RSC Advances



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PAPER

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Cite this: RSC Adv., 2017, 7, 25422

Received 25th March 2017 Accepted 6th May 2017

DOI: 10.1039/c7ra03483k

rsc.li/rsc-advances

Introduction

Since 1997, rechargeable lithium-ion batteries (LIBs) have been considered as one of the most attractive power sources for a wide range of applications, such as cellular phones, notebook computers, electric vehicles (EV) and hybrid electric vehicles (HEV), *etc.*, due to their high specific capacities, long cycle lives, high power densities and environmental friendliness.¹ At present, it is crucial to seek the state-of-the-art cathode materials, since their great importance in determining the whole performances of batteries.² Amongst the prospective cathode materials for large-scale next-generation LIBs applications, lithium transition-metal phosphates have aroused widespread concerns in recent decade, *e.g.* olivine LiFePO₄, because of its thermodynamically stable structure,³ high specific capacity (the

Large-scale synthesis of $Li_3V_2(PO_4)_3@C$ composites by a modified carbothermal reduction method as cathode material for lithium-ion batteries

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Carbon coated Li₃V₂(PO₄)₃ composites were prepared by a modified carbothermal reduction method. The method has the advantages of being simple and scalable, while the whole synthesis is devoid of any reducing/protecting gases, therefore, it can be directly carried out in muffle furnace. The obtained Li₃V₂(PO₄)₃@C composites have particles sizes from 500 nm to 3 μ m, and with homogeneous carbon coating layer thickness of about 7 nm. The battery based on this Li₃V₂(PO₄)₃@C composites cathode exhibits a specific discharge capacity of 130.9 mA h g⁻¹ at 0.1 C, and exhibits an initial specific discharge capacity of 102.8 mA h g⁻¹ even at 10 C and almost keeps 96.6% of initial capacity retention after 500 cycles. The performance enhancement can be directly ascribed to the robust structure merits of the Li₃V₂(PO₄)₃@C composites synthesized by the modified carbothermal reduction method. Furthermore, the as-developed method has the potential to be expanded to other lithium transition-metal phosphates (such as lithium iron phosphate), which conventionally need to be sintered in reductive or protective atmospheres.

theoretical capacity is 170 mA h g^{-1}) and abundance in nature. However, its low redox potential (only 3.4 V vs. Li/Li^+), as well as low gravimetric energy density, has become the obstacles for the practical application of LiFePO₄ cathode, although the great progress has been made. Therefore, tremendous efforts have been devoted to developing other lithium transition-metal phosphates with high operating voltages such as LiMnPO₄ (~4.1 V vs. Li/Li⁺),⁴ LiCoPO₄(~4.8 V vs. Li/Li⁺),⁵ LiNiPO₄(~5.2 V vs. Li/Li⁺),⁶ and Li₃V₂(PO₄)₃(~4.8 V vs. Li/Li⁺).⁷ Among them, monoclinic phase $Li_3V_2(PO_4)_3$ with the highest theoretical specific capacity (197 mA h g⁻¹ for complete extraction/ insertion of 3 Li⁺ in redox reactions at the voltage window of 3.0-4.8 V) has been the most widely studied system in recent years due to its high-power density, high operating voltage (4.8 V)8,9 and natural abundance,10,11 these advantages make it become highly prospective¹²⁻¹⁶ for high-energy LIBs.

However, the applications of $\text{Li}_3\text{V}_2(\text{PO}_4)_3$ cathode are still far, because of its low electronic conductivity, which will significantly limit electrochemical properties of the electrode. Tremendous efforts have been devoted to overcoming the deficiency, such as reduction of particle size and coating conductive carbon.^{17,18} The pathways for Li⁺ to transport across the particles can be enlarged and the distance can be shorted, moreover, the electronic contacts between electrode and electrolyte can be increased through decreasing the size.¹⁹ Nevertheless, it has negative effects on the morphology and structure of Li₃V₂(PO₄)₃ particles, and the crystallinity of Li₃V₂(PO₄)₃,²⁰ so it has not achieved a satisfactory level to be fully utilized.

