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High-performance, long lifetime chloride ion battery using a NiFe-Cl layered double hydroxide cathode†

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Chloride ion batteries (CIBs) are an example of a promising new emerging rechargeable battery technology, that exhibits large theoretical volumetric energy density performance and good safety. However, unsatisfactory capacity and poor cycling lifetime of the cathode currently hinder the development of CIBs. Herein, we report the use of an Ni²⁺Fe³⁺-based layered double hydroxide (LDH) intercalated by chloride ions as a promising cathode material for CIBs. [Ni₂Fe(OH)₆]Cl·1.37H₂O (NiFe–Cl LDH) exhibits a high maximum capacity of 350.6 mA h g⁻¹ and a long lifetime of over 800 cycles (at 101.1 mA h g⁻¹) at a current density of 100 mA g⁻¹, which is superior to most currently reported CIB cathodes. *In situ* X-ray absorption near-edge structure (XANES) and *ex situ* X-ray photoelectron spectroscopy (XPS) reveal the valency changes of the Fe²⁺/Fe³⁺ and Ni²⁺/Ni³⁺ redox pairs within the metal hydroxide layers of the LDH during electrochemcial cycling. *In situ* XRD reveals that 2D anion diffusion within the LDH results in only ~3% structural change. Oxygen K-edge soft X-ray absorption spectroscopy (SXAS) reveals the oxygen atoms within the MO₆ octahedra reversibly participate in the electrochemical reaction. In view of the extensive chemical variation, low-cost, and ease-of-preparation of LDH-based materials we regard LDHs as a promising materials platform for application as cathode materials in chloride ion batteries.

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Introduction

Over the past few decades, secondary power supplies represented by lithium-ion batteries (LIBs) have experienced revolutionary developments because of their high energy density and good cycling stability.¹⁻³ Recently, a new type of rechargeable battery has emerged and has attracted increasing attention from both academia and industry. Unlike reversible migration of metal cations in traditional batteries, battery-based energy storage whereby anions shuttle between two electrodes, providing more options and opportunities for electrochemical energy storage.⁴⁻⁸ The chloride ion battery (CIB) is a representative of this kind of battery, which shows high theoretical volumetric energy density (~2500 W h L⁻¹), comparable with that of Li–S batteries.^{9,10} To date, only a few cathode materials have been reported for CIBs and most of them are metal

chlorides/oxychlorides, such as CoCl₂, CuCl₂, FeOCl, BiOCl, and their hybrids.^{11–18} They normally suffer from unsatisfactory electrochemical performance and poor structural stability. Therefore, the key challenge in CIBs is to develop alternative cathode materials with competitive capacity, long cycle life and low-cost characteristics.

Layered double hydroxides (LDHs) are one type of important brucite-like anionic clay generally expressed as $[M^{II}_{1-x}M^{III}_{x}(OH)_{2}](A^{n-})_{x/n} \cdot mH_{2}O$ (M^{II} and M^{III} are divalent and trivalent metals respectively, A^{n-} is an exchangeable anion for compensating the positive charge of the bimetal hydroxide layers). 19-23 The tunability in both chemical composition and metal valency endows LDH materials with wide application potential in the fields of supercapacitors, LIBs anode materials, and electrocatalysts.24-30 In particular, LDH materials have large anion-exchange capacity with reversible anion uptake/release. Very recently, Young et al. used in situ techniques to observe the intercalation of anions into the interplanar space of an LDH under a positive bias and the de-intercalation under a reverse bias.31 Furthermore, the unique topochemical character and facile 2D diffusion pathways endow LDH materials with high structural stability and simultaneous facile anion intercalation/ de-intercalation ability. In our previous work, we reported for the first time that a CoFe LDH intercalated with chloride exhibited reversible chlorination/dechlorination with a high capacity of ~160 mA h g⁻¹.32 However, this capacity drops off

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dramatically after \sim 100 cycles, which probably arises as a result of local structure distortions or partial dissolution of Co from the LDH platelets.

In this work, Cl⁻ intercalated NiFe LDH was synthesized by a simple continuous process including hydrothermal and anion-exchange steps. When used as the cathode in a CIB, the NiFe-Cl LDH shows an ultrahigh initial capacity of 350.6 mA h g⁻¹ at a current density of 100 mA g⁻¹. The intercalated interlamellar chloride ions reversibly shuttle between the two electrodes which is accompanied by redox reactions of both the Ni²⁺/Ni³⁺ and Fe²⁺/Fe³⁺ in the LDH layers during electrochemical cycling. The layered structure of LDH exhibits less than a 3% volume variation during chloride insertion/deinsertion, that leads to a superior cycling life (reversible capacity of 101.1 mA h g⁻¹ after 800 cycles). In addition, a possible charge compensation mechanism involving oxygens in the MO₆ octahedra in the brucite-like sheets is discussed for the first time. Owing to the hypo-toxicity and low cost of nickel and facile synthetic procedure of NiFe-Cl LDH, these encouraging results present a range of opportunities for LDHs as a material platform for anion insertion electrochemistry and an exciting new frontier for the development of novel anion-based rechargeable batteries and large-scale energy-storage systems.

