Facile Synthesis of Reduced Graphene Oxide In-situ Wrapped MnTiO$_3$ Nanoparticles for Excellent Lithium Storage

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Abstract: The stable high-capacity anode has been an urgent demand for high energy density lithium-ion batteries (LIBs). Herein, a simple and effective strategy to synthesize high-performance reduced graphene oxide (rGO) in-situ wrapped MnTiO$_3$ nanoparticles (MnTiO$_3$@rGO) by Sol-Gel method is designed. The MnTiO$_3$ nanoparticles are uniformly dispersed and wrapped by few-layer graphene. Due to high conductivity of rGO, MnTiO$_3$@rGO nanoparticles show excellent rate performance, with a specific capacity of 286 mAh·g$^{-1}$ being displayed at the higher rate of 5.0 A·g$^{-1}$. Moreover, benefited from porous structure and flexible rGO shell, the MnTiO$_3$@rGO anode delivers a remarkable long-term cycling stability. The specific capacity maintains 441 mAh·g$^{-1}$ after 500 cycles at 0.5 A·g$^{-1}$, only losing 8.4%. Therefore, the results demonstrate that the facile synthetic strategy is highly desirable for improving the conductivity and stability of metal oxide anodes.

Key words: in-situ wrapped; reduced oxide graphene; MnTiO$_3$ nanoparticles; lithium-ion batteries anode

The transition metal oxides (MnO$^{[11]}$, Fe$_2$O$_3$$^{[2-3]}$, CoO$^{[4]}$, NiO$^{[5]}$, etc.) have been considered as promising anode materials for lithium-ion batteries (LIBs) due to their high theoretical capacity (~800 mAh·g$^{-1}$), wide availability, and low cost$^{[6-8]}$. However, the terrible cycle stability and poor rate capability of conversion-type anode is still impeding practical applications$^{[9]}$. Above mentioned problems are mainly due to the conversion-type transition metal oxides anode undergoing a large volume expansion with lithium ions insertion/extraction, rapid capacity degradation$^{[10]}$. The stable electrochemical performance of the electrode material is mainly related to the structure design and conductivity of the material. To obtain the excellent electrode material, many strategies have been applied to synthesize electrode materials with effectively reserved expansion space and improved electric conductivity, including the design of the unique hierarchical nanostructures materials (such as hollow$^{[11]}$, core-shell$^{[12]}$, yolk-shell structure$^{[13]}$ and various morphology, etc.) and composite materials$^{[14-16]}$. In terms of composite materials, carbonaceous materials, such as activated carbon$^{[17-18]}$, carbon nanotubes$^{[19]}$and graphene$^{[20]}$, etc., have been widely developed as coating shell or conductive additives. The carbon materials could buffer the volume change and enhance the electric transmission of electrode materials during the charging/discharging process$^{[21]}$.

Unfortunately, although the scale synthesis electrode of “A+B” mode electrode could improve the electrochemical performance, the effects were far from satisfactory$^{[22]}$. The weak interfacial adhesion between carbon shell and transition metal oxides electrode could not effectively protect the integrity of the material structure, leading to the relative low capacity$^{[23-26]}$. In addition, the in-situ growing graphene shell on the transition metal oxides surface is an interesting designing idea, including chemical vapor deposition (CVD) or plasma methods. The growing conductivity graphene has an excellent electric transmission and flexible mechanical nature, accommodating the volume expansion and providing excel-
lent conductivity layer during charging/discharging\cite{27}. However, the in-situ growing graphene is expensive and difficult to control, meanwhile this method requires the catalyst in electrode materials, such as Fe, Ni, and Cu, etc\cite{28}. So, it is difficult to have a low-cost and facile process for graphene coating transition metal oxides electrodes.

According to the previous reported works\cite{20,27}, the combination of TiO$_2$ and transition metal oxide can hinder the aggregation of metal particles and accommodate the volumetric expansion of the electrode material, improving the electrochemical performance. Thus, the MnTiO$_3$ becomes a promising anode material for LIBs by improving the electrochemical performances with the introduction of TiO$_2$. Although the poor conductivity of MnTiO$_3$ is still a hindrance to maintain excellent performance, carbon coating shell is an effective strategy to improve the conductivity and enhance the stability of semiconductor MnTiO$_3$ with cycling. Herein, a facile strategy to synthesize the reduced graphene oxide in-situ wrapped MnTiO$_3$ nanoparticles (MnTiO$_3$@rGO) by the Sol–Gel method was designed. The perfectly encapsulated few-layer rGO shell provides fast pathway for electron transportation, improving the lithium dynamics. The few-layer rGO with excellent mechanical nature refrains the aggregation of MnTiO$_3$ nanoparticles and suppresses fracture of particles and electrode stress during lithiation/delithiation\cite{29}. In addition, the solid electrolyte interface (SEI) film could grow uniformly on the graphene surface, preventing the side reactions occurrence of electrode. All the features induced by in-situ wrapped rGO resulted in excellent and stable electrochemical performances for MnTiO$_3$@rGO\cite{30}.

