CONTROLLABLE MORPHOLOGY OF SEA-URCHIN-LIKE NICKEL–COBALT CARBONATE HYDROXIDE AS A SUPERCAPACITOR ELECTRODE WITH BATTERY-LIKE BEHAVIOR

Nutthapong Poompiew, Prasit Pattananuwat,* and Pranut Potiyaraj*

ABSTRACT: Nickel–cobalt carbonate hydroxide with a three-dimensional (3D) sea-urchin-like structure was successfully developed by the hydrothermal process. The obtained structure enables the enhancement of charge/ion diffusion for the high-performance supercapacitor electrodes. The mole ratio of nickel to cobalt plays a vital role in the densely packed sea-urchin-like structure formation and electrochemical properties. At optimized nickel/cobalt mole ratio (1:2), the highest specific capacitance of 950.2 F g⁻¹ at 1 A g⁻¹ and the excellent cycling stability of 178.3% after 3000 charging/discharging cycles at 40 mV s⁻¹ are achieved. This nickel–cobalt carbonate hydroxide electrode yields an energy density in the range of 42.9–15.8 Wh kg⁻¹, with power density in the range of 285.0–2849.9 W kg⁻¹. The charge/discharge mechanism at the atomic level as monitored by time-resolved X-ray absorption spectroscopy (TR-XAS) indicates that the high capacitance behavior in a nickel–cobalt carbonate hydroxide is mainly dominated by cobalt carbonate hydroxide.

1. INTRODUCTION

Since electrical energy plays an important role in daily routines, requiring a fast charge feature, lightweight, high electrical storage capacity, and long-period usage.1,2 Unlike the battery such as Li-ion3,4 and hybrid type,5 supercapacitors (SCs) are the most attractive energy storage technology due to high power efficiency and long life cycles.6 Different redox-active materials such as polyaniline,7 polypyrrole,8 and metal oxide have been promoted as active materials for pseudocapacitors (PCs) due to high theoretical specific capacitance, high mechanical properties, and high stability.9 Transition-metal oxides with multiple oxidation states have been widely investigated for pseudocapacitor, such as bis(selanylidene)-rhodium,10 manganese dioxide,11 molybdenum disulfide,12 manganese hexacyanoferrate,13 titanium dioxide,14 vanadium oxide,15 copper and nickel hexacyanoferrate,16 cobalt sulfide,17 cobalt selenide,18,19 cobalt ferrate,20 cobalt oxide,21 nickel hydroxide,22 nickel cobaltite,23 etc. Many research efforts have been devoted to the nickel–cobalt complex due to high theoretical specific capacitance values, low cost, and long life cycles.24 Numerous attempts have been made to synthesize the nickel–cobalt complex such as direct oxidation and reduction,25 sol–gel method,24 electrodeposition,26 spraying process,27 chemical vapor deposition28 and hydrothermal process.29 The hydrothermal process is an efficient way to create a new class material of metal oxide with the defined morphologies. Supercritical water formed under high pressure provides an excellent reaction environment for the crystalization of metal oxide particles.30 For example, the flower-like structure of nickel–cobalt oxide was synthesized by the hydrothermal technique with hexamethylenetetramine (HMTA), yielding the specific capacitance of 750 F g⁻¹ at 1 A g⁻¹.31 Nanosheet structure of the nickel–cobalt layered double hydroxide was obtained by two steps of hydro/solvothermal process, demonstrating the high specific capacitance of 1497 F g⁻¹ at 5 mA cm⁻².32 Porous nickel–cobalt oxide nanowires via hydrothermal process yielded an excellent specific capacitance value of 1479 F g⁻¹ at 1 A g⁻¹.33

One important parameter for controlling the morphology and microstructure of the nickel/cobalt alloy complex is the modulating Ni/Co ratio. However, the nanostructured nickel/cobalt alloy complex synthesis is hampered by the numerous complicated steps required and the difficulty in controlling phase morphology. As a result, the findings of the morphology-controlled synthesis of the Ni–Co binary hydroxide for high-performance supercapacitors are still widely unknown. Furthermore, despite numerous studies of nickel and cobalt composites as SCs, the knowledge of the charge–discharge storage mechanisms remains hazy. To understand the behavior
at the atomic level, a charge/discharge mechanism analysis using time-resolved X-ray absorption spectroscopy (TR-XAS) is also proposed.

