Design and self-assembly of metal-organic framework-derived porous Co₃O₄ hierarchical structures for lithium-ion batteries

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Abstract

Porous metal oxides hierarchical structures with controlled morphologies have received great attention because of their promising applications in catalysis, energy storage, gas sensing, etc. Porous Co₃O₄ hierarchical structures with controlled morphologies were synthesized on the basis of a pyrolytic conversion of Co-based metal-organic frameworks (Co-MOFs), which were initially grown in solutions containing Co(NO₃)₂·6H₂O, 1,3,5-Benzenetricarboxylic acid and pyrazine as solute and N,N-dimethylformamide (DMF) as solvent under a solvothermal condition. Porous Co₃O₄ with twin hemispherical and flower-like structures were obtained with the assistance of PVP by adjusting the amount of pyrazine. The results of nitrogen adsorption–desorption indicate the BET surface area (22.6 m² g⁻¹) of twin hemispherical Co₃O₄ structures is lower than that (33.3 m² g⁻¹) of flower-like Co₃O₄ structures. However, the pore size of twin hemispherical Co₃O₄ structures is smaller, which is centered at about 2.5, 4.0 and 20.0 nm. The Co₃O₄ with twin hemispherical structures exhibit more excellent electrochemical performance as anode materials for lithium batteries than that of flower-like Co₃O₄ structures, which may be attributed to the smaller particle size and compact porous structures with suitable pore size.

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1. Introduction

Hierarchical micro- and nanostructures of porous metal oxides have attracted considerable attention due to their highly ordered architectures and well-defined physical properties, which show various potential applications in energy storage and conversion, adsorption, catalysis, gas sensing, etc. [1–4]. In past decades, tremendous efforts have been devoted to synthesize porous hierarchical structures involving the coating of a shell on removable or sacrificial templates based on different principles including the galvanic replacement [5,6], Kirkendall effect [7,8], ionic exchange [9,10], chemical etching [11,12], self-assembly [13], thermal decomposition [14,15], etc. Among the diverse synthetic methods, the self-sacrificial template strategy based on thermolysis of solid precursors is considered to be the most straightforward and representative one in fabricating porous hierarchical structures due to its low cost and effectiveness. For instance, 3D flower-like Co₃O₄ hierarchical microstructures were assembled by hexagonal porous nanoplates using facile surfactant-free hydrothermal and annealing processes [16]. Hierarchical porous ZnCo₂O₄ microspheres were successfully synthesized via a solvothermal method followed by an annealing process [17]. Hierarchical porous NiO nanoflowers were converted from flower-like α-Ni(OH)₂ nanostructures through calcinations procedure [18]. The design of novel and complex porous metal oxides hierarchical structures by a simple, low cost and

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effective method is still a very interesting topic in the field of advanced materials science, although it has been studied extensively.

Recently, metal-organic frameworks (MOFs) as sacrificial templates were devised to fabricate porous metal oxides or carbon nanostructures through thermal decomposition under controlled atmospheres [19–23]. MOFs are a class of organic–inorganic hybrid functional materials with high porosity and large specific surface area. It’s worthwhile to note that their pore sizes and morphologies can be easily tuned upon the selection of different metal ions and organic bridging ligands, and the porosity and long-range ordering of MOFs can offer a fast and convenient access to income and leave small molecules and ions in the transformation process. Recently, research emphasizing tuning and controlling of MOF crystal morphology and size becomes a newer way of tailoring MOF properties for specific functions, such as gas storage [24,25] and biomedical imaging [26].

