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Introduction

Electrochemical water splitting, a combination of two half-cell reactions consisting of the hydrogen evolution reaction at the cathode and the oxygen evolution reaction (OER) at the anode, provides an attractive path in various energy conversion¹⁻⁵ and storage systems.^{6,7} However, electrochemical water splitting usually suffers from significant efficiency loss and high overpotential in the oxidative half-cell reaction. The OER is complex and sluggish because the generation of one oxygen molecule involves the transfer of $4e^-$ and removal of $4H^+$ from water, which hinders practical application of overall water splitting.⁸⁻¹⁰ To date, the most efficient OER catalysts have been IrO_2 and RuO_2 ;¹¹⁻¹³ however, the scarcity of these noble metals has further hindered practical application of water splitting on

Phase segregated Cu_{2-x} Se/Ni₃Se₄ bimetallic selenide nanocrystals formed through the cation exchange reaction for active water oxidation precatalysts[†]

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Control over the composition and nanostructure of solid electrocatalysts is quite important for drastic improvement of their performance. The cation exchange reaction of nanocrystals (NCs) has been reported as the way to provide metastable crystal structures and complicated functional nanostructures that are not accessible by conventional synthetic methods. Herein we demonstrate the cation exchange-derived formation of metastable spinel Ni₃Se₄ NCs (sp-Ni₃Se₄) and phase segregated berzelianite Cu_{2-x}Se (ber-Cu_{2-x}Se)/sp-Ni₃Se₄ heterostructured NCs as active oxygen evolution reaction (OER) catalysts. A rare sp-Ni₃Se₄ phase was formed by cation exchange of ber-Cu_{2-x}Se NCs with Ni²⁺ ions, because both phases have the face-centered cubic (fcc) Se anion sublattice. Tuning the Ni : Cu molar ratio leads to the formation of Janus-type ber-Cu_{2-x}Se/sp-Ni₃Se₄ heterostructures exhibited high catalytic activities in the OER with small overpotentials of 250 and 230 mV at 10 mA cm⁻² in 0.1 M KOH, respectively. They were electrochemically oxidized during the OER to give hydroxides as the real active species. We anticipate that the cation exchange reaction could have enormous potential for the creation of novel heterostructured NCs showing superior catalytic performance.

a large scale. Therefore, it is imperative to develop an earthabundant and cost-effective OER catalyst with excellent electrocatalytic activity to make water splitting a viable energy conversion system.

Transition metal-based chalcogenides represent an alternative to the use of noble metals in water electrolysis.¹⁴ They exhibit diverse properties that depend on their composition and crystalline structures, leading to enhanced catalytic activity in the water splitting reaction. In particular, Ni-based electrocatalysts show great potential as alternative catalysts for the OER due to their unique 3d electron number and special e_o orbitals.15,16 Recently, catalysts of the nickel selenide group have gained attention because of their earth-abundance and high activity for water splitting. They have various crystal structures, such as hexagonal NiSe,¹⁷ cubic NiSe₂ (ref. 18-20) and rhombohedral Ni_3Se_2 (ref. 21), mediated by the slight difference in electronegativity between Ni and Se. Although nickel selenide systems have shown a favorably low onset potential and overpotential for the OER in alkaline media, they have practical limitations associated with their bulk-like morphologies. In this context, catalytic systems based on nanocrystals (NCs) are highly promising because of their large specific surface areas. In addition, homogeneous dispersions of catalyst NCs in a solvent

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can be used to modify various electrodes and photocatalysts by simple deposition or coating methods.^{22–24} However, it remains challenging to synthesize uniformly structured nickel selenide NCs with high electrocatalytic activity.

One of the ultimate goals of functional NC synthesis is to access arbitrarily defined structures with desired compositions. The cation exchange reaction has become a growing area in NC research, because the substitution of the original cations with other species enables us to synthesize metastable NCs that are not accessible by conventional one-pot synthesis.²⁵⁻²⁹ Furthermore, if specific procedures are employed in cation exchange reactions, the residual cations can be removed selectively and/ or partially, representing a new strategy for accessing structurally novel and multicomponent NCs. Recently, cation exchange reactions of Sn⁴⁺, Sn²⁺, In³⁺, Cd²⁺, Zn²⁺, Hg²⁺ and Pb²⁺ cations, using parent Cu_{2-x}Se NCs, have been developed.^{26,30-32} The exchange reactions of these cations give rise to ternary alloys with distinct nanoheterostructures, including core@shell, sandwich and Janus structures, making them potentially useful materials for solar energy harvesting and catalysis.33,34

Here we report the synthesis of a monodisperse spinel-type nickel selenide (sp-Ni₃Se₄) NC based OER electrocatalyst using the cation exchange reaction. In this work, we successfully replaced Cu^+ of berzelianite copper selenide (ber- $Cu_{2-x}Se$) as starting NCs with Ni²⁺ (Scheme 1). The formation of rare and metastable sp-Ni₃Se₄ is feasible because both ber-Cu_{2-x}Se and sp-Ni₃Se₄ phases contain an fcc Se anion sublattice, which is retained during the cation exchange reaction. We also achieved fine tuning of the Ni/Cu molar ratio in the NCs, resulting in the formation of partially cation-exchanged NCs with a Janus-type heterostructure composed of ber-Cu2-xSe and sp-Ni3Se4 domains arising from the large lattice mismatch, anisotropic reaction process and immiscibility of the two phases. These Nication-exchanged ber-Cu2-xSe NCs showed efficient electrocatalytic water oxidation activity. Specifically, the heterostructured ber-Cu_{2-x}Se/sp-Ni₃Se₄ NCs exhibited a lower overpotential (230 mV(a)10 mA cm⁻² in 0.1 M KOH) than pure sp-Ni₃Se₄ NCs. The characterization of NCs after the OER revealed that the NCs were electrochemically oxidized to hydroxides to serve as the real active species. Thus, these selenide NCs were found to behave as precatalysts for efficient water oxidation. Our work not only provides synthetic pathways to these novel-phased NCs but also supplements the existing modification methods for constructing high-performance catalysts.

