UC Irvine UC Irvine Previously Published Works

Title

Ultrafast Preparation of Nonequilibrium FeNi Spinels by Magnetic Induction Heating for Unprecedented Oxygen Evolution Electrocatalysis.

Permalink

https://escholarship.org/uc/item/3zf1w70h

Authors

Lu, Bingzhang Liu, Qiming Wang, Chunyang <u>et al.</u>

Publication Date

2022

DOI

10.34133/2022/9756983

Peer reviewed

Research Article

Ultrafast Preparation of Nonequilibrium FeNi Spinels by Magnetic Induction Heating for Unprecedented Oxygen Evolution Electrocatalysis

Bingzhang Lu^(b),¹ Qiming Liu^(b),¹ Chunyang Wang,² Zaheer Masood^(b),³ David J. Morris^(b),⁴ Forrest Nichols,¹ Rene Mercado^(b),¹ Peng Zhang^(b),⁴ Qingfeng Ge^(b),³ Huolin L. Xin^(b),² and Shaowei Chen^(b)

¹Department of Chemistry and Biochemistry, University of California, 1156 High Street, Santa Cruz, California 95064, USA ²Department of Physics and Astronomy, University of California, Irvine, California 92697, USA ³Department of Chemistry and Biochemistry, Southern Illinois University, Carbondale, Illinois 62901, USA

⁴Department of Chemistry, Dalhousie University, 6274 Coburg Road, Halifax, NS, Canada B3H 4R2

Correspondence should be addressed to Qingfeng Ge; qge@chem.siu.edu, Huolin L. Xin; huolinx@uci.edu, and Shaowei Chen; shaowei@ucsc.edu

Received 17 February 2022; Accepted 18 April 2022; Published 1 June 2022

Copyright © 2022 Bingzhang Lu et al. Exclusive Licensee Science and Technology Review Publishing House. Distributed under a Creative Commons Attribution License (CC BY 4.0).

Carbon-supported nanocomposites are attracting particular attention as high-performance, low-cost electrocatalysts for electrochemical water splitting. These are mostly prepared by pyrolysis and hydrothermal procedures that are time-consuming (from hours to days) and typically difficult to produce a nonequilibrium phase. Herein, for the first time ever, we exploit magnetic induction heating-quenching for ultrafast production of carbon-FeNi spinel oxide nanocomposites (within seconds), which exhibit an unprecedentedly high performance towards oxygen evolution reaction (OER), with an ultralow overpotential of only +260 mV to reach the high current density of 100 mA cm⁻². Experimental and theoretical studies show that the rapid heating and quenching process (ca. 10^3 K s⁻¹) impedes the Ni and Fe phase segregation and produces a Cl-rich surface, both contributing to the remarkable catalytic activity. Results from this study highlight the unique advantage of ultrafast heating/ quenching in the structural engineering of functional nanocomposites to achieve high electrocatalytic performance towards important electrochemical reactions.

1. Introduction

Design and engineering of low-cost, high-performance catalysts play an important role in the development and advancement of electrochemical energy technologies, such as fuel cells and water electrolyzers [1]. Up to now, a range of functional nanocomposites have been hailed as viable alternatives to the conventional, noble metal-based catalysts, such as doped carbon, and carbon-supported nanoparticles of metals, metal oxides, sulfides, phosphides, and selenides [2–4]. These materials are typically prepared via the "traditional" thermal methods based on pyrolysis and hydrothermal procedures [5–12]. While these methods are rather facile and effective in sample synthesis, they are energy and time-consuming [13, 14], and the slow heating ramp makes it difficult to produce a nonequilibrium phase within the samples, which might be critical in regulating the electronic structure and hence the electrocatalytic activity [15–18]. In a prior study [19], Holewinski et al. demonstrated that whereas Ag and Co metals were immiscible at equilibrium, AgCo alloy nanoparticles could be obtained by removing the sample from the tube furnace while hot and letting it cool down in ambient instead of following a traditional programmed cooling process and exhibited a remarkable electrocatalytic activity towards oxygen reduction reaction (ORR) unseen for either Ag or Co metals alone. Li et al. [20, 21] showed that the laser ablation method could be exploited for the synthesis of Ag and Ru nanoparticles for high-efficiency water splitting, due to the formation of plenty of stacking faults or grain boundaries that were

difficult to produce via conventional methods. Yao et al. [22] developed carbothermal shock synthesis to prepare highentropy alloy nanoparticles within seconds. Chen et al. [23] adopted flash joule heating to synthesize metastable 1 T phase of MoS_2 with S vacancies and observed a high efficiency towards hydrogen evolution reaction (HER). In another study [24], Zhang et al. developed a laser scribing method where metal oxide/graphene composites were prepared by simply lasing metal precursors loaded onto a graphene scaffold and exhibited a high performance towards oxygen evolution reaction (OER) likely due to the formation of surface defects. Li et al. [20, 25, 26] successfully used the laser ablation method to prepare a series of water splitting electrocatalysts with oxygen vacancies, faults, and unique morphologies.

Despite the progress, the toolbox for such sample synthesis has been limited, and the range of materials that can be produced and the extent of structural engineering remain narrow. Thus, further development of effective protocols for the synthesis of materials with unprecedented structures and properties is of both fundamental and technological significance [27, 28]. In electrochemical water splitting, OER has been recognized as a major bottleneck that limits the overall performance because of complex reaction pathways and sluggish electron-transfer kinetics [17], and FeNi (oxy) hydroxides and spinel oxides have been extensively studied as viable alternatives to the traditional, noble metal-based commercial catalysts [29-34], where manipulation of the occupation of the e_{σ} orbitals of the octahedral metals and/ or metal-oxygen covalency represents the leading strategies for further enhancement of the OER activity [30]. This is generally achieved by engineering the spinel components, heterometal doping, and introduction of oxygen vacancies [25, 31, 35, 36]. Phase segregation of Fe and Ni in the spinels has been believed to be the leading cause of the apparent loss of the electrocatalytic activity [37]. Yet, such segregation is inevitable for samples prepared via a "tedious" conventional thermal procedure as it is energetically favorable. In addition, residual heteroanions (e.g., Cl) adsorbed on or doped into the surface of Fe, Co, and Ni (hydro/oxyhydro) oxides may play a significant role in OER electrocatalysis [38-40]. Yet the impacts of such anion impurities have remained largely ignored, although most pyrolytically prepared spinel oxides are derived from iron and nickel chlorides. These issues can be addressed with the development of appropriate synthetic methods where nonequilibrium structures can be produced with reduced phase segregation and a remarkable concentration of anion impurities. This is the primary motivation of the present study.

Herein, we report an ultrafast heating/quenching method based on magnetic induction heating/rapid quenching (MIHRQ) [41] to prepare carbon-supported FeNi spinel composites (within seconds), which exhibit a clear mixing of the Ni and Fe phases and a Cl-rich surface, in contrast to the control samples that are prepared by prolonged heating and/or natural cooling to the ambient. In electrochemical measurements, the former displays an outstanding electrocatalytic performance towards OER, with an ultralow overpotential of only +260 mV to reach the high current density of 100 mA cm⁻², due to the formation of a nonequilibrium structure that is optimal for the adsorption of key reaction intermediates and eventual production of oxygen. The enhanced performance of the catalyst is also confirmed by results of first principle calculations. These results highlight the unique potential of MIHRQ in the deliberate production of nonequilibrium features in composite electrocatalysts. Notably, such an unprecedented tool can be readily extended to the preparation of a wide range of functional nanocomposites for diverse applications [42–47].

2. Results and Discussion

2.1. Magnetic Induction Heating-Rapid Quenching. The homemade MIHRQ apparatus is shown in Figure 1(a). A four-turn induction solenoid was twisted at a diameter of 5 cm, under which was placed a beaker containing ethanol and dry ice $(-78^{\circ}C)$ as the quenching agent. Experimentally, a calculated amount of the metal precursors (i.e., FeCl₃ and NiCl₂) was dropcast onto a piece of carbon paper $(1.5 \text{ cm} \times 0.5 \text{ cm})$, which was then sandwiched between two rectangular iron sheets $(2.5 \text{ cm} \times 2.5 \text{ cm} \times 0.01 \text{ cm})$. An iron nail was inserted into the center of the iron sheets and clamped to hold the assembly, which was placed in the center of the induction solenoid (Figure 1(b)). When a high frequency (30 kHz) current was passed to the solenoid, a strong magnetic field was produced, which instantly generated a strong Eddy current in the iron sheets and thus heated the sample rapidly to a high temperature. The induction current and time can be varied to control the heating temperature (Figure 1(c)). For example, a solenoid current of 200 A for a heating time of 4 s would generate a temperature of 200-300°C, which barely changed the color of the iron sheets (Movie S1). Yet, when the solenoid current was increased to 400 and 600 A, the temperature could reach ca. 600 and 1000°C, respectively, inducing a glowing color of the iron sheets from faint red to white (Figure 1(b), Movie S2). Such ultrafast heating (instant heating rate up to 10^3 Ks^{-1}) can not only drastically enhance the time efficiency of sample preparation but more importantly also be exploited for the kinetic control of the materials structures, in contrast to typical pyrolysis or hydrothermal synthesis, which is usually completed within the time frame of hours to even days. After a select period of heating (of the order of seconds), the sample would be dropped into the quenching solution below (Movie S1 and S2) or removed from the solenoid and cooled down naturally in the ambient. This offers an additional control of the materials structures, especially for the production of nonequilibrium features.

