness) was used as a cathode. The aqueous solution containing 6 M KOH and 0.2 M zinc acetate was used as the electrolyte. Fe_xNC@NiFe(OH)_x supported on carbon cloth and a gas diffusion layer were combined as the air cathode. The effective area of the air cathode was 1 × 1 cm². Galvanostatic discharge-charge cycling tests were evaluated using a LAND CT3001A battery test system at 10 mA cm⁻² with 5 min discharge and 5 min charge per cycle. For comparison, 20% Pt (0.8 mg cm⁻²) and RuO₂ (2.8 mg cm⁻²) were also loaded on the carbon cloth as the air cathode.

Results and discussion

As depicted in Fig. 1a, the rational fabrication strategy of the bifunctional Fe_xNC@NiFe(OH)_x electrode involves three steps: (1) the immobilization of FeFe(CN)₆ nano-seeds on carbon cloth by a wet-chemistry method; (2) carbonization facilitating the growth of N-doped carbon nanotubes (CNTs) catalyzed by

in situ evolved Fe/Fe₃C; and (3) electrodeposition of NiFe(OH)_x clusters over the CNTs. Details of the synthesis process are provided in the experimental section. For the first step, as shown in the scanning electron microscopy image (SEM, Fig. S1a and b†), uniform FeFe(CN)₆ nano-seeds are tightly anchored on the carbon cloth (CC) by facile solution precipitation. Subsequently, under high-temperature pyrolysis at 850 °C, the FeFe(CN)₆ precursors are decomposed, forming entangled CNTs on the CC (Fig. 1b and c). Transmission electron microscopy (TEM) images show that dark particles are enveloped in the middle or tips of the CNTs (Fig. 1d). The high-resolution TEM displays well-defined lattice fringes, which are consistent with the (211) and (110) crystalline planes of Fe/Fe₃C (Fig. S2a†). The average size of metal nanoparticles is around 30–40 nm (Fig. S2b†). The X-ray diffraction (XRD) patterns confirm the co-existence of Fe and Fe₃C (green curve), where the diffraction peaks are attributed to crystalline planes (102) of Fe₃C at 43.7° (JCPDS No. 35–0772) and (110) and (211) planes at 44.7° and 65.2° of metallic Fe (JCPDS No. 06–0696), respectively (Fig. 1e).

Raman spectra further probe the crystallinity of the carbons. The D band (1357 cm⁻¹) signal reflects the vibrational environment of defective/disordered carbon, and the G band (1560 cm⁻¹) shows the graphitic/crystalline carbon structure.^{67,68} As illustrated in Fig. S2c,† the intensity ratio of D to G of Fe_xNC is 0.58, indicating a high degree of crystallinity of the carbon tubes. The highly graphitized carbon endows 3D conductive and porous networks for electron transfer and ion transport. Energy-dispersive X-ray spectroscopy (EDS) elemental mapping images reveal the uniform distribution of Fe, N, and C in the nanotube matrix (Fig. S2d†), and their respective proportions have also been measured (Table S1†). From the elemental mapping images, apart from the signal of relatively large aggregates of Fe/Fe₃C, Fe element is also enriched on the carbon wall.

The facile electrodeposition technique is used to immobilize NiFe(OH)_x onto the as-prepared nanotubes. As revealed by the SEM images in Fig. 1f and g, platelet-like NiFe(OH)_x clusters are uniformly decorated on the surface of CNTs after rapid electrodeposition for only 5 min, denoted as Fe_xNC@NiFe(OH)_x. The hierarchical nanotubes provide a high surface area for the growth of the nanosheets and avert undesirable aggregation, as compared with electrodeposition of NiFe(OH)_x on pristine CC surface (Fig. S3a and b†). The schematic diagram shown in Fig. 1h manifests the merits of the integrated 3D electrode for oxygen electrocatalysis with high electron transfer, ion transport, and dual active components. XRD patterns further verify the amorphous nature of the deposited NiFe(OH)_x (Fig. 1e and Fig. S3c, d†). The amorphous NiFe(OH)_x exhibits better catalytic activity than the crystalline counterpart due to more unsaturated active sites.^{69–71} Fig. 1i shows the TEM and EDS mapping images of Fe_xNC@NiFe(OH)_x, revealing that nanoplates are tightly grown on the nanotubes with uniform dispersions of Ni, Fe, and O elements.