Results and discussion

Synthesis and characterization

NiFe-Cl nanoplatelets were prepared by a facile two-step process: preparation of NiFe-CO₃ LDH precursor via hydrothermal method followed by ion-exchange of the interlayer

CO₃²⁻ by Cl⁻ ions. X-ray diffraction (XRD) pattern (Fig. S1, ESI†) of the NiFe-CO₃ LDH sample with Ni/Fe ratio of 2/1 indicates a well-defined carbonate-intercalated LDH with a basal plane spacing of ~7.42 Å.33 After ion-exchange, all Bragg reflections in the XRD pattern can be indexed to a rhombohedral unit cell, reflections indexed as (003), (006), (110), and (113) were observed at $2\theta = 11.54^{\circ}$, 23.22° , 59.80° , and 61.14° respectively. The XRD showed no evidence for the presence of any crystalline impurity phases. The interlayer spacing of the NiFe-Cl LDH expands to 7.67 Å ($\Delta = 0.25$ Å; 3%), to accommodate the Cl⁻ anions between the layers (inset of Fig. 1a).34 Fourier transform infrared spectroscopy (FTIR) has been employed to provide additional evidence for the anion exchange (Fig. S2, ESI†). We find that the intensity of the absorption band at 1380 cm⁻¹, attributed to the ν_3 mode of CO_3^{2-35} significantly decreased in intensity after ion-exchange; (the weak feature remaining may be assigned to slight contamination from atmospheric CO₂). Scanning electron microscopy (SEM) image of the as-prepared NiFe-Cl LDH (Fig. 1b) reveals a typical plate-like morphology with plates ranging in size from \sim 100–200 nm, which is entirely consistent with the morphology of the NiFe-CO3 LDH precursor (Fig. S3, ESI†). The composition of the NiFe-CO₃ LDH precursor and NiFe-Cl LDH are determined to be Ni₂Fe(OH)₆(CO₃)_{1/} 2·1.56H2O and Ni2Fe(OH)6Cl·1.37H2O, respectively, based on the results of inductively coupled plasma-atomic emission spectrometer (ICP-AES) and thermogravimetry analysis (TGA) (Fig. S4, ESI†).

Fig. 1c-e shows the Ni 2p, Fe 2p and O 1s X-ray photoelectron spectroscopy (XPS) spectra of the NiFe-Cl LDH. The binding energies for the Ni 2p electrons (Fig. 1c) are similar to those of

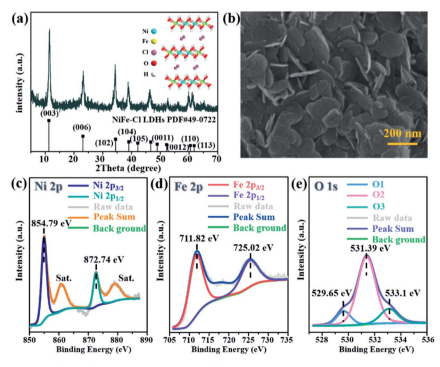


Fig. 1 (a) Powder XRD data of NiFe-Cl LDH (inset: structural model). (b) SEM image of the as-prepared NiFe-Cl LDH sample. XPS spectra of (c) Ni 2p, (d) Fe 2p and (e) O 1s for NiFe-Cl LDH.

Ni $^{2+}$ with Ni $2p_{3/2}$ and $2p_{1/2}$ peak located at 854.79 and 872.74 eV, respectively. 36 For Fe, two photoemissions are observed at 711.82 eV (Fe $2p_{3/2}$) and 725.02 eV (Fe $2p_{1/2}$) (Fig. 1d), indicating the presence of Fe $^{3+}$. 37 Fig. 1e presents that three different photoemission components in the O 1s region. The binding energies at 529.65 eV (O1) and 531.39 eV (O2) are assigned to the metal–oxygen (M–O) bonds and O–H bonds in LDH, respectively; the low intensity photoemission feature at 533.1 eV (O₃) may be ascribed to adsorbed O–O bonds from molecular oxygen. 38

