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Effects of amorphous and crystalline MoO₃ coatings on the Li-ion insertion behavior of a TiO₂ nanotube anode for lithium ion batteries†

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Amorphous and crystalline MoO_3 coatings are synthesized on anodic TiO_2 nanotube arrays by electrodeposition, as a self-standing, binder-free anode material in Li-ion batteries for enhancing the Li-ion insertion performance. The amorphous MoO_3 layer is uniform and conformal with a thickness of 10 nm, and is converted into crystalline nanoparticles via thermal treatment. Our results show that both the coated TiO_2 nanotubes deliver much higher areal capacities than bare nanotubes or a dense crystalline α - MoO_3 film, while the crystalline α - MoO_3 coating greatly increases the areal capacity of TiO_2 nanotubes compared to the amorphous. The results are obtained at 1340 μ A h cm⁻² initial capacity for nanotubes with a crystalline coating, 977 μ A h cm⁻² for those with an amorphous coating and 342 μ A h cm⁻² for the bare ones. The significant enhancement is due to a combination of MoO_3 with high specific capacity and TiO_2 nanotube arrays with large surface area allowing uniform MoO_3 deposition and rapid ionic transfer. Crystalline α - MoO_3 is better than amorphous MoO_3 and the coating medium is discussed in terms of chemical state, crystal defects, capacitive contributions and the charge–discharge kinetics in coated TiO_2 nanotube electrodes.

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Introduction

Rechargeable lithium-ion batteries (LIBs) have drawn tremendous interest in the past decades due to their high energy density, light weight, low self-discharge and long lifespan.^{1,2} At present, graphite is the most commonly-used anode material in commercial LIBs, but its drawback of easy formation of lithium dendrites, hinders the development of compact LIBs with higher operation voltage, larger power density and better cell safety.3 TiO₂ is a promising alternative to graphite because of its high safety and good cyclability, low self-discharge rates and small volume change (<3%).4 It is also attractive owing to its abundant resource, low cost, chemical stability and harmless properties. Self-organized TiO2 nanotube arrays formed by anodic oxidation could serve as an excellent host for electrons and Li⁺ ions, and their tight contact with the metallic substrate allows electrons to be collected directly and efficiently.5 However, the Li-ion insertion performance of anatase TiO2 nanotubes (\sim 168 mA h g⁻¹) is poor and needs to be improved. Several methods have been proposed for this using surface modifications with metal, 6,7 or other anode materials such as SnO₂,8 Cu₆Sn₅,9 NiO,10 Co₃O₄,11 Fe₂O₃,12 etc.

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Besides TiO₂, orthorhombic α-MoO₃ is also an excellent anode material for LIBs, with a high theoretical capacity of 1111 mA h g⁻¹, almost three times that of the graphite.¹³ Recently we reported α-MoO₃ deposited TiO₂ nanotube composites showing good Li-ion storage capacities and stable cycling performances.14 α-MoO₃ can be obtained by crystallization of amorphous MoO₃ that is well-known for its excellent electrochromic behavior during the insertion/desertion process of ions including Li⁺ and H⁺. 15-17 Although electroinsertion behaviors of amorphous and crystalline MoO3 are reported, 15,18 impacts of metal oxidation states and morphological evolution of MoO₃ coatings on Li-ion insertion of MoO3-deposited TiO2 nanotubes have not been investigated yet. It remains unclear if ordered TiO₂ nanotubes are qualified to act as a robust substrate to MoO₃ for its better performance in thin-film electrochromic devices and electrochemical batteries. Hence, we propose synthesis of amorphous MoO3 coatings onto highly-oriented TiO2 nanotubes by employing a controllable and facile electrodeposition method. Evolution of the surface coating is studied in detail in terms of morphological features, chemical state and electrochemical kinetics.

In this work, ordered anatase TiO₂ nanotubes were synthesized by potentiostatic anodization and subsequent thermal treatment. Electrodeposition of MoO₃ was performed on these nanotube arrays and bare Ti foils to obtain amorphous coatings that are transformed into the crystalline phase upon annealing. We compared the effects of amorphous and crystalline MoO₃ coatings on the Li-ion intercalation capacity of TiO₂ nanotubes.

