Enhancing capacitance behaviour of CoOOH nanostructures using transition metal dopants by ambient oxidation

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Cobalt hydrate and doped binary Co0.9M0.1OOH (M = Ni, Mn, Fe) nanorings of 100–300 nm were fabricated in solution through a facile ambient oxidation method. A transformation from Co0.9Ni0.1(OH)2 nanodiscs to hollow Co0.9Ni0.1OOH nanorings was observed with prolonged reaction time. Core-shell nanodiscs have elemental segregation with a Co(OH)2 core and Ni(OH)2 shell. Co0.9Ni0.1OOH nanorings displayed a higher electrochemical capacitance than Mn and Fe doped nanorings materials or materials with disc-like geometries.

Hollow inorganic nanomaterials have attracted intense research interest due to low mass density, high porosity and increased surface area1–3. These characteristics make them good candidate materials for high surface area devices such as lithium-ion batteries and gas sensors4–8. Nanorings are one of the hollow structures which have been successfully fabricated and exhibit extraordinary properties due to their ring-like morphology9–10. Divalent metal oxides or their hydrates’ nanorings are the most intensively studied hollow materials due to their high surface-to-volume ratio and special magnetic properties10–12. CoOOH is an active material used in conjunction with nickel hydroxide in Ni–H cells. Cobalt oxide is coated on the positive electrodes of rechargeable alkaline nickel batteries to achieve high electrochemical efficiency13. In addition, CoOOH is non-stoichiometric and the Co is at a higher oxidation state (+3) than in Co3O4 (the oxidation states are +4, +3, +2 and the average is +8/3)14,15. Usually the synthesis process for cobalt/iron hydrates contains a microwave irradiation/ultrasonic chiseling pre-treatment and a high temperature (140 °C–230 °C) post-hydrothermal reaction for few hours16–18 with low yield19. Cobalt hydroxide nanorings have been reported with polycrystalline structure after 10 h in a sealed autoclave at 180 °C20. Hexagonal Co3O4 nanorings have also been produced from precursor Co(OH)2 nanodiscs at 160 °C for 10 h with diameters of a few micrometres12. Co-Sn-O systems have also been reported using a hydrothermal method at high temperature via the Kirkendall effect through the diffusion of pores21. However, other doping such as Mn, or Fe into the Co(OH)2, is still not reported. A facile room temperature method is needed to prepare doped cobalt hydroxide nanorings with high yield.

Here, we will give detailed preparation and analysis on Co0.9Ni0.1OOH nanorings via ambient oxidation. Typical images of Co0.9Ni0.1OOH products under different reaction times will be discussed, revealing the shape transformation and growth mechanism. Electrochemical properties of CoOOH materials doped with different elements will also be presented.

Results and Discussion

Figure 1 shows the XRD profiles of the products reacted for 3 h (black) and for 72 h (red). The main diffraction peaks in the black spectrum can be attributed to the hexagonal trigonal β-Co(OH)2 structure (JCPDS file No. 30-0443, space group: P3-3m (166); lattice constants: a = 3.2 Å, c = 4.6 Å, α = 90°, β = 120°). While weak peaks in the spectrum of 3 h products can be attributed to the β-CoOOH structure (JCPDS file No. 73–1213, space group: Fd-3m (227); lattice constants: a = 4.7 Å, α = 35.5°). After reaction for 72 h, the peaks of CoOOH were observed

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mental distribution of the nanodiscs can be observed in the last part of Fig. 2g–k with a Co(OH)2 core (purplish color) wrapped by a Ni(OH)2 shell (azure color).