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Another feasible technique for enhancing electronic conductivity of $Li_3V_2(PO_4)_3$ cathode is coating conductive carbon. It has been extensively researched for its economical and practical features (such as high electronic conductivity, excellent chemistry/electrochemistry stability, low cost, etc.).21 It can modify the surface chemistry and form more efficient electron pathways,^{4,22,23} so the active materials can be largely utilized at high current rates,²⁴ and it can also alleviate the growing up and aggregation of Li₃V₂(PO₄)₃ particles during the high temperature calcination. Additionally, carbon can act as a reducing agent to reduce V5+ to V3+ during the reaction process.25 Meanwhile, various methods are used to synthesize Li₃V₂-(PO₄)₃@C composites, such as sol-gel method,²⁶ hydrothermal synthesis,27 freeze-drying method28 and rheological phase reaction synthesis,²⁹ etc. they all have great potential for manufacturing, but there are also some drawbacks of restricting their mass productions, for instance, sol-gel and hydrothermal method often demand low reaction temperatures, which will result in low crystallinity of materials, and the rheological phase reaction synthesis demand complicated operations and high cost. It is necessary to further optimize and modify these methods.

On the other hand, Li₃V₂(PO₄)₃@C composites can be commonly synthesized by conventional high temperature solid state reaction.^{30,31} Carbothermal reduction method,³² one of the conventional high temperature solid state reactions, has become one of the best choices for industrial production since the advantages of high yield and simple process. Nevertheless, large consumption of energy and enormous waste of protective gases or reductive gases (including H₂, it will bring the risk of explosion) are the drawbacks of this synthetic method. In this work, a modified carbothermal reduction method is employed to prepare $Li_3V_2(PO_4)_3$ (a)C composites, they can be large-scale synthesized without using reduction gas, and reducing the risk of hydrogen explosion in muffle furnace. Meanwhile, it has the merits of ease of using, energy conservation and low consumption. The chief merit among them is that it does not sacrifice its stable structure and excellent electrochemical performances in mass production. In addition, it has the potential to expand to other cathode of lithium transition-metal phosphates, which are also need to be sintered in reduction or protective atmosphere, such as lithium iron phosphate.

Experimental details

Synthesis

Li₃V₂(PO₄)₃@C composites were synthesized *via* a two-step carbothermal reduction approach including the pre-sintering and the subsequent heat treatment. Firstly, according to the molar ratio of Li₃V₂(PO₄)₃ ($N_{Li}: N_V: N_P = 3:2:3$), analytic grade CH₃COOLi·2H₂O (25.3 g), NH₄VO₃ (49.0 g), NH₄H₂PO₄· ·2H₂O (33.1 g) and sucrose (C₁₂H₂₂O₁₁, 43.2 g) were added into ethanol (100 ml). Secondly, the precursors were obtained after the mixtures were ball-milled for 12 h and dried for 12 h. Thirdly, the precursor powders were put into a smaller skittle pot, and wrap it with an aluminum foil before being tossed upside down into a larger one, and pour a certain amount of reduction powders which are the mixtures of carbon powders, iron powders and silica sands with the molar ratio of 1.5:1:2 into the extra space. Finally, skittle pot was put into muffle furnace to sinter. The precursors were pre-sintered at 450 °C for 8 h to carbonize sucrose and followed by subsequently sintering at 950 °C for 12 h to crystallize. After cooling down, Li_3V_2 (PO₄)₃@C composites were obtained.

Materials characterization

The surface chemistries of $\text{Li}_3\text{V}_2(\text{PO}_4)_3$ composites were characterized by the following techniques: the amount of residual carbon was determined by TG/DTA (DIAMOND TG/DTA) in flowing air with the temperature ranging from 30 to 800 °C at a heating rate of 10 °C min⁻¹. The crystalline phase of the synthesized materials was characterized by the X-ray diffractometer (XRD, BRUKERD8-A25) with 2 θ between 10° and 60°, at the scan rate of $0.02^{\circ} \cdot \text{sec}^{-1}$, the voltage and current were 40 kV and 40 mA, respectively. Raman scattering spectroscopy was recorded on a SPM-RAMAN (Ntegra upright, NT-MDT). The morphologies of the samples were observed by using scanning electron microscope (SEM, JSM-7100F) and the morphologies of surface carbon layers and rates of elements were obtained by transmission electron microscopy (HRTEM, Tecnai G2 20 U-Twin) coupled with an Energy Dispersive Spectrometer (EDS).