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Our results demonstrate that these TiO2 nanotubes could be an excellent substrate in thin film MoO3 electrodes for high performance LIBs.‡

Results and discussion

Fig. 1 presents SEM images of bare and MoO₃-coated TiO₂ nanotubes. The top-view (Fig. 1a) and cross-sectional images (Fig. S1†) show well-ordered TiO₂ nanotubes (B-TN) with the top end open at an average inner diameter of ~110 nm and a length up to 9 μ m. It can be seen from the inset of this figure that the hollow tubes are directionally arrayed with individual and smooth walls. Apparently, their inner and outer walls provide large surface area which offers numerous sites for MoO₃ deposition in order to form an effective hybrid structure. Fig. 1b gives SEM images of TiO2 nanotubes coated with 8 deposition cycles of MoO₃ (AMO-TN). Compared to B-TN, their walls are

‡ Experimental details: titanium foils (99.5 wt%, 250 µm thick) were purchased from Alfa Aesar, USA. Prior to electrochemical treatment, the Ti foils were degreased and rinsed by sonicating in ethanol and deionized water, and then covered by Scotch tape at their back. A two-electrode cell composed of a Ti foil as working electrode and a Pt foil as the counter electrode was assembled for anodization. TiO2 nanotubes (designated as B-TN) were synthesized in ethylene glycol (EG, Alfa Aesar) electrolytes containing 0.3 wt% NH₄F (MP Biomedicals, USA) and 5 vol% H₂O at 70 V for 40 min. After oxidation was completed, samples were immediately rinsed with deionized water, dried and annealed at 450 $^{\circ}\mathrm{C}$ for 1 h in air, to obtain crystalline TiO2 nanotubes. Later, electrochemical deposition of MoO3 onto these oxide nanotubes was carried out using a VersaSTAT 3F potentiostat/galvanostat with FRA (Princeton Applied Research, USA) with a three-electrode setup composed of anodized Ti as the working electrode, a Pt foil as the counter electrode and an Ag/AgCl reference electrode. The deposition was performed in 0.05 M (NH₄)₆Mo₇O₂₄ (Alfa Aesar) aqueous solution by cycling from 0 to -0.7 V with a step rate of 20 mV s⁻¹ for 8 or 16 cycles. The as-deposited TiO₂ nanotubes are designated as AMO-TN, which were then annealed in air at 450 °C for 1 h to obtain crystalline MoO₃-coated nanotubes (CMO-TN). For comparison, a MoO3 film (400MO) was grown by electrodeposition of MoO3 on Ti via 400 cycles and post thermal treatment at 450 °C for 1 h. The way to measure masses of TiO2 or MoO3 was described in our previous work.14 Areal masses of B-TN, AMO-TN, 400MO and weight ratios of MoO_3 are approximately 2.2 mg, 3.0 mg (26.7 wt%) and 0.27 mg. CMO-TN is considered to have the same composition and mass with AMO-TN.

A Hitachi S-4800 scanning electron microscope (SEM) and a Hitachi H9000NAR transmission electron microscope (TEM) were employed to observe the oxide films. Their crystallographic structure was determined by X-ray diffraction (XRD) using a Scintag XDS 2000 X-ray diffractometer with Cu K_{α} radiation at a scan rate of 2° min⁻¹. Chemical composition of deposited TiO₂ nanotubes was characterized by an energy dispersive X-ray spectroscope (EDX) with an XFlash detector (Bruker AXS) on SEM, and an X-ray photoelectron spectroscope (XPS) on a Hewlett-Packard 5950A spectrometer using a monochromatized Al K_{α} X-ray radiation source. All XPS spectra were calibrated according to the binding energy of the C1s peak at 284.8 eV.

CR2032 coin cells were assembled with bare and MoO_3 -deposited TiO_2 nanotubes acting as the working electrode. A lithium disc served as the counter and reference electrode, which was separated from these oxide nanotubes by a Celgard 20 μm-thick monolayer polyethylene membrane. Electrolytes were 1 M lithium hexafluorophosphate (LiPF₆) dissolved in a mixed solvent of ethylene carbonate and diethylene carbonate in 1:1 volume ratio (Novolyte Technologies, Inc). All the cells were assembled in an Ar-filled glove box with oxygen and moisture levels lower than 1 ppm. The galvanostatic charge-discharge cycling was carried out on LANHE automatic battery testers (China) at a current of 200–1000 μA cm⁻² for 10 or 30 cycles. AC impedance measurements were also conducted on these cells at a test potential of ~1.7 V (versus Li/Li⁺) from 100 kHz to 0.1 Hz with a perturbation of 5 mV applied.