Therefore we say the losing of H atom just in the symbolic meaning to discuss the two atomic models. More specifically, the transformation of Co(OH)2 to CoOOH there while the inner part maintains its structure. After reaction for 72 h, ring-like nanorings are formed. Figure 2 gives typical microstructures of products prepared in 20 °C for 3 h. A typical TEM image in Fig. 1a of a 220 nm outer diameter nanodisc shows that it has uniform thickness. The electron diffraction pattern inset of Fig. 2a indicates its hcp structure with lattice parameters of a = b = 3.2 Å which corresponds to (β-CO(OH)2). High resolution transmission electron microscope (HRTEM) images (Fig. 2b,c) from the border and inner space of the nanodisc indicate their single crystal structure. A lattice contraction of about 2% in the <010> direction of the nanodics can be detected, a change from 3.08 Å to 3.02 Å from average measurement results. This contraction can be attributed to the nanomaterials’ border effect or structural variation from Co(OH)2 (lattice parameter a is 3.18 Å) to Ni(OH)2 (lattice parameter a is 3.11 Å), which has a contraction from about 1.6% to 2.5% in bulk material. The lattice contraction can nearly be omitted in the nanorings, while 1–3 nm amorphous layers can be detected in the outer border of most rings. A further study on the elemental distributions on the room temperature nanodiscs is shown in Fig. 2d–k. It can be seen that the nickel is preferentially distributed in a border range of about 20–30 nm, while the cobalt has a lower concentration in the border compared to the centre. The mean concentration of Ni in the border is about 27.5% while it is 6.9% in the center. A schematic illustration of the elemental distribution of the nanodics can be observed in the last part of Fig. 2g–k with a Co(OH)2 core (purplish red color) wrapped by a Ni(OH)2 shell (azure color).

Figure 3 shows the morphologies, elemental distribution and valence state of nanorings provided from the following techniques: transmission electron microscopy (TEM), energy dispersive X-ray spectroscopy (EDX) and electron energy loss spectroscopy (EELS). In a typical low magnification TEM image in Fig. 3a, tens of rings are shown with diameters from 200–300 nm. Most of the nanorings maintain the hexagonal inner and outer border without any angle variation. The quantitative EDX results show that the ratio of Co/Ni is 87.7/12.3, which contains slightly more Ni than the starting Co/Ni ratio (9/1). A typical EDX map of one nanoring in Fig. 3b shows that the distributions of O, Co and Ni are uniform. An enlarged TEM image in Fig. 3c gives the detailed morphology of the nanorings and shows that the wall thickness of the nanorings is 30–50 nm. Although SEM observations of these nanoparticles suggest a single crystal nanoring, HRTEM results indicate an orientation related multicrystalline structure. The ring-like structure is not comprised of perfect single crystals. The inner edge of the hexagonal border is made of small nanoparticles aggregated continuously or quasi-continuously as in Fig. 3c. A thin porous structure can be seen in the centre of some ring-like structures. A HRTEM image in Fig. 3d indicates that the particles composing the hexagonal skeleton contain a single crystal structure with [001] facets of hcp CoOOH as a surface plane. Though the fast Fourier transform (FFT) of the HRTEM in the inset of Fig. 3d gives two sets of the diffraction patterns, the strong spots (indexed by the arrowhead) indicate that the main phase of the product is CoOOH, while it contains traces of the Co(OH)2 phase. The pre-peak in the EELS spectrum in Fig. 3e of the O edge in 535 eV indicates the existence of OH− bonds22. Since L1 and L2 lines in Co-L edge correspond to transitions from 2p1/2 to 3d10 3d3/2 and from 2p3/2 to 3d10 3d5/2 respectively, the integration of the L1/L2 ratio can provide the ratio of Co3+ inside it23. The L1/L2 ratio in the Co-L edge in our rings after calculation is about 2.2 which indicates that the products contain large quantities of trivalent cobalt. From the EELS results, we can conclude that these nanorings are mainly made up of CoOOH.

Figure 4 provides an SEM image of an intermediate state reacted for 24 h and a schematic illustration of shape evolution. In an intermediate state after reaction for 24 h as shown in Fig. 4a, some ring-like structures are formed while most of them maintain a bowl-like structure with a rough surface. The roughness of the surface can be explained by the atomic model in Fig. 4b. In Co(OH)2 (right part in Fig. 4b) the distance between Co-O layers is 2.66 Å, while it is 2.59 Å in CoOOH (left part in Fig. 4b) due to the loss of H atoms and an electron in the same time. In comparison of the chemical composition of Co(OH)2 and CoOOH, there is a H-atom removed, which in fact is referring to the removing a proton (H+ ion) and an electron as well and there forms no H(0) species. Therefore we say the losing of H atom just in the symbolic meaning to discuss the two atomic models. More surface area in the border is bare to the oxygen-filled environment than in the inner part, resulting in enhanced oxidation of Co(OH)2 to CoOOH there while the inner part maintains its structure. After reaction for 72 h, ring-like
structures form as shown in Fig. 1c. It indicates the formation of nanorings from the disks which first become porous and finally reach the ring-like structures.

