Flowery nickel–cobalt hydroxide via a solid–liquid sulphur ion grafting route and its application in hybrid supercapacitive storage†

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In our research, a two-step solid–liquid route was employed to fabricate flowery nickel–cobalt hydroxide with sulphur ion grafting (Ni1Co2–S). The utilization of NaOH/agar and Na2S/agar could efficiently retard the release rates of OH− or S2− ions at the solid–liquid interface due to strong bonding between agar hydrogel and these anions. Ni1Co2–S generally displays ultrathin flowery micro-frame, ultrathin internal nanosheets and expanded pore size. Besides, the introduction of suitable sulphide species into nickel–cobalt hydroxide could improve its conductivity due to the lower band gap of Ni–Co sulphide. The supercapacitive electrode Ni1Co2–S presented capacitance of 1317.8 F g−1 at 1 A g−1 and suitable rate performance (77.9% at 10 A g−1 and 59.3% at 20 A g−1). Furthermore, a hybrid supercapacitor (HSC) was developed utilizing positive Ni1Co2–S and negative activated carbon electrodes. As expected, the HSC device exhibited excellent specific capacitance (117.1 F g−1 at 1 A g−1), considerable energy densities (46.7 W h kg−1 at 0.845 kW kg−1 and 27.5 W h kg−1 even at 9 kW kg−1) and suitable cycling performance, which further illuminated the high energy storage capacity of Ni1Co2–S.

1 Introduction

Numerous electronic equipments such as electric vehicles, consumer electronics and portable devices have facilitated our daily lives, but some adverse issues such as low recyclability and energy storage capacity remain for their extensive applications. Therefore, substantial attention is focused on achieving efficient energy storage of supercapacitors (SCs) and their successful applications in various electronic equipments due to their high power deliveries, long lifespans and rapid charge/discharge rates.1–4 To fully promote energy density and enlarge the potential window, assembling hybrid SCs is a feasible strategy, wherein a redox electrode serves as the power source.5,6 Meanwhile, exploring efficient functional electrode materials plays a vital role for integrated performance of HSCs. As typical battery-type materials,7–9 a class of Ni/Co hydroxides represent multiple oxidation states, high theoretical values and long discharge platforms,10–12 which endow them with high potentials in supercapacitive materials.

The actual capacitance values of electrode materials are always lower than the theoretical values due to many factors such as specific surface area, porosity, conductivity, crystal structure and chemical composition.13–16 Thus, a rational structural design is highly desired to further improve electrochemical performances, especially while fabricating multi-level structures with large specific areas and optimizing components. On the one hand, versatile micro/nanostructures have been constructed and applied in catalysts,17 batteries,18 and supercapacitors.19 Sufficient interior space and large specific surface area expose more active areas and shorten contact distance and thus, the generation of ultrathin nanosheets in flowery structures enhances electrochemical performance. On the other hand, the conductivity or stability performance of Ni/Co hydroxides needs further improvement.20–22 Recently, many studies have reported different forms of S incorporated with Ni/Co hydroxides, because Ni/Co sulphides display lower band gap values than their hydroxides.23–25 For instance, Zhou and co-workers reported a distinct core–shell Ni3S2 nanorod@Ni(OH)2 nanosheet nanoarchitecture that presented an enhanced performance of 1037.5 F g−1 (at 5.1 A g−1) than monocomponent Ni(OH)2, because the hybrid structure-integrated Ni(OH)2 nanosheets with rich active sites and crystalline Ni3S2 nanorods promoted rapid electron transfer.23 Also, core–shell NiCo2S4@Co(OH)2 nanotube arrays were prepared, which showed lower polarization and improved reactivity compared to bare NiCo2S4 structures.24 Recently, a gas–solid reaction route was selected to fabricate novel metallic nickel hydroxide nanosheets utilizing H2S gas to treat β-Ni(OH)2 nanosheets at 110 °C. The introduction of sulphur into Ni(OH)2 efficiently promoted electrical conductivity, and the material exhibited metallic
property. Therefore, slight sulphur ion grafting route has been designed to improve the conductivities of hydroxide samples.