A series of samples were prepared with the MIHRQ setup at a controlled induction current (X) for a select period of time (Y) and denoted as FeNiO-X-Y (Figure 1(d)). Control samples were prepared in the same manner except for cooling in ambient, and referred to as FeNiO_{NC} -X-Y. Notably, MIHRQ can be extended to the preparation of a wide range of electrocatalysts beyond the FeNi spinel oxides (vide infra).

2.2. Structural Characterizations. High-angle annular dark field-scanning transmission electron microscopy (HAADF-

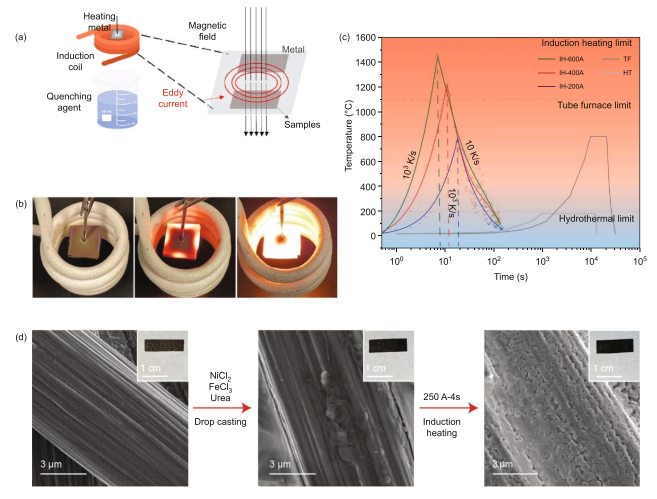


FIGURE 1: Sample preparation by MIHRQ. (a) Schematic illustration of magnetic induction heating-quenching for material preparation. (b) Photographs of induction heating with a solenoid current at (left) 200 A, (middle) 400 A, and (right) 600 A for 4 s, respectively. (c) Variations of temperature versus time of magnetic induction heating, traditional hydrothermal heating, and pyrolysis. The dash lines indicate the cases with the quenching process. 'IH,' 'TF,' and 'HT' are short for induction heating, tube furnace, and hydrothermal, respectively. (d) SEM images of the NiFeO-250-4 sample at different preparation stages. The insert images are the corresponding photographs of the electrodes. The carbon paper can be seen to become darkened after the deposition of the metal salt precursors, and subsequent induction heating and rapid quenching leads to a pitch-black appearance of the carbon paper, with particulates formed onto the carbon fibers.

STEM) measurements of the NiFeO-250-4 sample (Figure 1(d)) show the formation of a number of nanoparticles (dia. 20 to 100 nm) in irregular shapes (Figure 2(a)), with the structure consistent with Fe3-xNixO4-type spinel, as observed along the <111> zone axis (Figure 2(b) and Figure S1). The interatomic distance was estimated to be 0.816-0.824 nm (inset to Figure 2(b)), close to that of FeNi spinel oxide (0.835 nm) [48]. Additional STEM images acquired along the <112>, <100>, and<103> zone axes also confirmed the spinel lattice structure (Figure S2-S4). Furthermore, energy-dispersive X-ray spectroscopy- (EDS-) based elemental mapping studies clearly show an even distribution of Fe and Ni within the lattice (Figure 2(c)), suggesting atomic mixing of the Fe and Ni elements and no phase segregation. Notably, spindle-like FeNi oxide nanocrystals can also be found around these particles (Figures 2(d)-2(f) and Figure S5–S7), which feature a chlorine-rich surface (Figure 2(g)), with a Cl concentration of 12% in comparison to under 2% within the particles (Table S1). In fact, the atomic ratio of Fe:Ni:O:Cl in the nanospindles is estimated to be 1.9:1:4.9:1.1, while the overall ratio is close to 4.1:1:5.8:0.22, indicating that the spinel particles were Fe-rich oxide, while the nanospindles likely represent an intermediate phase between the precursors (metal chlorides) and the final spinel crystal (Table S1). Furthermore, it is noticeable that the Fe:Ni ratio is higher than the feeding ratio. This is likely because part of Ni was not fully converted into Ni oxide and washed away during the rapid quenching process.

For FeNiO-250-16 that was prepared via a longer heating time, the Fe:Ni:O:Cl ratio was estimated to be 6.6:1:8.6:0.028 (Table S1), indicative of the formation of a Fe-rich structure that was almost free of Cl. In fact, the nanospindle features, with the unique chlorine rich surface, can only be produced with a short heating time and rapid quenching process.

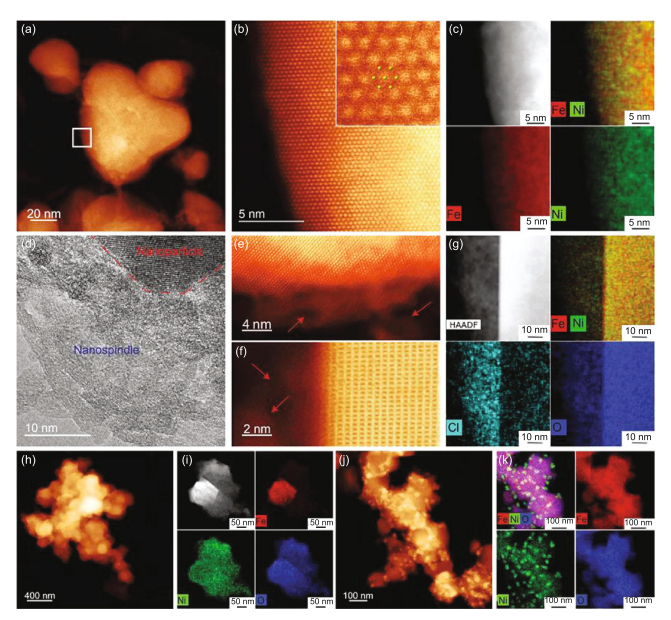


FIGURE 2: Electron microscopic studies of FeNi spinel oxides. (a) High-angle annular dark-filed scanning transmission electron microscopy (HAADF-STEM) image showing the morphology of FeNiO-250-4. (b) Atomic-resolution HAADF-STEM image and an enlarged image (inset) corresponding to the boxed region in (a) acquired along the <111> zone axis. (c) High-resolution energy-dispersive spectroscopy-(EDS-) based elemental maps of Fe and Ni in a FeNiO-250-4 particle, which features a FeNi spinel structure with no FeNi phase segregation. (d) High-resolution TEM (HRTEM) image of nanospindles in FeNiO-250-4. (e, f) HAADF-STEM images of nanospindles on the edge of FeNiO-250-4 (low-contrast regions), as highlighted by red arrows. (g) EDS mapping images of the interface between nanospindles and nanoparticles in FeNiO_{NC}-250-4, which show a Cl-rich surface of the nanospindles. (h) HAADF-STEM images and (i) EDS mapping images of FeNiO_{NC}-250-4, where natural cooling leads to the formation of nanoparticles aggregates and obvious FeNi phase segregation. (j) HAADF-STEM and (k) EDS mapping image of FeNiO-250-16, where prolonged heating (higher temperature) leads to the formation of a significant amount of metallic Ni nanoparticles on the FeNi spinel.