2. RESULT AND DISCUSSION

2.1. Morphology Study. Scanning electron microscopy (SEM) images of the as-prepared nickel−cobalt at different ratios of Ni to Co salt solution, as shown in Figure 1, show remarkable differences in surface morphology in each sample, indicating that the morphology of the as-prepared Ni−Co complex is sensitive to the mole ratio of Ni to Co for the hydrothermal process. As seen in Figure 1a, the as-prepared nickel carbonate hydroxide demonstrates the growth of three-dimensional (3D) flower-like microstructures assembled from nanoflake structures. Meanwhile, cobalt carbonate hydroxide exhibits a nanoflake-like structure in a rectangular shape as shown in Figure 1b. At a high ratio of Ni to Co (2 to 1), the Ni−Co complex shows a sea-urchin-like structure, originating from the densely packed nanoneedle in a 3D core structure (Figure 1c). The less-dense nanoneedle packing in 3D sea-urchin-like structure is observed at the Ni/Co ratio of 1:1 and 1:2 as seen in Figure 1d,e, respectively. The imperfect urchin-like shape of the Ni−Co complex occurs with the nanonoodle-like structure at a high ratio of Co to Ni (2 to 1), implying that the growth of needle shape in the core structure of sea urchin is disturbed with increasing cobalt content in the Ni/Co ratio.

Figure 1. Representative SEM micrographs of (a) pure Ni, (b) pure Co, (c) NiCo2/1, (d) NiCo1/1, and (e) NiCo1/2 scale at 1 μm and (f) elemental mapping of NiCo1/2 sample for (f-i) nickel, (f-ii) cobalt, (f-iii) oxygen, and (f-iv) carbon element.

The element composition analysis of samples at different Ni/Co ratios is further investigated. Figure 1f presents the element mapping of the Ni/Co1/2 sample, confirming the presence of Ni, Co, O, and C elemental distribution on the surface of the NiCo1/2 sample (Figure 1f-i–f-iv). Furthermore, the energy-dispersive X-ray spectrometry (EDS) for the Ni−Co complex oxide samples at different Ni-to-Co ratios were analyzed as shown in Table 1. The designed feeding for the Ni/Co mole ratios of 2:1, 1:1, and 1:2 samples reveal the actual Ni/Co ratios of 1:1, 1:1.5, and 1:3.5, respectively. This result indicates that the high reactivity of cobalt ion is superior to nickel ion in our experiment.

The surface area characteristic of all samples is investigated using the N₂ adsorption/desorption isotherms as shown in Figure 2. All samples can be categorized as an isotherm of type II with a hysteresis loop of type H3, indicating a typical macroporous structure of different sizes. The as-prepared NiCo1/2 exhibits the highest N₂ gas absorption, implying the

| sample          | Ni (%)   | Co (%)   |
|-----------------|---------|---------|
| NiCo2/1         | 13.12 ± 0.14 | 14.6 ± 0.13 |
| NiCo1/1         | 13.40 ± 0.16 | 20.18 ± 0.17 |
| NiCo1/2         | 5.01 ± 0.08  | 17.72 ± 0.13 |

Table 1. Percentage of Elemental Atom from EDS Results
highest surface area of the nanoneedle sea-urchin-like structure. The specific surface area and porosity parameter are calculated as shown in Table S2. Obviously, the specific surface area of NiCo1/2 exhibits the highest area of 24.84 m² g⁻¹ with the smallest average pore size of 155.3 Å. The Brunauer–Emmett–Teller (BET) results are in agreement with the SEM results, demonstrating that the decrease in needle density of 3D sea-urchin-like structure can facilitate an ion/charge diffusion pathways for electrochemical active surface area (ECSA).

2.2. Characterization. X-ray diffraction (XRD) technique was performed to investigate the crystal structure of the Ni–Co samples at different Ni/Co ratios, as shown in Figure 3.