The formation mechanism is illustrated in Fig. 4c. In the beginning, cobalt hydrate precipitates to form a disc-like core and this can be tested by EDX results on samples reacted for short times as shown in Figure S5 and Table S1 in the supplementary information. As time progresses, the loose layered borders and surfaces of β-Co(OH)₂ begin to lose some of their H atoms and an electron due to oxidation. The outer border tends to oxidize to a contracted CoOOH structure since more areas there are exposed to the oxygen. Then a Co(OH)₂/CoOOH boundary will be formed. Inner Co(OH)₂ layers adjacent to the outer CoOOH will then transform/diffuse to CoOOH structure by losing H atoms. Pores will then form in the centre, eventually creating a hollow ring-like structure.

The lowest pH value needed for precipitation can be calculated using the following

$$\text{pH} = 14 - \frac{1}{n} \left( 1 \ g[M^{+n}] - 1 \ gK_{sp} \right)$$

where n is the valence of the metal, M represents activity of ions of the metal and $K_{sp}$ is the solubility product constant. In a dilute solution, the activity of ions (also known as effective concentration) can be expressed as
\[ a_i = \gamma_i N_i \], where \( a_i \), \( \gamma_i \) and \( N_i \) represent activity, activity coefficient and concentration of the \( i \)th ion, respectively. This means that the activity \( a_i \) is monotonically dependent on the nominal concentration \( N_i \). Because \( \text{Co}^{2+} \) and \( \text{Ni}^{2+} \) have the same valence value, their activities in a dilute solution are very close \(^\text{24} \). It is reasonable to correlate these thermodynamic quantities with the nominal concentration in this case. Calculation results indicate that if the concentration of \( \text{Co}^{2+} \) is 0.018 M, the pH value needed is 7.53. Similarly, pH values for a concentration of 0.002 M Ni, Mn and Fe are 7.93, 8.87, and 7.85 respectively. This suggests that the \( \text{Co(OH)}_2 \) precipitate firstly forms disc-like structures due to its intrinsic hcp structure and provides borders and surfaces to which Ni/Mn/
Fe(OH)$_2$ can then adhere. That is the reason that no Nickel can be detected in the disc-like structures formed on short timescales while its concentration was found to have increased to 12–3% in ring-like structures (Figure S5 and Table S1). As predicted in the calculation above, Mn$^{2+}$ and Fe$^{2+}$ are possible doping metals with a higher pH value for precipitation needed than Co$^{2+}$. Typical morphologies and chemical properties of Co$_{0.9}$Mn$_{0.1}$OOH and Co$_{0.9}$Fe$_{0.1}$OOH nanorings are shown in Fig. 5 and supplementary files (Figure S6-S14). The diameter of Co$_{0.9}$Mn$_{0.1}$OOH nanorings is about 150 nm with an average cobalt atomic composition of 91% of metal content. However, the diameter of the Co$_{0.9}$Fe$_{0.1}$OOH is about 100 nm with an average cobalt atomic concentration of 89%.