thickened as evidenced by open entrances showing a reduced inner diameter of ~90 nm. In addition, their tube walls remain smooth with a uniform coating layer in a thickness of \sim 10 nm, and the precipitates there are believed to contain various molybdenum oxides, 15,19 On the other hand, the coating layer is roughened after heat treatment, thanks to spontaneous origination of particles anchored on the tube walls (CMO-TN, Fig. 1c). The tubular structure survives, leaving a narrower entrance and lower porosity, because the walls of TiO2 nanotubes are further thickened and the pore size is mostly 50 nm. Fig. 1d presents an oxide film (400MO) formed by electrodeposition of molybdenum oxides directly on Ti. Edged particles with a size up to hundreds of nanometers are aggregated to yield a dense film only with a few cracks. They are polycrystalline with an orthorhombic crystal structure (α phase).¹⁵ As a result, it is understood that the hollow nanotubes not only work as a support for MoO3 deposition, but offer numerous channels to let electrolytes readily infiltrate and Li⁺ ions rapidly transfer in a fast charge-discharge process, which is likely to achieve better Li-ion insertion behaviour of TiO2 anodes in LIBs.

MoO₃-deposited TiO₂ nanotubes are examined by TEM as well. Fig. 2a shows nanotubes from AMO-TN with a conformal coating layer of \sim 9.4 nm thickness on their outer wall. It is assumed that such a coating also exists on inner walls since the pores are quite wider than the tube gaps as shown in Fig. 1a, and thus electrolytes can easily access the inner walls for deposition. It is proved by reducing the inner diameter of TiO₂ nanotubes from 110 nm to 90 nm and to 30 nm, as the deposition cycles of amorphous MoO3 increase from zero to 8 and to 16 (Fig. 1a and b and S2a†). Direct deposition of MoO₃ inside the nanotube is also reported in previous work. 17,20 Upon heating many α-MoO₃ nanoparticles are further viewed on both the inner and outer walls due to the crystallization process (Fig. S2b†). The particles could be seen in Fig. 2b as well. When compared, there are fewer particles in Fig. 2a, suggesting the growth of α-MoO₃ particles on CMO-TN rather than AMO-TN and B-TN. In addition, co-existence of polycrystalline α-MoO₃ and anatase TiO2 is also confirmed by indexing these spots in a selected area electron diffraction ring pattern of CMO-TN, as seen in Fig. 2b. Three rings consisting of the spots are exampled. With their diameters, we obtain an interplanar spacing of \sim 0.355 nm for the anatase TiO₂ (101) plane, 0.271 nm for the α - MoO_3 (101) plane and ~ 0.185 nm for the α -MoO₃ (002) plane, but a great amount of interplanar spacings of anatase TiO2 and α -MoO₃ are too close to index other spots readily in the pattern. An EDX analysis is also made from TEM observation to claim the presence of Mo, Ti and O elements. An EDX quantitative analysis of Ti (59.14 at%) and Mo (12.67 at%) in the nanotubes provides an elemental ratio of 27.8 wt% for α -MoO₃ in CMO-TN, which is quite close to 26.7 wt% of the mass ratio obtained by weighing samples.

Both EDX and XPS techniques are employed to analyze the elemental distribution and chemical states of the deposited TiO₂ nanotubes. Fig. 3a shows the EDX mapping of AMO-TN in which elements including Mo, Ti and O are detected and well distributed in the TiO2 nanotube film, suggesting formation of a uniform molybdenum oxide coating on TiO₂ nanotube arrays.

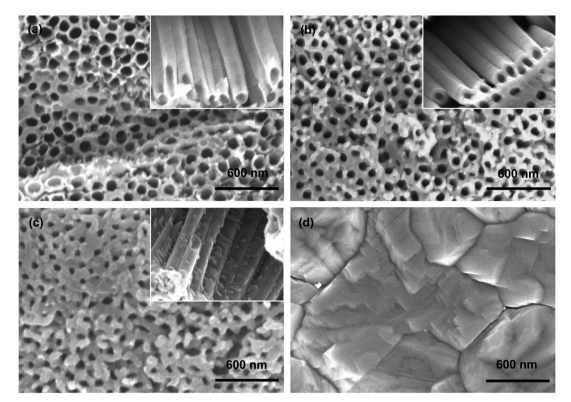


Fig. 1 SEM images of (a) B-TN, (b) AMO-TN, (c) CMO-TN and (d) 400MO.