Porous structures have been proven to have appropriate Li-ion diffusion lengths and large contact area with electrolytes [27,28]. Poixot et al. suggested that there are optimum particle and pore size of each metal oxide for the best electrochemical performance [29]. Co3O4 has been widely applied electrode materials in supercapacitors and lithium-ion battery [30]. Admittedly, specific surface areas of the obtained porous Co3O4 products are remarkably enlarged relative to solid particles. However, some formed porous Co3O4 products failed to maintain original morphologies of the precursor templates, probably due to a lack of suitable templates or optimal synthetic conditions. So it still remains a great challenge to synthesize porous Co3O4 with specific morphology and high specific surface area. In this study, Co-MOFs have been selected as a model MOFs to investigate the feasibility of MOFs as sacrificial templates for large-scale controllable preparation of porous metal oxides hierarchical structures. By controlling the reaction parameters, twin hemispherical and flower-like Co-MOFs hierarchical structures were synthesized and calcined to obtain porous Co3O4 hierarchical structures with corresponding morphologies at 500 °C for 30 min under air atmosphere. The results indicate that twin hemispherical Co3O4 hierarchical structures with smaller pore and particle size display excellent reversible capacity and good cycle performance, and the initial discharge capacity is as high as 1325.5 mA h g⁻¹.

2. Experimental procedure

2.1. Synthesis of Co-based metal-organic frameworks (Co-MOFs)

2.1.1. Synthesis of twin hemispherical Co-MOFs

In a typical solvothermal preparation procedure, 0.5 mmol cobalt nitrate hexahydrate, 0.5 mmol 1,3,5-Benzentricarboxylic acid, 0.9 mmol pyrazine and 1.31 g PVP (K85–95) were dissolved in 30 mL DMF. After magnetically stirred for 10 min, the homogeneous solution was transferred into a Teflon-lined stainless steel autoclave with 40 mL capacity, and placed in an oven at 150 °C for 24 h. Finally, the resulting purple powders were collected by centrifugation and washed with ethanol for 3 times and dried at 60 °C in a vacuum oven for 6 h. The prepared sample was designated as Co-MOFs-a.

2.1.2. Synthesis of novel flower-like Co-MOFs

The preparation procedure of the novel flower-like Co-MOFs was similar to that of the twin hemispherical Co-MOFs, which only changed the amount of pyrazine from 0.9 to 3 mmol. Finally, the light purple products were obtained, which were designated as Co-MOFs-b.

2.2. Synthesis of porous Co3O4 hierarchical structures

The above-synthesized Co-MOFs precursors were put into a ceramic crucible and then heated to 500 °C with a heating rate 1 °C min⁻¹, and maintained at 500 °C for 30 min under air atmosphere. The Co-MOFs-a and Co-MOFs-b were finally converted into two fluffy black powders, which were designated as Co3O4-a and Co3O4-b, respectively.

2.3. Characterization

2.3.1. Characterization of materials

Thermogravimetric analysis (TGA) of the Co-MOFs precursors were carried out on a Rigaku standard TG-DTG analyzer under air atmosphere in the temperature range of 25–550 °C with a heating rate of 15 °C min⁻¹. FTIR spectra of KBr powder pressed pellets were recorded on a Bruker Vector 22 spectrometer. X-ray powder diffraction (XRD) patterns were recorded on a MAX-RB X-ray diffractometer (Rigaku, Japan) equipped with graphite-monochromatized Cu Kα radiation (λ=1.54178 Å). The field-emission scanning electron microscopy (FESEM) images of the products were taken by using a field-emission scanning electron microscope (FESEM, JEOL JSM-7600F) and transmission electron microscope (TEM) and high-resolution TEM (HRTEM) images were taken on a JEM-2100F high-resolution transmission electron microscope at an acceleration voltage of 200 kV. Energy-dispersive X-ray (EDX) analysis was performed for the products using the energy-dispersive X-ray spectroscopy attached to the JSM-6700F. X-ray photoelectron spectroscopy (XPS) of the products was performed on a Perkin-Elmer model PHI 5600 system with a monochromatic Kα radiation (1486.6 eV) X-ray source. The nitrogen adsorption–desorption isotherms and textural properties were determined on a Micromeritics Instrument Corporation sorption analyzer (TriStar II 3020).