The electrodeposition time largely determines the amount of NiFe(OH)_x, which may, in turn, affect the catalytic activity

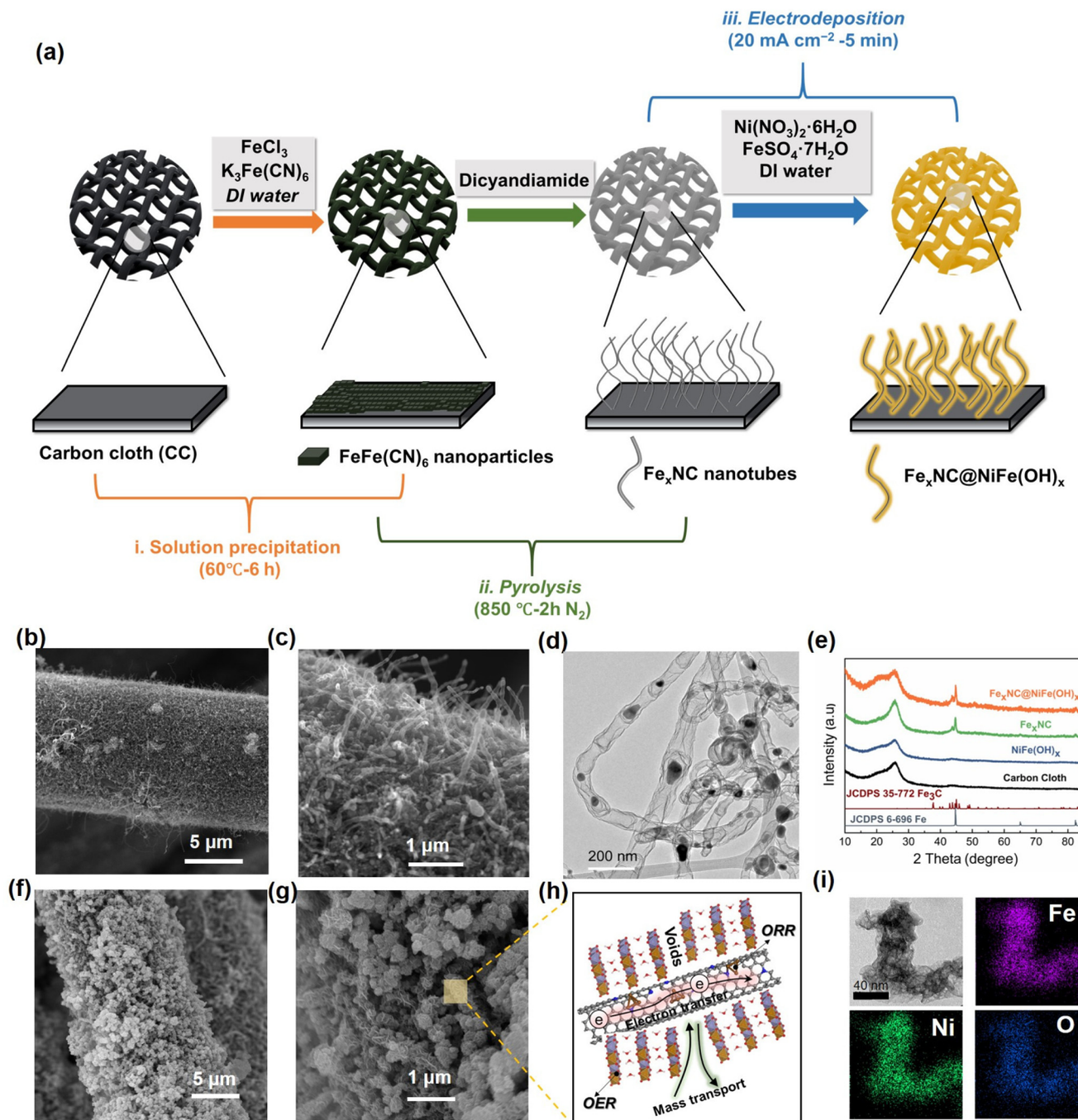


Fig. 1 (a) Schematic illustration of the synthesis route for $\text{Fe}_x\text{NC}@NiFe(OH)_x$. (b–d) SEM and TEM images of Fe_xNC . (e) XRD patterns of as-prepared electrodes. (f and g) SEM images of $\text{Fe}_x\text{NC}@NiFe(OH)_x$ and (h) the corresponding schematic diagram. (i) TEM image of $\text{Fe}_x\text{NC}@NiFe(OH)_x$ and EDS elemental mapping images.