Results and discussion

To obtain nickel selenide NCs by the cation exchange reaction, we chose ber- Cu_{2-x} Se NCs as the parent material. The ber-



Scheme 1 Schematic of the cation exchange reaction of $ber-Cu_{2-x}Se$ NCs with Ni cations.

 Cu_{2-x} Se NCs were synthesized using the procedure reported by Shen et al. with minor modifications (see the ESI for details[†]).³⁵ A transmission electron microscope (TEM) image of the resulting NCs shows the formation of monodisperse and cuboctahedral NCs with a size of 16.2 \pm 1.4 nm (Fig. 1a). The Xray diffraction (XRD) pattern of the NCs was well matched with that of the ber- Cu_{2-x} Se crystal structure (JCPDS 00-006-0680, Fig. 1e). The elemental analysis of the NCs using X-ray fluorescence spectroscopy (XRF) gave a Cu: Se molar ratio of 64:36, indicating the sub-stoichiometry of Cu in the ber- Cu_{2-x} Se NCs (x = 0.22). These ber- Cu_{2-x} Se NCs exhibited an intense near-infrared (NIR) absorption peak at around 1000 nm related to localized surface plasmon resonance (LSPR) arising from the Cu deficiency in their composition (Fig. S1[†]).³⁶ A high density of Cu vacancies in sub-stoichiometric copper chalcogenide NCs is generally expected to enhance the cation exchange reaction by vacancy diffusion.37

Then, the as-synthesized ber-Cu_{2-x}Se NCs underwent cation exchange with Ni acetate in the presence of tri-*n*-octylphosphine (TOP) at 140 °C. For Cu⁺ \rightarrow Ni²⁺ exchange, we used half the amount of Ni²⁺ relative to Cu⁺ in the ber-Cu_{2-x}Se NCs, as the reaction involves the replacement of two Cu⁺ cations with one Ni²⁺. In the ber-Cu_{2-x}Se NCs, Cu⁺ (chemical hardness: $\eta = 6.3$ eV) can be easily exchanged with harder metal cations when employing soft Lewis bases like TOP ($\eta = 6$ eV) to promote Cu⁺



Fig. 1 (a and c) TEM and (b and d) HRTEM images of (a and b) ber-Cu_{2-x}Se NCs and (c and d) sp-Ni₃Se₄ NCs. (e) XRD patterns of ber-Cu_{2-x}Se NCs and sp-Ni₃Se₄ NCs. The reference XRD pattern shows sp-Ni₃Se₄ (JCPDS 18-0889).

dissolution into the solvent.³⁸ Finally, we obtained monodisperse NCs of 11.7 \pm 1.3 nm size, *i.e.*, smaller than the original ber-Cu_{2-x}Se NCs (Fig. 1c).

When we monitored the change in the molar ratio of Cu, Ni and Se in the NCs during the cation exchange reaction by XRF, it was found that the cation exchange proceeded quite rapidly and was completed within 15 min (Table S1 and Fig. S2⁺). The rapid replacement of Cu⁺ with Ni²⁺ was also confirmed from the UVvis-NIR absorption spectra, which showed that the LSPR peak arising from the ber-Cu_{2-x}Se phase completely disappeared within 15 min (Fig. S1[†]). In the XRD analyses, the diffraction pattern of ber-Cu_{2-x}Se NCs also completely disappeared within 15 min, while a new set of peaks assignable to the sp-Ni₃Se₄ structure emerged (Fig. 1e). This sp-Ni₃Se₄ phase was maintained when prolonging the reaction time up to 60 min (Fig. S3[†]). In the high-resolution TEM (HRTEM) images in Fig. 1b and d, the ber- Cu_{2-x} Se NC seeds show a lattice spacing of 1.70 Å, which is consistent with the (311) lattice spacing of ber- Cu_{2-x} Se, while the cation-exchanged NCs exhibit a lattice spacing of 2.48 Å, which can be assigned to the (400) family of planes of sp-Ni₃Se₄ (additional HRTEM images of sp-Ni₃Se₄ NCs are shown in Fig. S4[†]).

Sp-Ni₃Se₄ was first discovered in 1964 as a natural mineral in Finland and named trüstedtite.³⁹ The crystal structure resembles that of polydymite, spinel Ni₃S₄, with a larger lattice constant (sp-Ni₃Se₄: a = 9.94 Å and Ni₃S₄: a = 9.48 Å, Table S2 and Fig. S5†). Although many researchers have reported the synthesis of nickel selenide crystals, most of them were rhombohedral Ni₃Se₂,^{40,41} hexagonal NiSe,^{42,43} monoclinic Ni₃Se₄ (ref. 44 and 45) or cubic NiSe₂ (ref. 46 and 47) (Table S3†), which are thermodynamically stable phases in the phase diagram of the nickel selenide series at RT.⁴⁸ On the other hand, spinel Ni₃Se₄ has been artificially obtained only under extreme conditions.⁴⁹