Electrochemical measurements

The electrochemical characterizations were carried out in two electrode coin cells (CR2025). The cells were assembled in a dry Ar-filled glove box according to the order of cathode electrode, electrolyte, separator, and lithium plate and nickel foam. Each component of cells would be introduced as followed: the cathode electrode contained 80 wt% of active material, 10 wt% of acetylene black and 10 wt% of PVDF (polyvinylidene fluoride), to provide sufficient Li^+ . The cathode slurry was prepared by thoroughly mixing a N-methyl-2-pyrrolidine solution with the active material, acetylene black, PVDF. Subsequently, the obtained slurry was deposited on aluminum foil (thickness of 20 µm) and rolling pressed before the entire assembly was dried in a vacuum oven at 120 °C overnight. The average loading of the electrode was 3.8-5.1 mg cm⁻².²¹ The electrolyte contributing to the diffusion of Li⁺ was made up of 1 M LiPF₆ dissolved in a solvent mixture of ethylene carbonate (EC), dimethyl carbonate (DMC) and ethyl methyl carbonate (EMC) with the volume ratio of 1:1:1. A porous polypropylene film (Celgard 2400) was used to protect inner circuit by only permitting the pass of Li⁺. Pure metallic lithium foil was served as the anode electrode, which was acted as counter electron and reference electrode. Nickel foam played the role of supporting stabilization of cells. The following electrochemical measurements were conducted. Electrochemical impedance spectroscopy (EIS) and cyclic voltammetry (CV) were analyzed on an electrochemical workstation (ZAHNER CIMPS-2). The impedance data were recorded with an AC voltage of 2 mV as the vibration voltage in the frequency from 100 mHz to 1 MHz. The CV measurements were performed in the potential window of 3.0-4.3 V with a scan rate of 0.2 mV s⁻¹. Galvanostatic cycling and rate capacity

performances were gauged on a Land automatic battery tester (LAND CT2001A) in the potential window of 3.0–4.3 V.

Results and discussion

Synthesis of the Li₃V₂(PO₄)₃@C composites

The $Li_3V_2(PO_4)_3$ (a)C composites are synthesized via the modified carbothermal reduction method, combining with the ballmilling procedure and the high temperature sintering reaction, the detailed fabrication process is schematically shown in Fig. 1. The reduction powders are used to build the reducing atmosphere in the whole process, instead of using reduction gas, that can reduce the risk of explosion. The carbon powders and iron powders are used to isolate from oxygen into the smaller skittle pot, so they have the effect of preventing precursors from being oxidized, the silica sands are used to prevent iron powders and carbon powders from being sintered into blocks at high temperature which can weak the ability of isolating oxygen, the aluminum foil has the functionality to further isolate oxygen, it can ensure that reduction powders will not mix with precursor powders simultaneously. What's more, they can be utilized repeatedly. The next prominent merit is hundreds of grams Li₃V₂(PO₄)₃@C composites can be synthesized in laboratory at a time, it is hundreds of times of production synthesized via conventional carbothermal reduction method. In general, all of these make it easier to operate and have greater output than conventional carbothermal reduction method, moreover, it can be easily scaled-up in manufacturing.

Physical characterizations of Li₃V₂(PO₄)₃@C composites

Fig. 2a shows the TG curves of $\text{Li}_3V_2(\text{PO}_4)_3$ @C composites measured from RT to 800 °C under an air flow. The weight loss is mainly due to the combustion of carbon and oxidation of vanadium elements in the composites (from V³⁺ to V⁵⁺). The weight of the carbon content is estimated to be about 4.18%.³³

Fig. 2b shows the XRD pattern of $Li_3V_2(PO_4)_3$ @C composites. The sharp diffraction peaks of the sample indicate that the



Fig. 1 Schematically formation process of carbon coated $Li_3V_2(PO_4)_3$ composites.

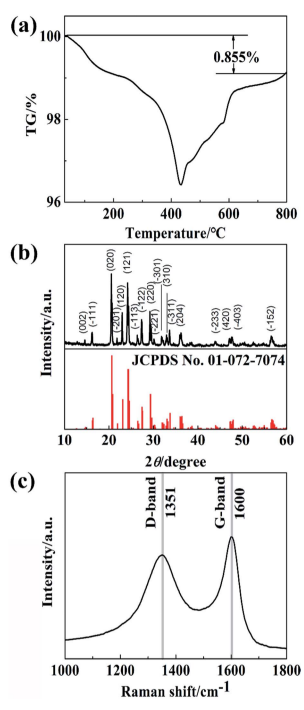


Fig. 2 (a) TG/DTA curves of the composites recorded from room temperature to 800 °C at a heating rate of 10 °C min⁻¹ under an air flow of 80 mL min⁻¹. (b) X-ray diffraction pattern of carbon coated Li₃V₂(PO₄)₃ materials. (c) Raman scattering spectrum of carbon coated Li₃V₂(PO₄)₃ materials.