The control sample, FeNiO_{NC}-250-4 that was produced by similar heating but natural cooling in the ambient, exhibited an obviously different morphology (Figure 2(h)), consisting of aggregates of nanoparticles into large chunks. In addition, significant phase segregation occurred within the sample (Figure 2(i)), where the elements of Ni and O appeared to be evenly distributed across the sample, whereas Fe was mostly confined within a small region, suggesting the growth of FeNiO spinel nanocrystals on a nickel oxide scaffold. This is consistent with the sample atomic ratio of Fe : Ni : O : Cl = 0.15 : 1 : 2.3 : 0.036. The fact that the sample was markedly richer in nickel was likely due to higher thermal volatility of the iron compounds [49], where the enhanced loss of Fe was facilitated by the relatively slow cooling (about 10 K s^{-1}). Similar phase segregation was also observed with a prolonged heating time (e.g., FeNiO-250-16), where the temperature could reach ca. 1000° C (Figures 2(j) and 2(k) and Figure S8). Note that such

nonhomogeneous segregation is actually the equilibrium state at high temperatures based on the phase diagram of Fe-Ni-O₂ (FToxid database and FactSage) [50], which can be rationally avoided by rapid heating and quenching as manifested with FeNiO-250-4.

The material structures were further characterized by Xray photoelectron spectroscopy (XPS) and X-ray absorption spectroscopy (XAS) measurements. Figure 3(a) depicts the high-resolution XPS scan of the Ni 2p electrons of FeNiO-250-4, where two peaks can be resolved at 855.9 and 856.9 eV for the Ni(II) $3p_{3/2}$ electrons, suggesting the formation of five and six oxygen-coordinated Ni atoms on the surface (i.e., Ni(OH)₂), respectively [51], since no NiO species (binding energy around 854.7 eV) was detected [52]. A single peak was resolved at 711.2 eV in the Fe 2p scan (Figure 3(b)), due to the Fe(III) 3p_{3/2} electrons [53], whereas three peaks were deconvoluted in the O 1s spectrum (Figure 3(d)) at 531.5 eV for hydroxide, 529.9 eV for metal-O, and 533.0 eV for C-O [54]. These observations are consistent with the formation of FeNiO spinel lattices (vide infra). Deconvolution of the Cl 2p spectrum (Figure S9) yields two peaks at 198.3 and 199.9 eV, due to the $2p_{3/2}$ and $2p_{1/2}$ electrons of metal-Cl, respectively [55]. Taken together, these results suggest that the FeNiO-250-4 sample surface was mostly terminated with OH and Cl groups, in good agreement with results from the TEM and EDS measurements.

The XPS spectra of the FeNiO_{NC}-250-4 and FeNiO-250-16 samples are shown in Figure S10–S12. The Fe 2p spectra showed only a rather insignificant difference among the sample series (Figure S11), likely because of the high thermal activity of FeCl₃ and the facile formation of Fe oxide. Additionally, FeNiO-250-4 and FeNiO_{NC}-250-4 exhibited a very similar Ni 2p profile with the Ni(II) $2p_{3/2}$ binding energy at 855.9 and 856.9 eV, which was somewhat lower for FeNiO-250-16 (855.6 and 856.75 eV, Figure S10). Furthermore, in the O 1 s spectra (Figure S12), the Ni(Fe)-O peaks (ca. 530 eV) of FeNiO_{NC}-250-4 and FeNiO-250-16 are significantly larger than that of FeNiO-250-4, indicating increasing contributions from bulk oxides.

Further oxidation state and structural insights were obtained by XAS measurements. Figure 3(d) depicts the Ni K-edge X-ray absorption near edge structure (XANES) spectra of the sample series, where the absorption edge intensity can be seen to vary in the order of Ni foil<FeNiO-250-16<FeNiO-250-4<FeNiO_{NC}-250-4<NiO, suggesting that the Ni valence state in the three FeNiO samples was in the intermediate between those of metallic Ni and Ni²⁺. A similar trend can be observed in the Fe K-edge XANES in Figure 3(e), where all samples show a clear deviation from that of Fe foil, with the absorption edge intensity varying in the order of Fe foil<FeNiO-250-16<FeNiO-250-4<FeNiO_{NC}-250-4<Fe₂O₃, confirming that the Fe valence state in the three FeNiO samples was in the intermediate between those of metallic Fe and Fe^{3+} . Importantly, the Ni and Fe elements of FeNiO-250-4 can be seen to possess an average oxidation state between those of FeNiO_{NC}-250-4 and FeNiO-250-16, likely due to reduced carbothermal effects by the rapid heating and quenching process [22].

Further insights into the bonding configurations of the metal centers were obtained from the extended X-ray absorption fine structure (EXAFS) results. Fitting of the FT-EXAFS data (Figure 3(f), Figure S13, and Table S2-S4) show that FeNiO-250-4 actually possessed Ni-O bonds with a bond length of 2.03 Å, somewhat smaller than those of FeNiO_{NC}-250-4 (2.05 Å) and rock salt NiO (2.09 Å) [56]. This is consistent with the phase segregation in FeNiO_{NC}-250-4 (vide ante). Meanwhile, the Ni-Cl path in FeNiO-250-4 was found to possess a coordination number (CN) of 3 and an average bond length of 2.40 Å, slightly larger than Ni-O (CN = 2.7). Due to the low cooling rate, a severe Cl loss occurred with FeNiO_{NC}-250-4 leading to a low CN of 1.5, while Ni-O showed a CN of 4.7, consistent with the absence of nanospindles in TEM measurements (Figure 2). The profile of FeNiO-250-16 is almost identical to that of Ni foil with a main peak at 2.13 Å for the Ni-Ni path. The Fe EXAFS profile of FeNiO-250-4 (Figure 3(g)) shows three major peaks at 1.38, 1.96, and 2.63 Å, due to Fe-O, Fe-Cl, and second-shell Fe-Fe/Ni bonds, respectively. Yet, the feature of Fe-Cl diminished in both $FeNiO_{NC}$ -250-4 and FeNiO-250-16. FeNiO-250-16 displayed a shorter Fe-Fe/Ni bond length (2.57 Å) than FeNiO-250-4 (3.01 Å) and FeNiO_{NC}-250-4 (3.04Å), suggesting a possible transition from spinel structure to metallic Fe [57].

In summary, results from these characterization measurements show that prolonged heating and slow cooling facilitated the O and Cl loss for the spinel samples. Prolonged heating also promoted phase segregation of Ni into rock salt NiO or metallic form. With a deliberate control of the heating time and cooling rate, two key nonequilibrium features of the FeNiO spinel nanoparticles can be achieved, minimal FeNi phase segregation and formation of a Cl-rich surface, both critical in OER electrocatalysis (details below).

2.3. Electrocatalytic Activity. The FeNiO-250-4 sample that possessed a unique FeNi oxide spinel with Cl-rich surface nanospindles exhibited a remarkably high activity towards OER. From the polarization curves in Figure 4(a), FeNiO-250-4 reached the high current density of $100 \,\mathrm{mA \, cm^{-2}}$ in 1 M KOH at an ultralow potential of +1.49 V vs. reversible hydrogen electrode (RHE) (corresponding to an overpotential, η_{100} , of only +260 mV), in comparison to +1.54 V for FeNiO_{NC}-250-4, +1.60 V for FeNiO-250-16, and +1.55 V for commercial 20% RuO2. FeNiO-250-4 also displayed a Tafel slope of only 25 mV dec⁻¹ markedly lower than the rest of the sample series, 39 mV dec⁻¹ for FeNiO-250-16, 48 mV dec⁻¹ for FeNiO_{NC}-250-4, and 58 mV dec⁻¹ for commercial RuO₂ (Figure 4(b)). In addition, at 100% iR compensation, FeNiO-250-4 can even produce an exceedingly high current density of 1 A cm⁻² at only +1.64 V (Figure S14), which represents an unprecedentedly high activity among the leading FeNi oxide-based OER electrocatalysts reported in recent studies (Table S5 and Movie S3).

