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Ming-Yu Ding

Wen-Jie Jiang

Tian-Qi Yu

Xiao-Yan Zhuo

Xiao-Jing Qin

Shi-Bin Yin Guangxi Key Laboratory of Electrochemical Energy Materials, School of Chemistry and Chemical Engineering, Guangxi University, Guangxi Nanning 530004, China, yinshibin@gxu.edu.cn

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ARTICLE

Electronically Modulated FeNi Composite by CeO_2 Porous Nanosheets for Water Splitting at Large Current Density

Ming-Yu Ding, Wen-Jie Jiang, Tian-Qi Yu, Xiao-Yan Zhuo, Xiao-Jing Qin, Shi-Bin Yin*

Guangxi Key Laboratory of Electrochemical Energy Materials, School of Chemistry and Chemical Engineering, Guangxi University, Guangxi Nanning, 530004, China

Abstract

Exploiting highly active and non-noble metal bifunctional catalysts at large current density is significant for the advancement of water electrolysis. In this work, CeO₂ electronically structure modulated FeNi bimetallic composite porous nanosheets in-situ grown on nickel foam ($NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF$) is synthesized. Electrochemical experiments show that the NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF exhibited the outstanding activities toward both oxygen and hydrogen evolution reactions (OER and HER) ($\eta_{1000} =$ 352 mV and $\eta_{1000} =$ 429 mV, respectively). When assembled into a two-electrode system for overall water splitting (OWS), it only needs a low cell voltage of 1.81 V to drive 100 mA \cdot cm⁻². And it can operate stably at ± 500 mA \cdot cm⁻² over 30 h toward OER, HER and OWS without significant activity changes. The reason could be assigned to the electronic modulating of $CeO₂$ on FeNi composite, which can boost the intrinsic activity and optimize the adsorption of reaction intermediates. Moreover, the porous nanosheets insitu grown on NF could enhance the contact of active site with electrolyte and facilitate the gas release, thus improving its chemical and mechanical stabilities. This study highlights a novel approach to design bifunctional non-noble metal catalysts for water splitting at large current density.

Keywords: FeNi composite; Hydrogen evolution reaction; Oxygen evolution reaction; Catalyst; Water splitting

1. Introduction

The development of hydrogen energy with zero pollution and high combustion calorific value will enable the transition from traditional energy to clean and sustainable hydrogen economy $[1-3]$ $[1-3]$ $[1-3]$ $[1-3]$ $[1-3]$. Water electrolysis driven by renewable energy (e.g., wind and solar) to produce H_2 is deemed a promising method. However, its practical application is hindered due to the slow reaction kinetics and high overpotential $[4,5]$ $[4,5]$ $[4,5]$ $[4,5]$. Although the above issues have been deeply studied, it is still difficult to meet the requirements of long-term and efficient operation at large current densities in industrial water electrolysis for hydrogen production [[6\]](#page-8-3). It is still challenging to design catalysts with abundant active sites and good durability/efficiency at large current density.

Among various catalysts, Fe-Ni-based composites have been considered as ideal catalysts for OER because of their high intrinsic activity and fast kinetics under alkaline conditions $[7-9]$ $[7-9]$ $[7-9]$. And there exists synergistic effect between Ni and Fe compared with single-component catalysts, which can effectively improve the intrinsic activity $[10-12]$ $[10-12]$ $[10-12]$. Meanwhile, Fe-Ni-based composites could form the FeNi double metal oxyhydroxide (NiFeOOH) during OER, thus $Fe³⁺$ occupies an octahedral position in NiFeOOH, and the Fe-O bond length is very short due to the influence of the surrounding edge-sharing octahedron (NiO_6) [\[13](#page-8-6)]. This structure can facilitate the adsorption of oxygen-containing intermediates during OER, therefore enhancing the reaction rate [[14\]](#page-8-7). Nevertheless, most Fe-Ni-based composite materials are only used for OER because of the weak bonding

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^{*} Corresponding author, Shi-Bin Yin, Tel: (86-771)3233718, E-mail address: yinshibin@gxu.edu.cn.

between hydrogen and Fe^{3+} [\[15](#page-8-8),[16](#page-8-9)]. Thereby designing advanced multifunctional catalysts is necessary.