A detailed XPS analysis reveals that the as-formed coating contains Mo^V oxides and Mo^{VI} oxides, since the $Mo3d_{5/2}$ peak at 232.0 eV and the $Mo3d_{3/2}$ peak at 235.2 eV correspond to Mo^V , and the $Mo3d_{5/2}$ peak at 236.2 eV $(\Delta E = 3.1 \text{ eV})$ correspond to Mo^{VI} (Fig. 3b). The fraction of $Mo^{VI}O_3$ is as high as 50% in the amorphous coating, based on the area ratio of 1 : 1 between Mo^{VI} peaks and Mo^V peaks. Upon annealing, the XPS peak shape shifting to high energies is in line with α -MoO $_3$ (Fig. 3c). The area ratio between the two peaks $(3d_{5/2}:3d_{3/2})$ was almost 3 : 2, as expected for the coatings on

CMO-TN which are dominantly composed of Mo^{VI}. A tiny peak at 234 eV tells us that a trace amount of molybdenum species (MoO_x) in lower oxidation states may reside. As for the O1s peak, a slight shift (0.5 eV) to a lower energy is observed (Fig. 3d), which is in agreement with previous studies.^{12,20} The peak shift is attributed to elimination of OH⁻ groups and oxidation of Mo^V ions in the coating on annealing.¹⁹ Our results indicate a uniform coating composed of Mo^V and Mo^{VI} oxides on TiO₂ nanotubes after electrodeposition, which turns to be an α -MoO₃-predominant film right after annealing at 450 °C. In

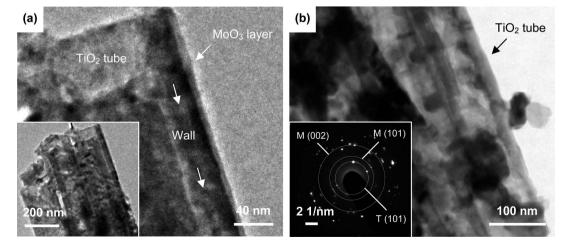


Fig. 2 (a) TEM images of AMO-TN, (b) TEM image of CMO-TN and an electron diffraction pattern taken from these nanotubes with distinguishable Miller indices of anatase TiO_2 (T(101)) and α -MoO₃ (M(101), M(002)).

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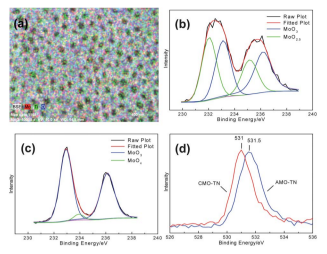


Fig. 3 (a) EDX mapping image of AMO-TN, high-resolution Mo3d XPS spectra taken from (b) AMO-TN, (c) CMO-TN, and (d) O1s XPS spectra taken from both samples.

summary, these results demonstrate that the MoO_3 -coated TiO_2 nanotube arrays have been successfully achieved by a facile electrodeposition in molybdenum salt solutions.

Anodic TiO₂ nanotubes are found to be in anatase phase after annealing at 450 °C by XRD and Raman. ^{14,21} Fig. 4 describes XRD patterns of MoO₃-coated TiO₂ nanotubes. Typical diffraction peaks for anatase TiO₂ are observed, including the (101) peak at $2\theta = 25.3^{\circ}$ and the (200) peak at $2\theta = 47.9^{\circ}$. Diffraction peaks of Ti are also found in both patterns, as evidenced by the (101) peak at $2\theta = 40.0^{\circ}$.

There are no XRD peaks found for molybdenum metal or oxides in the pattern of AMO-TN, suggesting deposition of amorphous MoO_3 layers onto anatase TiO_2 nanotubes. After heat treatment, the amorphous coating is gradually transformed to nanoparticles in the crystalline (α) phase, as evidenced by peaks (020), (040) and (021) of α -MoO₃ at $2\theta = 12.6^{\circ}$, 25.4° and 27.4°, respectively.

Based on above analyses, it is confirmed that electrodeposition and crystallization of $\alpha\text{-MoO}_3$ on anodic TiO_2 nanotube arrays results in a hybrid nanostructure. Fig. 5 vividly illustrates the morphological evolution of TiO_2 nanotubes with MoO_3

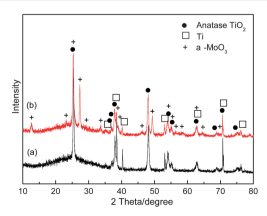


Fig. 4 X-ray diffraction patterns of (a) AMO-TN and (b) CMO-TN.