composites are well crystallized, which are in good agreement with that of monoclinic $\text{Li}_3\text{V}_2(\text{PO}_4)_3$ (JCPDS card no. 01-072-7074, space group of $P2_1/n$). The Miller indexes are added in the XRD pattern. There are no impure diffraction peaks, which suggests that the residual carbon in materials are in amorphous state. The refined lattice parameters of $\text{Li}_3\text{V}_2(\text{PO}_4)_3$ @C composites are a = 8.6049(5), b = 8.6056(3), c = 11.9920(8) Å

and $\beta = 90.813(8)^{\circ}$ with an $R_{wp} = 4.73\%$ and the cell parameters are consistent with those in previous reports.¹²

Fig. 2c exhibits the Raman scattering spectrum of the carbon coated $\text{Li}_3\text{V}_2(\text{PO}_4)_3$ particles with wavenumbers from 1000 to 1800 cm⁻¹. It can be observed that the characteristic carbon signatures at 1351 and 1600 cm⁻¹ corresponding to a disorder-induced phonon band (D-band) and the graphite band (G-band) respectively. The D-band and the G-band are associated with sp³-type carbon and sp²-type carbon, respectively. The I_D/I_G ratio provides useful information about the crystallinity of the carbon coated over $\text{Li}_3\text{V}_2(\text{PO}_4)_3$ particles. As previously reported,³⁴ the improvement of the electronic conductivity of the material is corresponding to the decline of I_D/I_G is 0.96, reflects the good electronic conductivity of the obtained samples.

Fig. 3 shows the SEM and TEM images with different magnifications of $\text{Li}_3\text{V}_2(\text{PO}_4)_3$ @C composites. The sizes of these small particles are about 500 nm to 3 µm, and they are uniformly distributed as the Fig. 3a shown. Fig. 3b shows the high-magnification SEM image of the composites, it reflects that the carbon is homogeneously coated on the surface of the particles, which are helpful to enlarge the pathways and shorten the distance for Li⁺ to transport across the Li₃V₂(PO₄)₃ particles. It can be clearly seen from Fig. 3c that the Li₃V₂(PO₄)₃ particles are indeed coated with a rough amorphous carbon layer. To further understand the microstructure of the composites and check the carbon coating on the particles, the magnified HRTEM images of the Li₃V₂(PO₄)₃@C composites are presented, Fig. 3d and e show the high resolution TEM images of

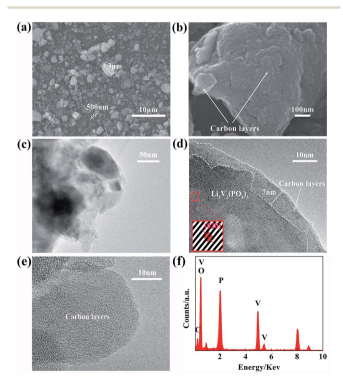


Fig. 3 SEM and TEM images of $Li_3V_2(PO_4)_3@C$ composites with different magnifications. (a and b) SEM images; (c-e) TEM and HRTEM images; (f) The EDS profile acquired from $Li_3V_2(PO_4)_3@C$ composites.

the composites, reveal that the average thickness of the carbon layer is 7 nm, the presence of carbon layers can well control the shape of composite particles and modify the surface chemistry without agglomeration.³⁵ The lattice fringes are clearly observed in the inset figure within Fig. 3d, it clearly reveals that regular lattice fringes of the (020) planes corresponding to the Li₃V₂(PO₄)₃ *d*-spacing value of 0.432 nm through the selected area electron diffraction scan. Fig. 3f shows EDS profile of Li₃-V₂(PO₄)₃@C composites. The EDS spectrum confirms the existence of C, V, O and P elements in the Li₃V₂(PO₄)₃@C composites, and further verifies a rough molar ratio of 2 : 3 for vanadium and phosphorus elements in accordance with the chemical formula. No obvious peaks corresponded to other impurity elements can be observed (the element Li cannot be identified by EDS detector).

