Large interlayer spacing vanadium oxide nanotubes as cathodes for high performance sodium ion batteries†

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Sodium ion batteries (SIBs), as a potential alternative to Li-ion batteries (LIBs), have attracted great attention from researchers. Herein, large interlayer spacing ferric ion substituted vanadium oxide nanotubes (Fe-VNTs) are fabricated using dodecylamine as a template via a facile hydrothermal treatment followed by a ferric ion substitution process. The distances between the adjacent layers of VNT, Fe-VNTs and orthorhombic V$_2$O$_5$ are 2.7 nm, 1.2 nm and 0.44 nm, respectively. The larger interlamellar spacing results in faster Na$^+$ diffusion reaction kinetics, and the insertion of ferric ion into vanadium oxide layers removes the organic templates between the vanadium oxide layers, leading to high conductivity and small electrochemical reaction resistance. Serving as the sodium ion battery cathode, Fe-VNTs display enhanced sodium storage performance over orthorhombic V$_2$O$_5$.

1. Introduction

In the past decades, much attention has given to energy production and storage technologies. Lithium-ion batteries (LIBs), as one of the most important energy-storage technologies, are the main candidates for large-scale energy storage systems (ESSs). The increasing requirement for lithium may cause the price of lithium to skyrocket due to its low abundance and uneven distribution. Hence, the discovery of cheaper alternatives to replace lithium has attracted great interest. By virtue of the wide availability and low-cost of sodium resources, sodium-ion batteries (SIBs) have emerged as one of the promising substitutes for LIBs. Meanwhile, SIBs have a similar ion intercalation mechanism.

Vanadium oxides, with unique open-layered structures, are considered as one of the most promising electrode materials for electrochemical energy storage due to its high theoretical capacity, abundance resource and low cost. Generally, the layered structures contain [VO$_6$] octahedral geometry and/or [VO$_5$] square pyramid with a long V=O bond and a short vanadyl bond, $V=\text{O}$. Various vanadium oxide layered structures are formed and exhibit promising sodium storage performance. The crystal structure of orthorhombic V$_2$O$_5$ can be described as layers of VO$_3$ square pyramids, with an interlayer spacing of 4.37 Å, exhibits the quite attractive electrochemical performance when used as cathode materials of LIBs. However, Na$^+$ ions (1.02 Å) are larger compared to Li$^+$ ions (0.76 Å) resulting in the kinetically sluggish of Na$^+$ insertion/extraction and low diffusion rate in the compact crystal structure. Therefore, rational design of cathode materials with larger diffusion channels/suitable lattice space to host Na$^+$ ions and promote rapid Na$^+$ insertion and extraction are of great importance. To enhance Na storage performance via controlling the interlayer spacing has been recognized in few works. Wang et al. synthesized single-crystalline bilayered V$_2$O$_5$ nanobelts with large (001) interlayer spacing (11.53 Å) can accommodate Na$^+$ intercalation and deintercalation. Additionally, Mai et al. synthesized V$_2$O$_5$-nH$_2$O xerogel with water molecules intercalated into layer structure which can enlarge interlayer spacing and enhance high initial capacity.

In this study, we successfully synthesize vanadium oxide nanotubes (VO$_2$-NTs) with super wide interlayer spacing using dodecylamine as templates through a simple hydrothermal treatment. Since organic templates do not possess good conductivity and will block the diffusion of Na$^+$ during the insertion-extraction. In order to improve the electrochemical performance, the following ferric ion substitution process was carried out to remove the organic template. This ferric ion substitution reaction can effectively remove the amine templates within VO$_2$-NTs, and maintain their tubular morphology and multiwalled structure. It is expected that the as-synthesized ferric ion substituted vanadium oxide nanotubes (Fe-VO$_2$-NTs) may exhibited faster diffusion reaction kinetics, and accommodated larger volume changes, benefiting for the sodium storage.

2. Experimental

2.1. Material preparation

2.1.1 Synthesis of VNTs. Vanadium oxide nanotubes (VNTs) were prepared as following: 0.51 g of V$_2$O$_5$ powder and...
0.52 g of dodecylamine (C12H25NH2) were dissolved in 40 ml distilled water under vigorous stirring for two days. The obtained solution was transferred to a 50 ml Teflon lined autoclave and heated at 180 °C for 5 days in an oven and then cooled to room temperature naturally. The final product was filtered, washed with de-ionized (DI) water and pure alcohol several times and then dried at 70 °C in vacuum for 12 h.

2.1.2 Synthesis of Fe-VNTs. VO2-NTs (500 mg) was dispersed in a mixture of distilled water and ethanol (10 ml and 80 ml respectively) by sonication for 2 h. Then, 5 g of FeCl3-6H2O was dissolved in 10 ml of de-ionized (DI) water. After that, the FeCl3-6H2O solution was slowly added into VNTs suspension and kept continuously stirring for 24 h. The precipitate was filtered, rinsed, and dried under vacuum at 80 °C for 12 h to obtain Fe-VNTs.