To quantify the electrochemical surface area (ECSA), voltametric measurements were also carried out in the non-faradaic region of +0.9 to +1.1 V (Figure S15), and FeNiO-250-4 can be seen to exhibit the highest ECSA of 23.06 cm^2 , as compared to 15.75 cm^2 for FeNiO_{NC}-250-4

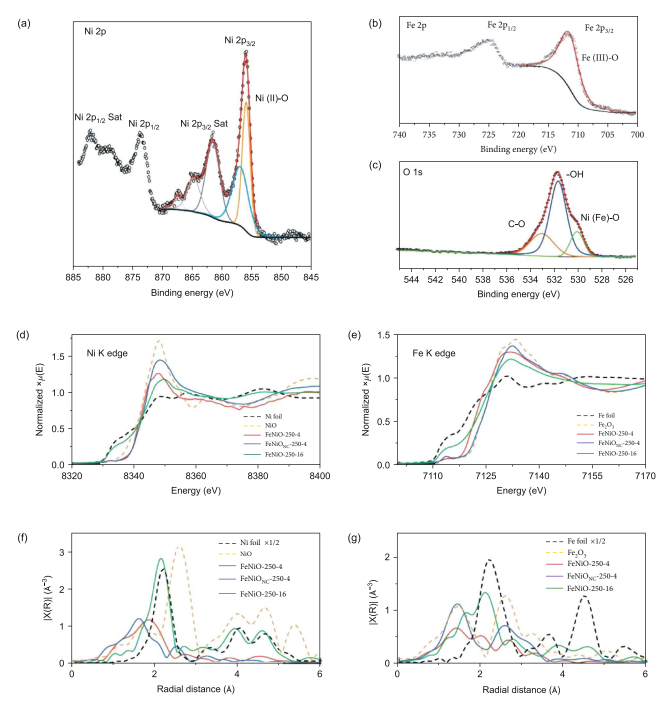


FIGURE 3: X-ray characterizations of FeNi spinel oxides. High-resolution X-ray photoelectron spectroscopy (XPS) scans of the (a) Ni 2p, (b) Fe 2p, and (c) O 1 s electrons of FeNiO-250-4. (d) Ni K-edge and (e) Fe K-edge X-ray absorption near-edge structure spectra (XANES) of FeNiO-250-4, FeNiO_{NC}-250-4, FeNiO-250-16, and reference samples (Ni/Fe foil, NiO, and Fe₂O₃), along with the corresponding Fourier transformed extended X-ray absorption fine structure spectra (FT-EXAFS) of (f) Ni and (g) Fe. Note that the Ni-Ni path (2.57 Å) in NiO is essentially absent in FeNiO-250-4 and FeNiO_{NC}-250-4, and FeNiO_{NC}-250-4 exhibits a first main peak at 1.58 Å, very close to the Ni-O bonds of the NiO reference (1.64 Å), in comparison to 1.83 Å for FeNiO-250-4, possibly because of strong interactions with Cl atoms.

and 17.06 cm² for FeNiO-250-16 (Table S6). When the OER currents were normalized to the respective ECSA, FeNiO-250-4 remained the best among the series (Figure S16a), reaching a current density of 10 mA cm^{-2} at +1.64 V. The corresponding turnover frequency (TOF) is depicted in Figure S16b at up to 0.21 s⁻¹ for FeNiO-250-4, which was

at least one order of magnitude higher than those of FeNiO $_{\rm NC}$ -250-4 and FeNiO-250-16.

In fact, the FeNiO-250-4 sample represents the optimal condition (Figure S17). It also shows excellent stability. At the applied potential of +1.53 V, over 80% of the initial current was retained even after 10 h's continuous operation

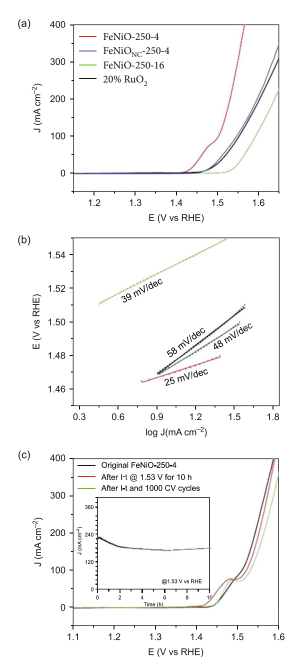


FIGURE 4: Electrochemical activities. (a) LSV curves and (b) the corresponding Tafel plots of FeNiO-250-4, FeNiO $_{\rm NC}$ -250-4, FeNiO-250-16, and 20% RuO $_2/C$ in 1 M KOH. (c) Polarization curves of FeNiO-250-4 in the first scan, after 10 h's stability tests at +1.53 V (the corresponding i-t curve is shown in the figure inset) and after additional 1000 CV cycles within the potential range of +1.20 to +1.65 V at the scan rate of 10 mV s⁻¹.

(Figure 4(c) inset), which is one of the best performances as compared to the state of the art [37]. The corresponding OER polarization curve showed an anodic shift of only 10 mV. When the electrode was subject to additional 1,000 cyclic voltammetric (CV) scans between +1.20 and +1.65 V, the subsequent polarization curve exhibited a further anodic shift of only 10 mV (Figure 4(c)).

Notably, XPS measurements of the electrocatalyst after the stability test (Figure S18) showed that the Fe:Ni ratio decreased to 1:1.8, as compared to that of the as-prepared sample (4.1:1, Table S1). This indicates that part of Fe was leached away during the prolonged OER process, which is commonly observed with relevant catalysts [37]. However, Cl remained detectable, and the relative ratio to Fe and Ni (1:3) is consistent with that of the pristine sample. Furthermore, in EDS-based elemental mapping analysis (Figure S19), a homogeneous distribution of Fe and Ni, along with Cl, can be readily observed after the stability tests; and the atomic ratio of Fe to Ni was estimated to be 1:2, consistent with that from XPS measurements. Taken together, these results suggest that the activity decay is largely due to the Fe loss instead of structural change.

2.4. Theoretical Study. To unravel the mechanistic origin of the remarkable OER activity observed above with FeNiO-250-4, slab models were built by chlorine substitution of the surface oxygen atom originally located between Fe_{oct} (octahedral site) and Ni_{td} (tetrahedral site) in $NiFe_2O_4$ (100) (Figure 5(a)), based on the structural features identified in the above experimental characterization. Free energy calculations indicate that adsorption of OH favors the Ni_{td} sites over the Fe_{oct} sites on the surface, consistent with the XPS results (Figure 3). Consequently, the NiFe₂O₄(100) surface with Cl substituting O and Ni_{td} binding an OH (Ni(OH) $Fe_2O_4(Cl)$ was used as the model catalyst (Figure 5(a)). Other structures, Ni(OH)Fe₂O₄ (Figure 5(b)), Fe(OH)Fe₂O₄(Cl) (Figure 5(c)), Fe(OH)Fe₂O₄ (Figure 5(d)), Ni(OH)NiO(Cl) (Figure 5(e)), and Ni(OH)NiO (Figure 5(f)), were constructed as comparative references.

A two-site (*-#) model was adopted to study the OER mechanism (details in the Supporting Information). From the free energy diagram in Figure 5(g), Ni(OH)Fe₂O₄(Cl) displays a thermodynamic overpotential of only 90 mV and stands out as the optimal catalyst for OER among all models. The potential-limiting step is the second OH adsorption with a reaction free energy (ΔG_2°) of 1.32 eV, while all other steps, including the first OH binding (ΔG_1°), O–O coupling (ΔG_3°) and O_2 release (ΔG_4°) , have a reaction free energy equal to or slightly lower than 1.23 eV. For comparison, Ni (OH)Fe₂O₄, with a similar structure but without Cl substitution, shows a very high reaction free energy for the O-O coupling step (ΔG_3°), whereas ΔG_1° , ΔG_2° , and ΔG_4° are all markedly below 1.23 eV. The high reaction free energy of O–O coupling accompanied by proton extraction (ΔG_3°) indicates that this step is the potential limiting step. These results suggest that incorporation of Cl onto the surface of NiFe₂O₄ spinel enhances the OER activity by facilitating O–O bond formation.

To confirm that the active site ensemble on Ni(OH) $Fe_2O_4(Cl)$ is unique and responsible for the enhanced activity, we also mapped out the potential energy profiles of OER on the monometal systems of Fe(OH)Fe₂O₄, Fe(OH) $Fe_2O_4(Cl)$, Ni(OH)NiO, and Ni(OH)NiO(Cl) (Figure 5(g)), where the O–O coupling step, with a respective reaction free energy of 2.81, 2.16, 2.20, and 2.03 eV, remains to be the potential limiting step. These results indicate that these

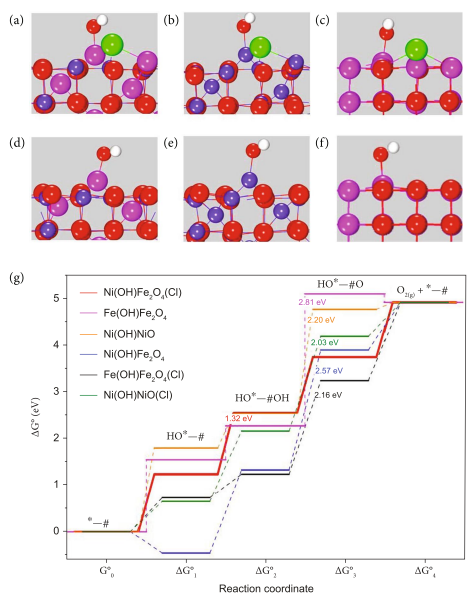


FIGURE 5: Computational studies of OER energetics. Structural models of (a) Ni(OH)Fe₂O₄(Cl), (b) Ni(OH)Fe₂O₄, (c) Fe(OH)Fe₂O₄(Cl), (d) Fe(OH)Fe₂O₄, (e) Ni(OH)NiO(Cl), and (f) Ni(OH)NiO. Free energy profile of OER on Ni(OH)Fe₂O₄, Ni(OH)Fe₂O₄(Cl), Fe(OH)Fe₂O₄, Fe (OH)Fe₂O₄(Cl), Ni(OH)NiO and Ni(OH)NiO(Cl). Color codes: green, Cl; red, O; dark blue, Fe; magenta, Ni; white, H.

monometal systems, even with Cl substitution, exhibited only a limited OER activity. Therefore, the remarkable activity of FeNiO-250-4 is most likely a synergistic effect of the formation of the metastable FeNi spinel phase and the incorporation of Cl in the surface by substituting surface oxygen atom.

