Facile synthesis of Nickel-iron based electrocatalyst anodes for efficient oxygen evolution reaction

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Abstract: In this research, we reported facile synthesis of efficient Ni3Fe electrocatalyst nanostructures deposited on conducting carbon fibers surface by a simple chemical bath deposition method at moderate temperature. The composition, phase and electrocatalytic property of as-prepared binder-free electrocatalyst was characterized by X-ray diffraction (XRD), field emission scanning electron microscopy (FE-SEM), and electrochemical measurements. The as-synthesized Ni3Fe@CF presented excellent performance and durability as water oxidation electrocatalyst in alkaline electrolyte owing to non-segregated deposition of Ni3Fe nanostructures on conducting surface of carbon, high electrochemical surface area, and the fast absorption and desorption of water molecules during oxygen evolution reaction due to the 3D architecture of conducting interlaced carbon fibers template.

Keywords: Chemical deposition, Ni3Fe nanostructures, carbon cloth fibers, electrochemical surface area, conductivity, oxygen evolution.

1. Introduction

Impeding global energy crises became the world extremely defensive over their natural resources and it is crucial to seek alternative energy conversion and storage systems such as water electrolysis, fuel cells and metal-air batteries [1]. Oxygen evolution reaction (OER) is bottleneck in water splitting and is process of generating molecular oxygen through electrochemical oxidation of water [2]. Nevertheless, OER process suffers from sluggish kinetics with a complicated multi-electron transfer step, resulting in a high overpotential requirement [3]. An effective electrocatalyst is needed in order to expedite the OER reaction to reduce the overpotential, and thus enhance energy conversion efficiency. Currently, noble metal oxides such as IrO2 or RuO2 are the benchmark water oxidation electrocatalysts in alkaline solutions, but their scarcity, high cost and poor stability restrict their implementation as water oxidation electrocatalyst at large scale [4–7]. In this regard, the exploit of earth-abundant, stable and highly efficient water oxidation electrocatalysts are highly needed.

A wide range of non-noble metal electrocatalysts have been developed with promising water oxidation performance including, transition-metal oxides, sulfides and nitrides [8–12]. Among these developed electrocatalysts, the nickel-iron based materials have attracted intensive attention because of their high water oxidation activity and stability [13]. However, they exclusively yield low catalytic activity owing to their insufficient intrinsic electrical conductivity and small surface area of the catalyst structures. Recently, combining highly conductive scaffold materials with inexpensive transition metal oxides have been used to enhance charge transport properties; thereby promoting catalytic activity. However, most of these fabricated materials exhibit buried active sites due to segregated morphology. Expensive synthetic protocols have been adopted to address these problems [14–16].

Herein, we report a simple strategy to develop binder-free nickel-iron based electrocatalyst films by adopting chemical bath deposition technique at moderate temperature. The developed Ni3Fe nanostructures deposited on carbon scaffold exhibited high OER performance with stability up to 6500s with the 89% activity...
retention in alkaline electrolyte under vigorous bubbling across its electrode surface corresponding to $J = 10$ mA cm$^{-2}$ (overpotential $= 320$V versus RHE). This improvement in the OER performance correlated with the significantly improved conductivity and the large electrochemical active surface area to provide more accessible catalytic active sites for the adsorption and desorption of water oxidation intermediates. We believe that this would be an attractive and facile strategy for the cost-effective development of efficient nickel-iron based water oxidation catalyst.

2. Materials and Methods

Isopropyl alcohol (HPLC, $\geq 99.5\%$), ethanol (AR, $\geq 99.7\%$), iron nitrate ($Fe(NO_3)_3\cdot 9H_2O$) (AR grade, 99.9%), nickel nitrate ($Ni(NO_3)_2\cdot 6H_2O$) (99.9%, Aladdin Industrial Co., Ltd.) and KOH (AR grade, 99%) were used in this experimental work. Double distilled water was used in preparing solutions and washing of electrodes.

2.1. Synthesis of t-CF

The cut pieces of CF ($11 cm^2$) were washed with isopropyl alcohol, ethanol and then water in succession, each for 5 minutes in water bath sonication and then oxidized by using previously reported modified Hummers method and named as $t-CF$ [17].

2.2. Synthesis of Ni$_3$Fe@t-CF

Typically, the Ni$_3$Fe@t-CF electrocatalyst was prepared by chemical bath deposition technique at room temperature. The deposition of Ni$_3$Fe nanostructures on t-CF surface was carried out using chemical bath solution containing $Ni(NO_3)_2\cdot 6H_2O$ (3 mM) and $Fe(NO_3)_3\cdot 9H_2O$ (1 mM). The chemical bath solution was continuously purged with Ar gas during electrodeposition to avoid the oxidation of $Fe^{2+}$ to $Fe^{3+}$. The deposition was carried by oil bath heating at 80$^\circ$C for 2 hours.