For comparison, crystalline V2O5 powder was also employed as a controlled group.

2.2. Characterizations

The morphology, structure and composition of the as-prepared samples were analyzed by field emission scanning electron microscopy (FESEM, Philips-XL-30FEG) and transmission electron microscopy (TEM, JEOL-1230), X-ray powder diffraction (XRD) patterns were obtained by using a RigakuD/Max-2500 diffractometer with Cu Kα radiation source (λ = 1.5406 Å). Fourier transform infrared spectroscopy (FTIR) was measured using a Bruker-TENSOR27 FTIR spectrometer over the range from 400 to 4000 cm⁻¹. The thermogravimetry analysis (TGA) and differential scanning calorimeter (DSC) were carried out on a SDT Q600 over the temperature range from 50 to 800 °C using a heating rate of 10 °C min⁻¹ under air atmosphere. XPS experiments were carried out on a RBD upgraded PHI-5000C ESCA system (Perkin-Elmer) with Mg Kα radiation (hν = 1253.6 eV). Binding energies were calibrated by using the containment carbon (C1s = 284.6 eV).

2.3. Battery assembly and electrochemical performance measurements

The SIBs performance for the as-made samples was evaluated by using 2032-type coin cells with a voltage window of 1.5–3.5 V. The working electrodes were prepared by mixing 70 wt% active materials, 20 wt% carbon black as a conducting agent and 10 wt% poly (vinylidene fluoride) (PVDF) as a binder. N-Methylpyrrolidone (NMP) solvent was added to the mixture to form a slurry, which was uniformly pasted on aluminum foils. The coated electrodes were dried in vacuum at 120 °C for 10 h and then cut into disks. The coin cells can be assembled with a sodium metal slice, electrolyte (1 M NaClO4 in ethylene carbonate (EC) and diethyl carbonate (DEC) (volume ratio of 1 : 1) with 5 wt% fluoroethylene carbonate (FEC) additive), a glass fiber separator (Whatman) and a slice of the as-made sample in Ar-filled dry glovebox. For comparison, crystalline V2O5 powder was also used as electrode to assemble 2032-type coin cells via similar procedures. Cyclic voltammograms (CV) were collected usingCHI660C (Chenghua, Shanghai) electrochemical workstation at a scan rate of 0.5 mV s⁻¹. Galvanostatic charge/discharge tests were performed by using LAND cell-testing system. An AC signal of 5 mV was employed to measure electrochemical impedance spectra (EIS) within the

Fig. 1 Low- and high-magnification FE-SEM images of the (a and c) VNTs (b and d) Fe-VNTs.
frequency range from 0.01 Hz to 100 kHz. All tests were carried out at the room temperature.

3. Results and discussion

Fig. 1 shows the FE-SEM images of VO\textsubscript{x}-NTs and Fe-VO\textsubscript{x}-NTs. Clearly, both of the samples exhibit the hollow tubular structure with a diameter from 50 to 130 nm. The tubular morphology of VO\textsubscript{x}-NTs are under preservation during ferric ions substitution process.

Fig. 2 represents the HRTEM images of VO\textsubscript{x}-NTs and Fe-VO\textsubscript{x}-NTs at different magnifications, respectively. Both two samples exhibit a well-defined multilayered structure with an open-ended tube. The Fe-VO\textsubscript{x}-NTs show larger inner diameter (50–80 nm) than that of VO\textsubscript{x}-NTs (30–50 nm). The removal of ferric ion and Fig. 2b shows the interlayer spacing of VO\textsubscript{x}-NTs are expanded to be 2.7 nm under dodecylamine embedding. The dark fringes are VO\textsubscript{x} layers and the light fringes represent dodecylamine templates. To improve electrochemical performance, the removal of amine templates is necessary. As shown in Fig. 2d, the distance between VO\textsubscript{x} layers within Fe-VO\textsubscript{x}-NTs decreases obviously from 2.7 to 1.2 nm, in agreement with the TEM results. The shrinkage of interlayer spacing results from the effective removal of dodecylamine templates. It can be calculated that the distance between adjacent layers are changed from 2.7 to 1.2 nm, in agreement with the TEM results. The typical (hk0) diffraction peaks are at same position after ferric ion substitution process which indicate two-dimensional structure of VO\textsubscript{x} layers remain unchanged.