Electrochemical performance

The electrochemical characteristics of the NiFe-Cl LDH cathode were carried out in a Li/Bpy14Cl-PC/NiFe-Cl LDH CIB system, using metallic Li as the anode and 0.5 M 1-butyl-1methylpyrrolidinium chloride (Bpy_{1.4}Cl) in propylene carbonate (PC) as electrolyte. To optimize the Ni/Fe ratio to achieve the best energy storage performance, different Ni_xFe-Cl LDH samples (x = 2, 3, 4 and 5) were synthesized (Fig. S5, ESI†) and the NiFe LDH with Ni/Fe = 2 was selected as the optimized sample (Fig. S6, ESI†). Therefore, our further study was performed using NiFe LDH sample with Ni/Fe ratio of 2. Fig. 2a displays the cyclic voltammetry (CV) profiles of the initial several cycles in a voltage range of 1.2-3.0 V (vs. Li/Li⁺) with a scan rate of 0.2 mV⁻¹. The battery was firstly discharged to 1.2 V (vs. Li/Li⁺) from open circuit voltage. Two pairs of broad redox events at 2.12/2.45 V and 1.93/1.55 V were observed. These current-voltage profiles maintain a similar shape during the subsequent cycles except for an apparent polarization during the first discharging process. The broad redox feature is caused by either multi-step reaction processes or multi-element redox reactions during the charge-discharge (discussed later). Galvanostatic discharge/charge curves (Fig. 2b) at different current densities show simple sloping profiles without an obvious plateau, suggesting that the NiFe–Cl LDH cathode undergoes no significant structural phase transformations but only anion intercalation/de-intercalation. Maximum discharge capacities of 350.1, 140.9 and 87.5 mA h g $^{-1}$ were obtained at 100, 250 and 500 mA g $^{-1}$, respectively.

The cycling stability of the NiFe-Cl LDH electrode operated at a current density of 100 mA g⁻¹ is shown in Fig. 2c. Surprisingly, a high discharge specific capacity of 101.1 mA h g⁻¹ was obtained after 800 cycles with a coulombic efficiency of 90.3%, which demonstrated that the cell is capable of delivering outstanding cycling stability. Some inevitable side reactions involving either the solvent and/or electrolyte resulted in coulombic efficiencies of more than 100% in the initial cycles. The slowly decreasing coulombic efficiency may be due to the gradual accumulation of non-conducting LiCl on the anode or side reactions involving electrolyte decomposition during cycling. For comparison, the specific capacity and cyclic stability data of other CIB cathode materials reported works to date are summarized in Fig. 2d and Table S1.† To the best of our knowledge, the NiFe-Cl LDH outperforms most of the previously reported CIB cathodes, 11-18,32 exhibiting superior Clstorage capability.

Evolution of structure and composition

In order to probe the evolution of the host structure and Cl⁻ storage mechanism for NiFe–Cl LDH, an *in situ* XRD study of the cathode was performed during a whole charge/discharge process at a constant rate of 50 mA g⁻¹ (Fig. 3a). No new phases except the start and final LDHs were detected during the electrochemical process (Fig. 3b), this demonstrates a clean topochemical phase transformation and no electrolyte intercalation. During the charging process, the (003) Bragg reflection (Fig. 3c) exhbits a monotonic shift toward the lower 2θ values, implying that the basal spacing of NiFe–Cl LDH decreases

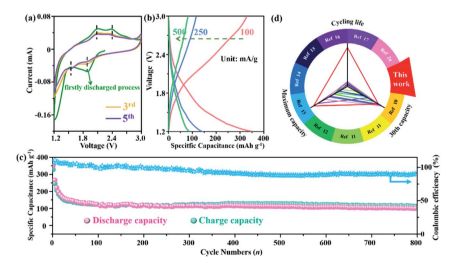


Fig. 2 Electrochemical performance of the NiFe–Cl LDH based CIB. (a) CV curves at a scan rate of 0.2 mV s^{-1} . (b) Galvanostatic discharge/charge curves (current density: 100 mA g^{-1}) at a voltage range from 1.2-3.0 V. (c) Cycle performances and corresponding coulombic efficiency at a current density of 100 mA g^{-1} within 800 cycles. (d) Discharge specific capacity and cycling life of NiFe–Cl LDH material compared with other reported cathodes of CIBs.

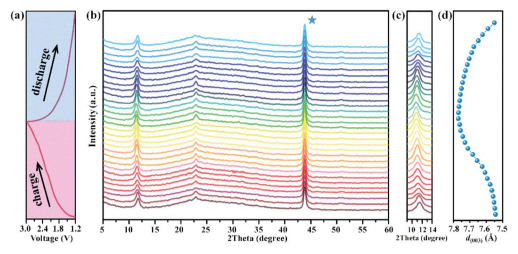


Fig. 3 (a) Typical galvanostatic charge/discharge curves. (b) In situ XRD patterns collected during the whole charge/discharge process of NiFe-CI LDH electrode. The asterisk represents peaks from the current collector (stainless steel). (c) Partial enlargement for the diffraction of (003) plane. (d) Basal spacing (d_{003}) of the NiFe-Cl LDH in different charge-discharge states.