The typical cyclic voltammograms of the Co$_{0.9}$Ni$_{0.1}$OOH nanodiscs (black lines) and nanorings were acquired at a sweep rate of 5 mV s$^{-1}$ with a potential window of −0.2 to 0.7 V vs a standard calomel electrode (SCE) and are shown in Fig. 6a and The four peaks of the C-V curve in Fig. 6a can be attribute to transformation of Co$^{2+}$→Co$^{3+}$, Co$^{3+}$→Co$^{4+}$, Co$^{4+}$→Co$^{3+}$ and Co$^{3+}$→Co$^{2+}$

There is a distinct increase in the redox peak of the nanorings compared to that of the nanodiscs. The strong redox peak in the nanorings indicates that the capacitance characteristics are mainly governed by Faradaic reactions, which are very distinct from the rectangular shape of the electric double layer. The charge/discharge curves at current densities of 1 A g$^{-1}$ in Fig. 6b show that after the shape changes to nanorings, its capacitance increased from 211.5 to 439.7 F g$^{-1}$. The comparison of M doped (M = Ni,
Mn, Fe) nanorings in Fig. 6c,d indicates that Ni doped CoOOH nanorings exhibit better capacitance performance than Mn and Fe doped CoOOH nanorings. The enhanced capacitance can be attributed to the porosity of the surface of the nanorings. We must point out that for pure CoOOH, the nanoring morphology enhances the capacitance but the enhancement is not as significant as the Ni doped ones.19,25,26 This means that both the nanoring morphology and Ni doping can enhance the capacitance.

Conclusions
In conclusion, cobalt hydrate and doped Co0.9M0.1OOH (M = Ni, Mn, Fe) nanorings were successfully fabricated by shape evolution from Co0.9M0.1(OH)2 nanodiscs to Co0.9M0.1OOH nanorings via ambient oxidation in room temperature. The formation and evolution process of the Co0.9Ni0.1OOH system were illustrated. Ring-like structures formed due to the contraction in the surface of the nanodiscs during the oxidation process. Co0.9Ni0.1OOH nanorings displayed a higher capacitance than its disc-like materials and other systems. Nickel doped CoOOH nanorings displayed better capacitance performance than Mn and Fe doped CoOOH nanorings.

Methods
Synthesis of M-doped CoOOH. For the synthesis procedure, Ni doped systems were described as an example, while synthesis with different dopants (Mn, Fe and pure CoOOH materials) use similar methods. In a typical synthesis, CoCl2·6H2O and NiCl2·6H2O were dissolved in deionized water with a ratio of 9:1. A small amount of N2H4·H2O was added to the solutions. During the reaction, NaOH solution was added to tune the pH value to about 13.7. This solution was continually stirred for 3–72 h in room temperature (17–20°C) while exposed to the air. A pink brown precipitate in the beginning and then a brown precipitate was collected on the bottom of the bottle. The precipitates were then washed and dried for further characterisation.

Characterisation. An X-ray diffractometer (XRD) on a BRUKER-D8 ADVANCE diffractometer was used while employing a graphite monochromatized Cu Kα radiation flux at a scanning rate of 0.02° s⁻¹ in the 2θ range of 1090°. A Carl Zeiss Ultra scanning electron microscope (SEM) was used to image the morphologies of the Co-Ni hydrate nanoparticles. The SEM was able to perform energy dispersive X-ray spectroscopy (EDX) on the sample. A 300 kV FEI Titan transmission electron microscope (TEM) was also used for imaging and was capable of both electron energy loss spectroscopy (EELS) and EDX. In all cases a wide selection of individual oxide nanoparticles were examined to determine their average structure and morphology.
Electrode preparation and electrochemical tests. Electrochemical measurements were carried out in a 5 M aqueous KOH in a half-cell setup configuration at room temperature. Platinum wire and a standard calomel electrode (SCE) served as counter and reference electrodes, respectively. The working electrode was prepared by casting a paste containing CoO$_2$Mn$_3$OOH nanomaterials (8 mg), carbon black, and poly(vinylidenefluoride) (PVDF) in a weight ratio of 80:10:10 onto a 1 cm $\times$ 1 cm Ni foam (2 mm thick, 100 ppi, 95% porosity, purchased from Bitaxiang Co. Ltd., Kunshan, China). The electrodes were calendared and degassed in vacuum at 80 °C for 12 h. The resulting electrode was pressed at 5 MPa to form about 150 μm thick with 1 cm$^2$ area sheet for electrochemical tests. Cyclic voltammetry (CV) and charge/discharge experiments were obtained on a CHI 660D electrochemistry workstation. CV curves were recorded at a sweep rate of 5 mV s$^{-1}$.

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