2.3.2. Characterization of electrochemical performance

The electrochemical performance of Co3O4 after calcination was carried out using a coin cell (CR2016). The working electrodes consist of 70 wt% active materials (Co3O4-a or Co3O4-b), 20 wt% of conductivity agent (Super P) and 10 wt% of polymer binder (polyvinylidene fluoride, PVDF) mixed with N-methyl-2-pyrrolidone (NMP). After being stirred for 2 h, the slurry was pasted on a Cu foil using a doctor blade, and dried
at 120 °C for 12 h under vacuum to serve as the working electrode. The coin cells were assembled in an argon-filled glove box using lithium as the negative electrodes, Celgard 2500 as the separators, and 1 mol L⁻¹ LiPF₆ (dissolved in ethylene carbonate, dimethyl carbonate and ethyl methyl carbonate, with a 1:1:1 volume ratio) as the electrolytes. The galvanostatic charge and discharge tests were performed between 0.01 and 3.0 V at different current density on battery testing system (LAND CT2001A, Wuhan, China). Cyclic voltammetry (CV) measurements were carried out on a PARSTAT2273 electrochemical workstation (Princeton Applied Research, USA) at a scanning rate of 0.1 mV s⁻¹ in the potential range of 0–3 V. Electrochemical impedance spectroscopy (EIS) was performed using PARSTAT2273 electrochemical workstation; the sinusoidal excitation voltage applied to the cells was kept at 5 mV and with a frequency range of 100 kHz to 10 mHz.

3. Results and discussion

3.1. Morphological and structural characterization

To control the morphologies of porous Co₃O₄ hierarchical structures after thermal decomposition of Co-MOFs, cobalt complexes with the ligands of 1,3,5-benzentricarboxylic acid and pyrazine were synthesized, and different morphologies of Co-MOFs (Co-MOFs-a and Co-MOFs-b) were controlled by adjusting the amount of pyrazine. In order to understand the morphologies and structures of Co-MOFs, FESEM was employed to investigate the structures and morphologies of Co-MOFs-a and Co-MOFs-b in Fig. 1, which indicate that both Co-MOFs-a and Co-MOFs-b have uniformed morphologies with hierarchical structures. It is noteworthy that the Co-MOFs-a displays twin hemispherical structures with the average size of 2 μm, and the equatorial juncture between the crystal halves can be clearly observed from the FESEM image in Fig. 1(a). The high magnification FESEM image in Fig. 1(b) reveals that the large configuration assembled by small nanocrystals with the size of less 100 nm. When the amount of pyrazine is increased to 3 mmol from 0.9 mmol, Fig. 1(c) shows that Co-MOFs-b is relatively uniform flower-like structures, which is self-assembled by nanosheets. From the high magnification image in Fig. 1(d), the diameter of these novel flower-like structures is from 10 to 15 μm, and the thickness of nanosheets is about 50 nm.

The Fourier transform infrared (FTIR) spectrum of 1,3,5-benzenetricarboxylic acid, pyrazine, Co-MOFs-a and Co-MOFs-b were measured at room temperature for elucidating the structures and components of the Co-MOFs, as shown in Fig. 2(a). The strong peak around 1720 cm⁻¹ is assigned to C=O stretching vibration, and the characteristic peaks at 1606, 1454 and 1404 cm⁻¹ are attributed to the antisymmetric

Fig. 1. (a) and (b) Different magnification FESEM images of Co-MOFs-a; (c) and (d) different magnification FESEM images of Co-MOFs-b.
stretching vibration and symmetric stretching vibrations of the COO$^-$ of 1,3,5-benzentricarboxylic acid. In the FTIR spectra of Co-MOFs-a and Co-MOFs-b, absorption bands at about 1720 cm$^{-1}$ is disappeared, which is attributed to the deproto-
nation of 1,3,5-benzentricarboxylic acid due to the reaction with the metal ions [31]. The stretching vibration and sym-
metric stretching vibration peaks of the COO$^-$ for Co-MOFs-a
and Co-MOFs-b are moved to 1546 or 1549 cm$^{-1}$, 1439 or
1441 cm$^{-1}$, 1379 or 1385 cm$^{-1}$, which indicate that the
chelating or bridge bidentate have been formed between
COO$^-$ and Co$^{2+}$. In the FTIR spectra of pyrazine, the peak
at 1627 cm$^{-1}$ is due to C¼N stretching vibration. After the
formation of complexes, the peaks at about 1580 cm$^{-1}$ are
also ascribed to C¼N stretching vibration which indicate that
the two nitrogen atoms of pyrazine and cobalt ions formed
coordination bonds. Therefore, it is believed that Co-MOFs-a
and Co-MOFs-b might be Cobalt complexes with the ligands
of 1,3,5-benzentricarboxylic acid and pyrazine after solvother-
mal treatment.