and mass transfer properties. For comparison, shorter (0.3 min) and longer (10 min) periods of deposition time were also accessed, denoted as $\text{Fe}_x\text{NC}@NiFe(OH)_{x-0.3}$ and $\text{Fe}_x\text{NC}@NiFe(OH)_{x-10}$, respectively. Their morphologies are shown in Fig. S4,[†] revealing that the shorter time leads to loosely deposited layers on the nanotubes, whereas the longer deposition time causes excessive aggregation of deposits over the electrode surface.

X-ray photoelectron spectroscopy (XPS) is conducted to determine the surface composition and valence states. In the survey spectrum, $\text{Fe}_x\text{NC}@NiFe(OH)_x$ shows a similar pattern to pristine $NiFe(OH)_x$ due to the detection limit (~ 10 nm) of the surface detection technique (Fig. S5[†]). For Fe_xNC , the high-resolution N 1s spectrum shows five peaks: oxidized-N (402.8 eV), graphitic-N (401.3 eV), pyrrolic-N (400.3 eV), $\text{Fe}_x\text{-N}$ (399.4 eV), and pyridinic-N (398.4 eV) (Fig. 2a). The proportion of

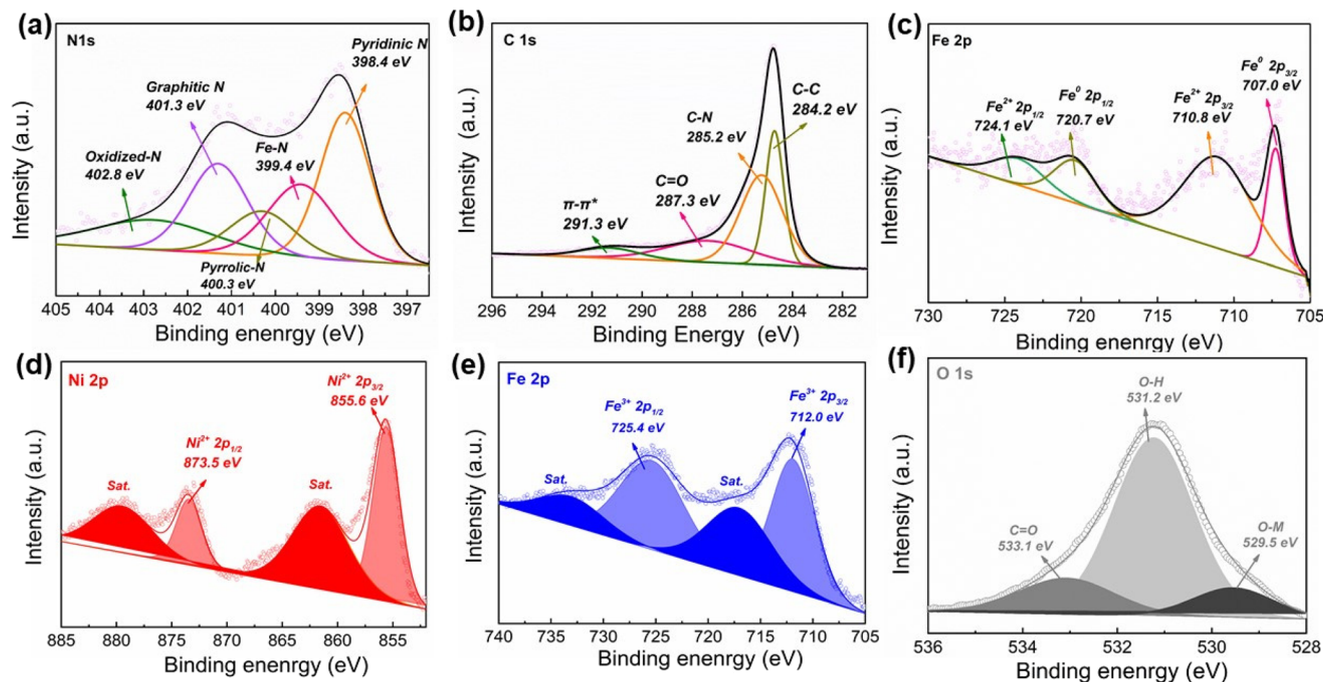


Fig. 2 High-resolution XPS spectra of Fe_xNC : (a) N 1s, (b) C 1s and (c) Fe 2p; and $\text{Fe}_x\text{NC}@NiFe(OH)_x$: (d) Ni 2p, (e) Fe 2p and (f) O 1s.

different nitrogen dopants is summarized in Table S2.† The rich nitrogen dopants in Fe_xNC have several benefits. Firstly, the pyridinic-N helps to create Lewis basic sites on the carbon surface and edge, which are attractive to O_2 molecules.⁷² Secondly, the high concentration of the Fe-N_x moiety also contributes to highly efficient sorption and coordination of the intermediate oxygen species during electrocatalysis.³⁹ Finally, the graphitic-N can donate their electrons to the π -conjugated sp^2 carbon and thus enhance electron mobility.^{73,74} The C 2p spectrum of Fe_xNC shows four peaks located at 291.3, 287.3, 285.2, and 284.2 eV, which are attributed to π - π^* , C=O, C-N and C-C bonds, respectively (Fig. 2b). The result further confirms the formation of N doped carbon moiety on the electrode. High-resolution Fe 2p spectrum is also deconvoluted into zero-valence Fe (707.0 and 720.7 eV) and Fe^{2+} (710.8 and 724.1 eV) (Fig. 2c). Fig. 2d and e demonstrate the high-resolution Ni 2p and Fe 2p spectra of $\text{Fe}_x\text{NC}@NiFe(OH)_x$, and their signals actually reflect the surface chemistry of top layer NiFe(OH)_x (Fig. S6†). In the Fe 2p spectra, the peaks located at 712.0 and 725.4 eV correspond to Fe 2p_{3/2} and Fe 2p_{1/2}, respectively, suggesting