According to previous studies, combination of $CeO₂$ with transition metals could enhance the intrinsic HER activity [[17](#page-8-10),[18\]](#page-8-11). $CeO₂$ as a water dissociation promoter exhibits excellent electronic/ ionic conductivity due to the abundant half-filled and empty d orbitals, thereby lowering the energy barrier for the formation of hydrogen intermediates [[19](#page-8-12),[20\]](#page-8-13). Chen et al. prepared a composite with $CeO₂$ modulating the electronic structure of Ni that is favorable for the transportations of electrons and ions to exhibit good HER activity [\[21\]](#page-8-14). Besides, $CeO₂$ is usually applied as OER catalysts because of its superior conductivity and oxygen storage ability accompanied by reversible exchange states between Ce^{3+} and Ce^{4+} [[22,](#page-8-15)[23\]](#page-8-16). Li and co-workers synthesized a FeOOH/CeO₂ hetero-layered nanotube, which exhibited better OER performance than single FeOOH component [\[24\]](#page-8-17).

The long-term stable operation of catalysts at large current density is another key factor for realizing industrial applications, since the vigorous reactions of anode and cathode will cause catalyst shedding and structural damage [[25](#page-8-18)[,26](#page-8-19)]. Morphology control and substrate selection are considered as efficient methods to enhance catalyst stability [\[27](#page-8-20),[28\]](#page-8-21). Among them, two-dimensional (2D) nanosheet in-situ grown on conductive substrate (NF) has the advantages of the stable combination between the active species and substrate, which is beneficial for accelerating electron and ion transfers [\[29](#page-9-0),[30\]](#page-9-1). As well as possessing large specific area to facilitate gas release, thus avoiding catalyst shedding and structural damage [\[31](#page-9-2)]. In our previous work [[32\]](#page-9-3), IrNi-FeNi₃ hybrid nanosheets are synthesized on NF via one-step solvothermal, which could maintain at 1000 mA \cdot cm⁻² for 100 h in industrial conditions.

Herein, $NiFe₂O₄-Fe₂₄N₁₀-CeO₂$ composite with porous nanosheets structure in-situ grown on NF ($NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF$) is designed and prepared by solvothermal and annealing treatments ([Fig. 1](#page-2-0)). It combines the advantages of $CeO₂$ in regulating electronic structure and self-supporting porous nanosheets, showing the feasibility of highefficiency, stable and low-cost water electrolysis at large current density. The electrochemical characterization demonstrates that it arrived at ± 1000 mA·cm⁻² only requiring overpotentials of 352 and 429 mV for OER and HER, respectively. Moreover, when used as a two-electrode system for water electrolysis, it needs a low voltage of 1.81 V to reach 100 mA \cdot cm⁻², which surpasses those of NiFe₂O₄-Fe₂₄N₁₀/NF (1.83 V), CeO₂/NF

Fig. 1. Schematic diagram for the synthesis of NiFe₂O₄-Fe₂₄N₁₀-CeO₂/ NF.

(1.94 V) and precursor (1.89 V), and it could also operate stably at 500 mA \cdot cm⁻² for 30 h.

2. Experimental section

2.1. Materials

All reagents were of analytical grade and used directly without further purification, including ferric nitrate nonahydrate (Fe(NO₃)₃.9H₂O), nickel foam (NF), urea (H_2NCONH_2) , cerium nitrate hexahydrate $(Ce(NO₃)₃·6H₂O)$, potassium hydroxide (KOH), hydrochloric acid (HCl) and absolute ethanol (C_2H_5OH).