The as-prepared pristine Ni exhibits the strong diffraction peak at 12.8, 25.8, 33.2, 36.4, and 59.5°, according to the (040), (260), (221), (071), and (0131) crystal planes of nickel carbonate hydroxide (PDF no. 00-029-0868), respectively.35 The as-prepared pristine Co reveals the diffraction peaks at 9.8, 17.3, 24.0, and 34.6° corresponding to PDF no. 00-048-0083, ascribed to cobalt carbonate hydroxide.36 For all diffraction patterns of the as-prepared Ni–Co samples, the XRD patterns reveal the diffraction peak of both nickel carbonate hydroxide and cobalt carbonate hydroxide. Compared with pristine Ni and Co, nickel carbonate hydroxide and cobalt carbonate hydroxide exhibit a gradual shift in the XRD pattern, resulting from the crystal intercalation of cobalt carbonate hydroxide and nickel carbonate hydroxide. Furthermore, the intensity of each characteristic peak relatively depends on the amount of initial Ni²⁺ and Co²⁺. In other words, as the Ni/Co ratio decreases, the peak of nickel carbonate hydroxide becomes smaller and narrower.

The further chemical compositions analysis of the Ni/Co ratio at 1:2 is investigated by X-ray photoelectron spectroscopy (XPS) technique. Figure 4a reveals the survey XPS spectra of nickel—cobalt carbonate hydroxide sample at ratio of Ni/Co of 1:2, confirming the presence of Ni, Co, O, and C atoms. Two spin-orbitals of Ni 2p at Ni 2p½ and Ni 2p½, with two satellite peaks (denoted as “Sat.”) are identified with binding energy at 856.6 and 874.2 eV, respectively, described as Ni²⁺ (seen in Figure 4b).35 Two spin-orbitals peaks of Co 2p½ and Co 2p½, are illustrated in Figure 4c. The peaks at 798.5 and 782.9 eV correspond with Co 2p½ and 2p½ of Co³⁺. In addition, the peaks at 792.7 and 776.1 eV are attributed to Co 2p½ and 2p½ of Co⁴⁺.37 Two species of oxygen are observed in Figure 4d. The higher intensity peak at 533.0 eV can be ascribed as a typical metal carbonate species while the peak at 532.1 eV is designated to hydroxyl groups.38 The deconvoluted Ni 2p, Co 2p, and C 1s spectra (Figure 4e) are observed at 283.6 and 285.3 eV, corresponding with the characteristic binding energy of Ni carbonate and Co carbonate species, respectively. The extra peak at 287.6 and 289.0 is ascribed to the carbon hydroxyl and carbonate species, respectively.34,35,37 Thus, these results suggest the main valence state of nickel—cobalt complex is Ni²⁺, Co³⁺, and Co⁴⁺, which is in good agreement with the previous report of the mixed nickel—cobalt carbonate hydroxide.

In addition, the comparative XPS spectra of the mixed nickel—cobalt carbonate hydroxide samples at different Ni-to-Co ratios were observed. The peak positions of Ni 2p for all mixed Ni–Co carbonate hydroxide samples reveal the almost same (Figure 5a—c), implying the similar chemical compositions of Ni²⁺. However, for Co 2p pattern, the presence of the peak intensities at 776.1 eV for Co 2p½ and 792.7 eV for Co 2p½ is manifestly increased with decreasing Ni/Co ratio (Figure 5e,f). This result suggests the high activity of the formation of Co(III) carbonate at a high cobalt concentration. The difference in peak intensities are also recognized, corresponding to the Ni/Co ratio in each sample. It can be seen that the peak intensity ratio of Ni 2p/Co 2p tends to decrease with increasing cobalt content (Table S1).