We compared the crystal structure of ber-Cu_{2-x}Se and sp- Ni_3Se_4 . Ber-Cu_{2-x}Se has a lattice constant of 5.69 Å and two kinds of Cu sites: Cu1 occupying every tetrahedral coordination site and Cu2 occupying nontetrahedral coordination sites. On the other hand, sp-Ni₃Se₄ also contains two kinds of Ni sites. While the first of these, Ni1, occupies one-eighth of tetrahedral coordination sites, the other, Ni2, occupies half of the octahedral coordination sites (Table S2[†]). Focusing on the Se arrangement, ber-Cu_{2-x}Se has an fcc Se anionic sublattice with a lattice constant of 5.69 Å (same as that of a unit cell), while sp-Ni₃Se₄ has a distorted quasi-fcc Se sublattice with a lattice constant of ~4.97 Å on average (half that of a unit cell) as shown in Fig. 2b and S6.[†] This indicates that the fcc Se sublattice is maintained with a slight distortion during the cation exchange process. In a fcc Se unit cell, sp-Ni₃Se₄ has one Ni1 and two Ni2 sites on average, indicating that one-eighth of Cu1 sites are replaced with Ni cations, and seven-eighth of Cu1 sites and all of Cu2 sites become empty after cation exchange (Table S4[†]). Further Ni cations newly occupy the octahedral Ni2 sites (Fig. S7[†]). As a typical spinel structure, Ni2 is located at only a half of the octahedral sites in the lattice, which disproportionates the size of Se octahedral units (Fig. S8[†]). As a result, forceful insertion of Ni2 sites induces the local distortion of the



Fig. 2 (a) Perspective images of ber-Cu_{2-x}Se (left, $2 \times 2 \times 2 = 8$ unit cells) and sp-Ni₃Se₄ (right, 1 unit cell) along the <110> zone axis. Cu1, Cu2, Ni1 and Ni2 sites are coloured in dark blue, light blue, purple and orange, respectively. (b) Se sublattice (pseudo) unit cell of ber-Cu_{2-x}Se and sp-Ni₃Se₄. Only Cu1 sites in ber-Cu_{2-x}Se are shown for simplification. Black dashed lines of sp-Ni₃Se₄ highlight the size of the sp-Ni₃Se₄ unit cell. (c) Se pseudo-unit cell of a common nickel selenide system.

Se configuration in perfect cubic ber- Cu_{2-x} Se, while the comprehensive fcc Se sublattice is preserved.

Replacement of Cu⁺ with half the amount of smaller Ni²⁺ to transform the ber-Cu_{2-x}Se lattice into the sp-Ni₃Se₄ lattice requires a volume shrinkage of 33%. Based on the comparative TEM images, the actual volume shrinkage of the NCs was calculated to be 48%. This further volume reduction can be explained by etching with TOP and OLAm during the reaction, because the size of the ber-Cu_{2-x}Se NCs decreased from 16 \pm 1.3 nm to 10 \pm 1.6 nm in the reaction even without Ni acetate. (Fig. S9†).

According to a few number of reports of sp-Ni₃Se₄ synthesis, we speculate that sp-Ni₃Se₄ is metastable. To better understand the stability of the nickel selenide phases, we monitored the transition of the crystal phases under annealing in an Ar atmosphere (Fig. S10†). Although the spinel structure of the Ni₃Se₄ NCs was retained at 100 °C, annealing at >200 °C caused the phase transition to the hexagonal NiSe phase, which confirms that sp-Ni₃Se₄ is the less stable nickel selenide species.

To reveal the reason why such a rare and metastable sp- Ni_3Se_4 phase formed, we compared the Se sublattice structure with other common Ni selenides (Fig. 2c). Rhombohedral

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 Ni_3Se_2 and hexagonal NiSe have bcc and hcp Se sublattices, respectively. Although monoclinic Ni_3Se_4 has the same stoichiometry as sp- Ni_3Se_4 , its Se sublattice has an hcp configuration with a small distortion. Cubic $NiSe_2$ has an fcc conformation; however, it is constructed from Se–Se dimer components, which is totally different from the usual fcc Se sublattice. This comparison shows that only sp- Ni_3Se_4 has an fcc Se sublattice which is the same as that of ber- $Cu_{2-x}Se$. This indicates that Se sublattice preservation preferentially occurs during cation exchange rather than the formation of a more thermodynamically preferable structure, even if the product (sp- Ni_3Se_4) is less stable.

Binary transition metal selenides often exceed monometallic selenides in OER activity due to the synergetic effect between two metals. The cation exchange method can facilely provide NCs with bimetallic composition at the intermediate stage of the reaction. However, in general, the cation exchange reaction proceeds too fast to obtain such partially exchanged NCs. Because the stability constant of Ni(acac)₂ (log K = 6.1) is larger than that of Ni acetate (log K = 1.8),⁵⁰ using Ni(acac)₂ as a Ni²⁺ precursor was expected to slow down the cation exchange kinetics. TEM images of the NCs obtained by cation exchange using Ni(acac)₂ at different reaction times showed monodisperse 11 nm NCs, which is similar to the case of Ni acetate. The Ni/Cu molar ratio during the reaction increased much more slowly than the case of Ni acetate as expected (Table S5 and Fig. S11[†]). Thanks to this slow reaction, we succeeded in obtaining partially cation-exchanged NCs with tunable Ni : Cu molar ratios from 10:50 to 40:20 (Table S5 and Fig. S11†). Fig. 3e shows the XRD patterns of partially cation-exchanged NCs after different reaction times. The patterns show a combination of the ber-Cu_{2-x}Se and the sp-Ni₃Se₄ phases without other Cu_xNi_ySe_z-related solid solution phases. As the reaction



proceeded, the peak intensities of the sp-Ni₃Se₄ phase increased, whereas those of ber-Cu_{2-x}Se decreased and almost disappeared at 60 min. This slow cation exchange also induced the gradual disappearance of LSPR absorption of the ber-Cu_{2-x}Se phase (Fig. S12†).