2.2. Syntheses of $NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF,$ $NiFe₂O₄-Fe₂₄N₁₀/NF$ and $CeO₂/NF$

NF (1.0 cm \times 6.0 cm) was sonicated with 1.0 mol \cdot L⁻¹ HCl solution to reduce the influence of oxide layer on the reaction, followed by repeated cleaning with absolute ethanol and DI water for several times, respectively. Fe($NO₃$)₃ $·9H₂O$ (252.75 mg) , $Ce(NO₃)₃·6H₂O$ (271.5 mg) and H2NCONH2 (363.75 mg) were added into 30 mL DI water, then the solution and pretreated NF were transferred into 50 mL Teflon-lined autoclave for 12 h heating treatment at 120 \degree C. Subsequently, the obtained precursors were annealed at 400 \degree C for 2 h under nitrogen (N_2) environment and then bubbled with ammonia (NH_3) for additional 2 h $NiFe₂O₄-Fe₂₄N₁₀/NF$ and $CeO₂/NF$ were also prepared by the similar procedure without Ce and Fe sources, respectively. To load the commercial Pt/C or 40 wt% IrO $_2$ /C catalyst on Ni foam, 5.0 mg of catalyst powders were dispersed in 1.0 mL $(20.0 \mu L)$ 5.0 wt% Nafion and 0.98 mL ethanol) solution. The suspension was sonicated for 30 min to prepare a homogeneous ink. The NF with a fixed area was drop-casted with $40 \mu L$ of the catalyst ink.

2.3. Material characterizations

The morphology of the above samples was investigated using scanning electron microscope (SEM, SU8220, Hitachi Corp.). Transmission electron microscopic (TEM) and elemental distribution observations were performed on Titan ETEM (TALOS 200X, FEI Corp.). X-ray diffraction (XRD, Rigaku Corp., with a copper K-edge X-ray) was utilized to determine the structure of the specimens. X-ray photoelectron spectroscopic (XPS) measurement was performed on an instrument (ESCALAB-250Xi, Thermo Fisher) with a monochromatic Al (300 W) K_{α} X-ray line source.

2.4. Electrochemical measurements

Electrochemical test was performed in a threeelectrode system (Interface 1010E, Gamry, USA) at 30 °C in N_2 saturated 1.0 mol·L⁻¹ KOH solution. Hg/HgO electrode and graphite rod were used as the reference and counter electrode, respectively. All potentials were calibrated against the reversible hydrogen electrode (RHE) according to Nernst equation ($E_{\text{RHE}} = E_{\text{Hg/HgO}} + 0.0591 \times pH + 0.098$). Cyclic Voltammetric (CV) activation at a scan rate of $100 \text{ mV} \cdot \text{s}^{-1}$ was performed for 10 cycles prior to conduct the linear sweep voltammetric (LSV) test at a scan rate of 5 mV \cdot s⁻¹ for characterizing the OER, HER and OWS activities of each sample. Unless otherwise specified, all the polarization curves were iR -corrected, where i is the experimentally measured current and R is the solution resistance. Electrochemical impedance spectroscopic (EIS) measurements were performed in the frequency range from 100 kHz to 0.1 Hz, and the polarization voltages were -0.20 V (HER) and 1.55 V (OER). The Tafel plots were obtained *via* Tafel equation: $\eta = a$ $\phi + b \log |j|$, where a, b and j are intercept, Tafel slop and current density, respectively.

3. Results and discussion

The microscopic morphologies of precursor and $NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF$ were observed by SEM. The nanosheets structures of precursor are smooth and vertically distributed on NF (Fig. S1a, b). The $NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF with rough and porous$ structure is formed after annealing at 400 \degree C in $NH₃$ atmosphere (Fig. S1c, d), which could increase the active specific surface area of the catalyst to expose abundant active sites [[33\]](#page-9-4). Meanwhile, this structure facilitates the contact between electrolyte and active sites, as well as bubbles escape, thereby improving the catalytic efficiency. Fig. S2 shows that the morphology of $NiFe₂O₄-Fe₂₄N₁₀/NF$

consisted of nanosheets and agglomerated nanoparticles, with only a few nanosheets growing on the agglomerated nanoparticles. The reason for this phenomenon could be that $NiFe₂O₄$ is a nanoparticle with uneven size, which is prone to agglomeration, and the introduction of $CeO₂$ avoids the sintering of NiFe₂O₄.