the dominant oxidation state of the Fe element is trivalent. The deconvoluted peak at 855.6 eV is assigned to Ni 2p_{3/2}, while the peak at 873.5 eV is consistent with Ni 2p_{1/2}. These two peaks are ascribed to Ni²⁺ in Ni(OH)₂. In addition, the O 1s spectrum manifests three characteristic peaks, located at 529.5 (metal-oxygen bond), 531.2 (hydroxyl in oxyhydroxide), and 533.1 eV (adventitious functional group, C=O) (Fig. 2f). Moreover, the stoichiometric composition of Ni to Fe is close to 1 (Table S1 and S3†).

Electrochemical tests are performed in 0.1 M KOH electrolyte using the conventional three-electrode assembly. As shown

in Fig. 3a and Table S4,† the $\text{Fe}_x\text{NC}@NiFe(OH)_x$ electrode demonstrates an overpotential of 264 mV at 10 mA cm⁻² for the OER, which is much lower than that of Fe_xNC (377 mV) and noble RuO₂ (330 mV). The half-wave potential ($E_{1/2}$) of ORR on $\text{Fe}_x\text{NC}@NiFe(OH)_x$ is as high as 0.858 V (vs. RHE), higher than that of the individual components. It is noted that neither individual Fe_xNC nor NiFe(OH)_x shows bifunctional properties towards OER and ORR. A performance indicator, potential difference ΔE ($E_{j=10} - E_{1/2}$), is calculated to further compare the overall oxygen electrocatalytic activity. As shown in Fig. S7,† the Tafel slope of the $\text{Fe}_x\text{NC}@NiFe(OH)_x$ electrode is 109 mV dec⁻¹, lower than that of individual Fe_xNC (127 mV dec⁻¹), NiFe(OH)_x (151 mV dec⁻¹), and noble Pt (116 mV dec⁻¹), indicating its fast kinetics for ORR. Further dissection of OER reaction kinetics by comparing the Tafel slopes is performed among the different electrodes (Fig. 3c). $\text{Fe}_x\text{NC}@NiFe(OH)_x$ possesses the smallest Tafel slope (60.9 mV Dec⁻¹) than that of NiFe(OH)_x (85.4 mV Dec⁻¹), Fe_xNC (84.6 mV Dec⁻¹), and RuO₂ (104.6 mV Dec⁻¹). For ORR, we realize that Fe_xNC functions as the actual active sites, but the composite electrode also shows a lower onset overpotential and half-wave potential than that of Fe_xNC , which arose our attention to analyze the role of outer NiFe(OH)_x clusters. In response, contact angle tests are carried out. As noted in Fig. 3d and Fig. S8,† the surface of the Fe_xNC electrode is hydrophobic with a large contact angle of 129°, which would inhibit the access of dissolved oxygen molecules for ORR. In comparison, significant improvement of hydrophilicity is observed on the composite electrode after the incorpor-