continuously upon Cl intercalation. In contrast, a gradual recoverly back to higher angle for the (003) Bragg reflection was observed upon discharging to 1.2 V due of de-intercalation of Cl from NiFe-Cl LDH. Energy-dispersive X-ray spectroscopy (EDS) (Fig. S7 and S8, ESI†) analysis of the cathode material shows the accumulation and release of Cl ions, further demonstrating the reversible shuttling of Cl⁻ ions during the charge/discharge process. The basal spacing of the NiFe-Cl LDH expanded from 7.54 Å to 7.76 Å upon charging to 3.0 V (Fig. 3d) and then reversibly contracted to 7.55 Å when the electrode was fully discharged.

Investigation of Cl⁻ storage mechanism

In situ X-ray absorption spectroscopy (XAS) data at the Fe K-edge and Ni K-edge at different charging and discharging states were recorded in order to further investigate the change in the oxidation states and local structure of NiFe-Cl LDH during the

electrochemical process. Fig. 4a and b present the Fe K-edge XAS spectra compared with Fe(OH)3 as a reference. Upon charging, the Fe K-edge absorption clearly shifts toward higher energy as a function of the degree of Cl intercalation, which reflects the change in the oxidation state of Fe. During discharge, reversible changes were observed, the Fe K-edge absorption profile completed returned to original sprectum following extraction of Cl⁻ ions, indicating the decrease of Fe valence in the NiFe-Cl LDH cathode. In addition, the Ni^{2+/3+} electrochemically active redox pair also participates in the charge/discharge cycles as obserbed in the Ni K-edge spectra (Fig. 4c and d) (using $Ni(OH)_2$ as the reference). However, unlike the spectral changes for Fe, the shift of Ni K-edge absorption is not as significant, and the intensity of the white edges shows a systematic change. We conclude that the detailed changes of the Ni ions are more complicated than the Fe centres. The evolution of the valence states of the two metals within the LDH

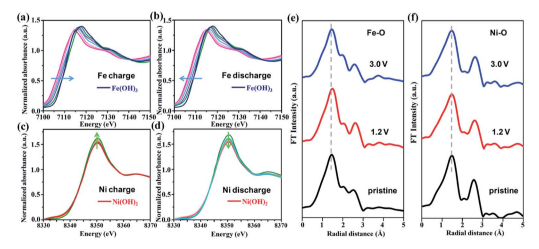


Fig. 4 In situ XANES spectra of the Fe K-edge and Ni K-edge collected during (a), (c) charge and (b), (d) discharge process. Fourier transform magnitude of the EXAFS spectra for (e) Fe K-edge and (f) Ni K-edge collected at different charge/discharge states.

host layer during the electrochemical process can also be highlighted by the first-order derivative of the XANES spectra (Fig. S9 and S10, ESI \dagger).

The evolution of the Fe or Ni local coordination was investigated by extended X-ray absorption fine structure (EXAFS) (Fig. 4e and f). The Fourier transformation of the weighted EXAFS spectrum for both Fe and Ni K-edge under fully charged or discharged states are very similar to the pristine sample, suggesting excellent structural stability of the NiFe–Cl LDH cathode. Furthermore, we note the change in unit cell volume of the NiFe–Cl LDH cathode is less than $\sim\!3\%$ throughout the entire electrochemical cycle, implying a high structural reversibility which may in part account for the remarkable cycling stability of the CIB. In addition, SEM image of the NiFe–Cl LDH cathode after 800 cycles (Fig. S11, ESI†) displays that the platelike morphology is well maintained without active material agglomeration, particle expansion or pulverization.

Although both Ni and Fe cations undergo reversible redox reactions during electrochemical cycling, they seem to have different charge compensation mechanisms and it appears that these perturb the electronic structure of the Ni atom more significantly than that of the Fe atom. To gain more insight, a series of O K-edge soft XAS were recorded over the whole electrochemical cycle. This technique is very sensitive to the electronic structure around oxygen and metal-oxygen bonding. 39,40 Fig. 5a shows the O K-edge spectra of NiFe-Cl LDH during the charging process. The first sharp peak around 533.5 eV corresponds to the transition of O 1s electrons to a hybridized state of the metal 3d with O 2p orbitals, while the broad higher energy peaks above 539 eV can be assigned to the transitions to hybridized states of oxygen 2p and metal 4s/4p orbitals. It was observed that upon increasing voltage, the first peak intensities decreased systematically and an additional well-resolved shoulder peak (marked by a vertical arrow) in a lower energy region at ~532.5 eV gradually emerged.