In addition, the NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF also shows the nanosheets morphology [\(Fig. 2](#page-4-0)a). In [Fig. 2b](#page-4-0), high-resolution TEM (HRTEM) images demonstrate the lattice distances of 0.251, 0.230 and 0.271 nm, which are derived from NiFe₂O₄(311), $Fe₂₄N₁₀(220)$ and $CeO₂(200)$ planes, respectively. And [Fig. 2](#page-4-0)c displays the lattice spacings of 0.208, 0.251, 0.230 and 0.271 nm, which could be assigned to NiFe₂O₄(400), NiFe₂O₄(311), Fe₂₄N₁₀(220) and $CeO₂(200)$, respectively. The above crystal planes are also observed in the XRD patterns ([Fig. 3](#page-5-0)a). As shown in Fig. $2d-i$, the HAADF-STEM and elemental mappings images demonstrate the homogeneous distributions of Ni, Fe, Ce, O and N on $NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF nanosheets.$

XRD was used to analyze the crystal structure of the prepared powder samples. The peaks located at 30.3° , 35.7° , 57.4° and 63.0° are derived from the (200), (311), (400) and (511) planes of NiFe₂O₄ (PDF#74-2081) in [Fig. 3a](#page-5-0). The reaction process could be as follows: The Fe^{3+} undergoes hydrolysis reaction to generate H⁺, and then H⁺ and NO₃ etch the NF to generate Ni^{2+} [\[34](#page-9-5)]. Subsequently, H_2O , Ni^{2+} and Fe³⁺ react to form NiFe₂O₄ in-situ on the surface of NF. The characteristic peaks at 39.1° (201), 41.5° (002), 44.5° (221) and 58.3° (222) belong to $Fe_{24}N_{10}$ (PDF#73-2103). The diffraction peaks of 28.5° , 33.1° , 47.5° and 56.3° are assigned to the (111), (200), (220) and (311) crystal planes of cubic fluorite $CeO₂$ (PDF#89-8436). XRD results indicate that the $NiFe₂O₄-Fe₂₄N₁₀-CeO₂$ is composed of three phases of NiFe₂O₄, Fe₂₄N₁₀ and CeO₂. Besides, the XRD patterns of NiFe₂O₄- $Fe₂₄N₁₀/NF$ and $CeO₂/NF$ are presented in Fig. S3.

Then, to study the discrepancy of surface chemical states and compositions between $NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF$, the XPS data of $NiFe₂O₄-Fe₂₄N₁₀/NF$ and $CeO₂/NF$ were obtained [\(Fig. 3](#page-5-0)b-d). The survey spectrum further proves that $NiFe₂O4-Fe₂₄N₁₀-CeO₂/NF contained Ni, Fe,$ Ce, N and O elements (Fig. S4a). For fine spectrum of Ni 2p in NiFe₂O₄-Fe₂4N₁₀-CeO₂/NF ([Fig. 3](#page-5-0)b), the peaks at 855.1 and 872.4 eV can be attributed to $Ni²⁺$, and the peaks at 857.0 and 874.3 eV correspond to $Ni³⁺$. The high-resolution Fe 2p spectrum consists of Fe²⁺(710.5/723.6 eV) and Fe³⁺(713.8/ 726.9 eV) ([Fig. 3c](#page-5-0)). Significantly, the Ni 2p and Fe 2p of NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF display a positive shift (Δ = 0.2 eV) relative to that of NiFe₂O₄-

Fig. 2. (a-c) TEM and HRTEM images of NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF, (d-i) HAADF-STEM and elemental mappings images.