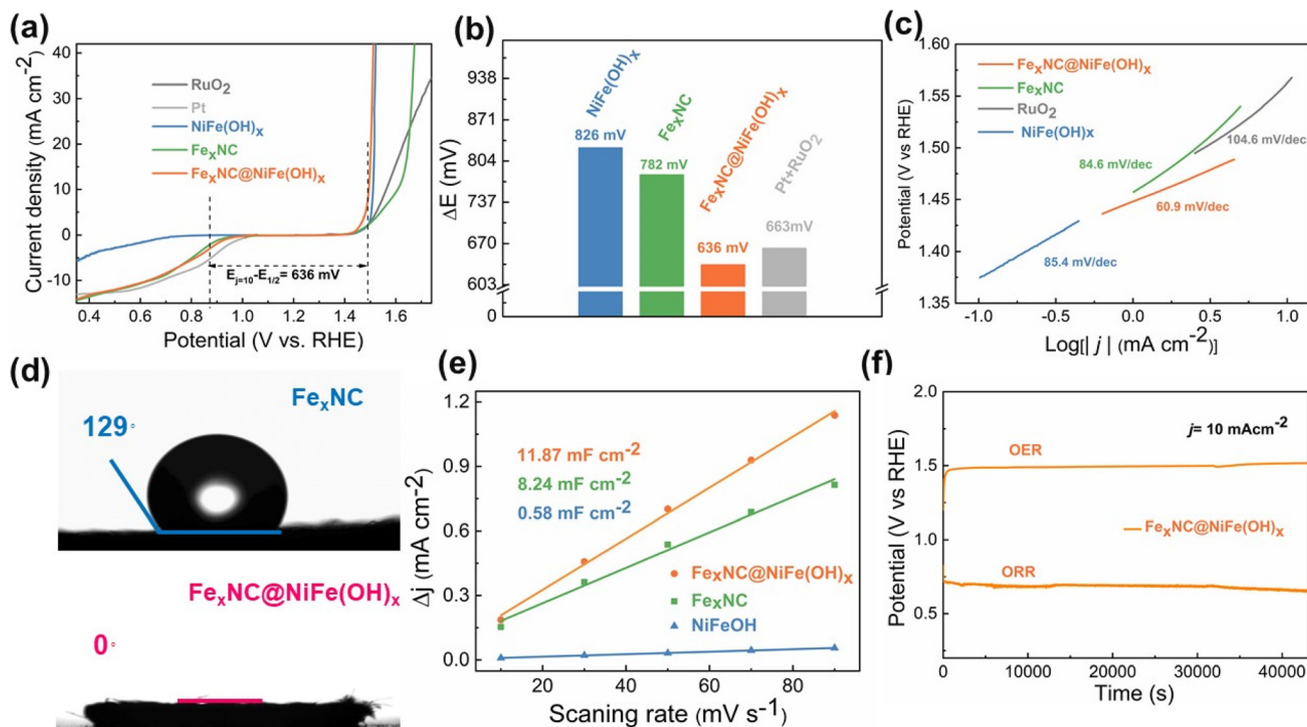


Fig. 3 (a) The polarization curves of different electrodes. (b) Potential difference (ΔE) of NiFe(OH)_x , Fe_xNC , $\text{Fe}_x\text{NC@NiFe(OH)}_x$, and $\text{Pt} + \text{RuO}_2$. (c) OER Tafel slopes. (d) Contact angle evaluation on Fe_xNC and $\text{Fe}_x\text{NC@NiFe(OH)}_x$. (e) ECSA results of different electrodes. (f) The voltage-time curves of $\text{Fe}_x\text{NC@NiFe(OH)}_x$ at 10 mA cm^{-2} .