Interestingly, under the electrochemical operational conditions of the CIB, the variations in O K-edge spectra were totally reversible as shown in Fig. 5b. According to previous reports, ^{41,42} the intensity of the pre-edge feature in the O K-edge spectra depends on the number of "d" electronic holes states in the metal 3d orbital hybridized with the O 2p orbital and the covalency of the M–O bond. These observations are consistent with Cl⁻ intercalation giving rise to an increased effective nuclear charge of the metal cations, decreased M–O bond covalency, indicating that the oxygen ions of the MO₆ octahedra in the LDH participate in the electrochemical reaction to some extent. Combined with the XANES measurements, we speculate that this kind of partial electron compensation between metal and oxygen ligand is more likely to occur in the Ni–O bonds.

In order to further identify the charge compensation mechanism during the electrochemically driven Cl intercalation (deintercalation) ex situ XPS measurement was perfromed. The Fe 2p core spectra (Fig. 6a) shows Fe 2p_{3/2} and Fe 2p_{1/2} photemission feature, this feature may be deconvoluted into two pairs of resonances at 710.13/712.61 eV and 723.17/725.94 eV, that may be ascribed to the binding energies for Fe2+ and Fe3+ respectively. Upon charging, the Fe3+ signals increase at the expense of the Fe²⁺ signals. When charged from 1.2 to 3.0 V, the ratio of the peak integrals for Fe³⁺/Fe²⁺ increases from 0.68 to 2.26 (Fig. 6c), as a result of insertion of Cl⁻. During discharge, Cl deintercalation resulted in a decrease of Fe³⁺/Fe²⁺ peak intergal ratio; when the NiFe-Cl electrode was discharged to 1.2 V, the ratio of Fe^{3+}/Fe^{2+} peak integrals (\sim 0.63) almost fully returned to its initial value before cycling. As shown in Fig. 6b, the main Ni 2p_{3/2} and Ni 2p_{1/2} photoemission peaks were divided into two components centred at 854.82/856.73 eV and 873.12/874.59 eV, which are assigned to Ni²⁺ and Ni³⁺ respectively. The ratio of Ni³⁺/Ni²⁺ peak intergals changed from 0.56 to 1.40 (Fig. 6d) during the charging process. The XPS data reveal that the peak intergal ratio change for Ni³⁺/Ni²⁺ is smaller than

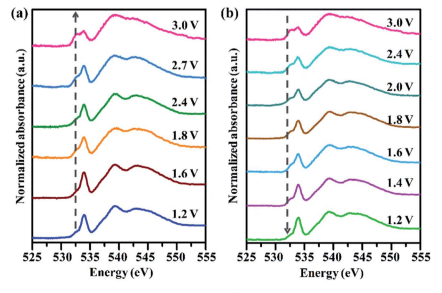


Fig. 5 Normalized oxygen K-edge soft X-ray absorption spectra of NiFe-Cl LDH cathode in a whole electrochemical cycle: (a) charge and (b) discharge.

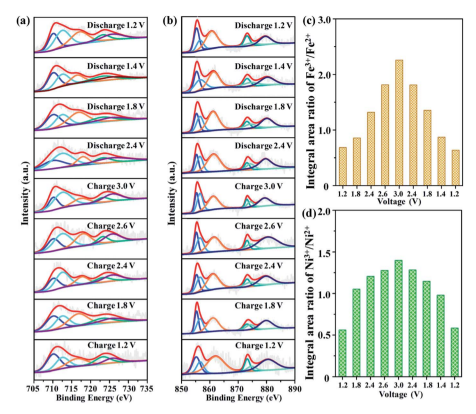


Fig. 6 Ex situ XPS analysis on (a) Fe 2p region and (b) Ni 2p region of NiFe-Cl LDH electrode in a whole charge/discharge cycle. Relative ratio of peak integrals for (c) Fe³⁺/Fe²⁺ and (d) Ni³⁺/Ni²⁺ photoemission

that of Fe³⁺/Fe²⁺ during the electrochemical cycle. The excellent oxidation/reduction reversibility of the LDH host gives the NiFe-Cl LDH a high specific capacity as an cathode material for CIB.

In addition to the investigation on the cathode, we aslo performed an XPS analysis on the Li anode, as shown in Fig. S12 (ESI†). The binding energy of lithium metal standard appears at 54.85 eV. This peak shifts to a higher binding energy of 55.23 eV during the electrochemical cycle. This is attributed to the migration of chloride ions to the surface of Li anode, leading to the formation of LiCl. After fully charging to 3.0 V, the Li 1s peak returns to the position of 54.87 eV, demonstrating a reversible uptake/release of Cl⁻ anions from the Li anode. Therefore, the half reaction of the anode can be described as Li + Cl \longleftrightarrow LiCl + e⁻.