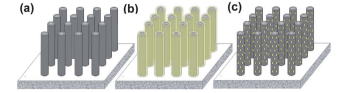


Fig. 5 Schematic showing (a) B-TN, (b) AMO-TN and (c) CMO-TN.

coatings. Initially, the formed nanotubes are well-arrayed on the Ti substrate by anodization and are then converted into anatase by annealing (Fig. 5a). Owing to electrodeposition of MoO₃, both the inner and outer walls of nanotubes are wrapped with a layer of amorphous MoO₃ to achieve a core–shell structure (Fig. 5b). Heat treatment induces crystallization within the amorphous layer, producing an amount of $\alpha\text{-MoO}_3$ nanoparticles (Fig. 5c) and yielding a TiO₂ nanotube supported $\alpha\text{-MoO}_3$ nanoparticle electrode.

Fig. 6 displays cycling performances of MoO₃ film, bare and deposited TiO2 nanotubes over 10 electrochemical cycles at a large current density of 800 μA cm⁻² (compared to the commonly-used current density of 50-200 µA cm⁻²)8-12,14 in a potential range of 0.05-3.0 V versus Li/Li⁺. Clearly, a capacity drop occurs after the initial cycle for all anodes, probably due to some fractions of Li⁺ ions becoming trapped inside the oxide from irreversible formation of a molybdenum bronze such as $\operatorname{Li}_{x+2}\operatorname{Mo}_{1-x}^{VI}\operatorname{Mo}_{x-z}^{VI}\operatorname{Mo}_{z}^{IV}\operatorname{O}_{3-y-z}$, or due to decomposition of organic solvent to form a SEI layer on the surface of the electrodes.15 Later, the charge-discharge capacities tend to stabilize. Apparently, all TiO2 nanotube films deliver significantly higher areal capacities (330–1300 μA h cm⁻²) than the dense pure MoO_3 film ($\sim 100 \, \mu A \, h \, cm^{-2}$), indicating a porous structure truly superior to a dense one for electrode materials. In addition, areal capacities of both MoO₃-coated TiO₂ nanotubes are larger than that of the bare one. For example, CMO-TN exhibits an initial charge capacity of 1340 μA h cm⁻² and a final charge capacity of 1136 μ A h cm⁻² over 10 cycles, more than three times that of B-TN (342 μ A h cm⁻² and 319 μ A h cm⁻²). Additionally,

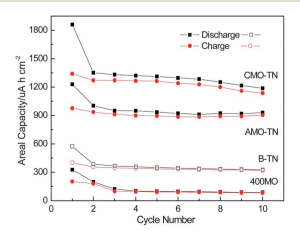


Fig. 6 Areal discharge (square symbol) and charge (round symbol) capacities of B-TN, 400MO, AMO-TN and CMO-TN over 10 cycles measured at a current of $800~\mu\text{A}$ cm⁻².

AMO-TN shows an initial charge capacity of 977 μA h cm⁻² and a final charge capacity of 905 μA h cm⁻², greater than B-TN but lower than CMO-TN. Specific capacities of these α-MoO₃ and TiO₂ anodes are obtained as well, among which CMO-TN delivers the highest and B-TN gives the lowest due to an intrinsically low capacity of anatase TiO₂ (~168 mA h g⁻¹).²² Typical specific charge capacities are 430 mA h g⁻¹ for CMO-TN, 340 mA h g^{-1} for 400MO, 290 mA h g $^{-1}$ for AMO-TN and 150 mA h g⁻¹ for B-TN (Fig. S3†). Obviously, the porous MoO₃/TiO₂ nanotube hybrid structure is surely able to provide better Li-ion insertion properties than a dense α-MoO₃ film electrode (400MO) and even a conventional α-MoO₃ powder electrode (with binder and conductive carbon black) showing capacities of \sim 125 mA h g⁻¹ with similar current and potential parameters for battery testing, mostly due to full utilization of nanosized α -MoO₃ and rapid Li-ion transfer inside a porous electrode. 13,18 Moreover, CMO-TN with both more areal and specific capacity than 400MO highlights nanoparticles superiority to submicron particles deposited onto TiO₂ nanotube arrays for the first time. That is, it has been reported that NiO or Co₃O₄ particles in submicron size deposited to anodic TiO2 nanotubes could improve their areal capacity rather than specific capacity compared to pure NiO or Co3O4 films made by same electrodeposition, since they are too large to spread effectively on the inner and outer walls of the nanotubes.11 It is also worth pointing out that CMO-TN and AMO-TN anodes deliver higher areal capacities than B-TN and better rate stability than 400MO within a wide current range from 200 to 1000 μ A cm⁻² (Fig. S4†). In brief, the remarkable improvement in Li-ion intercalation capacities of TiO2 nanotube arrays is attributed to the addition of MoO₃ with high specific capacity onto host TiO₂ material.¹⁴ It is also found that crystalline α-MoO₃ nanoparticles are capable of accommodating more Li⁺ ions than amorphous MoO₃ layers based on a comparison between AMO-TN and CMO-TN. The discrepancy is related to the crystal structure and chemical states of molybdenum oxides, capacitive factors and electrochemical kinetics. 15,18 Additionally, the TiO2 tubular structure plays an important role in this capacity boost as well. It allows uniform MoO3 deposition onto nanotubes and rapid mass transport within the electrode, favourable for α-MoO₃ coatings