For the synthesis of electrode materials, various methods such as the microwave method, solvothermal/hydrothermal method, electro-deposition reaction and calcination treatment have been reported. In spite of great improvements in the energy conversion of SCs, high costs and unsuitable synthesis processes hinder their further applications. Up to now, few studies have been reported on the NaOH/agar and Na2S/agar hydrogel synthetic method. Herein, we used a two-step solid–liquid method to obtain Ni–Co hydroxide and Ni–Co hydroxide with S2⁻ grating. The distinct agar hydrogel method has the following advantages: (1) the preparation of agar hydrogel only needs a short heat treatment time of 10 min, and the subsequent reactions are carried out at room temperature, which saves energy consumption; (2) there is controllable ionic diffusion in the solid–liquid interface owing to strong bonding between hydrogel and anions (OH⁻ or S2⁻ ions); (3) facile reaction equipments scale up sample preparation. In the hydrogel-assisted reaction, the flowery nickel–cobalt hydroxide via sulphur ions grafting not only introduces small amount of sulphide species, but also expands the pore size, which greatly stimulates charge transport. As anticipated, the Ni1Co2–S electrode and the corresponding hybrid supercapacitor (HSC) device deliver excellent capacitances, appropriate rate performances and cycling stabilities.

2 Experimental section

2.1 Preparation of NaOH/agar and Na2S/agar hydrogel

First, 12 mmol sodium hydroxide (NaOH) and 0.5 g agar powder were transferred to a 100 mL beaker. Then, 10 mL deionized water was added to the beaker. The mixture was homogeneously dispersed via consistent ultrasonication and then heated in a water bath at a temperature of 85 °C for 10 min. The obtained mixture was placed at room temperature to form NaOH/agar hydrogel. Similarly, 0.7 g agar was added to 20 mL 0.03 mol L⁻¹ sodium sulphide (Na₂S) aqueous solution, and the mixture was homogeneously dispersed, heated at 85 °C for 10 min and cooled to room temperature. The obtained hydrogel was marked as Na₂S/agar hydrogel.

2.2 Preparation of Ni1Co2, Ni1Co2-AN, Ni2Co1, Co3 and Ni3

First, 1.5 mmol nickel nitrate hexahydrate (Ni(NO₃)₂·6H₂O), 3 mmol cobalt hexahydrate (Co(NO₃)₂·6H₂O) and 1.5 mmol ammonium chloride (NH₄Cl) were added to a solution containing 15 mL deionized water and 50 mL N,N-dimethylformamide (DMF). The mixture was stirred for several minutes and gradually dropped on NaOH/agar hydrogel. After that, the beaker was placed for 24 h in a dark place without any heat treatment. The precipitate generated was collected, washed using abundant water and ethanol, and heated to dry. The above-mentioned sample was denoted as Ni1Co2. In the absence of NH₄Cl, the obtained precipitate was denoted as Ni1Co2-AN. Similarly, the molar ratios of Ni(NO₃)₂·6H₂O and Co(NO₃)₂·6H₂O were adjusted from 3 : 0, 2 : 1 and 0 : 3, and the other conditions were not changed; hence, the corresponding precipitates were marked as Ni3, Ni2Co1 and Co3.

2.3 Preparation of Ni1Co2–S, Ni1Co2-AN–S, Ni2Co1–S, Ni3–S and Co3–S

First, 50 mg Ni1Co2 precursor was dispersed in 40 mL deionized water with ultrasonicication to form Ni1Co2 suspension. The above-mentioned suspension was dropped on Na₂S/agar hydrogel and reacted at room temperature for 24 h. The precipitate generated was collected, washed using abundant water and ethanol, and heated to dry. The as-obtained composite was marked as Ni1Co2–S. The Ni1Co2 precursor was converted to Ni1Co2–AN, Ni2Co1, Ni3 and Co3 in turn; the samples obtained were denoted as Ni1Co2–AN–S, Ni2Co1–S, Ni3–S and Co3–S, respectively.