TGA of the Co-MOFs-a and Co-MOFs-b was carried out in
air atmosphere to measure the weight loss occurring during
pyrolysis, which is shown in Fig. 2(b). The results indicate that
the weight loss of Co-MOFs-a includes a slow weight loss
(23.3%) from 20 to 375 °C and a fast weight loss (51.5%) from
375 to 388 °C. Little weight loss before 150 °C indicates that
there are little amount of crystal water in Co-MOFs-a. The
weight loss from 150 °C to 375 °C is attributed to the release
of DMF. From 375 to 388 °C, the weight loss can be ascribed
to the decomposition of complex and subsequent oxidation of
ligands. However, the weight loss of Co-MOFs-b may be
divided four parts. The weight loss (12.9%) before 100 °C is
the desorption of crystal water, and the weight loss (3.9%)
from 100 to 200 °C is mainly attributed to the release of
coordinated water; and the weight loss (34.0%) from 200 to
283 °C is ascribed to the removal of DMF. From 375 to
393 °C, the weight loss can be attributed to the decomposition
of complex and subsequent oxidation of ligands. The above
results indicate that Co-MOFs-b has higher porosity. The
residual weight ratio of Co-MOFs-b is higher than that of
Co-MOFs-a, which might be due to the fact that the numbers
of 1,3,5-benzentricarboxylic acid and pyrazine ligands are
different in Co-MOFs-a and Co-MOFs-b. The XRD results
in Fig. 2(c) also prove that Co-MOFs-a and Co-MOFs-b are different complexes although no standard PDF cards match with the XRD pattern of Co-MOFs-a and Co-MOFs-b. With increasing the thermal decomposition temperature, there are no remarkable weight losses beyond 393 °C, indicating that Co-MOFs have decomposed completely. Hence, in order to yield pure and high crystalline Co₃O₄ phase, Co-MOFs were calcined at 500 °C for 30 min.

By calcining the Co-MOFs precursors at 500 °C for 30 min in air, fluffy black powders (Co₃O₄-a and Co₃O₄-b) were obtained. To ascertain the components and structures of the products after calcination, Fig. 2(d) shows the XRD patterns to investigate the crystal phases of Co₃O₄-a, Co₃O₄-b. All the diffraction peaks of Co₃O₄-a and Co₃O₄-b can be readily indexed to f.c.c phase Co₃O₄ (JCPDS card no. 42-1467). No other impurity peaks are detected such as CoO and Co₂O₃, revealing that the Co-MOFs precursors were converted to crystalline Co₃O₄ completely. Additionally, all the diffraction peaks of Co₃O₄-a and Co₃O₄-b are significantly broaden in width and weaken in intensity, which suggests that the as-obtained Co₃O₄ are essentially constructed with a large number of nanosized crystallites.

X-ray photoelectron spectra (XPS) have often been used for surface characterization of various materials, and unambiguous results are readily obtained when the various surface components each contain unique elemental markers. Here, in order to analyze the surface component of Co₃O₄-a and Co₃O₄-b, XPS of the products were measured, which are shown in Fig. 3. It can be found that the peaks on the full patterns are mainly attributed to C1s (285.6 eV), O1s (529.2 eV) and Co2p (794.5 and 779.5 eV) and their corresponding Auger peaks in Fig. 3a, indicating the existence of carbon, oxygen, and cobalt element. However, the lower peak for C1s binding energy suggests that the products consist of C elements, which is from a small amount of residue of the decomposition of organic ligands. Additionally, it is also found that the Co2p and O1s XPS peaks of the two Co₃O₄ samples are analogous for their position and intensity. Fig. 3(b) shows that the Co2p XPS spectra present two major peaks at 794.5 and 779.5 eV, which can be attributed to Co2p1/2 and Co2p3/2 of the Co₃O₄-a and Co₃O₄-b. The absence of prominent shakeup satellite peaks in the Co2p spectra further indicates the formation of the Co₃O₄ phase. The deconvoluted O1s spectrum in the inset of Fig. 3(a) exhibit two peaks at 529.8 and 531.1 eV, which can be assigned to the lattice oxygen of spinel Co₃O₄ [32].