ation of the hydrophilic NiFe(OH)_x , showing good wetting behaviour. Therefore, the improved wettability of the composite electrode favours the intimate contact between the ORR active sites and oxygen in the electrolyte. Furthermore, the hierarchical pseudo-3D structure constituted by the tubular carbon and NiFe(OH)_x platelets largely increases the electrochemically active surface area (ECSA) to 11.87 mF cm^{-2} , compared with that of the single component (8.24 mF cm^{-2} for Fe_xNC and 0.58 mF cm^{-2} for NiFe(OH)_x), shown in Fig. 3e and Fig. S9,[†] thus endowing more accessible active sites for the catalytic reactions. This point can be further confirmed by the BET assessment using nitrogen as the probe gas. As shown in Fig. S10 and Table S5,[†] among the samples investigated, the composite catalyst electrode possesses a relatively large surface area ($5.88 \text{ m}^2 \text{ g}^{-1}$) and pore volume ($0.041 \text{ cm}^3 \text{ g}^{-1}$), characterized by a range of interconnected porosities from micropores to relatively large mesopores. The results further indicate the intrinsically larger surface area and hierarchical porosities of the composite catalyst, exposing more active sites and facilitating mass transport to enhance electrocatalysis. The long-term durability of the $\text{Fe}_x\text{NC@NiFe(OH)}_x$ electrode towards oxygen electrocatalysis is further evaluated. There is negligible decay for both OER and ORR, suggesting the remarkable stability of the catalyst (Fig. 3f), even superior to the noble metals (Fig. S11[†]). We also carried out a series of tests to unveil the changes in structure and composition after the electrochemical tests. XRD results indicate no notable phase change

after the reactions except for the emergence of a broad peak at 34.8° on the post-OER electrode, corresponding to $\text{Ni}_{1.43}\text{Fe}_{1.7}\text{O}_4$ (JCPDS: 80-072) (Fig. S12[†]).⁵³ TEM results also reveal a clear crystalline lattice on the edge of the post-OER electrode (Fig. S13a and b[†]), while the NiFe(OH)_x remains amorphous after long-term ORR (Fig. S13c and d[†]). Moreover, in the XPS spectra of the post-OER electrode (Fig. S14a–d[†]), the fitted metal–oxygen peak in O 1s becomes stronger than that of the initial electrode, implying the formation of more metal–oxygen bonds, supporting the emergence of the metal oxide.

There is a trade-off between the catalytic activity and mass transfer capability upon deposition of NiFe(OH)_x . As shown in Fig. S15a,[†] the electrode prepared from the shorter electrodeposition time ($\text{Fe}_x\text{NC@NiFe(OH)}_{x-0.3}$) leads to a large overpotential of 285 mV towards OER at 10 mA cm^{-2} due to the insufficient OER catalytic active sites. While the electrodeposition time becomes long, the electrode ($\text{Fe}_x\text{NC@NiFe(OH)}_{x-10}$) shows deteriorated ORR performance because its surface is packed with a rather dense NiFe(OH)_x layer, impeding the access of electrolyte to the inner Fe_xNC (Fig. S15b[†]). Hence, a moderate deposition time of 5 min balances the manifold properties, namely, catalytic activity, electrical conductivity, and mass transport, to yield reasonable bifunctional characteristics (Fig. S15c and d[†]). Therefore, Fe_xNC is not densely entrapped inside NiFe(OH)_x in our optimized electrode. On the one hand, the 2D NiFe(OH)_x nanoplates are vertically grown on Fe_xNC

and thus leave large voids for the electrolyte penetration to active sites for ORR. On the other hand, as the reactants of ORR come from the dissolved oxygen in water, the incorporation of a hydrophilic NiFe(OH)_x layer favors the diffusion of the electrolyte into the active sites.