2.4 General characterization

On the basis of frequently used characterization technologies such as field emission scanning electron microscopy (FESEM, JEOL6700F) and transmission electron microscopy (TEM, JEM-2100F), superficial microtopographies and internal structures of our samples were studied in detail. Besides, an accelerating voltage of 200 kV was employed in the selected area electron diffraction (SAED) measurement. Wide angle X-ray diffraction (XRD, D8 ADVANCE) was also employed to analyze the phase components of our samples utilizing Cu-Kα radiation with a scanning range from 10° to 80°. Furthermore, X-ray photoelectron spectroscopy (XPS, ESCAlab220i-XL, Al-Kα source) and Raman microspectrometry (Renishaw, inVia-Reflex, 532 nm) tests were performed to gain insights into the valence states of different elements and chemical bonds of our samples. Furthermore, inductively coupled plasma-optical emission spectrometry (ICP-OES, SPECTRO CIROS VISION) was used to analyze the accurate mole ratio of metal anions. Fourier transform infrared (FTIR) spectrometry was carried out to study the chemical bonds in our samples. For obtaining the Barrett–Joyner–Halenda (BJH) mode, a Micromeritics ASAP-2020M apparatus was operated.

2.5 Electrode preparation and electrochemical measurements

The preparatory method of positive electrodes based on Ni–Co nanocomposites was as follows: first, we ground active materials poly(vinylidene fluoride) (PVDF) and acetylene black (AB) in a mass ratio of 85 : 5 : 10 for several minutes. The mixture was added to a suitable amount of N-methyl-2-pyrrolidone (NMP) solvent and stirred constantly to generate a uniformly distributed ropy slurry. Then, the ropy slurry was uniformly applied on a porous Ni foam (effective area of 1 cm² was controlled) and dried at 110 °C for 6 h in a vacuum oven. A series of electrodes were prepared including Ni1Co2, Ni1Co2-AN, Ni2Co1, Ni3, Co3, Ni1Co2–S, Ni1Co2–AN–S, Ni2Co1–S, Ni3–S and Co3–S. For effective comparison, we controlled the mass loadings of the as-mentioned electrodes to 1.2 ± 0.2 mg cm⁻². An active carbon (AC) electrode was also prepared utilizing...
a similar method except that we changed the mass ratio of the mixture (AC : PVDF : AB = 90 : 5 : 5). The mass loading of the AC electrode was controlled in the range of 2.9 ± 0.1 mg cm⁻². In the Ni₁Co₂//AC HSC device, we utilized an Ni₁Co₂ electrode as the positive electrode and AC positive electrode as the negative electrode. The total mass loading of the Ni₁Co₂–S/AC device was controlled in the range 4.1 ± 0.3 mg.

3 Results and discussion

3.1 Characterization of morphology, microstructure and composition

The overall fabrication procedure of flowery Ni₁Co₂ precursor is displayed in Fig. 1. OH⁻ ions were slowly transferred through the solid–liquid interface due to strong bonding between OH⁻ ions and agar hydrogel. These OH⁻ ions were slowly incorporated with Ni²⁺ and Co²⁺ ions, thus generating binary z-(Ni–Co) hydroxide. After that, the Na₂S/agar hydrogel slowly provided S²⁻ ions for partial incorporation with metal ions from nickel–cobalt hydroxide. A small amount of sulphur ions was grafted on Ni₁Co₂ hydroxide (Ni₁Co₂–S).