To understand the enhanced OER activity in Ni(OH) $Fe_2O_4(Cl)$, we tracked the charge density differences and compared the bond distances of the O*-#OH species on $Fe(OH)Fe_2O_4$, $Fe(OH)Fe_2O_4(Cl)$, Ni(OH)Fe_2O_4, and Ni(OH)Fe_2O_4(Cl). For O*-#OH adsorbed on Fe(OH) Fe_2O_4 (Figure 6(a)), there is no observable charge density redistribution at the Fe₁ site (adjacent to Cl) but a significant electron depletion is observed on Fe_2 (away from Cl). With the introduction of Cl (Figure 6(b)), charge redistribution at Fe₁ is clearly visible whereas the electron density

redistribution at Fe₂ is minimal. The electron density redistribution is believed to stabilize the O–O species on Fe(OH) $Fe_2O_4(Cl)$ as a result of losing the proton by O-O-H. Replacing Fe with Ni (Figure 6(c)) results in a relatively uniform charge density redistribution at both Fe1 and Fe2 sites and further stabilized the O-O species on Ni(OH)Fe2O4. Therefore, the presence of Cl and Ni (Figure 6(d)) strengthens bonding between the O*-#OH species and lowers the reaction free energy of the O-O coupling step. Charge redistribution at the Fe₁ and Fe₂ sites is also reflected in part in the decrease of Bader charge of the O-O pair, which was -1.2|e|, -1.14|e|, -1.04|e|, and -1.0|e| for the O-O pair adsorbed on Fe(OH)Fe₂O₄, Fe(OH)Fe₂O₄ (Cl), Ni(OH)Fe₂O₄, and Ni(OH)Fe₂O₄(Cl), respectively. A decreased negative charge value indicates an increase of acidity of O*-#OH, which benefits the proton transfer from

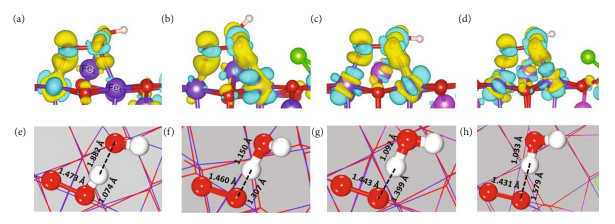


FIGURE 6: Charge density difference and evolution of bond lengths of two-site models. (a–d) Charge density difference isosurface with a value of 0.01. Light blue corresponds to negative and yellow to positive; (e–h) evolution of bond distances of HO*—#O species: (a, e) $Fe(OH)Fe_2O_4$, (b, f) $Fe(OH)Fe_2O_4(Cl)$, (c, g) $Ni(OH)Fe_2O_4$, and (d, h) $Ni(OH)Fe_2O_4(Cl)$. Fe_1 is next to Cl, and Fe_2 is away. Color codes: green, Cl; red, O; dark blue, Fe; magenta, Ni; white, H.

O*-#OH to the OH adsorbed on Nitd or Fetd (Figures 6(e)-6(h)). A complete proton transfer from HO*-#O species to the OH facilitates the O-O bond formation, and the O-H distance in HO*-#O increases from 1.074 Å on Fe $(OH)Fe_2O_4$ to 1.307 Å on Fe $(OH)Fe_2O_4(Cl)$, further to 1.399 Å on Ni(OH)Fe₂O₄, and finally to 1.579 Å on Ni $(OH)Fe_2O_4(Cl)$ (Figures 6(e)-6(h)). Correspondingly, this increasingly detached proton approaches the OH on Nitd or Fe_{td} to form H₂O. The loss of H from HO*-#O also strengthens the O-O bond, as the bond distance decreases from 1.473 Å on $Fe(OH)Fe_2O_4$ to 1.431 Å on Ni(OH) $Fe_2O_4(Cl)$. In summary, the presence of Ni and Cl in the catalyst synergistically stabilizes the O-O species while facilitates proton transfer from HO*-#O to the adjacent OH, resulting in a much reduced free energy barrier for the O-O coupling step.

From the Fe₁ density of state (DOS) plots (Figure S20), it is evident that the presence of Cl in Fe(OH)Fe₂O₄(Cl) and Ni(OH)Fe₂O₄(Cl) shifts the occupied d states closer to the Fermi level. For Ni(OH)Fe₂O₄(Cl), these states remain occupied up to the O–O coupling step (Figure S20m– S20p). The upward shift of the occupied d states results in an increased reactivity of the Cl and Ni containing catalysts.

3. Conclusions

In summary, the MIHRQ method was successfully developed and exploited for the ultrafast fabrication of metal oxide spinel nanostructures. Using NiCl₂ and FeCl₃ as the precursors, FeNi oxide spinels were obtained by heating at controlled currents within seconds and exhibited an even mixing of the Ni and Fe elements and a Cl-rich surface, in sharp contrast to samples prepared at prolonged heating and/or natural cooling in the ambient. The best sample FeNiO-250-4 needed an overpotential of only +260 mV to reach the high current density of 100 mA cm⁻² and exhibited significant stability in alkaline media. Such a remarkable activity was attributed to the unique metastable structure that facilitated the adsorption of key reaction intermediates and O-O coupling, a major limiting step in OER. Results from this study highlight the unique advantages of MIHRQ in the rapid production of unprecedented material structures that are unattainable in conventional thermal processes for enhanced electrocatalytic performance and potential applications in the structural engineering of a diverse range of materials (Figure S21).

4. Materials and Methods

4.1. Sample Preparation. Carbon paper (TGP-H-060 from Toray Industries, Inc.) was cut into $1 \text{ cm} \times 1.5 \text{ cm}$ pieces. A solution was prepared by dissolving 40 mg of NiCl₂•6H₂O, 10 mg of FeCl₃, and 0.8 g of urea into 10 mL of water (supplied with a Barnstead Nanopure Water System, 18.2 M Ω cm). 100 mL of the solution was dropcast onto the carbon paper, which was then dried at ambient temperature and sandwiched between two iron sheets of $2.5 \text{ cm} \times 2.5$ $cm \times 0.01$ cm. The assembly was then placed in the center of a four-turn induction coil with a diameter of 5 cm, and magnetic induction heating was carried out at a controlled current (X = 100 - 600 A) for a select heating time (Y = 2 - 16 s), when the sample was dropped into an ethanol-dry ice solution (-78°C) placed underneath the induction coil for rapid quenching (caution: the ethanol must be fully cooled down by dry ice, or it will catch fire). The temperature was measured with an industrial infrared laser thermometer (BTMETER, BT-1500). The corresponding sample was denoted as FeNiO-X-Y.

Control samples were also prepared by removing the sample assembly from the heater and being cooled down in ambient to room temperature. These samples were referred to as $FeNiO_{NC}$ -X-Y.

4.2. Structural Characterizations. (Scanning) transmission electron microscopy (S)TEM experiments ware conducted with a transmission electron microscope equipped with an X-FEG field-emission source, operated at 200 keV. To perform the high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) imaging and energy-dispersive X-ray spectroscopy (EDS) analysis, the samples were first sonicated, dispersed in ethanol, and then deposited onto copper grids for TEM characterization. Scanning electron microscopy (SEM) studies were carried out on FEI Quanta 3D FEG dual beam instrument. X-ray photoelectron spectroscopy (XPS) measurements were performed with a Phi 5400/XPS instrument equipped with an Al K_{α} source operated at 350 W and 10⁻⁹ Torr. X-ray diffraction (XRD) patterns were acquired with a Bruker D8 Advance diffractometer with Cu K_{α} radiation ($\lambda = 0.15418$ nm).