The excellent oxygen electrocatalytic activity of the composite electrode is further demonstrated in ZAB to showcase practical applications (Fig. S16†). The Fe_xNC@NiFe(OH)_x electrode can be directly used as the air cathode without the need for any binders, simplifying the component structure and ensuring hierarchical pseudo-3D structural integrity. Therefore, the polarization curve of the corresponding ZAB displays a higher discharging voltage and lower charging voltage than other cathode counterparts, implying better energy storage and energy output performance (Fig. 4a). The relatively small voltage gap with the composite electrode agrees well with the above-mentioned bifunctional ability for both OER and ORR (Fig. 4b).

As a result, the cell with the bifunctional electrode can achieve a power density of 85.1 mW cm⁻², which is higher

than that of Fe_xNC (77.9 mW cm⁻²), NiFe(OH)_x (70.1 mW cm⁻²) and is even comparable with that of the Pt/C//RuO₂ counterpart (84.9 mW cm⁻²) (Fig. S17†). The rate capability test further reveals that the ZAB with the composite electrode (ZAB_{CE}) show a higher and more stable discharge plateau at each current density than other counterparts (Fig. 4c). The assembled ZAB shows a stable open circuit voltage of 1.44 V over 50 000 s without severe self-discharge (Fig. 4d). Moreover, the specific capacity of 785.2 mA h g_{Zn}⁻¹ is achieved on ZAB_{CE}, which is larger than that of noble metal (721.6 mA h g_{Zn}⁻¹) (Fig. 4e). The battery performance is consistent with the above results, further confirming the merits of our rationally designed composite electrode for bifunctional oxygen electrocatalysis. Finally, the long-time cyclic performance of ZAB is accessed (Fig. 4g). It is evident that ZAB_{CE} shows a very stable lifespan over 350 h (2000 cycles) at the operating current density of 10 mA cm⁻², outperforming those with noble metals and single-component electrodes (Fig. S18–S20†), whose cells encounter significant polarization upon cycling. The perform-

