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Facile preparation of Mn₃O₄ hollow microspheres via reduction of pentachloropyridine and their performance in lithium-ion batteries

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 Mn_3O_4 hollow microspheres have been facilely prepared *via* a green synthesis of 2,3,5,6-tetrachloropyridine reduced from pentachloropyridine by manganese. The specific hollow microspheres were made by a H_2 gas bubble-templating method presenting a high specific surface area (87.1 m² g⁻¹) and a big total pore volume (0.2030 cm³ g⁻¹). The Mn_3O_4 hollow microspheres as an anode material demonstrate a good electrochemical performance, with a high reversible capacity of 646.9 mA h g⁻¹ after 240 cycles at a current density of 200 mA g⁻¹. The good cycling performance is attributed to numerous mesopores, high specific surface area and big total pore volume, which can offer good electrical contact and conductivity as well as accommodate the mechanism strains. In addition, the yield and selectivity of 2,3,5,6-tetrachloropyridine achieved up to 99.2% and 99.5%, respectively.

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Introduction

Recently, Mn₃O₄ have drawn considerable interest in many fields including lithium-ion battery (LIB),1-4 catalysis,5,6 ion exchange,7 molecular adsorption, magnetic applications8 and supercapacitors,9 etc. Particularly in LIBs applications, Mn₃O₄ has been regarded as a remarkably attractive anode material, owing to its relatively low cost, environmentally friendly nature, and abundant natural reserves.3,4 Hence, Mn3O4 with various nanostructures have been explored through different routes to improve the electrochemical performance of LIBs. Consequently, engineering hollow nanostructure or mesoporous architectures have been proposed as an effective approach for enhancing electrochemical performance because of its unique advantages. For instance, the large surface area contributes to a large electron/Li-ion contact area between the electrolyte and electrode solid; the pores provide flexible and fast transport pathways for the electrolyte ions and free space to alleviate the strains caused by significant volume change from lithium-ion insertion/extraction.10-13 D. Pasero synthesized pure Mn₃O₄ which have a capacity of 200 mA h g⁻¹, and a cobalt-doped sample of Mn₃O₄ exhibited a capacity of 400 mA h g⁻¹. 14 Wang synthesized carbon layer coated Mn₃O₄ nanorods which retained a capacity of 473 mA h g⁻¹ after 50 cycles at a current density of 40 mA g^{-1} . Deepak P. Dubal reported a mesoporous stacked Mn₃O₄ nano sheets exhibited a capacity of about 400 mA h g^{-1} at 0.1C. Wang and Du prepared an order-aligned Mn_3O_4 anode retaining a capacity of 494 mA h g⁻¹ over 100 cycles.3 Gao found that sponge-like Mn₃O₄ nanomaterial exhibited a high capacity of 800 mA h g⁻¹ after 40 charge/ discharge cycles.4 Despite achieving better performance, it should be noted that the methods used to prepare mesoporous architectures Mn₃O₄ nanomaterials are coprecipitation, electrospinning, hydrothermal/solvothermal and sol-gel techniques, etc., 1-4,13-18 which always include high-temperature processing and excessive consumption of organic agents leading to low yields. On the other hand, high purity Mn source used in these methods, such as MnCl₂, Mn(CH₃COO)₂, and KMnO₄, are obtained after certain processes from manganese metal, which will increase the cost of Mn₃O₄ production to some extent. In consideration of practical applications, it is highly desirable to explore a simple method to produce Mn₃O₄ from manganese metal with good control of morphology, mild operating conditions, lower cost and high through-put.

Even directly using manganese metal as raw materials to produce Mn₃O₄, the costs are still too high. To solve this problem, we turned our attention to better utilization of the high-grad chemical energy in metallic manganese, in which way will the cost significantly be reduced by the generation of high-value by-product. The reducibility of manganese is stronger than that of zinc attributing to higher reduction potential of Mn/Mn²⁺ with 1.185. Zinc is widely applied in organic reduction such as hydrodeoxygenation and hydrodechlorination.¹⁹⁻²¹ Additionally, plenty of developed theories on the mechanism of these organic reductions have suggested that zero valent metal just serves as a donor of electrons (reducing agent) to reduce H⁺ into active hydrogen which is the primary to finish the

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reduction process.²²⁻²⁵ Guided by the above theories and practical application, it is considered feasible to substitute zinc with manganese as reductant in hydrodechlorination. For example, it can be used in hydrodechlorination of pentachloropyridine (PCP) for preparing 2,3,5,6-tetrachloropyridine (TECP), a valuable commercial product as an important intermediate which is useful in manufacture agrochemical,26 such as chlorpyriphos,27 triclopyricarb,28 triclopyr and their derivatives.29,30 Meanwhile, common problems in organic reductions using zinc dust as reducing agent,31 such as poisonous solvent (acetonitrile) used, equipment corrosion by using concentrated hydrochloric acid, mass of concentrated acid wastewater containing Zn²⁺ yielded, can also be hopefully solved. Moreover, generated Mn²⁺ can easily transform into Mn₃O₄.16 And the organic solvents, necessarily used in organics involved reaction, is possibly beneficial to the formation of mesoporous architectures, because of their dispersion effect on nanocrystals. 32,33

Herein, a facile preparation of two products Mn₃O₄ hollow microspheres material and TECP reduced from PCP by manganese with a low temperature reaction using water as hydrogen source and ethanol as inexpensive, non-toxic solvent followed by a simple aeration step in the same pot, is reported. The specific hollow microspheres, consisted of Mn₃O₄ nanoparticles possessing high specific surface area of about 87.1 m² g⁻¹, self-assembled around the H₂ gas bubble templates into macro- and mesoporous structures. In addition, the as-prepared Mn₃O₄ hollow microspheres were used as an anode material for a lithium-ion battery to investigate its electrochemical properties, such as capacity, rate capability, cycle life, etc. Detailed characterization confirms that the Mn₃O₄ hollow spheres shows a comparable performance. The strategy of Mn₃O₄ preparation, substituting zinc with manganese can possibly be extended to other organic reductions using zinc dust as reducing agent.