As shown in Fig. 2a and S1a and b,† Ni₁Co₂ displayed well-proportioned flowery clusters (diameter up to several micrometers) and ultrathin internal nanosheets. In comparison, the Ni₁Co₂–AN sample showed irregular nanoflowers with the coexistence of small sheets and large sheets (in Fig. S2a and b†), because there was no chelation between NH₄⁺ and OH⁻ ions. Furthermore, the mole ratio of Ni(NO₃)₂ : Co(NO₃)₂ was adjusted from 3 : 0, 2 : 1 to 0 : 3 to optimize the proportion. Staked nanoflowers with insufficient space (Ni₂Co₁ or Co₃) and conglobulated nanoparticles (Ni₃) are exhibited in Fig. S2c–h† indicating that various mole ratios of metal nitrates could significantly influence the ionic reaction and tune the microtopography. On the basis of the accurate ICP-OES data in Table S1,† the Ni : Co ratios of Ni₁Co₂, Ni₁Co₂–AN, Ni₃, Ni₂Co₁ and Co₃ were 1 : 2.27, 1 : 2.26, 3 : 0, 1.86 : 1 and 3 : 0, respectively, which matched the dosages of metal nitrates, and this indicated a uniform co-precipitation reaction.

Subsequently, we utilized Na₂S/agar hydrogel to grate S²⁻ ions on Ni₁Co₂ and obtain Ni₁Co₂–S. The lower solubilities of metal sulphides than those of hydroxides caused ion exchanges between OH⁻ and S²⁻ ions. Fig. S3† reveals the colour change from green (Ni₁Co₂) to grey green (Ni₁Co₂–S), which supported the occurrence of S²⁻ ion grating processes. In Fig. 2b and S1c and d† we can see that the pore sizes of Ni₁Co₂–S extended with the partial damage of pristine flowery frames. Furthermore, the BJH modes of Ni₁Co₂ and Ni₁Co₂–S were also obtained to study the details of pore size distribution. As shown in Fig. S4,† the main pore size of Ni₁Co₂ was concentrated in a narrow range from 1.7 to 7.5 nm, whereas that of Ni₁Co₂–S exhibited a wider range from 1.7 to 50 nm. This clear broadening of pore size further revealed a structural change after S²⁻ ion grating. Fig. 2c displays the transparent structure of Ni₁Co₂–S, and Fig. S5† presents nanosheet thicknesses (mainly ranging from 2.2 to 5.3 nm), and these results revealed the ultrathin property of Ni₁Co₂–S. In the inset of Fig. 2c, the diffraction circles reveal the polycrystalline property of Ni₁Co₂–S. In Fig. 2d, the lattice fringes of 0.258, 0.263 and 0.266 nm resulted from the (102) planes of nickel–cobalt hydroxide. Notably, many internal pores and defects could be observed in Fig. 2d such as the red elliptical area, which probably could be ascribed to the grating effect of S²⁻ ion. Furthermore, the elemental mapping of Ni₁Co₂–S is provided to illuminate element distribution. As depicted in Fig. 2e, Ni, Co, S and O uniformly existed in the flowery Ni₁Co₂–S architecture. Based on energy dispersive spectrometry (EDS) data shown in Fig. S6,† the mole ratio S : O was found to be 1 : 5.14, indicating that
Ni1Co2–S was composed of Ni–Co hydroxides with slight S\(^2-\) grating. Also, the mole ratio of Ni : Co reached 1 : 1.89, which also generally coincided with the ratio of 1 : 2 (dosage of metal nitrates).