2.3. Electrochemical Properties Results. The comparative CV study has been used to evaluate the influence of the Ni/Co ion ratio on the electrochemical characteristics of the sample electrodes. As seen in Figure 6a, all sample electrodes exhibit the pair redox peak of a reversible Faradic redox reaction, confirming that the charge storage mechanism of nickel–cobalt carbonate hydroxide is mainly conducted by pseudocapacitance behaviors. Obviously, the diverse redox characteristic and electrochemical performance of all of the as-prepared electrodes reveal different Faradic characteristics, resulting from the distinct proportional ratio of Ni to CO in nickel–cobalt carbonate hydroxide.39 For pristine nickel oxide electrodes, the two obvious peaks of oxidation and reduction
reaction peaks are observed at the corresponding peak potentials of 0.28 and 0.18 V, respectively. This result may occur from the formation of the Ni$_2$(OH)$_2$CO$_3$ $\rightleftharpoons$ NiOOH during the CV cycling in KOH aqueous solution. The pure cobalt carbonate hydroxide electrode reveals one anodic peak at 0.05 V and two cathodic peaks at $-0.15$ and 0.35 V, caused by the phase transformation of Co(CO$_3$)$_{0.5}$(OH) $\rightleftharpoons$ CoOOH $\rightleftharpoons$ CoO$_2$. CV curves of mixed nickel−cobalt carbonate hydroxide electrodes exhibit a similar shape but shift in their redox position, indicating that the capacitive response related to the ratio of M−O, M−O−OH, and M carbonate hydroxide (M represents Ni or Co) in composite electrodes. The plausible redox mechanisms of nickel and cobalt carbonate hydroxide are illustrated in eqs 1−3.

\[
\text{Ni}_2(\text{OH})_2\text{CO}_3 + 4\text{OH}^- \quad \rightleftharpoons \quad \text{NiOOH} + \text{NiCO}_3 + 2\text{H}_2\text{O} + 2\text{e}^- 
\]

(1)

\[
2\text{Co(CO}_3)_{0.5}(\text{OH}) + \text{OH}^- \quad \rightleftharpoons \quad \text{CoOOH} + \text{CoCO}_3 + \text{H}_2\text{O} + \text{e}^- 
\]

(2)

\[
\text{CoOOH} + \text{OH}^- \rightleftharpoons \text{CoO}_2 + \text{H}_2\text{O} + \text{e}^- 
\]

(3)
Principally, the cyclic CV curve area is proportional to the electrochemically active surface area, indicating a sufficient number of electroactive sites at heterointerfaces during redox reactions. The precursor ratio of Ni to Co salt for the hydrothermal process is the key factor in influencing the electrocapacitance properties of the sample electrodes. Manifestly, the nickel−cobalt carbonate hydroxide electrode at the Ni-to-Co ratio of 1:2 demonstrates the highest CV area, implying the highest electrochemical active surface area relative to the other electrode samples. In contrast, the pure nickel carbonate hydroxide electrode reveals the lowest CV area, implying the lowest capacitance performance. The enhanced performance of NiCo1/2 due to the presence of the mixed nickel−cobalt carbonate hydroxide may be associated with the increase in conductivity related to electrochemical properties. The highest current density of the CV curve for NiCo1/2 is ascribed to the conversion of M (OH)\textsubscript{2} CO\textsubscript{3} to M−O−OH and M−CO\textsubscript{3}, which has gained more electron-transfer reaction as mentioned in eqs 1−3. The calculated specific capacitance by CV of nickel−cobalt carbonate hydroxide at different Ni-to-Co ratios of 1:0, 2:1, 1:1, 1:2, and 0:1 are 19.3, 153.7, 480.0, 735.33, and 329.33 F g\textsuperscript{−1} at 10 mV s\textsuperscript{−1}, respectively. Figure 6b shows the specific capacitance calculated by the CV curve of nickel−cobalt at different ratios as a function of the scan rate from 10 to 100 mV s\textsuperscript{−1}. It is clear that the nickel−cobalt carbonate hydroxide electrode at a ratio of 1:2 also exhibits high rate performance, maintaining the high specific capacitance of more than 400 F g\textsuperscript{−1} even at a high scan rate of 100 mV s\textsuperscript{−1}.