Conclusions

We have prepared phase pure $2:1 \text{ Ni}^{2+}: \text{Fe}^{3+} \text{ LDH}$ intercalated with chloride using hydrothermal synthesis followed by anionexchange. We have demonstrated it is a highly promising cathode material in a rechargeable CIB. Upon electrochemical cycling a high capacity cathode is observed. Our NiFe-Cl LDH containing CIB exhibits an ultralong cycle lifetime with stable reversible capacity of 101.1 mA h g^{-1} after 800 cycles at a current density of 100 mA g^{-1} , which is superior to most of the currently reported CIB cathodes. The topochemical transformation characteristics plus the available interlayer 2D diffusion pathway endow NiFe-Cl LDH with high structural stability (<3%

unit cell change) during facile Cl⁻ intercalation/deintercalation. The mechanism for charge compensation during Cl intercalation/deintercalation has been extensively studied. Reversible redox changes to the Fe and Ni cations is observed as are changes in the polarization of the Ni-O bonds during electrochemical cycling. In view of the extensive chemical variation, low-cost, and ease-of-preparation, LDH-based materials can be regarded as a promising materials platform for application in large-scale, safe energy storage devices.

Experimental section

Reagents and materials

FeCl₃·6H₂O, NiCl₂·6H₂O, NaCl, triethylamine (TEA, Sigma-Aldrich), urea and N-methyl-2-pyrrolidone (NMP) were of analytical grade and used without further purification. 1-Butyl-1-methylpyrrolidinium chloride (Bpy1.4Cl, 99%, loLiTech) and propylene carbonate (PC, anhydrous, Sigma-Aldrich) were purchased from Shanghai (China) Ailei Chemical Co. Ltd. Super P and polyvinylidene fluoride (PVDF) were obtained from Shenzhen (China) Kejing Co. Ltd.

Synthesis of NiFe-CO₃ LDH

TEA and urea were chosen as the chelating agent and ammonium releasing reagent during the hydrothermal process, respectively. NiCl₂·6H₂O, FeCl₃·6H₂O, and TEA were completely dissolved in 400 mL deionized water, in which the total concentration of metal cation was 20 mM, and the ratio of $\rm Ni^{2^+}/Fe^{3^+}$ was 2 : 1. The concentration of TEA was 2.5 times that of Fe^{3^+}. Then, 7 mmol urea was added into the solution. The above mixture was stirred for 1 h and then poured into a sealed stainless-steel Teflon-lined autoclave (100 mL) at 150 °C for 48 h. The autoclave was then cooled down to room temperature and the powder NiFe–CO $_3$ LDH sample was obtained by washing with deionized water and ethanol several times.

Preparation of NiFe-Cl LDH

Intercalation of Cl $^-$ in the NiFe–CO $_3$ LDH precursor was conducted in a high concentration of aqueous NaCl. A flask containing 100 mL deionized water was added with 100 mg asprepared NiFe–CO $_3$ LDH, 0.5 mol NaCl and 41.7 μ L HCl, and was then mechanically stirred under N $_2$ atmosphere for 48 h. The NiFe–Cl LDH sample was obtained successfully after centrifuging with deionized water three times and dried at vacuum oven at 60 °C for 12 h.

Electrodes preparation

The NiFe–Cl LDH was mixed with Super P and PVDF with the weight ratio of 7:2:1 in a mortar for 60 min, using NMP as the solvent. Then, the slurry was coated on the stainless-steel foil through a 100 μ m blade followed by drying under vacuum oven at 110 °C for 12 h. The resulting stainless-steel foil was cut into pieces with a diameter of 12 mm and the loading mass of active materials was approximate \sim 1.5 mg cm $^{-2}$. The electrolyte used in our experiments was 0.5 M Bpy_{1,4}Cl in PC solution. Glass fibre disks (Whatman) were served as the separator and the Li metal was chosen as the anode. Before the electrochemical tests, the battery had to rest for 12 h to reach the equilibrium state.