2.3. Characterizations

FE-SEM was carried out using a Hitachi S4800 with a 10kV accelerating voltage. XRD was performed on a PANanalytical XPert powder diffractometer at operating voltage of 40kV and operating current of 40mA. All electrochemical measurements were carried out in a 3-electrode cell in 1 M KOH electrolyte solution (pH $\sim 13.6$) using CHI 760 electrochemical workstation. As-prepared Ni$_3$Fe@t-CF, t-CF and CF samples were used directly as the working electrodes. All measured potentials were calibrated to RHE using the following Equation (1);

$$E_{RHE} = E_{SCE} + 0.244V + 0.059 \times pH.$$ (1)

Linear sweep voltammetry (LSV) polarization and cyclic voltammetry (CV) curves in non-Faradaic potential region (0.10.2 V versus RHE) were recorded at a scan rate of 10 mV s$^{-1}$. Tafel slopes were derived from the corresponding OER polarization curves with $iR$ compensation in alkaline solution for Ni$_3$Fe@t-CF, t-CF and CF using following Equation (2);

$$\eta = a + blog(j),$$ (2)

where $\eta$ is the overpotential, whose value can be calculated from Equation (3);

$$E(\eta) = E(SCE) + 0.244 + (0.059 \times pH)1.23,$$ (3)

where $b$ is the Tafel slope measured in mV/dec and $j$ is the current density. The chronoampermetric measurements were conducted under the same above described experimental conditions but without $iR$ drop compensation.

The electrochemical active surface area (ECSA) of samples CF, t-CF and Ni$_3$Fe@t-CF were calculated from the electrochemical double layer capacitance ($C_{dl}$). The scan-rate ($v$) dependent capacitive current, $I_c$ is associated with the $C_{dl}$ according to relation:

$$ECSA = C_{dl} = I_c / v,$$ (4)
where \( I_c = \frac{I_{anodic}}{I_{cathodic}} \), calculated in the middle of potential domain as a function of scan rate \( v \), that yielded a linear slope equal to \( 2 \times C_{dl} \).

Electrochemical impedance spectroscopy (EIS) was used to evaluate the electrochemical response of the synthesized OER electrocatalysts towards an applied alternating potential and to find the resistance in the transfer of charges \( (R_{CT}) \). The EIS measurements were conducted at 1.44 V versus RHE in the frequency range 100 kHz to 0.1 Hz with 10 mV amplitude of sinusoidal wave.

3. Results

3.1. X-ray Diffraction (XRD)

Phase determination of the CF, t-CF and \( Ni_3Fe@t-CF \) was carried out by XRD analysis and their XRD patterns as shown in Figure 1. Figure 1(a) shows the graphite nature of CF and t-CF (JCPDS card no. 03 − 1185). While the characteristic diffraction peaks of graphite at 25.60 and 430 are less intense in t-CF in comparison to CF, demonstrates its less ordered crystalline structure. Figure 1(b) shows the intense diffraction peaks at 44.90, 52.390 and 76.730 that correspond to \((111)\), \((200)\) and \((220)\) planes of crystalline \( Ni_3Fe \) and matches with JCPDS card no. 48 − 0419.

3.2. Field-Emission Scanning Electron Microscope (FE-SEM)

Figure 2(a) shows the FE-SEM image of pristine CF containing 3-dimensional framework of interlaced carbon fibers with void spaces between them. The high magnification image of the red squared area in Figure 2(a) shows the smooth surface of the CF. While the Figure 2(d) shows the high magnification image of the red squared area in Figure 2(c) that presents the dispersion of highly porous nickel-iron alloy nanostructures with high surface area on the smooth surface of CF. The high surface area with more exposed active sites, electrolyte penetration and fast adsorption and desorption of the generated gas bubbles facilitate OER activity.

3.3. Linear Sweep Voltammetry (LSV)

The LSV polarization curves of pristine CF, t − CF and \( Ni_3Fe@t-CF \) are shown in Figure 3. The OER performance was evaluated from onset and the overpotentials needed to drive current density of 10 mA cm\(^{-2}\) measured from corresponding LSV curves in alkaline electrolyte using standard 3-electrode electrochemical system. The pristine CF, t − CF and \( Ni_3Fe@t-CF \) exhibit the onset potentials of 1.6, 1.57 and 1.48 V versus RHE, respectively. The significantly lower overpotential needed for 10 mA cm\(^{-2}\) was 320 V for \( Ni_3Fe@t-CF \). While both pristine CF and t − CF exhibit negligible OER activity with very small current densities of 2 mA cm\(^{-2}\) and 1.7 mA cm\(^{-2}\), respectively at 1.7 V versus RHE. The large surface area is resulting from the highly porous and rough surface of \( Ni_3Fe@t-CF \) electrocatalyst as evident in its SEM images. While the efficient intrinsic OER activity is owing to more exposed active sites for the OER reactants.