The GCD results of all sample electrodes at a current density of 1 A g\textsuperscript{−1} with the potential window ranging from −0.2 to 0.37 V are presented in Figure 6c. The GCD evidence of all samples displays a plateau curve with low internal resistance (IR drop), indicating a good capacitance behavior. Among them, the nickel−cobalt carbonate hydroxide electrode at the ratio of 1:2 demonstrates the longest plateau discharging time, indicating the highest specific capacitance. An increase in the cobalt-to-nickel ratio leads to a decrease in the internal resistance in electrode samples, indicating that electrical capacitance improves as the cobalt component increases. The calculated specific capacitance from the GCD results of electrode samples at different Ni-to-Co ratios of 1:0, 2:1, 1:1, 1:2, and 0:1 are 2.4, 93.5, 584.1, 950.2, and 117.0 F g\textsuperscript{−1} at a current density 1 A g\textsuperscript{−1}, respectively. Further investigation of the rate performance of electrode samples at different current densities ranging from 1 to 10 A g\textsuperscript{−1} is shown in Figure 6d. These results show a good agreement with CV results, demonstrating the high performance of nickel−cobalt carbonate hydroxide electrode at the Ni/Co ratio of 1:2 is superior to that of other electrodes even at high current density (>400 F g\textsuperscript{−1} at 10 A g\textsuperscript{−1}). On the basis of these evidences along with the SEM and BET results, the obtained heterostructure of...
urchin-like structure with the nanonoodle-like structure of samples at Ni/Co ratio of 1:2 is the optimized combination morphology for the coexistence of nickel and cobalt carbonate hydroxide (Figure 1e) with the highest BET surface area, yielding the highest capacitance. Meanwhile, the large urchin-like structure of electrode samples at the Ni/Co ratios of 2:1 (Figure 1c) and 1:1 (Figure 1d) reveals the presence of highly dense packing and agglomerate structure, which decrease the ion diffusion and charge transfer, resulting in a low redox surface activity in the inner active site. Thus, the great improvement of redox capacitance behavior of electrode samples at the Ni/Co ratio of 1:2 can be explained as follows: (i) highly electroactive site-enriched cobalt carbonate hydroxide incorporated with nickel carbonate hydroxide for faradic reaction and (ii) the perfect densely packed sea-urchin-like structure provides the shortcut pathway for ion diffusion.

To further emphasize the electrochemical storage mechanism of nickel−cobalt carbonate hydroxide electrode at the Ni/Co ratio of 1:2, CV techniques of the electrode sample are complemented at different scan rates from 10 to 60 mV s⁻¹, as shown in Figure 7a. It is seen that the pair redox peaks of the CV curve reveal a slight shift in the higher potential with the increasing scan rate. The logarithmic relationship between the peak currents (i) and scan rate (ν) is calculated using the power law to exemplify the charge storage mechanism as in eqs 4 and 5.

\[ \log(i) = b \log(\nu) + \log(a) \]  

\[ i = a \nu^b \]  

where the b value prefers the slope of log(i) and log(ν) plots. The charge/discharge process is controlled by ionic diffusion when the b value is close to 0.5, While the b value reaches 1, the electrochemical performance of the electrode is dominated by pseudocapacitive behavior. From the fitting curves of the NiCo1/2 electrode in Figure 7b, the calculated b of the oxidation and reduction peaks are 0.7 and 0.71, respectively, indicating that the charge/discharge mechanism of the NiCo1/2 electrode is majorly controlled by ion diffusion participating in the pseudocapacitive behavior. The rate performance contribution mechanism of this electrode is calculated to compare the capacitive contribution with the relation of eq 6.

\[ i = k_1\nu + k_2\nu^{1/2} \]  

With the plotting of \( i/\nu^{1/2} \) vs \( \nu^{1/2} \), \( k_1 \) and \( k_2 \) are calculated from the slope and the y-axis interception, respectively. As eq 6, the term of the diffusion-controlled behavior is referred to as \( k_2\nu^{1/2} \) and that of the capacitive contribution behavior as \( k_1\nu \). Figure 7c presents the mechanism contribution of NiCo1/2 electrodes at different scan rates. It is clear that as the scan rate increases, the pseudocapacitive contribution is increased. The pseudocapacitive contribution of NiCo1/2 is 30.0, 37.8, 46.2, and 51.3% at 10, 20, 40, and 60 mV s⁻¹, respectively.
confirming that the pseudocapacitive characteristic is participating in charge storage at a high rate for the nickel–cobalt carbonate hydroxide electrode.46