In order to investigate the effect of Co-MOFs as sacrificial templates for the morphologies of the products, FESEM were employed to measure the morphologies and structures of Co₃O₄-a and Co₃O₄-b. Typical morphologies are shown in Fig. 4(a) and (b). Obviously, the calcined products retain the similar sizes and shapes of the Co-MOFs-a and Co-MOFs-b precursors. Fig. 4(a) shows that Co₃O₄-a is sphere-like shape, which is assembled by many uniform nanoparticles with the diameter of about 50 nm. Fig. 4(b) displays that Co₃O₄-b is still flower-like shape by assembly of nanosheets, and the inset of Fig. 4(b) reveals that the surface of nanosheets becomes fairly rough compared to the nanosheets of Co-MOFs precursors, and the rough nanosheets are assembled by smaller nanoparticles which is less than 100 nm. The EDX spectrum in Fig. 4(c) and (d) further confirm that the two samples all contain Co and O elements, and no other elements are detected, which are consistent with the results of XRD. The results provide an efficient way to facilitate the formation of the tunable morphologies of Co₃O₄.

More structural information of Co₃O₄-a and Co₃O₄-b were investigated by TEM and HRTEM characterization. As shown in Fig. 5(a), the projection profile of Co₃O₄-a remains twin hemispherical shape, agreeing well with the morphology in the FESEM observation. The high magnification TEM image in Fig. 5(b) shows the Co₃O₄ twin hemispherical architecture is porous structure, which is composed of numerous nanoparticles. The reason may be the successive release and loss of CO₂, N₂O₅ and H₂O during the thermal decomposition of precursors. The TEM image of scattered nanoparticles in the inset of Fig. 5(b) indicates that the size of nanoparticles is from 23 to 45 nm.

As evidence, the HRTEM image (Fig. 5(c)) shows that these nanoparticles are attached with each other in various orientations, despite displaying clear lattice fringes.
with spacings of 0.46 nm and 0.24 nm, which correspond to the (220) and (311) planes, respectively. The selected area electronic diffraction (SAED) pattern recorded from the whole twin hemispherical architecture (inset of Fig. 5(c)) shows a series of concentric rings, indicating the twin hemispherical architecture is polycrystalline structure. Fig. 5(d) displays individual flower-like structure with the diameter of 8 μm. The high magnification TEM image in Fig. 5(e) shows that the nanosheets are porous structures which are assembled by nanoparticles. The sizes of particles are between 50 and 100 nm. The HRTEM image also displays clear lattice fringes with spacings of 0.46 nm and 0.24 nm in Fig. 5(f), which correspond to the (220) and (311) planes. The SAED pattern from the nanosheets (inset of Fig. 5(f)) shows a series of concentric rings, indicating the nanosheets are polycrystalline structures.

Nitrogen absorption–desorption isotherms were performed to obtain information about the specific surface areas and pore sizes of the calcined samples at 77 K. Fig. 6(a) and (c) indicates that Co₃O₄-a and Co₃O₄-b display typical IV adsorption–desorption isotherms with distinct hysteresis loop at a relative pressures of 0.68–1.0 and 0.79–1.0, respectively, which suggest the presence of mesoporous structures [33]. The Brunauer–Emmett–Teller (BET) specific surface area of Co₃O₄-a and Co₃O₄-b were calculated to be 22.6, 33.3 m² g⁻¹, respectively. The pore size distribution derived from the adsorption branch using the BJH method shows that the pore size of Co₃O₄-a is centered at about 2.5, 4.0 and 20.0 nm in Fig. 6(b), and the pore size of Co₃O₄-b is centered at about 2.45 and 21.0 nm in Fig. 6(d). According to the morphology and structure characteristic, the pore structures can be attributed to the decomposition process that the ligands were removed and resulted in the formation of small pores.