Fe₂₄N₁₀/NF. While the Ce 3d in NiFe₂O₄-Fe₂₄N₁₀-CeO2/NF exhibits a distinguished negative binding energy shift ($\Delta = 0.3$ eV) than that of CeO₂/NF ([Fig. 3d](#page-5-0)). These phenomena confirm that $CeO₂$ acts as an electron acceptor to induce electron transfer from the two-phase $NiFe₂O₄-Fe₂₄N₁₀$ to $CeO₂$ due to the electronic interaction between $CeO₂$ and NiFe₂O₄-Fe₂₄N₁₀ [[35,](#page-9-6)[36](#page-9-7)], thereby changing the internal electronic structure [[37\]](#page-9-8). This will optimize the adsorption energy of reaction intermediates, thus improving catalytic performance [[38,](#page-9-9)[39](#page-9-10)]. The N 1s peaks of 397.6 and 399.4 eV correspond to Fe-N and N-H (Fig. S4b). It is reported that $Fe-N$ bonds can reversibly exchange oxygen reaction [[40\]](#page-9-11). The appearance of Fe-N is attributed to the $Fe₂₄N₁₀$ phase, which is generated by the coordination reaction of Fe atoms with N atoms under an ammonia atmosphere. Besides, the O 1s peak is presented in Fig. S4c.

Then electrochemical measurements were performed to assess the activity of catalysts. [Fig. 4](#page-6-0)a displays that $NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF only re$ quires low overpotential of 274 mV to reach 100 mA \cdot cm⁻² toward OER, which is better than $NiFe₂O₄-Fe₂₄N₁₀/NF$ (206 mV), CeO₂/NF (361 mV), NF (342 mV) and IrO₂/C/NF (353 mV). The reason may be that the introduction of $CeO₂$ can change the electronic structure and enhance the adsorption of intermediates, thereby speeding up the reaction rate. Obviously, the potential of NiFe₂O₄-
Fe₂₄N₁₀-CeO₂/NF relative to precursor $Fe₂₄N₁₀ - CeO₂/NF$ relative to $(E_{100} = 1.54 \text{ V})$ shifts significantly to a more negative direction, possibly benefitting from the porous structure formed by calcination to expose more active sites. Furthermore, $NiFe₂O₄-Fe₂₄N₁₀-CeO₂/$ NF still exhibits the best activity among all catalysts, even in harsh environment of large current density with severe reaction, suggesting that it exhibits significant promise for commercial applications. Among them, the activity of $IrO₂/C/NF$ is worse than that of NF due to the introduction of more Nafion binders with non-conductive properties in the preparation of $IrO₂/C/NF$.
Tafel slopes of NiFe₂O₄-Fe₂₄N

Tafel slopes of $NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF$ $(62.34 \text{ mV} \cdot \text{dec}^{-1})$, NiFe₂O₄-Fe₂₄N₁₀/NF (70.56) mV \cdot dec $^{-1}$), CeO₂/NF (125.47 mV \cdot dec $^{-1}$), IrO₂/C/NF (116.09 mV \cdot dec⁻¹), NF (104.49 mV \cdot dec⁻¹) and precursor (88.49 mV \cdot dec⁻¹) were calculated by polarization curves ([Fig. 4](#page-6-0)b). The minimum Tafel value of $NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF implies a kinetic advan$ tage relative to other prepared catalysts. It can be interpreted that the micro/nanoarray structure

Fig. 3. (a) XRD pattern of NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF; XPS spectra of (b) Ni 2p and (c) Fe 2p in NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF and NiFe₂O₄-Fe₂₄N₁₀/ NF, and (d) Ce 3d in NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF and CeO₂/NF.

formed by the rough nanosheets facilitates the mass transport and exposes abundant active sites, thereby accelerating the reaction kinetics. Besides, according to literatures, $NiFe₂O₄$ is prone to be agglomerated due to its inhomogeneous size of nanoparticles. The introduction of $CeO₂$ prevents NiFe₂O₄ nanoparticles from agglomeration and facilitates the adsorption of water molecules [[41\]](#page-9-12). In [Fig. 4b](#page-6-0), $NiFe₂O₄-Fe₂₄N₁₀$ - $CeO₂/NF$ exhibits the smallest charge transfer resistance (R_{ct}) observed by EIS, this is because that the interaction between $CeO₂$ and NiFe₂O₄-Fe₂₄N₁₀ accelerates electron transfer.