To evaluate the resistive nature of electrode samples, the comparative study of electrode samples at different Ni/Co ratios using electrochemical impedance spectroscopic analysis is investigated as shown in Figure 8a. The Nyquist plot is typically used to examine the terms of solution resistance ($R_s$), charge-transfer resistance ($R_{ct}$), and charge/ion mechanism diffusion in electrode materials. Nyquist plots of all electrode samples reveal two parts: a semicircular response between high–intermediate frequency and a linear response at low frequency. In the region of high frequency, the initial resistance value of the semicircle region is referred to as the contact resistance between the interaction of active materials and electrolytes on the surface. The arc line intersection of semicircle portion from high–intermediate frequencies indicates the total resistance of electrode sample, corresponding to electron-transfer-limited processes. A relative linear range from middle to low frequency through phase angle corresponds with ion/charge diffusion into porous and interlayer of materials.47 Obviously, the increase of cobalt-to-nickel ratio trends to reduce the solution resistance ($R_s$) component in the low-frequency region, indicating the improvement of electrical capacitance behavior. Similarly, the semicircle region also tends to decrease with increasing cobalt proportion. Obviously, at Ni-to-Co ratio of 1:2, the electrode sample exhibits the lowest initial resistance value and impedance resistance value through phase angle, implying the highest ion/charge diffusion. 

Figure 8b demonstrates the comparative capacitance retention of all electrode samples from 1 to 3000 cycles. It is seen that pristine nickel carbonate hydroxide can preserve only 30.1% after 3000 cycles. Meanwhile, cobalt carbonate hydroxide exhibits retention constant of 100% after 500 cycles and a slight decrease to 95.4% after 3000 cycles. Thus, cobalt carbonate hydroxide is proven to possess a highly stable electrochemical performance compared with nickel carbonate hydroxide. Interestingly, all the nickel–cobalt carbonate complex samples demonstrate better development of specific capacitance with increasing CV cycles than that of pristine nickel hydroxide and cobalt carbonate hydroxide. Especially, the retention of nickel–cobalt carbonate hydroxide at the Ni/Co ratio of 1:2 can increase up to 178.3% after 3000 cycles, while the Ni/Co ratio of 2:1 and 1:1 exhibit high stability at 101.7 and 98.6% after 3000 cycles. As the CV

Figure 8. (a) Nyquist plots at a frequency range of 0.01–10 000 Hz of different Ni–Co carbonate hydroxides, (b) percent retention of different Ni/Co ratios as a function of cycle number at a scan rate 40 mV s$^{-1}$, (c) cyclic voltammograms of NiCo1/2 after 3000 cycles, and (d) ragone plots of NiCo1/2 and recent research for comparison.
evidence supports cycle stability for the nickel–cobalt carbonate hydroxide at the Ni/Co ratio of 1:2 in Figure 8c, the current density of pair redox peak is dramatically increased with the CV cycles, suggesting that the development of phase crystallinity of the nickel–cobalt carbonate hydroxide during the charging–discharging cycles. This phenomenon is attributed to the formation of intermediate cobalt carbonate hydroxide species during the redox reaction, which increases the phase crystallinity of cobalt carbonate hydroxide, leading to an increase in faradic reaction. Furthermore, the performance of the nickel–cobalt carbonate hydroxide electrode at the Ni/Co ratio of 1:2 is compared with those of the other nickel–cobalt SCs sources as seen in the ragone plot of Figure 8d. The energy density and power density are calculated as eqs 8,9. The nickel–cobalt carbonate hydroxide electrode at the Ni/Co ratio of 1:2 can yield an energy density in the range of 42.9–15.8 Wh kg−1, while the power density values are in the range of 285.0–2849.9 W kg−1. In addition, the energy density and power density of the NiCo1/2 symmetric cell are 35.4 Wh kg−1 at 6797.25 W kg−1. This result demonstrates a much higher value than those of conventional capacitors and many reports of nickel–cobalt materials. A comparison of the electrochemical performance of the NiCo1:2 sample with other reports is also illustrated in Table 2. Apparently, our study shows an outstanding electrocapacitance performance compared to other reports.