In Fig. 3, the main peaks of Ni3 are directly indexed to \(\alpha\)-Ni hydroxide with turbostratic disorder, and the peaks of Co3 also present the property of \(\alpha\)-Co hydroxide.\(^{33,34}\) Meanwhile, the peaks of Ni1Co2, Ni1Co2-AN and Ni2Co1 are analogous with those of Ni3 and Co3, indicating the formation of binary \(\alpha\)-(Ni–Co) hydroxide. To further study the structures of these hydroxides, their FTIR patterns are displayed in Fig. S7a.\(^{†}\) In samples of Ni1Co2, Ni1Co2-AN, Ni2Co1, Ni3 and Co3, a series of broad peaks centred at 3400–3500 cm\(^{-1}\) were assigned to the hydrogen bonding with intercalated water molecules. Besides, the peaks at 1000–1500 cm\(^{-1}\) were ascribed to intercalated NO\(_3^−\) ions, and the peaks at 400–700 cm\(^{-1}\) resulted from M (Ni/Co)–O–H bending and M–O stretching vibrations. These FTIR results demonstrated the \(\alpha\)-hydroxide features of Ni1Co2, Ni1Co2-AN, Ni2Co1, Ni3 and Co3.\(^{33–35}\) Furthermore, the Raman spectra of these samples supported the chemical bonds of hydroxide precursors. In Fig. S8,\(^{†}\) the Raman peaks of Co3 located at 184.6, 464.0, 507.6, 599.8 and 666.2 cm\(^{-1}\) coincided with the features of \(\alpha\)-Co(OH)\(_2\).\(^{36–38}\) In the Raman pattern of Ni3, the peaks at 304.2, 400–600, and 1093.9 cm\(^{-1}\) corresponded to Ni–O stretching, E-type vibration of the Ni–OH lattice and hydroxyl group vibration mode of the Ni(OH)\(_2\) phase.\(^{39,40}\) In contrast, Ni1Co2, Ni1Co2-AN and Ni2Co1 maintained the peak locations of Ni3 and Co3 with some peak shifts, indicating the existence of binary Ni–Co hydroxide. Therefore, we successfully fabricated a series of unitary \(\alpha\)-(Ni/Co) or binary Ni–Co hydroxides.

In the XRD pattern shown in Fig. 4a, Ni1Co2–S displays similar peaks to Ni1Co2, demonstrating that Ni1Co2–S mainly contains \(\alpha\)-(Ni–Co) hydroxide. Compared to peaks from Ni1Co2, a clear negative shift (ca. 0.6 to 0.9°) can be observed in the XRD pattern of Ni1Co2–S because the slight S\(^2-\) grating reaction extends the internal planar spacing and pore size. No clear signals of Ni–Co sulphide can be detected because of weak crystallinity and low content of sulphide at room-temperature. By comparing the FTIR and Raman patterns of Ni1Co2 and Ni1Co2–S (in Fig. S7b and 4b\(^{†}\)), we further ascertain the \(\alpha\)-(Ni–Co) hydroxide component in Ni1Co2–S.

We utilized the XPS patterns of Ni1Co2 and Ni1Co2–S to evaluate the variation before and after S\(^2-\) grafting. All spectra were calibrated based on the C 1s benchmark of 284.6 eV. In the Ni 2p spectra of Ni1Co2 (Fig. 4c), the peaks at 856.33 and 873.95 eV corresponded to Ni 2p\(_{3/2}\) and 2p\(_{1/2}\) with a separation of 17.62 eV, which was in line with the feature of the Ni(OH)\(_2\) phase.\(^{41,42}\) In Fig. 4d, Ni1Co2 displays Co 2p\(_{3/2}\) and 2p\(_{1/2}\) peaks at 781.51 and 797.29 eV with a separation of 15.78 eV, indicating the co-existence of divalent and trivalent states of Co.\(^{43,44}\) The divalent and trivalent states of Co co-existed in \(\alpha\)-(Ni–Co) hydroxides. For Ni1Co2–S, the energy-bond separations of Ni 2p and Co 2p levels were 17.67 and 16.05 eV, respectively, verifying slight reduction of partial cobalt ions during the ion exchange process. In Fig. 4e, the peaks at 162.63 and 163.77 eV represented S 2p\(_{3/2}\) and 2p\(_{1/2}\) levels, respectively, and a satellite peak at 168.89 eV was ascribed to S\(_4\)O\(_6\)\(^{2-}\) ions with high oxidation states on the surface. Similar S 2p spectra could be found in many reported nickel/cobalt sulfides,\(^{45–47}\) which suggested...
successful grating of $S^{2−}$ ions in Ni1Co2–S. In contrast to the observations for Ni1Co2, Ni 2p, Co 2p and O 1s levels in the XPS spectrum of Ni1Co2–S clearly shifted to more positive levels, which was mainly due to electronic interactions involving S and Ni/Co components.\textsuperscript{25,48}