Scanning TEM (STEM)–energy-dispersive X-ray spectroscopy (EDS) elemental mapping was carried out to reveal the spatial distributions of the elements Cu, Ni and Se in the partially cation-exchanged NCs (Fig. 4a–d). Cu and Ni were concentrated at specific regions in each individual NC, while Se was distributed throughout each NC. This means that the ber-Cu_{2-x}Se and sp-Ni₃Se₄ phases were anisotropically segregated to form Janustype heterostructured NCs (HNCs), which is consistent with the XRD results. In the HRTEM image of partially exchanged NCs (Fig. 4e), we can see a clear boundary between domains with lattice spacings of ber-Cu_{2-x}Se ($\{002\}$, 2.86 Å) and sp-Ni₃Se₄



Fig. 3 (a–d) TEM images of ber- $Cu_{2-x}Se/sp-Ni_3Se_4$ HNCs at different cation exchange reaction times of (a) 15 min, (b) 30 min, (c) 45 min and (d) 60 min. (e) XRD patterns of ber- $Cu_{2-x}Se/sp-Ni_3Se_4$ HNCs at different cation exchange reaction times. The red-star symbols and reference XRD pattern indicate ber- $Cu_{2-x}Se$ and sp-Ni₃Se₄ (JCPDS 18-0889), respectively.

Fig. 4 (a–d) STEM-EDS elemental mapping for each element and (e) HRTEM image of ber-Cu_{2-x}Se/sp-Ni₃Se₄ HNCs (Ni : Cu : Se = 22 : 37 : 41 mol% in XRF). (f) Schematic atomic arrangement of the heterointerface between ber-Cu_{2-x}Se and sp-Ni₃Se₄ phases along with the [1–10] zone axis. Cu1, Ni1 and Ni2 sites are coloured in dark blue, purple and orange, respectively.

({004}, 2.48 Å). The fact that these two planes are parallel indicates the epitaxial relationship between ber-Cu_{2-x}Se and sp-Ni₃Se₄ with a large lattice mismatch of ~13% (Fig. 4f). Additional HRTEM images (Fig. S13†) also exhibit an epitaxial relationship between two phases. These observations indicate that these two phases share a continuous Se sublattice and the Se framework is maintained during the cation exchange reaction.

The formation of Janus-type HNCs through the cation exchange of copper chalcogenide NCs has been recently reported.⁵¹ Typical examples are the formation of Cu_{2-x}Se/CdSe and Cu_{2-x}Se/ZnSe HNCs,³⁷ which are structurally similar to the as-synthesized ber-Cu_{2-x}Se/sp-Ni₃Se₄ HNCs. The phase segregation can be explained as a consequence of the preferential starting sites and propagation paths for cation exchange within the NCs. The cation exchange reaction observed herein is also likely to have been initiated at the most energetically favored positions, like Cu vacancies. In addition, considering that no data for the Cu-Ni-Se ternary alloy can be found in the phase diagrams,⁵² the ber-Cu_{2-x}Se and sp-Ni₃Se₄ phases seem to be immiscible, which thermodynamically drives their phase segregation. As an exception, spinel CuNi₂Se₄ was very recently found as a natural mineral in 2017 (ref. 53) in the Andes in Bolivia and named Nickeltyrrellite.54 However, it took several millions of years to mineralize CuNi2Se4 under extreme conditions of continuous volcanic hydrothermal activity with repeated pressure and heating. Considering that there is almost no other literature of Cu-Ni-Se ternary systems, it can be concluded that ber-Cu_{2-r}Se and sp-Ni₃Se₄ are basically immiscible. Such a large lattice mismatch, preferable reaction site and pathway, and immiscibility of ber-Cu_{2-x}Se and sp-Ni₃Se₄ phases result in the formation of a Janus-type heterostructure with a single interface which is favorable to minimize the lattice strain and the interfacial area.

The sp-Ni₃Se₄ based NCs produced by the cation exchange reaction are expected to have high OER catalytic activity, as observed in bulk transition metal selenides electrocatalysts. The OER activity of the ber-Cu_{2-x}Se NCs, sp-Ni₃Se₄ NCs and ber- $Cu_{2-x}Se/sp-Ni_3Se_4$ (Ni : Cu : Se = 34 : 31 : 35 mol% in XRF) HNCs was evaluated in 0.1 M KOH electrolyte (pH 13). The working electrodes were prepared by the simple deposition of NCs on carbon paper. Fig. 5a shows the iR-corrected polarization curves. While pure ber-Cu_{2-x}Se NCs showed a high overpotential of 568 mV at 10 mA cm⁻², the fully cation-exchanged sp-Ni₃Se₄ NCs exhibited a much lower overpotential of 250 mV at 10 mA $\rm cm^{-2}$, which is comparable to the reported values for metal chalcogenide systems (Table S6^{\dagger}). Interestingly, the ber-Cu_{2-x}Se/sp-Ni₃Se₄ HNCs showed an even better catalytic activity with a lower overpotential of 230 mV at 10 mA cm⁻². For comparison, a simple mixture of ber-Cu_{2-x}Se NCs and sp-Ni₃Se₄ NCs was also applied to the OER; however, it has lower activity than sp-Ni₃Se₄ NCs. A scanning electron microscopy (SEM) image of the electrode of mixed NCs showed segregation of each kind of NCs (Fig. S14[†]), suggesting that the homogeneous distribution of ber- Cu_{2-x} Se and sp-Ni₃Se₄ phases contributes to the synergetic improvement of catalysis. Fig. 5b shows the Tafel slopes of the catalysts. It is generally accepted that a smaller Tafel slope indicates faster OER kinetics.55 The ber-Cu_{2-x}Se NCs had a high Tafel slope of 949 mV



Fig. 5 (a) Cyclic voltammograms of ber- $Cu_{2-x}Se$, sp-Ni₃Se₄, the mixture of ber- $Cu_{2-x}Se$ and sp-Ni₃Se₄ NCs and ber- $Cu_{2-x}Se$ /sp-Ni₃Se₄ HNCs in 0.1 M KOH. (b) Tafel plots of sp-Ni₃Se₄ NCs and ber- $Cu_{2-x}Se$ /sp-Ni₃Se₄ HNCs. (c) TOF calculations at an overpotential of 350 mV and (d) long-term durable operation with a constant potential of 1.5 V vs. RHE. (e) Capacitive current density versus scan rate and (f) Nyquist plots of the ber- $Cu_{2-x}Se$ NC, sp-Ni₃Se₄ NC and ber- $Cu_{2-x}Se$ /sp-Ni₃Se₄ HNC-loaded electrodes.

dec⁻¹ (not shown), implying their poor intrinsic OER kinetics. The sp-Ni₃Se₄ NCs and ber-Cu_{2-x}Se/sp-Ni₃Se₄ HNCs exhibited Tafel slopes of 57 and 48 mV dec⁻¹, respectively, revealing that the latter had the fastest OER kinetics.