Fig. 7a shows galvanostatic discharge–charge plots of B-TN, AMO-TN and CMO-TN on the first two cycles at a current of 800 μ A cm⁻². It is observed that the charge–discharge capacity of the two deposited samples is fairly higher than that of bare one. Two voltage plateaus appear near 1.7 V and 2.1 V in the curves of the three anodes. The one at 1.7 V in the negative scan corresponds to Li-ion insertion into TiO₂, while the other in the positive scan accords with Li-ion desertion from TiO₂. The total reaction for Li-ion insertion to/extraction from TiO₂ is described by:²²

to exert its Li-ion insertion/desertion nature at high rates.

$$TiO_2 + xLi^+ + xe^- \leftrightarrow Li_xTiO_2$$
 (1)

For anatase TiO_2 , x could be up to 0.5 ($Li_{0.5}TiO_2$), corresponding to a capacity of 168 mA h g⁻¹.

As the cell is discharged below 1.0 V, the capacity of B-TN continues to arise slightly, as shown by a decline slope of its plot in Fig. 7a. Besides a little extra Li-ion insertion to TiO2, an important reason for the increment could be the decomposition of electrolyte components at the TiO₂ electrodes in which Li⁺ ions may participate to cause a few irreversible capacities. 23,24 In addition, more prolonged, slowly declining slopes below 0.7 V in the discharge plots are observed for both AMO-TN and CMO-TN, suggesting a large quantity of Li⁺ ions intercalated into the MoO₃-coated TiO₂ nanotubes rather than the bare ones at low potentials. Such a difference is again confirmed by their differential capacity plots derived from this figure. Fig. 7b shows a pair of oxidation (at 2.08 V) and reduction peaks (at 1.7 V) in the charge-discharge process of TiO2, suggesting a one-stage lithiation mechanism in anatase TiO₂ according to eqn (1). It is also seen that in the second cycle the reduction and oxidation peaks for B-TN are weakened, due to Li-ion loss in the previous cycle. However, the peak positions almost remain unchanged, showing good reversibility of the Li-ion insertion/desertion reaction on TiO₂ nanotubes. On the contrary, the lithiation process in MoO3-deposited samples appears greatly different. Fig. 7c displays the differential capacity curves of AMO-TN. In this figure, the peaks at 1.5 V and 2.11 V in the first cycle mainly corresponds to Li-ion insertion/desertion on TiO2. As the potential goes down to 0.05 V, the plot inclines distinctly and even produces a small peak at \sim 0.28 V, claiming more Li⁺ ions insert into AMO-TN than into B-TN. It indeed happens that Li⁺ ions are inserted into the amorphous MoO₃ layer in the negative scan, because of much larger |dQ/dV| values for AMO-TN than for B-TN. The reduction peak shifts to 1.68 V in the second cycle, thanks to foregoing the activation process and reduced electrode polarization. Differential plots of CMO-TN are present in Fig. 7d where a pair of high reduction and oxidation peaks are seen at 1.59 V and 2.07 V in the first cycle, plus a sharp peak at 0.24 V. Li-ion insertion to α-MoO₃ is believed to be involved in this event, since long discharge plateaus and corresponding reduction peaks appear at 0.42 V and 0.32 V for 400MO (Fig. S5 \dagger). Undoubtedly, |dQ/dV| values continue to rise for CMO-TN, a sign of increased storage of Li⁺ ions. As the battery testing proceeds, its reduction peak moves to 1.7 V but its oxidation peak remains unchanged, so AMO-TN displays a wider separation of reduction and oxidation peaks of TiO2 (0.43 V) than B-TN (0.38 V) and CMO-TN (0.37 V), probably due to larger electrode polarization from slower Li⁺ diffusion in the amorphous MoO₃ coating. 18 Additionally, it is worth pointing out that a broad plateau appears between 1 V and 2 V in the positive scan of AMO-TN and CMO-TN, suggesting a great number of Li⁺ ions extracted out of the MoO₃ within the potential window, which is in accord with linearly increasing slopes of their charge plots in Fig. 7a. The plateau or slope is a typical electrochemical characteristic of MoO3 according to Fig. S5.† The differential capacity curves of the three samples within this potential window are enlarged and put as insets in Fig. 7b-d. The curve starts from 20 μA h cm⁻² V⁻¹ at 1 V and gradually goes up to 240 μ A h cm⁻² V⁻¹ at 2 V for B-TN, but from 170 to 510 $\mu A\,h\,cm^{-2}\,V^{-1}$ for AMO-TN, and from 260 to 740 $\mu A\,h$ cm⁻² V⁻¹ for CMO-TN. Since TiO₂ nanotubes serve as the