Chronopotentiometry (CP) method is adopted to evaluate the OER stability of NiFe₂O₄-Fe₂₄N₁₀- $CeO₂/NF$ [\(Fig. 4d](#page-6-0)). It can keep working at 500 mA \cdot cm $^{-2}$ for 30 h (potential change ca. 20 mV), indicating good OER stability. Fig. S5 shows the microstructure changes of catalysts during prolonged O_2 release, the porous nanosheets morphology could be well-remained under harsh conditions. The good OER stability benefits from

the porous nanosheets and the regulation of $CeO₂$, which can alleviate the catalyst agglomerate.

HER activity of $NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF was$ also evaluated. Although Pt/C/NF exhibited excellent intrinsic activity with the overpotential of 23 mV at -10 mA \cdot cm⁻² [\(Fig. 5](#page-6-1)a). However, it is fabricated with the assistance of Nafion binder, which cannot operate stably at large current density due to violent bubble release, thus leading to the mechanical shedding of active material. $NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF only needs an over-potential of 429 mV to stably reach$ potential of 429 mV to stably reach -1000 mA \cdot cm⁻², which is superior to NiFe₂O₄- $Fe₂₄N₁₀/NF$ (458 mV), NF (483 mV), precursor (458 mV) and $CeO₂/NF$. This excellent HER activity could be attributed to the synergy between $NiFe₂O₄-Fe₂₄N₁₀$ and $CeO₂$ optimizes the catalytic activity. As shown in [Fig. 5\(](#page-6-1)b and c), except for Pt/ C/NF, NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF displays smaller Tafel slope and R_{ct} than those of other samples, which implies that the introductions of $CeO₂$ and

Fig. 4. OER performance tests. (a) LSV curves, (b) Tafel slopes, (c) Nyquist curves, (d) chronopotentiometric curve at 500 mA·cm⁻².

Fig. 5. HER performance tests. (a) LSV curves, (b) Tafel slopes, (c) Nyquist curves, and (d) chronopotentiometric curve at –500 mA·cm⁻².

Fig. 6. (a) Overall water splitting LSV curves of NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF, NiFe₂O₄-Fe₂₄N₁₀/NF, CeO₂/NF, precursors and NF; (b) chronopotentiometric curve of NiFe2O₄-Fe₂₄N₁₀-CeO₂/NF at 500 mA·cm⁻² .

porous nanosheets can speed up the electron and mass transfers.

Besides, CP method was also employed to assess the HER stability [\(Fig. 5](#page-6-1)d). The potential changed by only 10 mV after $NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF$ was continuously operated for 30 h at -500 mA \cdot cm⁻², indicating the gratified HER durability. It is also demonstrated by the morphology of NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF after long-term H_2 evolution tests, that is, the morphology of porous nanosheets still preserved well (Fig. S6).

Electrochemical surface area (ECSA) is a significant parameter to evaluate the catalysts, which can be calculated by the following formula:

$$
ECSA = C_{\rm dl}/C_{\rm s} \tag{1}
$$

In Fig. S7, C_{dI} value was estimated by cyclic voltammetry (CV) method in the non-Faraday region $(0.8-1.0 \text{ V})$ with the scan rates from 20 to 100 mV \cdot s⁻¹. The NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF exhibits the highest $C_{\rm dl}$ value (22.69 mF \cdot cm $^{-2}$), which is 1.5, 5.9 and 10.8 times higher than those of $CeO₂/$ NF, $NiFe₂O₄-Fe₂₄N₁₀/NF$ and precursor. The above results clearly demonstrate that the introduction of $CeO₂$ can increase the ECSA and effective active sites, thereby enhancing the kinetics of the water electrolysis reaction. The reason could be that the presence of $CeO₂$ avoids the problem of easy agglomeration of $NiFe₂O₄$, thereby exposing active sites. Meanwhile, the porous nanosheet structure of NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF after annealing can facilitate the contact between electrolyte and active sites, and thus enriching the activity.