Electrochemical performances of $\text{Li}_3\text{V}_2(\text{PO}_4)_3$ (a)C composites cathode

To understand the property of $\text{Li}_3\text{V}_2(\text{PO}_4)_3$ @C composites synthesized by the modified carbothermal reduction method more deeply, tests are conducted in more details.

Fig. 4a displays the galvanostatic charge-discharge potential profiles of the sample between 3.0 and 4.3 V. At the constant

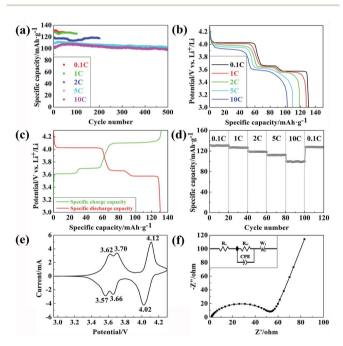


Fig. 4 (a) Nyquist plots of $Li_3V_2(PO_4)_3$ @C composites cathode measured in an AC voltage as a vibration of 2 mV in the frequency range from 100 mHz to 1 MHz and equivalent circuit of the $Li_3V_2(PO_4)_3$ @C composites cathode; (b) CV curves of $Li_3V_2(PO_4)_3$ @C composites cathode at 0.2 mV s⁻¹ in the potential range of 3.0-4.3 V; (c) cycle performance of the $Li_3V_2(PO_4)_3$ @C composites cathode at 0.1 initial discharge potential curves of the $Li_3V_2(PO_4)_3$ @C composites cathode at different rates (0.1C, 1C, 2C, 5C, 10C) in the potential range of 3.0-4.3 V; (e) initial charge-discharge potential curves of the $Li_3V_2(PO_4)_3$ @C composites cathode at 0.1C in the potential range of 3.0-4.3 V; (f) rate performance at various C-rates from 0.1C to 10C each for 20 cycles for the $Li_3V_2(PO_4)_3$ @C composites cathode.

current densities of 0.1 C, 1 C, 2 C, 5 C and 10 C (1C = 133 mA g^{-1}), the Li₃V₂(PO₄)₃@C composites cathode exhibits the initial specific discharge capacities of 130.9, 127.7, 118.6, 109.8 and 102.8 mA h g^{-1} (based on the removal of two Li⁺), corresponding to 98.4%, 96%, 89.2%, 82.6% and 77.3% of the theoretical specific capacity (133 mA h g^{-1}), respectively, which are higher than the previous reported.³⁶ When the sample is cycled at a current density of 10 C, the specific discharge capacity of the sample is about 98.5 mA h g^{-1} even after 500 cycles as Fig. 4a shown, and it occupies almost 96.6% of initial capacity. It proves that Li₃V₂(PO₄)₃@C composites synthesized by the modified carbothermal reduction method can obtain good capacities retention, even at high C-rates.

The results of Fig. 4b show that $\text{Li}_3V_2(\text{PO}_4)_3$ @C composites cathode deliver the specific discharge capacities of 130.9 mA h g⁻¹ (0.1 C), 127.7 mA h g⁻¹ (1 C), 118.6 mA h g⁻¹ (2 C), 109.8 mA h g⁻¹ (5 C) and 102.8 mA h g⁻¹ (10 C). Obviously, the discharge voltage platforms keep better when the discharge current density is lower. On the contrary, it becomes less conspicuous. It implies that charging/discharging in high current density will limit amount of the Li⁺ being participated in insertion/de-insertion action, which will reduce the capacity and reversibility of the battery. It explains well why the capacities in smaller current densities are higher than in larger current densities.

Subsequently, the initial charge–discharge potential profiles of the $\text{Li}_3\text{V}_2(\text{PO}_4)_3$ @C composites cathode at a current rate of 0.1 C in potential window of 3.0–4.3 V are presented in Fig. 4c, the three clear charge plateaus are 3.62, 3.71 and 4.11 V respectively, and three clear discharge plateaus are 3.56, 3.67 and 4.02 V respectively. In addition, the initial Coulomb efficiency is 99%, which embodies an excellent reversibility of the battery.

Fig. 4d describes the rate capability test of $Li_3V_2(PO_4)_3$ @C composites cathode at C-rates ranging from 0.1C to 10C, and back to 0.1C. It exhibits a specific discharge capacity of 130.2 mA h g⁻¹ at 0.1 C and still remains 99.8 mA h g⁻¹ at 10 C, which corresponds to 76.7% of initial specific discharge capacity. When back to 0.1 C, the specific discharge capacity is 128.1 mA h g⁻¹, and the $Li_3V_2(PO_4)_3$ @C composites cathode retains 98.4% of initial specific discharge capacity. It is explained that the structure of $Li_3V_2(PO_4)_3$ @C composites are stable and the formation of dead Li^+ are temporary, which generated at high rate performance, so the capacities and reversibility of the battery will decline at the larger current densities, and will return when back to 0.1 C.