Materials characterization

X-ray diffraction (XRD) patterns of LDH samples and electrodes were recorded by a Rigaku XRD-6000 diffractometer with a scanning rate of 10° min⁻¹, which was operated at 40 kV and 40 mA with Cu K α radiation ($\lambda = 0.15418$ nm). The morphology of the as-prepared samples was obtained via a Zeiss Supra 55 field emission scanning electron microscope (SEM), while the composition of substance was explored by an EDX attachment (Oxford Instrument Isis 300). The Fourier transform infrared (FT-IR) spectra were acquired through a Vector 22 (Bruker) spectrophotometer with a 2 cm⁻¹ resolution. The inductively coupled plasma (ICP) emission spectroscopy (Shimadzu ICPS-7500) was used for the metal elementary compositions of products. Thermogravimetric analyzer (TGA) was carried out on a PCT-1A thermal analysis system at an air atmosphere with a heating rate of 10 °C min⁻¹ and a range from 25 °C to 800 °C. X-ray photoelectron spectra (XPS) were tested by a Thermo VG Escalab 250 X-ray photoelectron spectrometer at a pressure of about 2×10^{-9} Pa with Al 193 K α X-rays as the excitation source. For the ex situ XPS measurement, the cathode was entirely charged or discharged to different states of voltage at 100 mA g⁻¹. In order to eliminate the influence of residual electrolyte, the electrodes were washed with PC several times, preserved in a vacuum glass box to avoid the contamination of moisture and oxygen. X-ray absorption spectroscopy (XAS) measurements of Ni and Fe were carried out at beamline 1W1B while O element was measured at beamline 4W7B, all of which located on the Beijing Synchrotron Radiation Facility (BSRF), Institute of High Energy Physics (IHEP), Chinese Academy of Sciences (CAS). And the IFFEFIT 1.2.11 data analysis package (Athena, Artemis, Atoms, and FEFF6) was used for data analysis. Al foil, instead of stainless-steel foil, was used as current collector for the in situ XAS test using a home-made cell, which has a high-precision 18 mm diameter sandwich geometry and is equipped with one high-purity Be window. Cyclic voltammetry (CV) measurements were performed on a CHI 660 electrochemical workstation at different scanning rates while a Land BT2000 battery tester was used to test the cycling and chargedischarge performance at 100 mA g⁻¹ within 1.2-3.0 V. All of the electrochemical tests were performed at 25 °C.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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Notes and references

- 1 S. Kim, H. Oguchi, N. Toyama, T. Sato, S. Takagi, T. Otomo, D. Arunkumar, N. Kuwata, J. Kawamura and S. Orimo, *Nat. Commun.*, 2019, **10**, 1081–1089.
- 2 M. Chen, X. Ma, B. Chen, R. Arsenault, P. Karlson, N. Simon and Y. Wang, *Joule*, 2019, **9**, 2622–2646.
- 3 J. Liu, Z. Bao, Y. Cui, E. J. Dufek, J. B. Goodenough, P. Khalifah, Q. Li, B. Y. Liaw, P. Liu, A. Manthiram, Y. S. Meng, V. R. Subramanian, M. F. Toney, V. V. Viswanathan, M. S. Whittingham, J. Xiao, W. Xu, J. Yang, X. Yang and J. Zhang, *Nat. Energy*, 2019, 4, 180–186.
- 4 V. K. Davis, C. M. Bates, K. Omichi, B. M. Savoie, N. Momčilović, Q. Xu, W. J. Wolf, M. A. Webb, K. J. Billings, N. H. Chou, S. Alayoglu, R. K. McKenney, I. M. Darolles, N. G. Nair, A. Hightower, D. Rosenberg, M. Ahmed, C. J. Brooks, T. F. Miller, R. H. Grubbs and S. C. Jones, *Science*, 2018, 362, 1144–1148.
- 5 F. Gschwind and J. Bastien, *J. Mater. Chem. A*, 2015, **3**, 5628-
- 6 X. Hu, F. Chen, S. Wang, Q. Ru, B. Chu, C. Wei, Y. Shi, Z. Ye, Y. Chu, X. Hou and L. Sun, ACS Appl. Mater. Interfaces, 2019, 11, 9144–9148.
- 7 C. Chen, T. Yu, M. Yang, X. Zhao and X. Shen, *Adv. Sci.*, 2019, 6, 1802130.
- 8 F. Chen, Z. Leong and H. Yang, *Energy Storage Materials*, 2017, 7, 189–194.