3.4. Tafel Slope

The Tafel plots for all the materials were generated from the LSV polarization curves by plotting the log of current \((\log I)\) against the applied potentials \((V_{SCE})\). The Tafel slopes \((b)\) were extracted by fitting straight line portion of the Tafel plots to the Tafel equation (Equation (2)). Figure 4 shows the Tafel slopes of CF, t − CF and \( Ni_3Fe@t-CF \) derived from their respective LSV polarization curves to show the influence of the applied
Figure 2. SEM image of (a) pristine CF, (b) high resolution SEM image of the area squared in (a), (c) $Ni_3Fe@t – CF$ and (d) high resolution SEM image of the area squared in (c).

Figure 3. LSV curves (with $iR$ correction) of CF (black curve), $t – CF$ (purple curve) and $Ni_3Fe@t – CF$ (red curve) in alkaline electrolyte.

OER overpotential on the current density ($I$) to evaluate their OER kinetics. The Tafel slope for $Ni_3Fe@t – CF$ was about $65 \text{ mV/dec}$ that is comparable to the previously reported nickel-iron based electrocatalysts [17].

Figure 4. Tafel slopes of CF (black curve), $t – CF$ (purple curve) and $Ni_3Fe@t – CF$ (red curve) in alkaline electrolyte.
3.5. Electrochemical active surface area (ECSA) measurements

Figure 4(a-c) show the CV curves for pristine CF, t – CF and Ni$_3$Fe@t – CF in the non-Faradaic potential region from 0.1 to 0.2V in alkaline solution. Where for these CV measurements, CV curves were run at multiple scan rates ($v$) of 20, 40, 60, 80, 100, 150, 200, 250, 300 and 350mVs$^{-1}$. The plot of capacitive current ($i_c$) at the middle of the applied potential window versus $v$ gives $C_{dl}$ that has direct relation with ECSA as given in equation (4). Figure 5(d) shows that Ni$_3$Fe@t – CF exhibits the highest ECSA of 1.7mFcm$^{-2}$ in comparison to t – CF (0.87mFcm$^{-2}$) and CF (0.44mFcm$^{-2}$). The larger ECSA is resulting from the highly porous and rough surface of Ni$_3$Fe@t – CF as evident in its SEM images. The efficient intrinsic OER activity is owing to more accessible active sites of this electrocatalyst for the reactant species.

![Figure 5. CV curves in non-Faradaic region (a) CF, (b) t – CF and Ni$_3$Fe@t – CF. (d) Calculated $C_{dl}$ plots of CF (black curve), t – CF (blue line) and Ni$_3$Fe@t – CF (red line).](image)

3.6. Electrochemical Impedance Spectroscopy (EIS)

Further, electrochemical impedance spectroscopy (EIS) measurements were carried out to study the kinetics of interfacial electrocatalytic behaviour of CF, t – CF and Ni$_3$Fe@t – CF. The obtained Nyquist plots and charge transfer resistances ($R_{CT}$) are shown in the Figure 6. These plots showing two semicircles corresponding to RCT at the electrocatalyst/electrolyte interface and in the electrocatalyst electrode, in the low and high frequency regions, respectively. The semicircle for the Ni$_3$Fe@t – CF in the high frequency range indicates the enhanced charge transport with very less $R_{CT}$ 0.9$\omega$ in comparison to CF and t – CF. It reveals the good conductivity and the good interfacial contact between the 3D conducting framework of CF and the Ni$_3$Fe electrocatalyst.

3.7. Chronoamperometric Measurements

Durability is an important feature to evaluate OER activity for practical applications at large scale. The stability of the as-synthesized Ni$_3$Fe@t – CF electrocatalyst was recorded over 6500s in alkaline electrolyte at 1.54V versus RHE (without iR drop compensation). Figure 7 shows good stability of Ni$_3$Fe@t – CF electrocatalyst by preserving a relative 89% of the initial current density of 10mAcm$^{-2}$ suggesting its good operational stability in the alkaline solution.

4. Conclusion

In this work, we have used a facile and effective strategy for the synthesis of Ni$_3$Fe nanostructures deposited on the 3D conducting surface of carbon at moderate temperature as an efficient OER catalyst in alkaline electrolyte. The high OER performance of the as-synthesized Ni$_3$Fe@t – CF sample is attributed to
Figure 6. Nyquist plots of CF (black curve), t – CF (purple curve) and Ni₃Fe@t – CF (red curve) in alkaline electrolyte.

Figure 7. Chronoamperometric test of Ni₃Fe@t – CF at 1.54 V versus RHE.

the high electrochemical active surface area as inferred from its FE-SEM images, low charge transfer resistance due to intimate contact between electrocatalyst nanostructures and the 3D conducting template. Considering the high performance of our synthesized binder-free electrocatalyst, this work will undeniably provide a prospect for the development of inexpensive and highly active OER electrocatalyst films by adopting this simple synthetic technique.

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References