Fig. 4e displays the CV curve of the sample at the scan rate of 0.2 mV s^{-1} in the potential window of 3.0–4.3 V. The CV curve is composed of three apparent redox couples, due to the insertion/ de-insertion of Li⁺ with only two of them being removed, which means that all of the three couples of current peaks are assigned to the V³⁺/V⁴⁺ redox couples, and it corresponds to the intercalation and extraction of Li⁺ as the stoichiometric ranges: x = 0.0-0.5, 0.5–1.0 and 1.0–2.0 in Li_{3–x}V₂(PO₄)₃@C, respectively. The three oxidation peaks of Li₃V₂(PO₄)₃@C composites cathode are located at about 3.62, 3.70 and 4.12 V respectively, and the three reduction peaks are located at 3.57, 3.66 and 4.02 V respectively, which are in agreement with the three

couples of charge/discharge plateaus in Fig. 4c. The extraction and intercalation potentials are similar to previous report.³⁷ The well-defined sharper oxidation/reduction current peaks of the sample are ascribed to the two-phase reaction mechanism, and the broad peak potentials corresponding to the solid-solution behaviors and small potential intervals show low electrochemical polarization reflect excellent reversibility of $\text{Li}_3\text{V}_2(-\text{PO}_4)_3$ @C composites cathode in the charge/discharge processes.³⁸

EIS measurements are performed on the Li₃V₂(PO₄)₃@C composites cathode. Firstly, the coins are galvanostatic charged/discharged for at least 5 cycles to ensure that the SEI films on the surface of the active particles are fully formed, and the coins are then measured with an AC vibration voltage of 2 mV in the frequency range from 100 mHz to 1 MHz. Nyquist plots of the Li₃V₂(PO₄)₃@C composites cathode and an equivalent circuit are shown in Fig. 4f. The EIS data are fitted by using Zview2 software. As seen in the figure, an intercept at high frequency on the Z' axis represents R_e of battery, which contains the resistance of electrolyte, separator and electrode. A depressed semicircle in the high-middle frequency region is attributed to the charge transfer resistance, which is associated with the Li^+ diffusion in the $\text{Li}_3\text{V}_2(\text{PO}_4)_3$ @C particles, and the straight line in the low frequency region stands for the diffusions controlled by Warburg. R_{ct} and W₀ are used to denote them in the equivalent circuit. A constant phase element CPE is placed to represent the double layer capacitance and passivation film capacitance,^{39,40} and the capacitance resistance is so small that it is negligible. R_{ct} of the Li₃V₂(PO₄)₃@C composites cathode is 55 ohm, smaller than previous report,41 which is in favor of rapid electrochemical reaction, reflects the excellent electrochemical reaction of the composites cathode. It is explained that the Li_3V_2 (PO₄)₃@C composites synthesized by the modified carbothermal reduction method indeed has better structure and good electronic conductivity, which contributes to the diffusion of these Li⁺ in the Li₃V₂(PO₄)₃@C particles.

Among repeated observations, $Li_3V_2(PO_4)_3$ cathode materials with conductive coating in this literature as Fig. 1 shown, possess both outstanding cycling stability and rate capability. The quality of the Li₃V₂(PO₄)₃@C composites synthesized by the modified carbothermal reduction method are evidently remarkable. The exceptional performance can be ascribed to the following structural factors. Firstly, the high quality of crystallization, good dispersion and small size of the $Li_3V_2(PO_4)_3$ phase, so it allows the full access of its maximum capacity. Secondly, the carbon coating layer functions as a shell which accommodates the Li₃V₂(PO₄)₃ particles just like eggs packed in the shell. With such affection of coated layer, the stress generated due to volume change of Li₃V₂(PO₄)₃ caused by the Li⁺ de-insertion/insertion during the charging/discharging processes will become much localized, thus the phase change and/or decomposition of the particles will be thoroughly eliminated, and the stability of the composites structure will be highly improved upon repeated charging/discharging, the morphology and size of these particles are also greatly reserved even after the deep cycling. Thirdly, the conductive and continuous carbon matrix of the surface carbon coating can enhance the efficient electronic connection between $Li_3V_2(PO_4)_3$ particles. In general, the superior electrochemical performances of $Li_3V_2(PO_4)_3$ composites cathode in this literature is benefit from the core-shell structure of $Li_3V_2(PO_4)_3$ composites synthesized by the modified carbothermal reduction method.