3.2. Electrochemical performances

Such porous Co₃O₄ hierarchical structures can not only shorten diffusion length for Li⁺, but also provide adequate buffer space for volume expansion to enhance lithium performance during the cycling process [34]. In order to prove the applicability of the porous Co₃O₄ hierarchical structures in lithium ion batteries, the electrochemical properties with respect to Li insertion/extraction were investigated, as shown in Fig. 7. In the charge–discharge curves of Co₃O₄-a and Co₃O₄-b in the voltage of 0.001–3.0 V (vs. Li⁺/Li) at a constant current density of 100 mA g⁻¹ at ambient temperature in Fig. 7(a) and (b), it can be clearly seen that there are two potential plateaus which is similar to the previous report [35]. As shown in Fig. 7(a), the initial discharge capacity of Co₃O₄-a is as high as 1325.5 mA h g⁻¹, which is higher than
the theoretical capacity (890 mA h g \(^{-1}\)). The first charge capacity is approximately 1003.5 mA h g \(^{-1}\), and the coulombic efficiency is about 75.7%. The irreversible capacity in the first cycle is usually ascribed to the decomposition of the electrolyte to form SEI layers and interfacial lithium storage, which is common for most anode materials [36]. The reversible capacities of the electrode in the 2nd, 5th and 10th cycles are about 965 mA h g \(^{-1}\), 840.6 mA h g \(^{-1}\), 749 mA h g \(^{-1}\), respectively, and the coulombic efficiencies are all above 1, which indicate that the porous hierarchical Co\(_3\)O\(_4\)-a electrode has a much higher capacity as well as good capacity retention. Furthermore, its coulombic efficiencies of Co\(_3\)O\(_4\)-a and
Co$_3$O$_4$-b are almost all above 100% except for the first charge–discharge cycles. It might be due to the fact that in the discharge process more Li involve in the active material of Co$_3$O$_4$, but in the charge process not all of Li are removed from Co$_3$O$_4$. The minor irreversibility of active material results in the columbic efficiencies of both samples higher than 1. Compared with the twin hemispherical porous Co$_3$O$_4$ structures, the flower-like porous Co$_3$O$_4$ structures (Co$_3$O$_4$-b) hinder more lithium cations to involve in the lithiation/delithiation processes of Co$_3$O$_4$, and present relatively low capacities in Fig. 7(b). Therefore, the morphology and structure of Co$_3$O$_4$ play crucial roles in the high performance anode materials for lithium batteries. The cycling performances of Co$_3$O$_4$-a and Co$_3$O$_4$-b at a current density of 100 mA g$^{-1}$ are shown in Fig. 7(c). The capacities of the Co$_3$O$_4$-a and Co$_3$O$_4$-b electrodes achieved a maximum initial capacities of 1325.5 and 1243 mA h g$^{-1}$, respectively. After 90 cycles, the reversible capacity of Co$_3$O$_4$-a retains 470.3 mA h g$^{-1}$, which is lower than the value (529.2 mA h g$^{-1}$) of Co$_3$O$_4$-b capacity. Even so, the values are all higher than the commercial graphite anode (372 mA h g$^{-1}$). The excellent performance of porous Co$_3$O$_4$ hierarchical structures are probably attributed to that the small nanoparticle subunits and porous structures facilitate the electrolyte penetration and Li$^+$ ions transport in the electrode and the hierarchical porous structures could efficiently buffer the stress caused by the volume variation during the charge–discharge process. Notably, twin hemispheric porous Co$_3$O$_4$ structures show better capacity retention and stability due to the maximization of the beneficial effects arising from the compact porous hierarchical structures, which could better withstand the repeated volume expansion/contraction and facilitate the lithium ions insertion with reduced diffusion length [37].