To investigate the valence state of the as-prepared Ni–Co carbonate hydroxide samples at different Ni/CO ratios, further analysis by X-ray absorption near-edge structure (XANES) technique is employed. The individual K energy of nickel and cobalt is separately monitored with NiO, CoO, and Co3O4 standard, representing Ni2+, Co2+, and Co3+ ion, respectively. The absorption K-edge of the as-prepared nickel carbonate hydroxide samples is nearly close to that of NiO standard, suggesting the same valence of Ni+2. All nickel–cobalt carbonate hydroxide samples exhibit nearly close to K-edge energy of NiO standard, indicating that all nickel–cobalt carbonate hydroxide samples exhibit Ni2+ ion in nickel part. In other words, the formation of Ni valence states does not interfere with the incorporation of cobalt during the hydrothermal process. Furthermore, the white line peak intensities of the NiCo1:2 sample with other reports is also decreasing nickel contents, implying the increase in the local disorder of the nearest neighbors of the Ni bulk.
For the absorption K-edge of cobalt (Figure 9b), the K-edge energy of the as-prepared cobalt carbonate hydroxide is nearly close to that of the Co$_3$O$_4$ standard, suggesting the valence of Co$^{2+}$ and Co$^{3+}$. Meanwhile, all as-prepared nickel–cobalt carbonate hydroxide samples at different Ni/Co ratios exhibit the shift of K-edge energy and are close to CoO when it is hydrothermally synthesized with nickel. These results indicate the formation of diverse Co-valence states of +2 and +3. The K-edge energy also shows a lower shift in intensity with cobalt content, implying the Co-valence state is nearly close to Co$^{2+}$.  

Figure 10. (a) Schematic of XANES collecting point on cyclic voltammetry curve at 1 mV s$^{-1}$ and the experimental cell of TR-XAS measuring; comparison of in situ XANES collected of NiCo1/2 on CV measuring. full cycle in (b) Co and (c) Ni part. (d) Charge and (e) discharge of Co part. (f) Charge and (g) discharge of Ni part.
The white line peak intensity of cobalt bulk for the Ni–Co carbonate hydroxide complex reveals an increase with cobalt content, implying the decrease in the local disorder of the nearest-neighbor atom of the Co bulk.54

To understand the charge–discharge mechanism, the optimized sample at Ni/Co ratio of 1:2 is further investigated the change of valence behavior during CV cycling at 1 mV s−1. Figure 10a shows that the asymmetrical cell of nickel–cobalt carbonate hydroxide is fabricated and monitored with the XANES technique at each CV measurement point. The in situ XANES spectra of Co and Ni signal for an overall charge–discharge cycle (as well as charge–discharge) are monitored as revealed in Figure 10b,c, respectively. The separated plots for Co- and Ni-charging from −0.3 to 0.45 V are presented in Figure 10d,e. However, no significant change of valence behavior during CV cycling at 1 mV s−1 at a power density of 285.0 mV s−1 at a scan rate of 40 mV s−1. The electrochemical impedance spectroscopy (EIS). The three-electrode system was used in a 6 M KOH electrolyte support solution and configured with active materials, platinum plate, and Ag/AgCl as the working electrodes, counter electrodes, and reference electrodes, respectively. CV was employed with a potential of −0.3 to 0.45 V at scan rates from 10 to 100 mV s−1. GCD was examined at −0.2 to 0.37 V at the current density from 1 to 10 A g−1. EIS was performed in 1 M KOH at the voltage of 5 mV over a frequency range from 0.01 Hz to 100 kHz. In addition, the symmetrical cell using a CR2032 coin cell was fabricated to measure the cycle stability performance. Two electrodes of NiCo1/2 are sandwiched together with a glass fiber separator and 6 M KOH as electrolyte. The cycle stability testing was conducted by CV technique at a scan rate of 40 mV s−1 for 3000 cycles using a potentiostat/galvanostat instrument (Metrohm Autolab, PGSTAT204, Netherlands) to investigate cyclic voltammetry (CV), galvanostatic charge–discharge (GCD), and electrochemical impedance spectroscopy (EIS). The electrode samples were measured the electrochemical properties by the potentiostat/galvanostat instrument (Metrohm Autolab, PGSTAT204, Netherlands) to investigate cyclic voltammetry (CV), galvanostatic charge–discharge (GCD), and electrochemical impedance spectroscopy (EIS). The three-electrode system was used in a 6 M KOH electrolyte support solution and configured with active materials, platinum plate, and Ag/AgCl as the working electrodes, counter electrodes, and reference electrodes, respectively. CV was employed with a potential of −0.3 to 0.45 V at scan rates from 10 to 100 mV s−1. GCD was examined at −0.2 to 0.37 V at the current density from 1 to 10 A g−1. EIS was performed in 1 M KOH at the voltage of 5 mV over a frequency range from 0.01 Hz to 100 kHz. In addition, the symmetrical cell using a CR2032 coin cell was fabricated to measure the cycle stability performance. Two electrodes of NiCo1/2 are sandwiched together with a glass fiber separator and 6 M KOH as electrolyte. The cycle stability testing was conducted by CV technique at a scan rate of 40 mV s−1 for 3000 cycles using a potentiostat/galvanostat instrument (Metrohm Autolab, PGSTAT204, Netherlands). The specific capacitance (Csp) is calculated from the GCD result following eq 7. The energy density (E) and the power density (P) are calculated from eqs 8 and 9.