To further estimate the inherent OER activity of the ber- Cu_{2-x} Se/sp-Ni₃Se₄ HNCs, the turnover frequency (TOF) at an overpotential of 0.35 V was calculated under the assumption that all the metal content in the NCs was catalytically active (Fig. 5c). While the TOF was 0.32×10^{-3} s⁻¹ for the ber-Cu_{2-x}Se NCs, this was enhanced by ~ 40 times to 12.98×10^{-3} s⁻¹ and $13.91 \times 10^{-3} \text{ s}^{-1}$ for sp-Ni₃Se₄ NCs and ber-Cu_{2-x}Se/sp-Ni₃Se₄ HNCs, respectively, which are comparable to or even surpass the TOFs reported for metal chalcogenide catalysts.56-58 Alternatively, assuming that only the Ni ions work as active sites, the TOF of the ber-Cu_{2-x}Se/sp-Ni₃Se₄ HNCs was calculated to be 27 $\times 10^{-3}$ s⁻¹, which is twice that of pure sp-Ni₃Se₄ NCs. After 500 cyclic voltammetry (CV) sweeps, the overpotential at 10 mA cm⁻² increased from the initial sweep by only 22 mV (from 230 to 252 mV) for the ber-Cu_{2-x}Se/sp-Ni₃Se₄ HNCs and by 15 mV (from 250 to 265 mV) for the sp-Ni₃Se₄ NCs (Fig. S15[†]), which illustrated their high durability as catalysts. Moreover, the sp-Ni₃Se₄ NCs and ber-Cu_{2-x}Se/sp-Ni₃Se₄ HNCs also exhibited excellent stability in long-term continuous operation at 1.5 vs. the reversible hydrogen electrode (RHE), in which the current densities were retained at more than 10 mA cm⁻² for 12 h without obvious degradation (Fig. 5d).

The electrochemically active surface area (ECSA) was determined from the electrochemical double-layer capacitance (C_{dl}) in the potential range in which the current response was nonfaradic. The ber-Cu_{2-x}Se/sp-Ni₃Se₄ HNCs and sp-Ni₃Se₄ NCs showed 6-times and 3-times higher C_{dl} values than the ber- Cu_{2-x} Se NCs, respectively, as shown in Fig. 5e and S16.[†] To clarify the reason for the different ECSA values of these NCs, the differences in NC morphologies before and after the OER were microscopically investigated (Fig. S17†). The SEM images showed that the ber-Cu_{2-x}Se NCs on carbon paper were transformed into larger solid particles of 50-100 nm size after the OER. The sp-Ni₃Se₄ NCs were also fused into aggregates after the OER. For the ber-Cu_{2-x}Se/sp-Ni₃Se₄ HNCs, the NC size was maintained, with only slight contact among NCs, during the OER, providing the largest surface area. The reason for this shape retention is still not fully understood. We speculate that the immiscibility of the ber-Cu_{2-x}Se and sp-Ni₃Se₄ phases prevents both species of metal ions from migrating to form larger agglomerates. XRF analyses of each NC catalyst after the OER demonstrated that the molar proportion of Se decreased, compared with the as-synthesized NCs, suggesting that Se dissolution occurred during OER operation (Fig. S17[†]). Surface oxidation of the bulk metal selenide catalyst during the OER has often been reported recently, resulting in the formation of micropores to increase the number of active sites.⁵⁹ For NCs, however, surface oxidation triggers the fusion of NCs, decreasing the merit of the large relative surface area. Therefore, suppressing such a fusion to maintain the nanoparticulate form is apparently advantageous to improve the overall activity.

The catalytic enhancement was further confirmed by electrochemical impedance spectroscopy (EIS) analyses, as shown in Fig. 5f. We fitted Nyquist and Bode plots using the equivalent circuit model which contains three kinds of resistances to reproduce the multistep kinetics in the OER (Fig. S18 and Table S7†).⁶⁰ For each catalyst, R_{ads} (resistance related to the ease of reaction intermediate adsorption on the surface) is much larger

than $R_{\rm CT}$ (charge transfer resistance), which suggests that OER catalysis is dominated by the formation of chemical intermediates. Ni₃Se₄ and ber-Cu_{2-x}Se/sp-Ni₃Se₄ HNCs have much smaller $R_{\rm ads}$ than ber-Cu_{2-x}Se, illustrating drastic improvement of OER activity after Ni cation exchange. Ber-Cu_{2-x}Se/sp-Ni₃Se₄ HNCs have not only the smallest $R_{\rm ads}$ but also the largest capacitance ($C_{\rm ads}$), which suggests that ber-Cu_{2-x}Se/sp-Ni₃Se₄ HNCs possess a larger number of preferential active sites for OER intermediates.

From the above results, the coexistence of ber- $Cu_{2-x}Se$ and sp-Ni₃Se₄ phases in a single NC seems to cause a synergetic effect to promote the OER. In the cyclic voltammogram, the ber- $Cu_{2-x}Se/sp$ -Ni₃Se₄ HNCs have a larger Ni redox current area around 1.3 V *vs.* RHE than the pure sp-Ni₃Se₄ NCs (Fig. 5a), meaning that more Ni³⁺ species were generated than in the sp-Ni₃Se₄ NCs, probably because of their larger ECSA. It is well known that in higher valence states, Ni acts as a more efficient OER active site.⁶¹ Thus, the presence of the ber- $Cu_{2-x}Se$ phase supports the generation of both a larger ECSA and larger number of Ni³⁺ species in the sp-Ni₃Se₄ phase.