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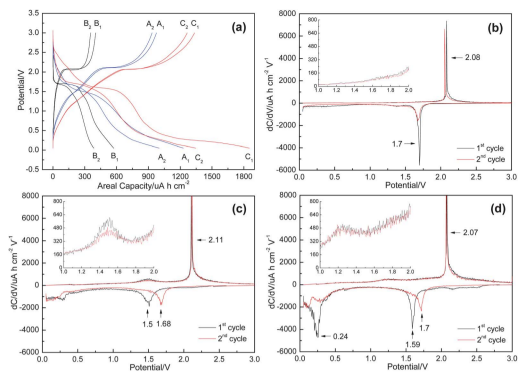


Fig. 7 (a) Charge-discharge profiles at the 1^{st} and 2^{nd} cycle of B-TN (B₁, B₂), AMO-TN (A₁, A₂) and CMO-TN (C₁, C₂), and their according differential capacity curves of (b) B-TN, (c) AMO-TN and (d) CMO-TN measured at a current of 800 μ A cm⁻². Insets: enlarged differential charge capacity curves within a potential range of 1.0 to 2.0 V

substrate for both amorphous and crystalline MoO₃ coatings, it confirms that α-MoO₃ is able to accommodate more Li⁺ ions than amorphous MoO₃, which is in good agreement with previous literature. 15,18

A two-stage lithiation process has been proposed for MoO₃.²⁵ As the potential is above 1.25 V, the reaction of Li-ion insertion into MoO₃ on the TiO₂ anode can be represented by:²⁵

$$MoO_3 + xLi^+ + xe^- \leftrightarrow Li_xMoO_3$$
 (2)

For α -MoO₃, x could be between 1 and 1.5 in the first stage, and Mo^{VI} is reduced to Mo^V and further to Mo^{IV}. α-MoO₃ has an orthorhombic structure containing a group of regularly distorted MoO₆ octahedra. They share edges to form chains that are crosslinked through oxygen atoms to yield layers. Li⁺ ions are inserted into the interlayer spacing between these octahedral layers and intralayers.26,27 The second stage occurs at potentials below 0.7 V here, and the according reaction can be expressed by:25

$$\text{Li}_x \text{MoO}_3 + y \text{Li}^+ + y \text{e}^- \leftrightarrow \text{Mo} + 3 \text{Li}_2 \text{O}$$
 (3)

It can be seen from eqn (3) that in lower voltage regions Li⁺ ions react with the solid solution (LixMoO3) to form Mo metal and Li₂O oxides. Li₂O is believed to be naturally irreversible, but the presence of nanosized Mo particles induces a reversible reaction of Li₂O during charging, 13,25 and thus a huge loss of active material is prohibited.