1 Experimental

1.1 Preparation of MnTiO$_3$ nanoparticles

The tetrabutyl titanate (TBT) and manganese titanate were purchased from Alladin, and ethylene glycol (EG) was purchased from Sinopharm Chemical Reagent Co., Ltd. The typical synthetic procedure for MnTiO$_3$@rGO nanoparticles composites is described as follows. 150 mL EG and 120 mg GO were added into a three-mouth flask with ultrasonic stirring, and nitrogen was used as inert gas. 3.4 mL TBT and 2.45 g manganese acetate were dissolved under vigorous stirring to form a red solution, and kept stirring overnight. Ultimately, the resultant precipitate was collected by centrifugation. The precipitate was annealed at 1073 K for 2 h in Ar atmosphere, obtaining a black powder of mesoporous MnTiO$_3$@rGO nanoparticles. The MnTiO$_3$ nanoparticles were obtained following the same procedure without GO.

1.2 Characterization of Samples

Samples measurements: the crystalline structure and morphology of the samples were characterized by X-ray diffraction measurement (XRD, Bruker D8 ADVANCE, CuKa radiation), and transmission electron microscopy (TEM, JEM 2100F), respectively. Nitrogen adsorption/desorption isotherms were measured using a Micrometitics Tristar 3000 system. The pore-size distributions were calculated from desorption branches of isotherms by the Barrett–Joyner–Halenda (BJH) method.

1.3 Electrochemical testing

The electrochemical performances of the as-obtained samples were evaluated by a 2016-type coin cell. The working electrode was composed of 80wt% of active material, 10wt% of conductivity agent (acetylene black), and 10wt% of binder (PVDF, Aldrich) solvating onto the Cu foil and then dried at 343 K overnight. The cells were assembled in an argon-filled glove box (moisture and oxygen contents below 0.1x10$^{-6}$) by using the electrode as the working electrode, lithium metal as the counter electrode, Celgard 2600 as separator, and 1 mol/L LiPF$_6$ (dissolved in ethylene carbonate and dimethyl carbonate with a 1 : 1 volume ratio) as the electrolyte. The galvanostatic discharge/charge tests were performed in the CT2001A battery testing system (Land®, China) at 298 K with a voltage window of 0.01 V–3.0 V. The cyclic voltammetry (CV) and EIS tests were conducted on a CHI electrochemical workstation.

2 Results and discussion

The MnTiO$_3$@rGO nanoparticles were synthesized by reacting tetrabutyl titanate with manganese acetate in the graphene oxide (GO) solution, followed by calcination in Ar atmosphere. From the SEM image (Fig. 1(a)), the surface of mesoporous MnTiO$_3$@rGO nanoparticles are closely bound together with graphene shell. The optical photograph of MnTiO$_3$@rGO powder is exhibited, emphasizing the capacity of scale preparation. A uniform size of ~15 nm of mesoporous MnTiO$_3$@rGO nanoparticles confirmed by the TEM images (Fig. 1(b)). As shown in the high-resolution TEM (HRTEM) image (Fig. 1(c)), the lattice spacing of the MnTiO$_3$ nanoparticle is measured to be ~0.28 nm, which agrees well with the d-spacing of the pyrophanite MnTiO$_3$ (104) plane. The MnTiO$_3$@rGO nanoparticle with a perfectly wrapped graphene shell around the highly crystalline core structure can be clearly observed. The in-situ wrapped rGO shell reveals a few-layer structure of about 2–4 layers graphene. As shown in the Fig. 1(d), the selected-area electron diffraction (SAED) pattern of MnTiO$_3$@rGO displayed the ring-like mode, indicating polycrystalline
Fig. 1  (a) SEM image with inset showing the optical photograph of MnTiO@rGO powder, (b) TEM image, (c) HRTEM image, and (d) SAED pattern, (e) high-angle annular dark field (HAADF) of MnTiO@rGO, and EDS mapping of C, Mn, Ti, and O elements which indicates uniformly elemental distribution in the MnTiO@rGO structure. The ring-like pattern can be clearly assigned to the diffractions of the (012), (104), (024), and (116) planes, respectively, which is consistent with the aforementioned XRD results. The high-angle annular dark field (HAADF) of MnTiO3@rGO was investigated by scanning transmission electron microscopy (STEM) (Fig. 1(e)). The energy dispersed spectrum (EDS) of MnTiO3@rGO shows the existence of C, Mn, Ti, and O elements and respective contents (Table 1). Corresponding elemental mapping results indicate the homogeneous distribution of C, Mn, Ti, and O elements.