As displayed in Fig. S8, the activity of catalyst was optimized by annealing at different temperatures (300, 400, 500, 600 and 700 °C) under NH₃ atmosphere. The sample prepared at $400\degree\text{C}$ has the superior OER and HER activities as well as the fastest kinetics. The annealing temperature will change the chemical composition and microscopic morphology of the catalyst, thereby affecting the catalytic activity.

In [Fig. 6](#page-7-0)a, considering the good HER and OER performances of $NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF,$ it was utilized as an anode and a cathode for assembling a water electrolyzer $(NiFe₂O₄-Fe₂₄N₁₀-CeO₂/$ $NF\vert\vert NiFe_2O_4-Fe_{24}N_{10}-CeO_2/NF$). The $NiFe_2O_4 Fe₂₄N₁₀-CeO₂/NF$ also displays good performance with a potential of 1.81 V at 100 mA \cdot cm⁻² for OWS, which is superb to $NiFe₂O₄-Fe₂₄N₁₀/NF$ (1.83 V), $CeO₂/NF$ (1.96 V), precursor (1.89 V) and NF (1.90 V). The OWS stability test of NiFe₂O₄- $Fe₂₄N₁₀-CeO₂/NF$ demonstrates that it can continuously operate for 30 h at 500 mA \cdot cm⁻² with voltage rise of only 0.2 V ([Fig. 6](#page-7-0)b), indicating its potential for industrial application.

4. Conclusions

In summary, the $CeO₂$ electronic structure modulated FeNi bimetallic composite porous nanosheets in-situ grown on 3D NF are prepared through a solvothermal and annealing method. Benefiting from the electronic interaction of $CeO₂$ modulating the electronic structure of FeNi composite and self-supporting porous nanosheets, the prepared $NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF$ exhibited excellent water electrolysis activity with a low potential of 1.81 V to arrive at 100 mA \cdot cm⁻² for OWS. It also displays good stability at 500 mA \cdot cm⁻² for 30 h with no significant potential augment. These results indicate that the $NiFe₂O₄-Fe₂₄N₁₀-CeO₂/NF$ meets the requirements of high efficiency and prolonged stable operation at large current

density. Hence, this work has implications for designing bifunctional catalysts applied in industrial water electrolysis hydrogen production.

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CeO₂ 电子调控 FeNi 纳米片大电流密度电解水催化剂

丁明宇, 蒋文杰, 余天琦, 卓小燕, 覃晓静, 尹诗斌* 广西电化学能源材料重点实验室, 广西大学化学化工学院, 广西 南宁 530004

摘要

开发高活性的大电流密度非贵金属双功能催化剂对于电解水制氢的发展意义重大。本文通过水 热法和高温退火处理制备了自生长在泡沫镍上的 CeO2 电子调控的 FeNi 双金属复合物多孔纳米片 (NiFe2O4-Fe24N10-CeO2/NF)。电化学测试结果表明, NiFe2O4-Fe24N10-CeO2/NF 在 1.0 mol·L⁻¹ KOH 电解液中具有出色的析氧和析氢反应(OER 和 HER)活性, 在±1000 mA·cm⁻² 电流密度下所需的过 电位分别为 352 mV 和 429 mV。将其组装成电解水 (OWS) 两电极体系, 只需 1.81 V 的电池电压 就能达到 100 mA·cm⁻² 的电流密度。对于 OER、HER 和 OWS, 可以在±500 mA·cm⁻² 的电流密度 下稳定运行 30 小时,其优异的大电流密度催化性能可以归功于 CeO2 对于 FeNi 复合物的电子结构 调控增强了催化剂的本征活性和反应中间体的吸附。原位生长在泡沫镍(NF)上的多孔纳米片可以 增强活性位点与电解质的接触,并利于气体产物的释放,从而提高其化学稳定性和机械稳定性。本 工作为制备双功能非贵金属电解水催化剂提供了一种新思路。

关键词: FeNi 复合物; 析氢反应; 析氧反应; 催化剂; 电解水