3.2 Electrochemical capacitive behaviors of various electrodes

To select an efficient hydroxide precursor, we investigated the performances of Ni1Co2, Ni1Co2-AN, Ni3, Ni2Co1 and Co3 in an alkaline three-electrode system. Fig. 5a shows the CV curves of Ni1Co2, Ni1Co2-AN, Ni3, Ni2Co1 and Co3 at 5 mV s\textsuperscript{−1}. The main contents of Ni1Co2, Ni1Co2-AN, Ni3, Ni2Co1 and Co3 were unitary Ni/Co or binary Ni–Co hydroxides. The energy storage mechanism could be described as follows:\textsuperscript{49–54}

\begin{align}
\text{Ni(OH)}_2 + \text{OH}^- & \rightarrow \text{NiOOH} + e^- \tag{1} \\
\text{Co(OH)}_2 + \text{OH}^- & \rightarrow \text{CoOOH} + e^- \tag{2} \\
\text{CoOOH} + \text{OH}^- & \rightarrow \text{CoO}_2 + e^- \tag{3}
\end{align}

A smaller potential disparity ($\Delta E_{\text{a,c}}$, the disparity between the anode and cathode peak positions) is presented in the CV curve of Ni1Co2, suggesting slight electrode polarization.\textsuperscript{35,56} Additionally, Ni1Co2 delivers longer discharge time, as shown in Fig. 5b, and better capacitance performance, as shown in Fig. 5c, thus highlighting the enhanced performance of the Ni1Co2 electrode. These results are mainly ascribed to the ultrathin flowery structure, providing more active contact sites for charge transfer. In the EIS spectra (Fig. 5d), the intersections of semicircles on the real axis, semicircle diameter at high frequency and slope of a line in the low-frequency region are employed to evaluate the equivalent series resistance ($R_s$), charge-transfer resistance ($R_{ct}$) and diffusion resistance ($Z_d$).\textsuperscript{27,58} The $R_s$ values of Ni1Co2, Ni1Co2-AN, Ni2Co1, Co3 and Ni3 electrodes are 2.90, 2.05, 4.38, 1.74 and 1.65 $\Omega$, respectively. In comparison, even though Ni1Co2 possesses higher $R_s$ values, the smaller semicircle diameter and steeper slope indicate lower $R_s$, $R_{ct}$ and $Z_d$ values, thus revealing enhanced performance.

To further promote the performance of Ni1Co2, controllable $S^{2−}$ grafting was carried out to obtain Ni1Co2–S. The CV curves of Ni1Co2 and Ni1Co2–S at various scanning rates from 5 to 50 mV s\textsuperscript{−1} are presented in Fig. 6a–c. Compared to that of Ni1Co2, the extended CV area of Ni1Co2–S revealed enhanced capacitance. Interestingly, compared to the observations for Ni1Co2, there was a pair of anodic peaks that disappeared for Ni1Co2–S electrode. This peak disappearance actually resulted due to closer location of various pairs of oxidation and reduction peaks, leading to overlapping.\textsuperscript{59,60} In the CV curves of

![Fig. 5](image-url) (a) CV curves at 5 mV s\textsuperscript{−1}, (b) GCD curves at 5 A g\textsuperscript{−1}, (c) specific capacitances at various current densities and (d) EIS spectra of Ni1Co2, Ni1Co2-AN, Ni2Co1, Co3 and Ni3 electrodes.