4.3. X-Ray Absorption Spectroscopy (XAS) Studies. Fe and Ni K-edge XAS data was collected from the CLS@APS Sector 20-BM beamline at the Advanced Photon Source (operating at 7.0 GeV) in Argonne National Labs, Chicago, IL, USA. Samples were enclosed within a Kapton tape and measured in fluorescence mode simultaneously with each element foil reference. All measurements were conducted at room temperature and ambient pressure. EXAFS data was transformed and normalized into k- and R-space using the Athena program following conventional procedures. A k weighting of 2 was used to obtain all FT-EXAFS spectra. The k-range used for each sample is as follows for Fe: $3.1-9.2 \text{ Å}^{-1}$ for FeNiO-250-4, 2.1–9.1 Å⁻¹ for FeNiO_{NC}-250-4, and 3.3–12.7 Å⁻¹ for FeNiO-250-16. For Ni, the k-range used was as follows: $3.0-8.9 \text{ Å}^{-1}$ for FeNiO-250-4, 2.9–12.2 \AA^{-1} for FeNiO_{NC}-250-4, and 2.6–14.4 \AA^{-1} for FeNiO-250-16. The R-range used for Fe is as follows: 1.0-3.7 Å for FeNiO-250-4, 1.0-3.6 Å for FeNiO_{NC}-250-4, and 1.0-3.4 Å for FeNiO-250-16. The R-range used for Ni is as follows: 1.0-3.5 Å for FeNiO-250-4, 1.0–3.0 Å for FeNiO_{NC}-250-4, and 1.0-3.0 Å for FeNiO-250-16. Self-consistent multiplescattering calculations were performed using the FEFF6 program to obtain the scattering amplitudes and phase-shift functions used to fit various scattering paths with the Artemis program. In the fitting of each sample, the E_0 values were correlated together to minimize the number of independent values, allowing reliable fitting results to be obtained. The σ^2 values were also correlated for some samples.

4.4. Electrochemistry. Electrochemical measurements were carried out with a CHI 700e electrochemical workstation in a three-electrode configuration. The prepared carbon paper was fixed onto a graphite electrode holder, with an exposed surface area of 1 cm^2 . A platinum wire was adopted as the counter electrode and a Ag/AgCl in saturated KCl as the reference electrode. The reference electrode was calibrated against a reversible hydrogen electrode (RHE), and all potentials in the present study were referenced to this RHE.

4.5. DFT computation. Spin-polarized density functional theory (DFT) calculations were carried out using the VASP (Vienna Ab-Initio Simulation Package) code [58]. Projector-augmented wave (PAW) method [59] with the Perdew-Burke-Ernzerhof (PBE) exchange-correlation functional [60] was used in all calculations. On-site Coulomb interactions were corrected within the DFT+U framework based on Dudarev's approximation [61]. U_{eff} = 4.20 [62] and 6.40 [63] for Fe and Ni, respectively, were used [64]. Plane-wave basis set with a 400 eV energy cutoff provides a

balance of accuracy and computational cost. Either quasi-Newton scheme or conjugate gradient algorithm implemented in VASP was used to relax structure until forces are converged to less than -0.03 eV Å⁻¹ on unconstrained atoms and selfconsistent convergence until 10⁻⁵ eV. The Gaussian smearing with a σ value of 0.05 was used to minimize entropy contribution to free energy. The bulk structure of NiFe₂O₄ was taken from JCPDS (JCPDS Card No. 10-0325) and optimized in (4 × 4 × 1) k-point grid sampling of the surface Brillouin zone. The optimized lattice constant of 8.37 Å was found in close agreement with the experimental value of 8.35 Å.

A supercell consisting of five layers of NiFe₂O₄ with an exposed (100) surface was constructed from the optimized bulk structure. A vacuum space of 14 Å in *z*-direction was inserted between the slabs, and the atoms in the top three layers were allowed to relax while those in the bottom two layers were fixed at the corresponding bulk position during structural optimization.

Additional details are included in the Supporting Information.

Data Availability

The data used to support the findings of this study are included within the article.

Conflicts of Interest

The authors declare that there is no conflict of interest regarding the publication of this article.

Authors' Contributions

S.W. Chen conceived the project and supervised the work. B.Z. Lu prepared the samples and carried out the majority of the experimental work. Q.M. Liu assisted in sample preparation, structural characterization, and results analysis. F. Nichols carried out the initial TEM studies. R. Mercado conducted the XPS measurements. C.Y. Wang conducted the HAADF-STEM and EDS measurements under the supervision of H.L. Xin. The XAS studies were carried out by D. Morris under the supervision of P. Zhang. DFT work was performed by Z. Masood under the supervision of Q.F. Ge. The paper was drafted by B.Z. Lu, and all authors discussed the results and commented on the manuscript. S.W. Chen finalized the paper. Bingzhang Lu and Qiming Liu contributed equally to this work.

Acknowledgments

This work was supported by grants from the National Science Foundation (CHE-1900235 and CHE-2003685, S.W.C. and CHE-1900401, H.L.X.). Part of the TEM and XPS work was carried out at the National Center for Electron Microscopy and Molecular Foundry, Lawrence Berkeley National Laboratory, which is supported by the Office of Science, Office of Basic Energy Sciences, of U.S. Department of Energy under Contract No. DE-AC02-05CH11231, as part of a user project. The XAS work used resources of the Advanced Photon Source, a User Facility operated for the U.S. Department of Energy (DOE) Office of Science by Argonne National Laboratory and was supported by the DOE under contract No. DE-AC02-06CH11357 and the Canadian Light Source and its funding partners. This research also used resources of the Center for Functional Nanomaterials (CFN), which is a U.S. Department of Energy Office of Science User Facility, at Brookhaven National Laboratory under Contract No. DE-SC0012704. The authors also thank Mr. Jeremy Barnett for the assistance in sample preparation and data acquisition of X-ray diffraction measurements in the UCSC X-ray Facility which was funded by the National Science Foundation (MRI-1126845).

Supplementary Materials

Additional computational details. Figure S1: crystal structure of FeNi oxide spinel, where blue, pink, and red indicate Fe, Ni, and O atoms, respectively. Figure S2: HAADF-STEM images of FeNi oxide spinel in FeNiO-250-4 along the [112] zone axis. Figure S3: HAADF-STEM images of FeNi oxide spinel in FeNiO-250-4 along the [100] zone axis. Figure S4: HAADF-STEM images of FeNi oxide spinel in FeNiO-250-4 along the [103] zone axis. Figure S5: TEM and electron diffraction studies. Figure S6: EDS-based elemental mapping analysis. Figure S7: EDS-based elemental mapping analysis. Figure S8: XRD patterns of the sample series. Figure S9: XPS spectrum of the Cl 2p electrons of FeNiO-250-4. Figure S10: high-resolution XPS spectra of the Ni 2p electrons of (a) FeNiO_{NC}-250-4 and (b) FeNiO-250-16. Figure S11: high-resolution XPS spectra of the Fe 2p electrons of (a) FeNiO_{NC}-250-4 and (b) FeNiO-250-16. Figure S12: high-resolution XPS scans of the O 1s electrons of (a) FeNiO_{NC}-250-4 and (b) FeNiO-250-16. Figure S13: EXAFS fitting results. Figure S14: LSV curves of FeNiO-250-4 in 1 M KOH at different levels of iR compensation. Figure S15: CV curves at difference scan rates for (a) FeNiO-250-4, (b) FeNiO_{NC}-250-4, and (c) FeNiO-250-16. (d) The variation of the double-layer charging current with potential scan rate. Figure S16: (a) LSV curves normalized to ECSA and (b) TOF curves of FeNiO-250-4, FeNiO_{NC}-250-4, and FeNiO-250-16. Figure S17: optimization of OER performance of FeNiO samples. Figure S18: highresolution XPS spectra of the (a) Ni 2p, (b) Fe 2p, (c) Cl 2p, (d) C 1 s, and (e) O 1 s electrons of FeNiO-250-4 after stability test. Figure S19: SEM image and the corresponding EDS-based elemental mapping analysis of FeNiO-250-4 after stability test. Figure S20: PDOS of the Fe1 atom. Figure S21: additional materials prepared by MIHRQ. Table S1: elemental contents of the sample series based on EDS measurements. Table S2: fitting results of the EXAFS data of FeNiO-250-4. Table S3: fitting results of the EXAFS data of FeNiO $_{
m NC}$ -250-4. Table S4: fitting results of the EXAFS data of FeNiO-250-16. Table S5: comparison with relevant FeNi oxide-based OER catalysts reported in the literature. Table S6: electrochemical surface areas of the sample series. Movie S1: magnetic induction heating at a solenoid current of 200 A for a heating time of 4 s. Movie S2: magnetic induction heating at a solenoid current of 600 A for a heating time of 4 s. Movie S3: oxygen evolution catalyzed by FeNiO-250-4. (Supplementary Materials)