To further reveal the origin of the activity difference among the three kinds of NCs, X-ray photoluminescence spectroscopy (XPS) measurements were performed. First, we investigated the XPS data of the NCs before the OER (Fig. 6a–d). In the Cu $2p_{3/2}$ region, the ber-Cu_{2-x}Se NCs showed a peak at 932.3 eV assigned to Cu^+ in ber- Cu_{2-x} Se, with a minor peak at higher binding energy assigned to Cu^{2+} in CuO (Fig. 6a). After the cation exchange, Ni $2p_{3/2}$ peaks of Ni^{2+/3+} in sp-Ni₃Se₄ and Ni²⁺ from NiO and $Ni(OH)_2$ newly emerged (Fig. 6b). Simultaneously, new Se 3d peaks appeared at \sim 58.6 eV resulting from SeO_x formation (Fig. 6c), and O 1s peaks also become strong after cation exchange (Fig. 6d). These results indicate that the cation exchange reaction caused surface oxidation of NCs, probably during the purification step in air. We would like to claim that the main phases are selenides, because all XRD peaks are assigned to selenides (Fig. 3e), and no oxide layers were



Fig. 6 XPS spectra of NCs (a–d) before and (e–h) after the OER. XPS spectra of ber- Cu_{2-x} Se NCs (red line), sp-Ni₃Se₄ NCs (green line) and ber- Cu_{2-x} Se/sp-Ni₃Se₄ HNCs (purple line) loaded on carbon paper electrodes before and after OER experiments; (a and e) Cu $2p_{3/2}$, (band f) Ni $2p_{3/2}$, (c and g) Se 3d and (d and h) O1s regions.

observed in HRTEM images. Therefore, the amorphous oxide layers are considered to be thin on the NCs. Second, we further conducted XPS measurements of all NCs after the OER (after 500 cycles CV, Fig. 6e-h). In all samples, the Se 3d peaks completely disappeared because of the Se dissolution during the OER (Fig. 6g), which is consistent with the decreased Se molar ratio after the OER (Fig. S17^{\dagger}). For the ber-Cu_{2-x}Se NCs, the Cu $2p_{3/2}$ peak was shifted from 932.3 eV (before the OER) to 933.9 eV, suggesting the oxidation of Cu^+ to Cu^{2+} and the formation of CuO (Fig. 6e). The sp-Ni₃Se₄ NCs showed a positive shift of the Ni 2p_{3/2} peak to 857.0 eV which can be assigned to the formation of Ni^{3+} in NiOOH (Fig. 6f). In the ber-Cu_{2-x}Se/sp- Ni_3Se_4 HNCs, the Cu $2p_{3/2}$ peak at around ~935.0 eV suggests the formation of CuO and Cu(OH)2, and Ni 2p3/2 peak at 856.8 eV and O 1s peak at 531.3 eV arise from NiOOH. The ber-Cu_{2-x}Se/sp-Ni₃Se₄ HNCs have higher binding energy of the Cu $2p_{3/2}$ peak (934.3 eV) than pure $Cu_{2-x}Se$ NCs (933.9 eV), implying that the higher valence state of Cu was stabilized by the coexistence of NiOOH species. It has been reported that highly oxidized Cu species can act as efficient OER catalysts,62 which suggests that the high valence state of Cu in (Cu/Ni)OOH might also catalyze the OER. However, additional experiments are required to confirm this hypothesis. Selected area electron diffraction (SAED) measurements of ber-Cu_{2-x}Se/sp-Ni₃Se₄ NCs were also carried out to further confirm the change of crystal structures during the OER (Fig. S19†). Immediately after immersing the NCs in 0.1 M KOH, the crystalline feature of ber-Cu_{2-r}Se and sp-Ni₃Se₄ was retained. However, the NCs become less crystalline after 500 CV cycles, indicating that the oxidation of NCs is electrochemically induced. Such a process can be considered as the activation step, as we can see an increase of current density in the first 50 CV cycles (Fig. S20[†]).

Conclusions

We have developed the selective synthesis of sp-Ni₃Se₄ NCs and novel Janus-type ber-Cu_{2-x}Se/sp-Ni₃Se₄ HNCs by the cation exchange reaction, and demonstrated their high OER catalytic activity. A rare sp-Ni₃Se₄ phase was selectively formed because of the preservation of the fcc Se sublattice of ber- $Cu_{2-x}Se$. Careful control of the progress of the cation exchange reaction selectively provided the Janus-like heterostructure as a result of lattice mismatch, anisotropic reaction process and immiscibility of the two phases. The ber-Cu_{2-x}Se/sp-Ni₃Se₄ HNCs exhibited exceptional OER activity, manifesting a low overpotential of 230 mV at 10 mA cm^{-2} , even better than that of the sp-Ni₃Se₄ NCs (250 mV). This outperforms the values for other Ni-Se systems and the most commonly reported solid-solution catalysts. The NCs were activated by in situ electrochemical oxidation, and their hydroxide species work as the real active species. Such an enhanced activity of the HNCs arose from the coexistence of ber-Cu_{2-x}Se and sp-Ni₃Se₄ phases in each individual NC, which provided synergistic benefits by increasing the ECSA and generating highly active Cu species. We believe that the utility of the cation exchange reaction might extend to other transition metal-based materials for highly active nextgeneration electrocatalysts. We also anticipate that the novel heterostructured NCs might be exploited in other materials systems for fabricating nanostructures with crystal phases that are not easily accessed by direct chemical synthesis.

Conflicts of interest

There are no conflicts to declare.