![Fig. 6](image-url) (a) CV curves of Ni1Co2 and Ni1Co2–S at 5 mV s\textsuperscript{−1}; (b) CV curves of Ni1Co2 at different scanning rates; (c) CV curves of Ni1Co2–S at different scanning rates; (d) linear fit of $\log I_{ap}$--$\log n$ scatters from three pairs of peaks in the CV curves of Ni1Co2 and Ni1Co2–S; (e) GCD curves of Ni1Co2 and Ni1Co2–S at 1 A g\textsuperscript{−1}; (f) GCD curves of the as-fabricated Ni1Co2–S at different current densities; (g) capacitance performances of Ni1Co2, Ni1Co2–S, Ni1Co2-AN–S, Ni2Co1–S, Co3–S and Ni3–S electrodes at various current densities; (h) 6000 cycle duration measurements of Ni1Co2 and Ni1Co2–S at the current density of 10 A g\textsuperscript{−1}.
Ni2Co1–S (Fig. S9†), there were many pairs of oxidation and reduction peaks, indicating incomplete overlapping of these peaks. Furthermore, the scatters of log $I_p$–log $v$ from the three pairs of oxidation peaks are fitted in Fig. 6d, wherein $I_p$ and $v$ stand for scanning rate and peak current, respectively. Based on a previous study, the relationship between $I_p$ and $v$ is in line with the formula $I = av^b$, wherein $a$ and $b$ represent empirical parameters; $b$ value of 0.5 is due to the diffusion-dominated reaction, whereas $b$ value of 1 originates from the ideal capacitive behaviour. In our study, the peak 1 and peak 2 from Ni1Co2 exhibited higher $b$ values of 0.63 and 0.84, respectively, and the peak 3 from Ni1Co2–S exhibited a lower $b$ value of 0.55, indicating that the capacitances of Ni1Co2 and Ni1Co2–S contained diffusion and capacitive contributions.

As shown in Fig. 6e and f, Ni1Co2–S displayed low discharge platform, which was in agreement with the result of CV curves. We further calculated the specific capacitances of our electrodes based on different GCD curves from 1 to 20 A g$^{-1}$. As shown in Fig. 6g, Ni1Co2–S possessed enhanced specific capacitances (1317.8 F g$^{-1}$ at 1 A g$^{-1}$ and 781.3 F g$^{-1}$ at 20 A g$^{-1}$) relative to those of Ni1Co2 (789.5 F g$^{-1}$ at 1 A g$^{-1}$ and 489.4 F g$^{-1}$ at 20 A g$^{-1}$). When the Ni1Co2–S electrode was at high current density of 10 or 20 A g$^{-1}$, it still displayed suitable capacitance retention of 77.9% or 59.3%, respectively. The electrochemical capacitances of Ni1Co2–AN–S, Ni2Co1–S, Ni3–S and Co3–S were also studied. Similarly, S$^{2−}$ ion grating enhanced the capacitance compared with that of the corresponding precursors. Ni1Co2–S displayed excellent capacitance and high rate performance. Therefore, the selection of precursor and further S$^{2−}$ ion grating efficiently promoted the comprehensive performances of our electrode materials.

Besides, long-term cycling duration is desired for reversible applications; hence, the cycling measurement of Ni1Co2–S was operated at a current density of 10 A g$^{-1}$. In Fig. 6h, after 6000 cycles, Ni1Co2–S presented a high capacitance of 804.3 F g$^{-1}$ and capacitance retention of 72.5%, which was higher than that of Ni1Co2 (58.8%). As shown in Fig. 7, Ni1Co2–S had lower semicircle diameter and steeper slope (lower $R_{ct}$ and $Z_w$), further highlighting the improved resistance performance. $R_{ct}$ and $Z_w$ significantly increased after 6000 cycles, probably due to the damage of internal nanosheets of Ni1Co2–S (in Fig. S10†). In view of the results mentioned above, the enhancement in the electrochemical performance of Ni1Co2–S was mainly due to the following factors: (1) suitable ratio of Ni : Co could tune the nanostructure and promote electrochemical performance; (2) S was introduced into Ni–Co hydroxide, and it greatly improved the conductivity; and (3) S$^{2−}$ ion grafting generated many defects and extended the pore size.