Conclusions

In summary, $Li_3V_2(PO_4)_3$ (a)C composites can be successfully synthesized by the modified carbothermal reduction method. This approach is facile and cost-saving since the $Li_3V_2(PO_4)_3$ (a)C composites are directly sintered without any protective gases or reductive gases, and can produce a large amount at a time, the production is hundreds of times of conventional carbothermal reduction method. The batteries based on this $Li_3V_2(PO_4)_3$ (a)C composites cathode exhibit remarkable reversibility and excellent rate capability, which can be directly ascribed to the robust structure merits of the Li₃V₂(PO₄)₃@C composites, it suggests that structure and electrochemical performance is not affected by mass production. In addition, ease of use, energy conservation and low consumption of this approach will greatly improve the efficiency of the industrial production, effectively solve the problems of the traditional method. What's more, this method can be easily extended to other high-performance cathode materials, and can be one of the most promising candidates for future industrial production.

Acknowledgements

This work was financially supported by the National Natural Science Foundation of China (Grant No. 61274073, 11474088 and 51562026), the Key Project of Education Department in Hubei Province (Grant No. D20151005).

Notes and references

- 1 J. M. Tarascon and M. Armand, *Nature*, 2001, **414**, 359–367.
- 2 M. S. Whittingham, Chem. Rev., 2004, 104, 4271-4302.
- 3 C. Masquelier and L. Croguennec, *Chem. Rev.*, 2013, **113**, 6552–6591.
- 4 L. H. Liao, J. Xie, S. C. Zhang, G. S. Cao and X. B. Zhao, *RSC Adv.*, 2015, 5, 99632–99639.
- 5 N. N. Bramnik, K. Nikolowski, C. Baehtz, K. G. Bramnik and H. Ehrenberg, *Chem. Mater.*, 2007, **19**, 908–915.
- 6 J. Wolfenstine and J. Allen, *J. Power Sources*, 2005, **142**, 389–390.
- 7 Q. Liu, L. B. Ren, C. J. Cong, F. Ding, F. X. Guo, D. W. Song, J. Guo, X. X. Shi and L. Q. Zhang, *Electrochim. Acta*, 2016, **187**, 264–276.
- 8 B. Kang and G. Ceder, Nature, 2009, 458, 190–193.
- 9 Y. Q. Qiao, J. P. Tu, X. L. Wang, D. Zhang, J. Y. Xiang, Y. J. Mai and C. D. Gu, *J. Power Sources*, 2011, **196**, 7715–7720.
- 10 J. Kim, J. K. Yoo, Y. S. Jung and K. Kang, *Adv. Energy Mater.*, 2013, **3**, 1004–1007.
- 11 J. Su, X. L. Wu, J. S. Lee, J. Kim and Y. G. Guo, *J. Mater. Chem. A*, 2013, **1**, 2508–2514.