\[
C_{sp} = \frac{I(t)}{m(\Delta V)} 
\]

(7)

\[
E = \frac{1}{2}C_{sp}(\Delta V)^2 
\]

(8)

\[
P = \frac{E}{T} 
\]

(9)

where \(I\) is the constant discharge current (A), \(t\) is the discharging time (s), \(m\) is the mass of active material sample (g), \(\Delta V\) is the window potential (V), and \(T\) is discharging time (h).

4.3. Preparation of Working Electrode Samples. Working electrode samples were prepared by mixing 80 wt % of active materials, 15 wt % of carbon black, and 5 wt % of poly(tetrafluoroethylene) (PTFE). The solid mixture was mixed with ethanol to obtain a slurry mixture and coated on carbon fiber paper (1 × 1 cm²). The total weight of the solid mixture was controlled at approximately 1.0 mg cm⁻² for all coated materials.

4.4. Electrochemical Characterization. The electrode samples were mixed with ethanol to obtain a slurry mixture and coated on carbon fiber paper (1 × 1 cm²). The total weight of the solid mixture was controlled at approximately 1.0 mg cm⁻² for all coated materials.
performed to investigate the chemical states of samples. Time-resolved X-ray absorption spectroscopy (TR-XAS, beamline 2.2 at Synchrotron Light Research Institute (SLRI), Thailand) was carried out with the absorption edge energy at 8333 and 7709 eV to detect nickel and cobalt elements, respectively. The transmission mode was performed to investigate the X-ray absorption near-edge structure (XANES) of sample side electrode during charging and discharging. The charge-discharge process was performed by CV technique at scan rate of 1 mV s$^{-1}$. Asymmetric supercapacitors (ASCs) assembled using two dissimilar electrode as the cathode, the carbon fiber paper as the anode, and filter paper separator with 6 M KOH electrolyte.

**ASSOCIATED CONTENT**

Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsomega.1c02139. Experimental details of the materials (XPS, BET, and EIS circuit fitting) (PDF)

**AUTHOR INFORMATION**

Corresponding Authors
Prasit Pattananuwat — Department of Materials Science, Faculty of Science, Chulalongkorn University, Bangkok 10330, Thailand; Research Unit of Advanced Materials for Energy Storage and Center of Excellence on Petrochemical and Materials Technology, Chulalongkorn University, Bangkok 10330, Thailand; Email: prasit.p@chula.ac.th

Pranut Potiyaraj — Department of Materials Science, Faculty of Science, Chulalongkorn University, Bangkok 10330, Thailand; Center of Excellence on Petrochemical and Materials Technology and Center of Excellence on Responsive Wearable Materials, Chulalongkorn University, Bangkok 10330, Thailand; Email: pranut.p@chula.ac.th

Author
Nutthapong Poompiew — Department of Materials Science, Faculty of Science, Chulalongkorn University, Bangkok 10330, Thailand

Complete contact information is available at: https://pubs.acs.org/10.1021/acsomega.1c02139

Notes
The authors declare no competing financial interest.

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