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References

- 1 N. S. Lewis and D. G. Nocera, *Proc. Natl. Acad. Sci. U. S. A.*, 2006, **103**, 15729–15735.
- 2 J. Barber, Chem. Soc. Rev., 2009, 38, 185-196.
- 3 D. Gust, T. A. Moore and A. L. Moore, *Acc. Chem. Res.*, 2009, **42**, 1890–1898.
- 4 T. Faunce, S. Styring, M. R. Wasielewski, G. W. Brudvig, A. W. Rutherford, J. Messinger, A. F. Lee, C. L. Hill, H. DeGroot, M. Fontecave, D. R. MacFarlane, B. Hankamer, D. G. Nocerea, D. M. Tiede, H. Dau, W. Hillier, L. Wang and S. Amal, *Energy Environ. Sci.*, 2013, 6, 1074–1076.
- 5 J. R. Swierk and T. E. Mallouk, *Chem. Soc. Rev.*, 2013, 42, 2357–2387.
- 6 T. R. Cook, D. K. Dogutan, S. Y. Reece, Y. Surendranath, T. S. Teets and D. G. Nocera, *Chem. Rev.*, 2010, **110**, 6474– 6502.
- 7 F. Cheng and J. Chen, Chem. Soc. Rev., 2012, 41, 2172-2192.
- 8 J. Suntivich, K. J. May, H. A. Gasteiger, J. B. Goodenough and Y. Shao-Horn, *Science*, 2011, **334**, 1383–1385.
- 9 M. W. Kanan and D. G. Nocera, *Science*, 2008, **321**, 1072–1075.
- 10 Y. Zhao, J. R. Swierk, J. D. Megiatto Jr, B. Sherman, J. Youngblood, D. Qin, D. M. Lentz, A. L. Moore, T. A. Moore, D. Gust and T. E. Mallouk, *Proc. Natl. Acad. Sci. U. S. A.*, 2012, **109**, 15612–15616.
- 11 Y. Lee, J. Suntivich, K. J. May, E. E. Perry and Y. Shao-Horn, *J. Phys. Chem. Lett.*, 2012, **3**, 399–404.
- 12 M. E. G. Lyons and S. Floquet, *Phys. Chem. Chem. Phys.*, 2011, 13, 5314–5335.
- 13 C. C. L. McCrory, S. Jung, J. C. Peters and T. F. Jaramillo, *J. Am. Chem. Soc.*, 2013, **135**, 16977–16987.
- 14 S. Anantharaj, S. R. Ede, K. Sakthikumar, K. Karthick, S. Mishra and S. Kundu, *ACS Catal.*, 2016, **6**, 8069–8097.
- 15 R. Subbaraman, D. Tripkovic, K. C. Chang, D. Strmenik, A. P. Paulikas, P. Hirunsit, M. Chan, J. Greeley, V. Stamenkovic and N. M. Markovic, *Nat. Mater.*, 2012, **11**, 550–557.

- 16 Y. Zhao, X. Jia, G. Chen, L. Shang, G. I. Waterhouse, L. Z. Wu,
 C. H. Tung, D. O'Hare and T. R. Zhang, *J. Am. Chem. Soc.*,
 2016, 138, 6517–6524.
- 17 S. Kukunuri, M. R. Krishnan and S. Sampath, *Phys. Chem. Chem. Phys.*, 2015, **17**, 23448–23459.
- 18 F. Wang, Y. Li, T. A. Shifa, K. Liu, F. Wang, Z. Wang, P. Xu, Q. Wang and J. He, *Angew. Chem.*, *Int. Ed.*, 2016, 55, 6919– 6924.
- 19 A. T. Swesi, J. Jasud, W. P. R. Liyanage, S. Umapathi, E. Bohannan, J. Medvedeva and M. Nath, *Sci. Rep.*, 2017, 7, 2401.
- 20 G. Nagaraju, S. M. Cha, C. Sekhar and J. S. Yu, Adv. Energy Mater., 2017, 7, 1601362.
- 21 Z. Zhuang, Q. Peng, J. Zhuang, X. Wang and Y. Li, *Chem.–Eur.* J., 2006, **12**, 211–217.
- M. Saruyama, S. Kim, T. Nishino, M. Sakamoto, M. Haruta,
 H. Kurata, S. Akiyama, T. Yamada, K. Domen and
 T. Teranishi, *Chem. Sci.*, 2018, 9, 4830–4836.
- 23 T. Yoshinaga, M. Saruyama, A. Xiong, Y. Ham, Y. Kuang, R. Niishiro, S. Akiyama, M. Sakamoto, T. Hisatomi, K. Domen and T. Teranishi, *Nanoscale*, 2018, **10**, 10420–10427.
- 24 S. Kim, T. Nishino, M. Saruyama, M. Sakamoto,
 H. Kobayashi, S. Akiyama, T. Yamada, K. Domen and
 T. Teranishi, *ChemNanoMat*, 2017, 3, 764–771.
- 25 J. B. Rivest and P. K. Jain, Chem. Soc. Rev., 2013, 42, 89-96.
- 26 H. Li, M. Zanella, A. Genovese, M. Povia, A. Falqui,
 C. Giannini and L. Manna, *Nano Lett.*, 2011, **11**, 4964–4970.
- 27 J. L. Fenton and R. E. Schaak, Angew. Chem., Int. Ed., 2017, 56, 6464–6467.
- 28 A. E. Powell, J. M. Hodges and R. E. Schaak, J. Am. Chem. Soc., 2016, 138, 471–474.
- 29 H.-L. Wu, R. Sato, A. Yamaguchi, M. Kimura, M. Haruta, H. Kurata and T. Teranishi, *Science*, 2016, **351**, 1306–1310.
- 30 L. D. Trizio, H. Li, A. Casu, A. Genovese, A. Sathya, G. C. Messina and L. Manna, *J. Am. Chem. Soc.*, 2014, 136, 16277–16284.
- 31 A. C. Berends, W. Stam, Q. A. Akkerman, J. D. Meeldijk, J. Lit and C. M. Donega, *Chem. Mater.*, 2018, **30**, 3836–3846.
- 32 R. Tu, Y. Xie, G. Bertoni, A. Lak, R. Gaspari, A. Rapallo,
 A. Cavalli, L. D. Trizio and L. Manna, *J. Am. Chem. Soc.*, 2016, 138, 7082–7090.
- 33 H. Tahara, M. Sakamoto, T. Teranishi and Y. Kanemitsu, *Phys. Rev. Lett.*, 2017, **119**, 247401.
- 34 Z. Lian, M. Sakamoto, J. J. M. Vequizo, C. S. K. Ranasinghe, A. Yamakata, T. Nagai, K. Kimoto, Y. Kobayashi, N. Tamai and T. Teranishi, *J. Am. Chem. Soc.*, 2019, **141**, 2446–2450.
- 35 H. Shen, H. Wang, H. Yuan, L. Ma and L. S. Li, *CrystEngComm*, 2012, **14**, 555–560.
- 36 D. Dorfs, T. Jartling, K. Miszta, N. C. Bigall, M. E. Kim, A. Genovese, A. Falqui, M. Povia and L. Manna, J. Am. Chem. Soc., 2011, 133, 11175–11180.
- 37 V. Lesnyak, R. Brescia, G. C. Messina and L. Manna, J. Am. Chem. Soc., 2015, 137, 9315–9323.