To understand the electrochemical performances, the cyclic voltamograms (CV) measurements were performed on Co$_3$O$_4$-a and Co$_3$O$_4$-b electrodes to characterize their electrochemical reactions in Li ion batteries. Fig. 8 shows the first three consecutive CV curves of the electrodes made from Co$_3$O$_4$-a and Co$_3$O$_4$-b recorded between 5 mV and 3 V at a scan rate of 0.1 mV s$^{-1}$. In the first cycle, the cathodic peaks are observed at around 1.10 V for Co$_3$O$_4$-a and at 0.70 and 0.33 V for Co$_3$O$_4$-b, which can be attributed to the electrochemical reduction (lithiation) reaction of Co$_3$O$_4$ with Li and formation of partially irreversible solid electrolyte interface (SEI) [38]. The main anodic peaks are observed at 2.10 V for Co$_3$O$_4$-a and at 1.64 V for Co$_3$O$_4$-b, which could be ascribed to the oxidation of CoO to Co$_3^{3+}$. Compared to the first cycle, the peak intensity and integral areas of the second and third

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cycles are reduced for Co$_3$O$_4$-a but are decreased more obviously for Co$_3$O$_4$-b, which indicate that the electrochemical reversibility of Co$_3$O$_4$-a was gradually built and highly stable SEI layer was formed after the initial cycle and much better than those for Co$_3$O$_4$-b. Additionally, cathodic peak about 1.1 V and anodic peak about 2.1 V shift to lower potential for Co$_3$O$_4$-a, which suggest drastic, lithium-driven, structural or textural modification during both the anodic and cathodic processes. Electrochemical impedance spectroscopy is carried out to identify the charge transfer resistance in the Co$_3$O$_4$-a and Co$_3$O$_4$-b electrode materials. The Nyquist plots of Co$_3$O$_4$-a and Co$_3$O$_4$-b in the frequency range from 100 kHz to 10 mHz are shown in Fig. 9. The plots consist of depressed semicircles in the high frequency regions and straight lines in the low frequency regions. Generally, the semicircle in the high frequency region represents the resistance of the lithium ion migration through the interface between the surface layer of the particles and electrolyte, while the linear region in the low
frequency region corresponds to the Warburg impedance related to the diffusion of the Li-ions in the Co$_3$O$_4$ electrodes. The inset of Fig. 9 shows that Co$_3$O$_4$-a and Co$_3$O$_4$-b have low high frequency resistance about 162 and 75 ohms, which suggests that the twin hemispherical and flower-like structures provide efficient lithium diffusion tunnels and increases the electrolyte/Co$_3$O$_4$ contact area. Based on above, it can be believed that the high capacity and good capacity retention of the Co$_3$O$_4$-a and Co$_3$O$_4$-b anode could probably be ascribed to the porous structures verified by HRTEM and BET measurements. Firstly, the small Co$_3$O$_4$ nanoparticles can shorten the diffusion length of Lithium ions insertion/extraction. Secondly, the porous structure can effectively buffer space for the volume expansion of Co$_3$O$_4$ during the cycling process so that improved the cycling stability.

4. Conclusion

A large scale and facile method was employed to synthesize Co-MOFs hierarchical structures with the assistance of PVP. Meanwhile, thermal decomposition was developed to successfully prepare intriguing twin hemispherical and novel flower-like porous Co$_3$O$_4$ hierarchical structures due to the successive release and loss of CO$_2$, N$_2$O$_5$ and H$_2$O, which was assembled by many uniform nanoparticles. The results indicate that Co$_3$O$_4$-a and Co$_3$O$_4$-b remain the twin hemispherical and flower-like hierarchical structures compared with Co-MOFs-a and Co-MOFs-b, which are attributed to Co-MOFs hierarchical structures as self-sacrificial templates. Such complex hierarchical structures might bring unique physical chemistry properties. The results demonstrate that these complex hierarchical structures, especially the twin hemispheric porous Co$_3$O$_4$ hierarchical structures, exhibit remarkable electrochemical performance as anode materials for lithium ion batteries. It can be anticipated that these complex hierarchical structures would find their promising applications in many areas.

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