References

- Z. W. Seh, J. Kibsgaard, C. F. Dickens, I. Chorkendorff, J. K. Nørskov, and T. F. Jaramillo, "Combining theory and experiment in electrocatalysis: insights into materials design," *Science*, vol. 355, no. 6321, 2017.
- [2] M. Shao, Q. Chang, J.-P. Dodelet, and R. Chenitz, "Recent advances in electrocatalysts for oxygen reduction reaction," *Chemical Reviews*, vol. 116, no. 6, pp. 3594–3657, 2016.
- [3] J. Zhu, L. Hu, P. Zhao, L. Y. S. Lee, and K. Y. Wong, "Recent advances in electrocatalytic hydrogen evolution using nanoparticles," *Chemical Reviews*, vol. 120, no. 2, pp. 851–918, 2020.
- [4] N.-T. Suen, S.-F. Hung, Q. Quan, N. Zhang, Y. J. Xu, and H. M. Chen, "Electrocatalysis for the oxygen evolution reaction: recent development and future perspectives," *Chemical Society Reviews*, vol. 46, no. 2, pp. 337–365, 2017.
- [5] Y. Peng, B. Lu, and S. Chen, "Carbon-supported single atom catalysts for electrochemical energy conversion and storage," *Advanced Materials*, vol. 30, no. 48, p. 1801995, 2018.
- [6] B. Lu, Q. Liu, and S. Chen, "Electrocatalysis of single-atom sites: impacts of atomic coordination," ACS Catalysis, vol. 10, no. 14, pp. 7584–7618, 2020.
- [7] C. Wang, N. M. Markovic, and V. R. Stamenkovic, "Advanced platinum alloy electrocatalysts for the oxygen reduction reaction," ACS Catalysis, vol. 2, no. 5, pp. 891–898, 2012.
- [8] S. Anantharaj, S. R. Ede, K. Sakthikumar, K. Karthick, S. Mishra, and S. Kundu, "Recent trends and perspectives in electrochemical water splitting with an emphasis on sulfide, selenide, and phosphide catalysts of Fe, Co, and Ni: a review," ACS Catalysis, vol. 6, no. 12, pp. 8069–8097, 2016.
- [9] Y. L. Zhu, Q. Lin, Y. J. Zhong, H. A. Tahini, Z. Shao, and H. Wang, "Metal oxide-based materials as an emerging family of hydrogen evolution electrocatalysts," *Energy & Environmental Science*, vol. 13, no. 10, pp. 3361–3392, 2020.
- [10] C. G. Kuai, Y. Zhang, D. Y. Wu et al., "Fully oxidized Ni-Fe layered double hydroxide with 100% exposed active sites for catalyzing oxygen evolution reaction," ACS Catalysis, vol. 9, no. 7, pp. 6027–6032, 2019.
- [11] M. R. Gao, W. C. Sheng, Z. B. Zhuang et al., "Efficient water oxidation using nanostructured alpha-nickel-hydroxide as an electrocatalyst," *Journal of the American Chemical Society*, vol. 136, no. 19, pp. 7077–7084, 2014.
- [12] R. D. Smith, M. S. Prévot, R. D. Fagan, S. Trudel, and C. P. Berlinguette, "Water oxidation catalysis: electrocatalytic response to metal stoichiometry in amorphous metal oxide films containing iron, cobalt, and nickel," *Journal of the American Chemical Society*, vol. 135, no. 31, pp. 11580– 11586, 2013.
- [13] B. Lu, Q. Liu, F. Nichols et al., "Oxygen reduction reaction catalyzed by carbon-supported platinum few-atom clusters: significant enhancement by doping of atomic cobalt," *Research*, vol. 2020, article 9167829, 2020.
- [14] B. Lu, T. J. Smart, D. Qin et al., "Nitrogen and iron-codoped carbon hollow nanotubules as high-performance catalysts toward oxygen reduction reaction: a combined experimental and theoretical study," *Chemistry of Materials*, vol. 29, no. 13, pp. 5617–5628, 2017.
- [15] M. Gong, Y. Li, H. Wang et al., "An advanced Ni-Fe layered double hydroxide electrocatalyst for water oxidation," *Journal* of the American Chemical Society, vol. 135, no. 23, pp. 8452– 8455, 2013.

- [16] Q. Xiang, F. Li, W. Chen et al., "In situ vertical growth of Fe–Ni layered double-hydroxide arrays on Fe–Ni alloy foil: interfacial layer enhanced electrocatalyst with small overpotential for oxygen evolution reaction," ACS Energy Letters, vol. 3, no. 10, pp. 2357–2365, 2018.
- [17] Q. Zhao, Z. H. Yan, C. C. Chen, and J. Chen, "Spinels: controlled preparation, oxygen reduction/evolution reaction application, and beyond," *Chemical Reviews*, vol. 117, no. 15, pp. 10121–10211, 2017.
- [18] R. Jiang, Y. Da, X. Han, Y. Chen, Y. Deng, and W. Hu, "Ultrafast synthesis for functional nanomaterials," *Cell Reports Physical Science*, vol. 2, no. 1, article 100302, 2021.
- [19] A. Holewinski, J.-C. Idrobo, and S. Linic, "High-performance Ag–Co alloy catalysts for electrochemical oxygen reduction," *Nature Chemistry*, vol. 6, no. 9, pp. 828–834, 2014.
- [20] Z. Li, J.-Y. Fu, Y. Feng, C. K. Dong, H. Liu, and X. W. du, "A silver catalyst activated by stacking faults for the hydrogen evolution reaction," *Nature Catalysis*, vol. 2, no. 12, pp. 1107–1114, 2019.
- [21] J.-Q. Wang, C. Xi, M. Wang et al., "Laser-generated grain boundaries in ruthenium nanoparticles for boosting oxygen evolution reaction," ACS Catalysis, vol. 10, no. 21, pp. 12575–12581, 2020.
- [22] Y. Yao, Z. Huang, P. Xie et al., "Carbothermal shock synthesis of high-entropy-alloy nanoparticles," *Science*, vol. 359, no. 6383, pp. 1489–1494, 2018.
- [23] W. Chen, Z. Wang, K. V. Bets et al., "Millisecond conversion of metastable 2D materials by flash joule heating," ACS Nano, vol. 15, no. 1, pp. 1282–1290, 2021.
- [24] J. Zhang, M. Ren, Y. Li, and J. M. Tour, "In situ synthesis of efficient water oxidation catalysts in laser-induced graphene," ACS Energy Letters, vol. 3, no. 3, pp. 677–683, 2018.
- [25] X. Yue, X. P. Qin, Y. D. Chen et al., "Constructing active sites from atomic-scale geometrical engineering in spinel oxide solid solutions for efficient and robust oxygen evolution reaction electrocatalysts," *Advanced Science*, vol. 8, no. 17, article 2101653, 2021.
- [26] C. Meng, M. Lin, X. Sun et al., "Laser synthesis of oxygen vacancy-modified CoOOH for highly efficient oxygen evolution," *Chemical Communications*, vol. 55, no. 20, pp. 2904– 2907, 2019.
- [27] H. Hu, Q. Li, L. Q. Li et al., "Laser irradiation of electrode materials for energy storage and conversion," *Matter*, vol. 3, no. 1, pp. 95–126, 2020.
- [28] K. M. Wyss, D. X. Luong, and J. M. Tour, "Large-scale syntheses of 2D materials: flash joule heating and other methods," *Advanced Materials*, vol. 34, no. 8, article 2106970, 2022.
- [29] F. Ran and S. Chen, Advanced Nanomaterials for Electrochemical-Based Energy Conversion and Storage, Elsevier, 2019.
- [30] Y. Zhou, S. N. Sun, C. Wei et al., "Significance of engineering the octahedral units to promote the oxygen evolution reaction of spinel oxides," *Advanced Materials*, vol. 31, no. 41, article 1902509, 2019.
- [31] X. K. Chen, X. H. Zhang, L. Z. Zhuang et al., "Multiple vacancies on (111) facets of single-crystal NiFe₂O₄ spinel boost electrocatalytic oxygen evolution reaction," *Chemistry, an Asian Journal*, vol. 15, no. 23, pp. 3995–3999, 2020.
- [32] B.-Q. Li, S.-Y. Zhang, C. Tang, X. Cui, and Q. Zhang, "Anionic regulated NiFe (oxy)sulfide electrocatalysts for water oxidation," *Small*, vol. 13, no. 25, p. 1700610, 2017.