- 9 F. Gschwind, H. Euchner and G. Rodriguez-Garcia, Eur. J. Inorg. Chem., 2017, 2784-2799.
- 10 X. Zhao, Z. Zhao-Karger, M. Fichtner and X. Shen, Angew. Chem., Int. Ed., 2020, 59, 5902-5949.
- 11 X. Zhao, S. Ren, B. Michael and M. Fichtner, J. Power Sources, 2013, 245, 706-711.
- 12 X. Zhao, Z. Zhao-Karger, D. Wang and M. Fichtner, Angew. Chem., Int. Ed., 2013, 52, 13621-13624.
- 13 P. Gao, M. A. Reddy, X. Mu, T. Diemant, L. Zhang, Z. Z. Karger, V. S. K. Chakravadhanula, O. Clemens, R. J. Behm and M. Fichtner, Angew. Chem., Int. Ed., 2016, 55, 4285-4290.
- 14 X. Zhao, Z. Zhao, M. Yang, H. Xia, T. Yu and X. Shen, ACS Appl. Mater. Interfaces, 2017, 9, 2535-2540.
- 15 Z. Zhao, T. Yu, Y. Miao and X. Zhao, Electrochim. Acta, 2018, **270**, 30-36.
- 16 T. Yu, O. Li, X. Zhao, H. Xia, L. Ma, J. Wang, Y. Meng and X. Shen, ACS Energy Lett., 2017, 2, 2341-2348.
- 17 K. P. Lakshmi, K. J. Janas and M. M. Shaijumon, J. Power Sources, 2019, 433, 126685.
- 18 R. Yang, T. Yu and X. Zhao, J. Alloys Compd., 2019, 788, 407-
- 19 X. Gao, X. Liu, D. Wu, B. Qian, Z. Kou, Z. Pan, Y. Pang, L. Miao and J. Wang, Adv. Funct. Mater., 2019, 1903879.
- 20 J. Liang, R. Ma, N. Iyi, Y. Ebina, K. Takada and T. Sasaki, Chem. Mater., 2010, 22, 371-378.
- 21 Z. Pan, Y. Jiang, P. Yang, Z. Wu, W. Tian, L. Liu, Y. Song, Q. Gu, D. Sun and L. Hu, ACS Nano, 2018, 12, 2968-2979.
- 22 R. Gao and D. Yan, Adv. Energy Mater., 2019, 1900954.
- 23 J. Chen, X. Wang, J. Wang and P. S. Lee, Adv. Energy Mater., 2016, 6, 1501745.
- 24 X. Li, H. Wu, C. Guan, A. M. Elshahawy, Y. Dong, S. J. Pennycook and J. Wang, Small, 2018, 1803895.
- 25 J. Yang, C. Yu, X. Fan and J. Qiu, Adv. Energy Mater., 2014, 4, 1400761.

- 26 M. Tian, C. Liu, Z. G. Neale, J. Zheng, D. Long and G. Cao, ACS Appl. Mater. Interfaces, 2019, 11, 35977-35986.
- 27 L. Shi, Y. Chen, R. He, X. Chen and H. Song, Phys. Chem. Chem. Phys., 2018, 20, 16437-16443.
- 28 G. Chen, T. Wang, J. Zhang, P. Liu, H. Sun, X. Zhuang, M. Chen and X. Feng, Adv. Mater., 2018, 30, 1706279.
- 29 X. Zhang, Y. Zhao, Y. Zhao, R. Shi, G. I. N. Waterhouse and T. Zhang, Adv. Energy Mater., 2019, 9, 1900881.
- 30 C. Tang, H. Wang, H. Wang, Q. Zhang, G. Tian, J. Nie and F. Wei, Adv. Mater., 2015, 27, 4516-4522.
- 31 M. J. Young, T. Kiryutina, N. M. Bedford, T. J. Woehl and C. U. Segre, Sci. Rep., 2019, 9, 2462-2473.
- 32 Q. Yin, D. Rao, G. Zhang, Y. Zhao, J. Han, K. Lin, L. Zheng, J. Zhang, J. Zhou and M. Wei, Adv. Funct. Mater., 2019, 29,
- 33 Q. Yin, J. Luo, J. Zhang, S. Zhang, J. Han, Y. Lin, J. Zhou, L. Zheng and M. Wei, Adv. Funct. Mater., 2019, 1907448.
- 34 M. Xu and M. Wei, Adv. Funct. Mater., 2018, 28, 1802943.
- 35 L. Dang, H. Liang, J. Zhuo, B. K. Lamb, H. Sheng, Y. Yang and S. Jin, Chem. Mater., 2018, 30, 4321-4330.
- 36 Y. Wan, W. Zhao, Y. Tang, L. Li, H. Wang, Y. Cui, J. Gu, Y. Li and J. Shi, Appl. Catal., B, 2014, 148, 114-122.
- 37 S. Tiwari, R. Prakash, R. J. Choudhary and D. M. Phase, J. Phys. D: Appl. Phys., 2007, 40, 4943-4947.
- 38 D. Zhou, S. Wang, Y. Jia, X. Xiong, H. Yang, S. Liu, J. Tang, J. Zhang, D. Liu, L. Zheng, Y. Kuang, X. Sun and B. Liu, Angew. Chem., Int. Ed., 2019, 131, 736-740.
- 39 W.-S. Yoon, K.-B. Kim, M.-G. Kim, M.-K. Lee, H.-J. Shin, J.-M. Lee, J.-S. Lee and C.-H. Yo, J. Phys. Chem. B, 2002, 106, 2526-2532.
- 40 B. Li and D. Xia, Adv. Mater., 2017, 29, 1701054.
- 41 B. Li, H. Yan, Y. Zuo and D. Xia, Chem. Mater., 2017, 29, 2811.
- 42 B. Li, H. Yan, J. Ma, P. Yu, D. Xia, W. Huang, W. Chu and Z. Wu, Adv. Funct. Mater., 2014, 24, 5112-5118.