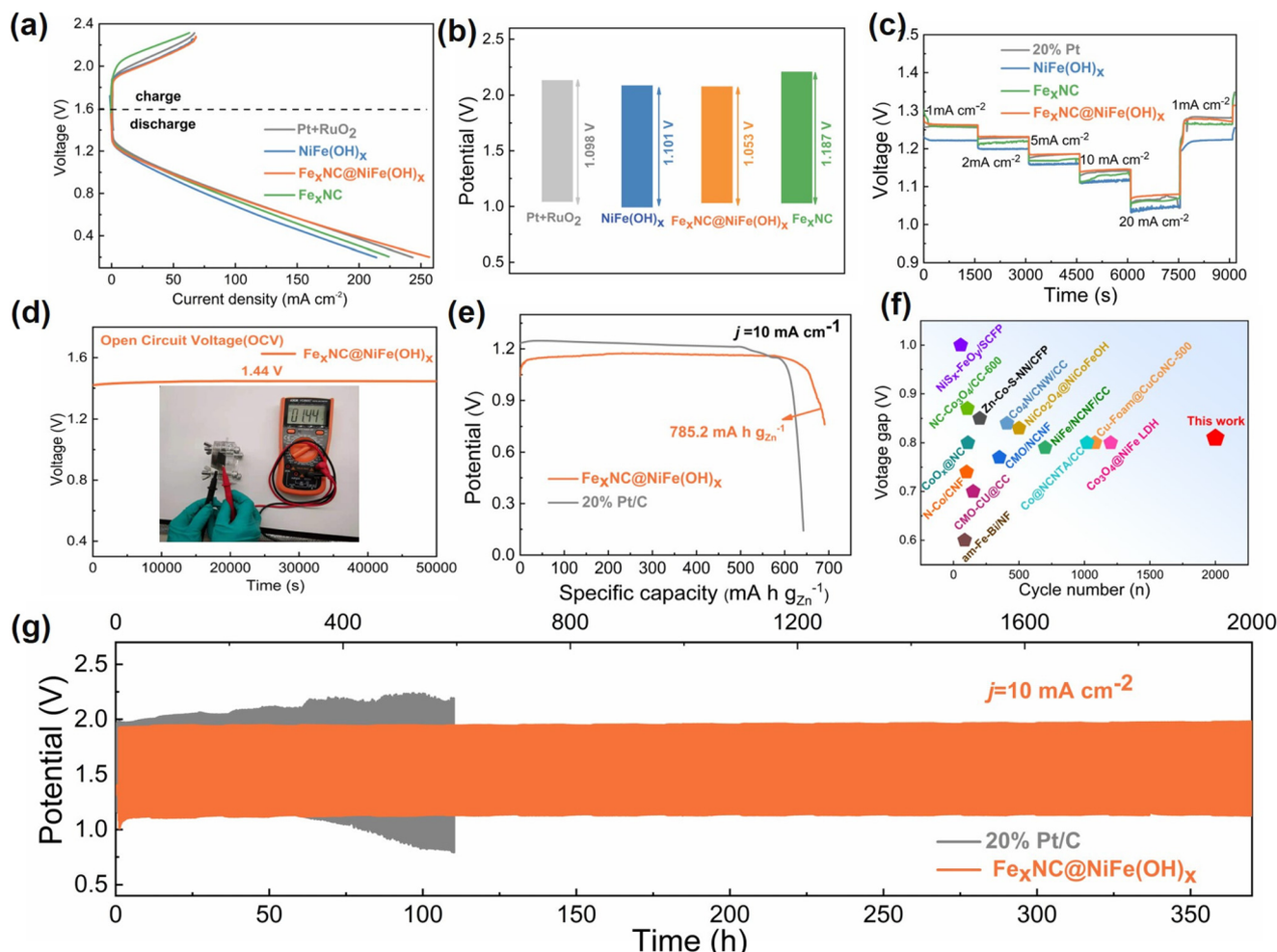


Fig. 4 (a) The charge/discharge polarization curves. (b) Voltage gaps at the current density of 40 mA cm⁻². (c) Galvanostatic discharge of Pt/C-ZAB, Fe_xNC-ZAB, NiFe(OH)_x-ZAB, and Fe_xNC@NiFe(OH)_x-ZAB under different current densities. (d) Open circuit potential, inset: optical image. (e) The specific capacity curves run at 10 mA cm⁻². (f) Performance comparison of ZABs using self-supported bifunctional electrodes operating at 10 mA cm⁻². (g) The discharge/charge cycle curves of Pt/C-ZAB and Fe_xNC@NiFe(OH)_x-ZAB at 10 mA cm⁻².

ance also favourably competes with other self-supported electrodes, as outlined in Fig. 4f and Tables S6, S7.† To summarize, our rationally designed hierarchical pseudo-3D composite electrode offers long durability for rechargeable ZABs.

Conclusions

A hierarchical pseudo-3D composite electrode supported on carbon cloth has been successfully fabricated with non-noble metal NiFe(OH)_x clusters grafted over Fe/Fe₃C- and N-codoped carbon nanotubes that are directly grown from carbon-cloth support. The composite electrode rationally integrates the active components for bifunctional properties, achieving an overpotential of 264 mV for OER at a current density of 10 mA cm⁻² and a half-wave potential of 0.858 V (vs. RHE) for ORR. From a series of characterizations, the enhanced performance is due to the following merits: (1) the inner conductive nitrogen-doped carbon nanotubes serve as the pathway for electron transfer, and the appropriate amount of outer NiFe(OH)_x clusters provide interconnected access for electrolyte penetration. (2) The outer layer NiFe(OH)_x tunes the hydrophilicity to cater to the electrode/electrolyte interface. (3) hierarchical pseudo-3D structure endows the composite electrode with an increased electrochemical surface for electrocatalysis. For practical applications, ZAB_{CE} shows a power density of 85.1 mW cm⁻² and a long cycle life of 350 h (2000 cycles), superior to those using noble metals. This work provides a low-cost alternative to noble metals for oxygen electrocatalysis and offers fresh insights underpinning the rational design of bifunctional oxygen electrodes.

Conflicts of interest

There are no conflicts to declare.

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