3.3 Electrochemical capacitive behaviour of HSC device
Considering that general three-electrode systems have narrow potential windows, we prepared an HSC device to enlarge the voltage using Ni1Co2–S and AC electrodes as positive and negative electrodes, respectively, as shown in Fig. 8a. The CV curves shown in Fig. 8b present large quasi-rectangular shapes with few weak redox peaks, which suggest the battery-type and electrical double-layer hybrid energy-storage mechanisms. Fig. 8c displays various linear GCD profiles with long discharge time, further highlighting its good capacitance performance. As depicted in Fig. 8d, Ni1Co2–S//AC delivered great specific capacitance values of 117.1, 106.3, 97.5, 82.3, 74.5 and 61.1 F g$^{-1}$ at various current densities of 1, 2, 3, 5, 7 and 10 A g$^{-1}$, respectively. Unfortunately, with the increase in current density, the specific capacity gradually faded, and the total voltage of the HSC also increased due to insufficient electrochemical reaction at high current density. Similar to the study of the three-electrode system, we also studied the cycling stability of Ni1Co2–S//AC with charge/discharge current density of 3 A g$^{-1}$ (in Fig. 8e). The initial electrochemical measurement contributed to gradual wetting between the electrode and electrolyte. The trapped OH$^−$ ions then gradually diffused out, and the contact area between the electrode and electrolyte was enlarged.
thus causing capacitance enhancement during the initial 1000 cycles. Subsequently, the capacitance of Ni1Co2–S//AC slowly faded and reached 82.4% after 7000 cycles at 3 A g\(^{-1}\), resulting from partial damage to the Ni1Co2–S structure.

The relationship between energy density and power density in Ni1Co2–S//AC is depicted in the Ragone diagram (Fig. 8f). The HSC device achieved a high energy density of 46.7 W h kg\(^{-1}\) at a power density of 0.845 kW kg\(^{-1}\) and also reached 27.5 W h kg\(^{-1}\) at 9 kW kg\(^{-1}\), which were comparable to the results of many peer HSC devices based on core–shell NiCoS\(_2\)@Co(OH)\(_2\) nanotube arrays,\(^{24}\) NiCoS\(_4\) nanosheets,\(^{44}\) and other cobalt/nickel hydroxide electrode materials.\(^{65–68}\) To trigger four light emitting diodes (LEDs, red, 5 mm), we further assembled two Ni1Co2–S//AC devices in series and charged them at 3 A g\(^{-1}\), as exhibited in Fig. 6g. As anticipated, the LED bulbs worked for a long duration of 2 min and displayed dim light after 5 min. These results suggested that Ni1Co2–S can serve as a promising energy storage material.

4 Conclusions

Herein, we designed a simple solid–liquid interface reaction to synthesize Ni1Co2–S using NaOH/agar and Na\(_2\)S/agar gels. Ni1Co2–S was mainly composed of α-(Ni–Co) hydroxide with S\(^2–\) ion grating. The slight ion exchange reaction between S\(^2–\) ions and Ni1Co2 introduced a small amount of sulphide to improve its conductivity and expanded the internal pore size, thus ensuring rapid charge transfer. Compared to other Ni1Co2, Ni1Co2–AN, Ni2Co1, Co3 and Ni3 electrodes, the Ni1Co2–S electrode delivered improved capacitance of 1317.8 F g\(^{-1}\) at 1 A g\(^{-1}\), suitable rate performance (77.9% at 10 A g\(^{-1}\) and 59.3% at 20 A g\(^{-1}\)) and improved cycling stability. The Ni1Co2–S//AC device also exhibited a high energy density of 46.7 W h kg\(^{-1}\) at 0.845 W kg\(^{-1}\) and long cycling stability, which further demonstrated the potential advantage of Ni1Co2–S material in supercapacitive storage.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was financially supported by the National Natural Science Foundation of China (Grant No. 51501068) and Graduate Innovation Fund of Jilin University (Grant No. 2017183).

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