- 38 J. Gui, M. Ji, J. Liu, M. Xu, J. Zhang and H. Zhu, Angew. Chem., Int. Ed., 2015, 127, 3754–3758.
- 39 Y. Vuorelainen, A. Huhma and A. Hakli, *Bull. Comm. Geol. Finl.*, 1964, 215, 114–125.
- 40 A. T. Swesi, J. Masud and M. Nath, *Energy Environ. Sci.*, 2016, 9, 1771–1782.
- 41 K. Xu, H. Ding, H. Lv, S. Tao, P. Chen, X. Wu, W. Chu, C. Wu and Y. Xie, *ACS Catal.*, 2016, 7, 310–315.
- 42 X. Li, G. Han, Y. Liu, B. Dong, W. Hu, X. Shang, Y. Chai and C. Liu, *ACS Appl. Mater. Interfaces*, 2016, **8**, 20057–20066.
- 43 S. Kukunuri, M. R. Krishnan and S. Sampath, *Phys. Chem. Chem. Phys.*, 2015, **17**, 23448–23459.
- 44 S. Ananthara, J. Kennedy and S. Kundu, ACS Appl. Mater. Interfaces, 2017, 9, 8714–8728.
- 45 J. Y. Zhang, X. Tian, T. He, S. Zaman, M. Miao, Y. Yan, K. Qi,
 Z. Dong, H. Liu and B. Y. Xia, *J. Mater. Chem. A*, 2018, 6, 15653–15658.
- 46 Z. Pu, Y. Luo, A. M. Asiri and X. Sun, ACS Appl. Mater. Interfaces, 2016, 8, 4718-4723.
- 47 H. Fan, H. Yu, X. Wu, Z. Yu, Y. Zhang, Z. Luo, H. Wang, Y. Guo, S. Madhavi and Q. Yan, *ACS Appl. Mater. Interfaces*, 2016, 8, 25261–25267.
- 48 H. Baker, *ASM Handbook, Alloy phase diagrams*, ASM International, 1992.
- 49 G. Wang, S. K. Lok, G. K. L. Wong and I. K. Sou, *Small*, 2011, 7, 1546–1551.
- 50 M. Misono, E. Ochiai, Y. Saito and Y. Yoneda, *J. Inorg. Nucl. Chem.*, 1967, **29**, 2685–2691.
- 51 L. D. Trizio and L. Manna, *Chem. Rev.*, 2016, **116**, 10852–10887.
- 52 P. Vilars, A. Prince and H. Okamoto, *Handbook of Ternary Alloy Phase Diagrams*, ASM International, 1995.
- 53 G. Grundmann and H. J. Föster, Minerals, 2017, 7, 68.
- 54 H. J. Förster, C. Ma, G. Grundmann, L. Bindi and C. J. Stanley, *Eur. J. Mineral.*, 2019, **31**, 197–202.
- 55 L. Liang, H. Cheng, F. Lei, J. Han, S. Gao, C. Wang and Y. Sun, *Angew. Chem., Int. Ed.*, 2015, **54**, 12004–12008.
- 56 L. Wu, Q. Li, C. H. Wu, H. Zhu, A. Mendoza-Garcia, B. Shem, J. Guo and S. J. Sun, *J. Am. Chem. Soc.*, 2015, **137**, 7071–7074.
- 57 Y. Shi, Y. Zhou, D. Yang, W. Xu, C. Wang, F. Wang, J. Xu, X. Xia and H. Chen, *J. Am. Chem. Soc.*, 2017, **139**, 15479– 15485.
- 58 Y. H. Fang and Z. P. Liu, ACS Catal., 2014, 4, 4364-4376.
- 59 B. R. Wygant, K. Kawashima and C. B. Mullins, ACS Energy Lett., 2018, 3, 2956–2966.
- 60 J. R. Swierk, S. Klaus, L. Trotochaud, A. T. Bell and D. Tilley, J. Phys. Chem. C, 2015, 119, 19022–19029.
- 61 M. Yoshida, Y. Mitsutomi, T. Mineo, M. Nagasaka,
 H. Yuzawa, N. Kosugi and H. Kondoh, *J. Phys. Chem. C*, 2015, **119**, 19279–19286.
- 62 S. Cui, X. Liu, Z. Sun and P. Du, *ACS Sustainable Chem. Eng.*, 2016, 4, 2593–2600.