- [33] X.-M. Liu, X. Cui, K. Dastafkan et al., "Recent advances in spinel-type electrocatalysts for bifunctional oxygen reduction and oxygen evolution reactions," *Journal of Energy Chemistry*, vol. 53, pp. 290–302, 2021.
- [34] C. Tang, H.-S. Wang, H.-F. Wang et al., "Spatially confined hybridization of nanometer-sized NiFe hydroxides into nitrogen-doped graphene frameworks leading to superior oxygen evolution reactivity," *Advanced Materials*, vol. 27, no. 30, pp. 4516–4522, 2015.
- [35] B. Lal and P. K. Rastogi, "Microwave assisted synthesis of chromium substituted nickel ferrite spinel for oxygen evolution reaction," *Orbital-the Electronic Journal of Chemistry*, vol. 12, no. 3, pp. 154–159, 2020.
- [36] D. Lim, H. Kong, N. Kim, C. Lim, W. S. Ahn, and S. H. Baeck, "Oxygen-deficient NiFe₂O₄Spinel nanoparticles as an enhanced electrocatalyst for the oxygen evolution reaction," *ChemNanoMat*, vol. 5, no. 10, pp. 1296–1302, 2019.
- [37] C. Kuai, Z. Xu, C. Xi et al., "Phase segregation reversibility in mixed-metal hydroxide water oxidation catalysts," *Nature Catalysis*, vol. 3, no. 9, pp. 743–753, 2020.
- [38] Y. Zhu, Q. Lin, Z. Wang et al., "Chlorine-anion doping induced multi-factor optimization in perovskties for boosting intrinsic oxygen evolution," *Journal of Energy Chemistry*, vol. 52, pp. 115–120, 2021.
- [39] J. Zhang, Y. Cui, L. Jia, B. He, K. Zhang, and L. Zhao, "Engineering anion defect in LaFeO_{2.85}Cl_{0.15} perovskite for boosting oxygen evolution reaction," *International Journal of Hydrogen Energy*, vol. 44, no. 44, pp. 24077–24085, 2019.
- [40] B. M. Hunter, W. Hieringer, J. R. Winkler, H. B. Gray, and A. M. Müller, "Effect of interlayer anions on [NiFe]-LDH nanosheet water oxidation activity," *Energy & Environmental Science*, vol. 9, no. 5, pp. 1734–1743, 2016.
- [41] V. Rudnev, D. Loveless, and R. L. Cook, Handbook of induction heating, CRC press, 2017.
- [42] T. Zhao, C. Xu, W. Ma et al., "Ultrafast growth of nanocrystalline graphene films by quenching and grain-size-dependent strength and bandgap opening," *Nature Communications*, vol. 10, no. 1, p. 4854, 2019.
- [43] C. Wu, F. Li, W. Chen, C. P. Veeramalai, P. C. Ooi, and T. Guo, "Electromagnetic induction heating for single crystal graphene growth: morphology control by rapid heating and quenching," *Scientific Reports*, vol. 5, no. 1, p. 9034, 2015.
- [44] R. Piner, H. Li, X. Kong et al., "Graphene synthesis via magnetic inductive heating of copper substrates," ACS Nano, vol. 7, no. 9, pp. 7495–7499, 2013.
- [45] M. Seifert, S. Drieschner, B. M. Blaschke, L. H. Hess, and J. A. Garrido, "Induction heating-assisted repeated growth and electrochemical transfer of graphene on millimeter-thick metal substrates," *Diamond and Related Materials*, vol. 47, pp. 46–52, 2014.
- [46] P. Sharma, N. Holliger, P. H. Pfromm, B. Liu, and V. Chikan, "Size-controlled synthesis of iron and iron oxide nanoparticles by the rapid inductive heating method," ACS Omega, vol. 5, no. 31, pp. 19853–19860, 2020.
- [47] X. B. Xiong, J. Y. Zhang, J. Ma, X. R. Zeng, H. Qian, and Y. Y. Li, "Fabrication of porous nickel (hydr)oxide film with rational pore size distribution on nickel foam by induction heating deposition for high- performance supercapacitors," *Materials Chemistry and Physics*, vol. 181, pp. 1–6, 2016.

- [48] R. Tiwari, M. De, H. S. Tewari, and S. K. Ghoshal, "Structural and magnetic properties of tailored NiFe₂O₄ nanostructures synthesized using auto-combustion method," *Results in Physics*, vol. 16, article 102916, 2020.
- [49] W. Niu, L. Li, X. Liu et al., "Mesoporous N-doped carbons prepared with thermally removable nanoparticle templates: an efficient electrocatalyst for oxygen reduction reaction," *Journal* of the American Chemical Society, vol. 137, no. 16, pp. 5555– 5562, 2015.
- [50] C. Bale, P. Chartrand, S. A. Degterov et al., "FactSage thermochemical software and databases," *Calphad - Computer Coupling of Phase Diagrams and Thermochemistry*, vol. 26, no. 2, pp. 189–228, 2002.
- [51] A. G. Marrani, V. Novelli, S. Sheehan, D. P. Dowling, and D. Dini, "Probing the redox states at the surface of electroactive nanoporous NiO thin films," ACS Applied Materials & Interfaces, vol. 6, no. 1, pp. 143–152, 2014.
- [52] A. P. Grosvenor, M. C. Biesinger, R. S. C. Smart, and N. S. McIntyre, "New interpretations of XPS spectra of nickel metal and oxides," *Surface Science*, vol. 600, no. 9, pp. 1771–1779, 2006.
- [53] T. Yamashita and P. Hayes, "Analysis of XPS spectra of Fe²⁺ and Fe³⁺ ions in oxide materials," *Applied Surface Science*, vol. 254, no. 8, pp. 2441–2449, 2008.
- [54] M. A. Peck and M. A. Langell, "Comparison of nanoscaled and bulk NiO structural and environmental characteristics by XRD, XAFS, and XPS," *Chemistry of Materials*, vol. 24, no. 23, pp. 4483–4490, 2012.
- [55] L. Liu, F. Tian, X. Wang, Z. Yang, and X. Wang, "Influence of NiCl₂ modification on the electrochemical performance of LiV3O8 cathode for lithium ion batteries," *Ionics*, vol. 19, no. 1, pp. 9–15, 2013.
- [56] C. Lin, Y. H. Zhao, H. J. Zhang et al., "Accelerated active phase transformation of NiO powered by Pt single atoms for enhanced oxygen evolution reaction," *Chemical Science*, vol. 9, no. 33, pp. 6803–6812, 2018.
- [57] M. H. Nilsen, C. Nordhei, A. L. Ramstad, D. G. Nicholson, M. Poliakoff, and A. Cabañas, "XAS (XANES and EXAFS) investigations of nanoparticulate ferrites synthesized continuously in near critical and supercritical water," *Journal of Physical Chemistry C*, vol. 111, no. 17, pp. 6252–6262, 2007.
- [58] G. Kresse and J. Furthmüller, "Efficient iterative schemes ab initio total-energy calculations using a plane-wave basis set," *Physical Review B*, vol. 54, no. 16, pp. 11169–11186, 1996.
- [59] G. Kresse and D. Joubert, "From ultrasoft pseudopotentials to the projector augmented-wave method," *Physical Review B*, vol. 59, no. 3, pp. 1758–1775, 1999.
- [60] J. P. Perdew, K. Burke, and M. Ernzerhof, "Generalized gradient approximation made simple," *Physical Review Letters*, vol. 77, no. 18, pp. 3865–3868, 1996.
- [61] S. Dudarev, G. Botton, S. Savrasov, C. J. Humphreys, and A. P. Sutton, "Electron-energy-loss spectra and the structural stability of nickel oxide: an LSDA+ U study," *Physical Review B*, vol. 57, no. 3, pp. 1505–1509, 1998.
- [62] G. Hautier, S. P. Ong, A. Jain, C. J. Moore, and G. Ceder, "Accuracy of density functional theory in predicting formation energies of ternary oxides from binary oxides and its implication on phase stability," *Physical Review B*, vol. 85, no. 15, article 155208, 2012.

- [63] L. Wang, T. Maxisch, and G. Ceder, "Oxidation energies of transition metal oxides within the GGA+U framework," *Physical Review B*, vol. 73, no. 19, article 195107, 2006.
- [64] L. Gao, X. Cui, Z. Wang et al., "Operando unraveling photothermal-promoted dynamic active-sites generation in NiFe₂O₄ for markedly enhanced oxygen evolution," *Proceedings of the National Academy of Sciences*, vol. 118, no. 7, article e2023421118, 2021.