To further investigate the intrinsic structure of as-obtained samples, the X-ray diffraction (XRD) pattern of the samples are presented in Fig. 2(a). The characteristic diffraction peaks within the 2θ range from 20° to 80° could be well indexed to the (012), (104), (110), (113), (024), (110), (018), (124), and (300) planes for pyrophanite MnTiO3 (JCPDS No.77-1858). The (002) diffraction peak of graphene exists at 26.5° indicating the GO fully reduced to rGO via the high temperature procedure. In addition, the characteristic peaks of other impurities such as MnO, and TiO2 do not appear, indicating the pure MnTiO3 phase.

The Raman spectra (Fig. 2(b)) demonstrated the existence of rGO and MnTiO3 in MnTiO3@rGO composites. Compared with MnTiO3, the stronger Raman peaks locating at 1340 cm⁻¹ and 1591 cm⁻¹ for MnTiO3@rGO are typical D and G bands, respectively, inducing the existence of rGO, which is consistent with the HRTEM result. The existence of D and G bands for MnTiO3 could be ascribed to glycol carbonized in the precursors during annealing. The presence of typical Raman-active modes of MnTiO3 is located at 153 cm⁻¹, 202 cm⁻¹, 232 cm⁻¹, 284 cm⁻¹, 338 cm⁻¹, 389 cm⁻¹, 458 cm⁻¹, 608 cm⁻¹, 705 cm⁻¹, and 760 cm⁻¹. The strong Raman absorption peak of MnTiO3 nanoparticles locates at 682 cm⁻¹, which is due to the high-frequency vibration mode of MnO6 octahedron. The phonon mode absorption peaks below 300 cm⁻¹ are caused by the lattice vibration. These results coincide with XRD pattern and TEM observation.

Nitrogen sorption isotherms of the mesoporous MnTiO3@rGO showed a type-IV curve (Fig. 2(c)). The specific surface area is calculated to be 91 m²·g⁻¹, which is higher than that of MnTiO3 (12 m²·g⁻¹). The corresponding pore size distribution of MnTiO3@rGO are mainly within 3 nm due to the accumulation of nanoparticles. The porous structure could reserve more space to accommodate the volume expansion of the electrode and improve lithium ions transportation during charging/discharging. In addition, the conductivity of MnTiO3 and MnTiO3@rGO is investigated by the two electrode methods, the results indicate that the electrical conductivity of the MnTiO3@rGO sample reaches 2.37 mS·cm⁻¹, which is higher than that of MnTiO3 due to the conductive rGO coating shell (Fig. 2(d)). These are beneficial for excellent electrochemical performance of MnTiO3@rGO anode.

| Materials   | C/wt% | O/wt% | Ti/wt% | Mn/wt% |
|-------------|-------|-------|--------|--------|
| MnTiO3@rGO  | 9.01  | 18.87 | 32.11  | 40.01  |
To evaluate the electrochemical performance of MnTiO₃@rGO electrode, cyclic voltammetry (CV) and galvanostatic cycling were carried out by assembling half-cells with lithium metal foils as counter and reference electrodes. All of the electrode mass consist of MnTiO₃ and rGO in the MnTiO₃@rGO electrode. The type of electrochemical reactions are probed by initial three CV with a voltage window of 0.01–3.0 V. The CV curves at a sweep rate of 0.2 mV·s⁻¹ was shown in Fig. 3(a).

In the first discharge process, the cathodic peak located around 0.72 V correlates with the formation of solid electrolyte interface (SEI) film and the others at 1.71 V and 0.38 V, indicating the lithium ions insertion into TiO₂ and the reduction of MnO to metal Mn [1, 31]. In the subsequent charge process, two broad anodic peaks occurring around 1.25 V and 1.75 V are correspond to the oxidation of Mn to Mn²⁺ and lithium ions extraction from TiO₂, respectively [32]. From the second cycle, there is only a broadened reduction peak in the range of 0.3 V–0.5 V due to the overlap and broadening of peaks [33-34].

Besides, the CV curves of the second and third cycles tend to overlap, demonstrating the excellent cycling performance. In Fig. 3(b), the rate capability of the MnTiO₃@rGO was investigated by increasing the current densities from 0.1 A·g⁻¹ to 5.0 A·g⁻¹. Compared with MnTiO₃, the rate performance of MnTiO₃@rGO electrode significantly outperformed that of MnTiO₃ anode. The corresponding charge/discharge curves are shown in Fig. 4(a). The MnTiO₃@rGO electrode performed excellently under high current densities of 2.0 A·g⁻¹ and 5.0 A·g⁻¹, with the specific capacity recorded as 342 mAh·g⁻¹ and 286 mAh·g⁻¹, respectively.