- 12 H. Huang, S. C. Yin, T. Kerr, N. Taylor and L. F. Nazar, *Adv. Mater.*, 2002, **14**, 1525–1528.
- 13 M. X. Jing, J. Q. Li, Z. C. Pi, H. A. Zhai, L. L. Chen, S. S. Yao, J. Xiang, X. Q. Shen, X. M. Xi and K. S. Xiao, *Electrochim. Acta*, 2016, 212, 898–904.
- 14 P. Li, L. Y. Shao, P. F. Wang, X. Zheng, H. X. Yu, S. S. Qian, M. Shui, N. B. Long and J. Shu, *Electrochim. Acta*, 2015, 180, 120–128.
- 15 C. C. Yang, S. H. Kung, S. J. Lin and W. C. Chien, *J. Power Sources*, 2014, **251**, 296–304.
- 16 L. Chen, B. Yan, H. Y. Wang, X. F. Jiang and G. Yang, J. Power Sources, 2015, 287, 316–322.
- 17 L. F. Shen, H. S. Li, E. Uchaker, X. G. Zhang and G. Z. Cao, *Nano Lett.*, 2012, **12**, 5673–5678.
- 18 H. Q. Li and H. S. Zhou, *Chem. Commun.*, 2012, **48**, 1201–1217.
- 19 N. Raver, Y. Chouinard, J. F. Maganan, S. Besner, M. Gauthier and M. Armand, *J. Power Sources*, 2001, 97–98, 503–707.
- 20 Y. G. Wang, P. He and H. S. Zhou, *Energy Environ. Sci.*, 2011, 4, 805–817.
- 21 L. F. Fei, L. Sun, W. Lu, M. Guo, H. T. Huang, J. P. Wang, H. L. W. Chan, S. S. Fan and Y. Wang, *Nanoscale*, 2014, 6, 12426–12433.
- 22 H. D. Liu, G. Yang, X. F. Zhang, P. Gao, L. Wang, J. H. Fang, J. Pinto and X. F. Jiang, *J. Mater. Chem.*, 2012, 22, 11039–11047.
- 23 H. Wu, G. H. Yu, L. J. Pan, N. Liu, M. T. McDowell, Z. N. Bao and Y. cui, *Nat. Commun.*, 2013, 4, 1943.
- 24 W. F. Mao, J. Yan, H. Xie, Z. Y. Tang and Q. Xu, *J. Power Sources*, 2013, 237, 167–171.
- 25 X. H. Rui, C. Li and C. H. Chen, *Electrochim. Acta*, 2009, 54, 3374–3380.
- 26 L. Chen, B. Yan, H. Y. Wang, X. F. Jiang and G. Yang, *J. Power Sources*, 2015, 287, 316–322.
- 27 F. Teng, Z. H. Hu, X. H. Ma, L. C. Zhang, C. X. Ding, Y. Yu and C. H. Chen, *Electrochim. Acta*, 2013, **91**, 43–49.
- 28 Y. Q. Qiao, X. L. Wang, Y. J. Mai, X. H. Xia, J. Zhang, C. D. Gu and J. P. Tu, *J. Alloys Compd.*, 2012, **536**, 132–137.
- 29 Y. Wu, X. M. Zhao, Z. H. Song, L. P. Lin, C. Q. Du and Z. Y. Tang, *J. Power Sources*, 2015, 274, 782–790.
- 30 P. Fu, Y. M. Zhao, Y. Z. Dong, X. N. An and G. P. Shen, *Electrochim. Acta*, 2006, 52, 1003–1008.
- 31 L. J. Wang, X. C. Zhou and Y. L. Guo, *J. Power Sources*, 2010, 195, 2844–2850.
- 32 Y. Q. Qiao, J. P. Tu, X. L. Wang and C. D. Gu, *J. Power Sources*, 2012, **199**, 287–292.
- 33 J. J. Chen and M. S. Whittingham, *Electrochem. Commun.*, 2006, 8, 855–858.
- 34 T. Muraliganth, A. V. Murugan and A. Manthiram, *Chem. Commun.*, 2009, **19**, 7360–7362.
- 35 W. He, X. D. Zhang, X. Y. Du, Y. Zhang, Y. Z. Yue, J. X. Shen and M. Li, *Electrochim. Acta*, 2013, **112**, 295–303.
- 36 X. Y. Du, W. He, X. D. Zhang, Y. Z. Yue, H. Liu, X. G. Zhang,
 D. D. Min, X. X. Ge and Y. Du, *J. Mater. Chem.*, 2012, 22, 5960–5969.
- 37 Q. Kuang, Y. M. Zhao, X. N. An, J. M. Liu, Y. Z. Dong and L. Chen, *Electrochim. Acta*, 2010, 55, 1575–1581.

- 38 Y. Z. Li, X. Liu and J. Yan, *Electrochim. Acta*, 2007, **53**, 474–479.
- 39 X. H. Rui, N. Yesibolati, S. R. Li, C. C. Yuan and C. H. Chen, *Solid State Ionics*, 2011, 187, 58–63.
- 40 C. X. Chang, J. F. Xiang, X. X. Shi, X. Y. Han, L. J. Yuan and J. T. Sun, *Electrochim. Acta*, 2008, **53**, 2232–2237.
- 41 Q. L. Wei, Q. Y. An, D. D. Chen, L. Q. Mai, S. Y. Chen,
 Y. L. Zhao, K. M. Hercule, L. Xu, A. Minhas-Khan and
 Q. J. Zhang, *Nano Lett.*, 2014, 14, 1042–1048.