Fig. 4 is the schematic illustration of the ferric ion substitution. For VNT, the protonated dodecylamine cations (C\textsubscript{12}H\textsubscript{25}NH\textsubscript{3}\textsuperscript{+}) is positively charged and intercalates the negatively charged VO\textsubscript{x} layers through electrostatic interaction. However, this interaction is broken, when more positively charged Fe\textsuperscript{3+} with smaller size insects into VO\textsubscript{x} layers. Meanwhile, the organic templates are removed out of the VO\textsubscript{x} layers and lead to the shrinkage of interlayer spacing.

The TG-DSC curves of samples are showed in Fig. 5. The two samples undergo significant thermal weightlessness owing to
the decomposition of organic templates. The weight loss of VO\textsubscript{x}-NTs and Fe-VO\textsubscript{x}-NTs are 52% and 23%, respectively. It can demonstrate that a large content of organic templates are removed by ferri ion substitution reaction. There are two endothermic peaks locate at 220°C and 380°C on DSC curve corresponding to the decomposition of dodecylamine and the crystallization of vanadium oxide, respectively.

The samples are further characterized by Fourier transform infrared (FTIR) spectra, X-ray photoelectron spectroscopy (XPS). For VO\textsubscript{x}-NTs, the peak at 575 and 797 and 1002 cm\(^{-1}\) are in Fig. 6 are corresponding to the vibrations of O–(V)\(_3\) bending, V–O–V vibration and stretching vibration of V=O.\(^{33,34}\) The absorption bands located at 2853 and 2927 cm\(^{-1}\) are attributed to the various C–H vibrations in the organic templates.\(^{33}\) Both of the samples have the board peak at 1623 and 3445 cm\(^{-1}\) which are assigned to H–O–H bending vibration and H–O stretching.\(^{33,36}\) Remarkably, there are some differences between two samples. The intensity of the peaks at 2927 cm\(^{-1}\) association with organic templates in Fe-VO\textsubscript{x}-NTs has weakened significantly, and the peak at 2853 cm\(^{-1}\) disappears (inset figure of Fig. 6), which indicates large amount of templates are removed after ferric ion substitution.
X-ray photoelectron spectroscopy (XPS) measurements is further performed to identify the elemental composition of two samples and study the valence change of vanadium. From XPS general spectra of VNTs and Fe-VNTs in Fig. 7a and b, the Fe-VNTs show weaker C1s peaks and new Fe2p peak which demonstrate the organic templates are effectively removed and the existence of ferric ions. The N1s peak corresponding to \(-\text{C-NH}_2\) group disappeared in Fig. 7b due to less organic templates within Fe-VNTs. The V2p\(_{3/2}\) are comprised of two parts locating at the banding energy of 517.7 and 516.5 eV which are consistent with the two formal reported values for V\(^{5+}\) and V\(^{4+}\).\(^{37,38}\) The V\(^{5+}/\)V\(^{4+}\) ratio of VNT and Fe-VNT are 0.93 and 1.13 (Fig. 7c and d), respectively. The V\(^{5+}\) concentration in samples is increased after ferric ion substitution, which may caused by the Fe\(^{3+}\) ions. To further proved it, the spectra of Fe2p is involved (Fig. 8). The peak at 711.4 eV is observed which is attributed to Fe\(^{2+}\) in addition to a strong peak at 712.7 eV is ascribed to Fe\(^{3+}\).\(^{39}\) The result indicates Fe\(^{3+}\) are partially reduced to Fe\(^{2+}\) in Fe-VNTs. It is clear that the strong oxidizing Fe\(^{3+}\) has oxidized some V\(^{4+}\) into V\(^{5+}\) which cause the appearance of Fe\(^{2+}\). Furthermore, the atomic ratio of Fe/V is 0.053/1, which was figured out by inductively coupled plasma (ICP) analysis.

Electrochemical performance of two samples are evaluated as a cathode material for NIBs by assembling the coin cell (2032-type). Fig. 9a shows the initial charge-discharge curves in 1.5–3.5 V at current density of 50 mA g\(^{-1}\). Both VNTs and Fe-VNTs exhibit smooth charge and discharge without obvious discharge plateau, manifesting amorphous features.\(^{17}\) The VNTs show a low first discharge specific capacity of 138 mA h g\(^{-1}\). By contrast, the Fe-VNTs deliver a higher discharge specific capacity of 185 mA h g\(^{-1}\). This might due to
the release of organic templates and the preintercalation of ferric ion. While the orthorhombic V\textsubscript{2}O\textsubscript{5} exhibits lowest discharge capacity of 76 mA h g\textsuperscript{-1} and worst reversible discharge capacity (Fig. S1†), originating from its compact crystal structure which may causes the slow sodium ion diffusion. Cyclic voltammetry (CV) (Fig. 9b) were carried out at a scan rate of 0.5 mV s\textsuperscript{-1}. Different from VNT which has no pronounced peaks (Fig. S3†), one cathodic peak at ~2.07 V and a pair of anodic peaks at ~2.61, 2.98 V appear in the CV curves of Fe-VNTs. Larger surrounded area of the Fe-VNTs demonstrates higher capacity and faster kinetic during electrochemical insertion/extraction than those of VNTs. Fig. 10 shows cycling performance of the samples at different discharge current density of 50 and 80 mA g\textsuperscript{-1}. By 50 cycles at 50 mA g\textsuperscript{-1} (Fig. 10a), the Fe-VNTs deliver a discharge specific capacity of 70 mA h g\textsuperscript{-1} which is higher than 25 mA h g\textsuperscript{-1} of VNTs. Fe-VNTs have a better cycling performance than VNTs. The cycling performance of orthorhombic V\textsubscript{2}O\textsubscript{5} has shown in Fig. S2,† the second discharge specific capacity drops from 76 to 11 mA h g\textsuperscript{-1} and remain below 10 mA h g\textsuperscript{-1} in the following cycling. The small interlayer spacing hinder the insertion/extraction of Na\textsuperscript{+}.