It is understood from Fig. 7c and d that the two stages could occur in both amorphous and crystalline MoO3 layers on TiO2 nanotubes, despite different capacities of AMO-TN and

CMO-TN. Several reasons are proposed to explain such discrepancies: first of all, the presence of molybdenum species in lower oxidation states in AMO-TN decreases chemical reactions of metal ion and Li⁺ ions according to eqn (2) and (3). Secondly, absorbed moisture and oxygen deficiencies in amorphous MoO3 cause a huge Li-ion loss when AMO-TN is subject to deep discharge conditions.15 Thirdly, distinct capacitive capabilities of MoO₃ are involved in the Li-ion storage behaviors of AMO-TN and CMO-TN electrodes. Both crystalline and amorphous MoO3 deliver redox pseudocapacitance but the former has a unique layered crystal structure in which extra Li+ ions can enter van der Waals gaps to create intercalation pseudo-capacitance.28,29 Hence, more capacitive contributions from α-MoO₃ help CMO-TN to deliver larger Li-ion storage capacities than AMO-TN at high rates. More importantly, the kinetics of Li⁺ ions at these oxide electrodes also play an important role in their electrochemical performance, which is studied by the electrochemical impedance spectroscopy (EIS) technique.

Fig. 8 gives Nyquist plots and their equivalent circuit for B-TN, 400MO, AMO-TN and CMO-TN after cells are discharged and charged for ten battery cycles. In this circuit, R_s is the ohmic resistance from the electrolyte, polymer separator and electrodes, corresponding to the intercept of the depressed semicircle in the high-to-medium frequency region with the Z-real axis. The semicircle is related to the Li-ion transfer process from the electrolyte into the TiO2 anode. Rct is the charge transfer resistance at the electrode/electrolyte interface, which could be inferred from the semicircle arc. The sloping line at the low Paper

Fig. 8 Nyquist plots and the equivalent circuit of half cells containing B-TN, 400MO, AMO-TN and CMO-TN anodes measured at a potential of 1.7 V after ten electrochemical cycles.

frequency range represents Li-ion diffusion in the bulk electrode, known as the Warburg impedance (W). From these plots, it is found that the semicircle diameter for 400MO is much longer than that for others, implying larger charge transfer resistance (R_{ct}) in the dense oxide film. B-TN gives the smallest $R_{\rm ct}$, and $R_{\rm ct}$ is clearly higher for CMO-TN than for AMO-TN with amorphous coatings. EIS analyses reveal that Li-ion transfer at the TiO₂ anode is hindered by MoO₃ oxide coatings partly because of reduced surface area of deposited nanotubes. Fortunately, MoO3 with the nature of good specific capacity accepts more Li⁺ ions to obtain a capacity increment in AMO-TN and CMO-TN compared to B-TN. Similarly, it is discovered that ionic transfer resistance is greater in CMO-TN than in AMO-TN, which is supposed to be related to decreased surface area or porosity of α-MoO₃ particle coated TiO₂ nanotubes after crystallization of as-deposited MoO₃ layers, as shown in Fig. 1b and c. However, α-MoO₃ with higher capacities than amorphous MoO₃ still succeeds in enhancing the Li-ion insertion properties of the TiO2 nanotube anode.

Conclusions

In summary, anatase TiO2 nanotube arrays have been synthesized by anodic oxidation of Ti in fluoride-containing EG electrolyte and annealing at 450 °C in air, for effective surface coating with MoO₃ via electrochemical deposition, in order to greatly enhance the Li-ion intercalation properties of TiO2 anodes. The uniform layer of amorphous MoO₃ on nanotubes is demonstrated by XRD, SEM and TEM. The as-deposited MoO₃ coating is converted to the crystalline (α) phase upon thermal treatment. All TiO₂ nanotubes deliver larger areal capacities than pure MoO₃ film, partly due to high surface area and fast Liion kinetics in porous electrodes. In addition, Li-ion storage capacities of TiO2 nanotubes are increased by the as-formed amorphous MoO₃ coatings, and further enlarged with a crystalline coating. The improved electrochemical property is due to the combination of MoO₃ with high specific capacity and ordered TiO2 nanotubes with large surface area for efficient deposition and fast Li-ion kinetics. Despite larger Li-ion transfer resistance, α -MoO₃ coated TiO₂ nanotubes possess the advantages of crystalline α -MoO₃ including more capacitive contributions and tighter chemical bonds with the TiO₂ substrate to achieve the best Li-ion storage performance. Our results demonstrate that crystalline TiO₂ nanotube arrays composited with transition oxides having more Li-ion storage capabilities could be a promising electrode material for high-performance LIBs.

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