To demonstrate excellent cycle stability, long cycle performance of as-obtained materials was compared [35]. Fig. 3(c) reveals the cycle stability and coulombic efficiency of MnTiO₃@rGO and MnTiO₃ electrode. Serious capacity attenuation with cycling occurs on the MnTiO₃ electrode. The poor conductivity of MnTiO₃ causes a considerable lithium kinetic obstacle for lithium ions extraction and particles are pulverized, which induces irreversible capacity. In addition, the rGO electrode shows the excellent cycling stability. This greatly improves the cycling stability of MnTiO₃@rGO. During the cycle testing, the coulombic efficiency of MnTiO₃@rGO is above 98.8%, and the specific capacity is maintained 441 mAh·g⁻¹ after 500 cycles, losing 8.4% referring to the initial capacity at a current density of 0.5 A·g⁻¹.

The excellent electrochemical performance mainly stems from the synergistic effect between the MnTiO₃ nanostructures and in-situ wrapped few-layer rGO shell. The porous MnTiO₃@rGO has a greater specific surface area than that of MnTiO₃, decreasing the transport
distance of lithium ions of electrochemical reactions. In addition, the rGO shell not only inhibits the aggregation of the MnTiO$_3$ nanoparticles, enhancing the lithium kinetics, but also helps the forming of a stable SEI film, avoiding the electrolyte decomposition. As shown in Fig. 4(b), the galvanostatic charge/discharge profiles of the MnTiO$_3$@rGO electrode with cycle numbers of 10th, 100th, 200th, 300th, 400th, and 500th were demonstrated. The low discharge platform is consistent with the CV results. Fig. 3(d) compares the Nyquist plots of electrodes of MnTiO$_3$@rGO and MnTiO$_3$ after three cycling. The semicircle at high-medium frequency and the straight line at low frequency are displayed, corresponding to charge transfer and diffusion, respectively. Apparently, the MnTiO$_3$@rGO electrode shows a much lower resistance than the MnTiO$_3$ electrode (24 vs. 46 $\Omega$). These largely ascribe to the highly conductive rGO shell, facilitating electron transfer from MnTiO$_3$ within the
whole electrode and thus decrease resistance.

3 Conclusion

In summary, a general and scale strategy was designed for in-situ wrapped of conductive few-layer rGO shell on MnTiO$_3$ materials via a facile Sol-Gel method. After the high temperature annealing, MnTiO$_3$ nanoparticles are tightly encapsulated with few-layers rGO. Benefiting from the conductive rGO and MnTiO$_3$ nanoparticles, the MnTiO$_3$@rGO electrode revealed the excellent rate performance that the specific capacity of 342 mAh$\cdot$g$^{-1}$ and 286 mAh$\cdot$g$^{-1}$ were performed under high current densities of 2.0 A$\cdot$g$^{-1}$, and 5.0 A$\cdot$g$^{-1}$. In addition, the porous structure and flexible rGO could accommodate the large volume change, maintaining the electrode stability during lithium/delithiation. The resultant MnTiO$_3$@rGO nanocomposite exhibited a reversible capacity of 484 mAh$\cdot$g$^{-1}$ at 0.5 A$\cdot$g$^{-1}$ after 500 cycles (only losing 8.4%). Interestingly, the facile and effective preparation method could be extended to synthesize high stability nano-structure metal oxide anode for LIBs.

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还原氧化石墨烯原位包覆纳米MnTiO₃颗粒的简易合成及储锂性能研究

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摘 要：对于高能量密度的锂离子电池而言，研究稳定、高容量负极材料的需求十分迫切。基于此，本工作设计了一种简单有效的溶胶–凝胶法，来合成高性能的被还原氧化石墨烯氧化物原位包覆的MnTiO₃纳米颗粒(MnTiO₃@rGO)。合成的MnTiO₃纳米粒子分散均匀，被少层的石墨烯包裹。由于还原氧化石墨烯的高电导率，MnTiO₃@rGO 作为锂离子电池负极表现出优异的倍率性能，MnTiO₃@rGO 在5.0 A·g⁻¹的高电流密度时，比容量为286 mAh·g⁻¹。此外，得益于MnTiO₃@rGO 的多孔结构和柔性的还原氧化石墨烯外层，MnTiO₃@rGO 负极具有显著的长期循环稳定性。在500 个循环后，比容量仍保持在441 mAh·g⁻¹，仅损失了8.4%。结果表明，该方法对提高金属氧化物负极的导电性和循环稳定性具有较高的应用价值。

关键词：原位包覆；还原氧化石墨烯；MnTiO₃纳米颗粒；锂离子电池负极

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