It is excepted that VNTs show better electrochemical performance than Fe-VNTs due to its larger interlayer spacing which allow more Na\textsuperscript{+} to insert. However, large amount amine templates within VNTs will impede the diffusion of Na\textsuperscript{+}. It can demonstrate the ferric ion substitution process facilitate electrochemical capacity and cycling performance.

To understand the increased electrochemical performance of Fe-VNTs, the electrochemical impedance spectra (EIS) were measured (Fig. 11a). The Nyquist plots consist of a compressed semicircle in the high-frequency region, which related to the charge transfer resistance (\(R_{ct}\)) of the electrode, and a sloped line in the low frequency region which corresponds to the Warburg impedance (\(W_\omega\)). In this equivalent circuit, \(R_e\) stands for all ohmic resistance, including the electrolyte and other part of the system. \(R_{ct}\) represents the charge-transfer impedance at the electrode/electrolyte interface. CPE and \(W_\omega\) are constant phase-angle element and Warburg impedance, respectively. The \(R_{ct}\) value of VNT is 1589.9 $\Omega$ and that of Fe-VNTs is 337.7 $\Omega$. By contrast, the orthorhombic V\textsubscript{2}O\textsubscript{5} delivers the \(R_{ct}\) value of 2325 $\Omega$ (Fig. S3†). Obviously, it demonstrates that Fe-VNTs possesses faster kinetics of electrochemical reaction and higher capacity, which owe to its large interlayer spacing boosting sodium insertion/extraction. The sodium ion diffusion coefficient value (\(D\)) is proportional to the reciprocal of squared value of Warburg factor \(\sigma(D_{Na} \propto 1/\sigma^2)\).\textsuperscript{40-42} \(\sigma\) is relative to \(Z_{re}\). From the slop of the lines in Fig. 11b \(\sigma\) can be obtained.\textsuperscript{43}

\[
Z_{re} = R_D + R_L + \sigma\omega^{-1/2}
\]

Na\textsuperscript{+} diffusion coefficient at room temperature can be indirectly evaluate by calculating \(\sigma\) values. As calculated, \(\sigma\) value of Fe-VNTs is 167 which is lower than 654 of VNT, indicating the Fe-VNTs possess better Na\textsuperscript{+} diffusion. The increase of sodium ion diffusion coefficient results from the release of organic templates which facilitate Na\textsuperscript{+} intercalation. To be specific, it can provides more open channels for facile Na\textsuperscript{+} intercalation and deintercalation. As discussed above, enhancement electrochemical performance of Fe-VNTs can be well explained by ferric ion substitution.

Fig. 8 XPS spectrum of Fe2p.

Fig. 9 (a) Galvanostatic charge–discharge profiles of first cycle of VNTs and Fe-VNTs between 1.5–3.5 V at 50 mA g\textsuperscript{-1}. (b) Cyclic voltammograms of VNTs and Fe-VNTs at scan rate of 0.5 mV s\textsuperscript{-1}.
4. Conclusions

The Fe-VNTs were prepared by a facile hydrothermal treatment and followed by ion substitution process. With the ferric ion preintercalation, the interlayer spacing is reduced from 2.7 to 1.2 nm. The enhanced electrochemical performance of Fe-VNTs are mainly attributed to the reasons as the following: (i) the organic templates which have no electrochemical contribution are effectively removed with its morphology and structure remain. (ii) The insertion of ferric ion lead to enhanced electronic conductivity and good charger transfer ability for reversible sodium storage capacity. (iii) The large interlayer space between Fe-VNTs can facility the kinetics of sodium ion intercalation/deintercalation. The novel ferric ion substituted vanadium oxide nanotubes demonstrate a great potential as cathode material for Na\(^+\) ion batteries application.

Conflicts of interest

There are no conflicts to declare.

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