2. Experimental

Synthesis of Mn₃O₄ hollow microspheres and 2,3,5,6tetrachloropyridine

In a typical process, 2 g pentachloropyridine (PCP) (Aladdin, reagent grade, 98%) was dissolved in 200 mL ethyl alcohol (Sinopharm Chemical Reagent Co., Ltd, analytic grade, 99%) with vigorously stirring in 40 °C. Then 0.6 g pure metallic manganese (Mn) dust (Aladdin, reagent grade, 99%, 200 meshes) was added into the mixture. Solution of 1.3 g ammonium acetate (NH4Ac) (Sinopharm Chemical Reagent Co., Ltd, analytic grade, 99%) dissolved in 5 g water was added drop-wise over a 2 hours' period. After agitation for about 4 hours under the protection of nitrogen, the slurry was heated up to 65 °C and adjusted the pH value close to 8.0 by addition 20% ammoniaque (NH₃ H₂O) solution. At the same time, fresh air was blown into the reaction for 1 h.

Then the mixture was centrifuged and the solids portion was washed several times with absolute ethanol to remove impurities and dried at 60 °C for 5 h in air to get the Mn₃O₄ hollow microspheres. 2,3,5,6-Tetrachloropyridine products was obtained by decompressing and distilling the centrifugal liquid, recovering the solvent, washing the residual in the cold water

and filtering. The acetate in washing water was then recovered and recycled.

2.2 Characterization and analysis

Transmission electron microscopy (TEM), high resolution transmission electron microscopy (HR-TEM), and selected area electron diffraction (SAED) measurements were obtained from a JEOL-2100F instrument at an acceleration voltage of 200 kV. Field emission scanning electron microscopy (FE-SEM) is performed with an ISM-7800F. X-ray diffraction (XRD) analyses were carried out on Bruker D8 diffractometer. Diffraction patterns were collected in the range of $2\Theta = 10-80^{\circ}$ under ambient conditions with a scanning rate of 8° min⁻¹. X-ray photoelectron spectroscopy (XPS) analysis was conducted by an ESCALAB 250 with Al K alpha radiation. Nitrogen adsorption-desorption isotherms were measured on a ASAP 2020 adsorption analyser.

Chloride pyridine were analysed by a high-performance liquid chromatography (HPLC) equipped with a C18-WR column (250 \times 4.6 mm, 5 μ m) and an UV detector. The mobile phase composed 90% methanol (HPLC-grade) and 10% ultrapure water, and the detective wavelength was 298 nm, with a flow rate of 1.0 mL min⁻¹ and an injection volume of 20 μ L.

2.3 Electrochemical investigation

The electrodes for electrochemical studies were prepared with 70 wt% active material of Mn₃O₄ hollow microspheres, 10% polyvinylidenefluoride (PVDF), 20 wt% conducting acetylene black binder in N-methyl pyrrolidinone, grinded for 30 min in an agate mortar, thus the slurry was coated on a piece of Cu foil, and the plate was dried at 80 °C under vacuum for 12 hours, the cells were fabricated by using lithium foil as the counter electrode and the reference electrode, Celgard 2300 microporous membrane as separator, and a solution of 1 mol L⁻¹ LiPF6 in a mixture of ethylene carbonate (EC) and dimethyl carbonate (DMC) with volume ratio of 1:1 as electrolyte. The assembly of the cell was prepared in an argon-filled humidity-free glove box. And the cells were cycled galvanostatically from 0.01 to 3.00 V at different current rates (Land CT2001A). Cyclic voltammograms (CV) were performed on a CHI-760 electrochemical workstation over a potential range 0.01 to 3.00 V at a scan rate of 0.40 mV s^{-1} .

3. Results and discussion

Characterization of Mn₃O₄ hollow microspheres

As shown in Fig. 1, all the diffraction peaks of the XRD pattern should be indexed to the spinel structure of Mn₃O₄ (JCPDS card of 24-0734, space group of $I4_1/amd$, a = 5.762, b = 5.762, c =9.47), which confirms that these products are pure Mn₃O₄. Relatively broad diffraction peaks indicate that Mn₃O₄ crystals are in small sizes. From the width of diffraction peaks and calculation by the Debye-Scherrer formula, an average particle size of Mn₃O₄ is around 9.7 nm.

The morphology of the Mn₃O₄ product was investigated by FE-SEM and TEM. The typical SEM image of the Mn₃O₄ product **RSC Advances**

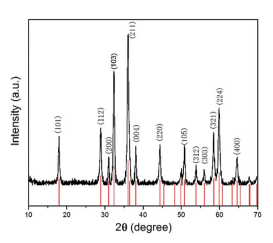


Fig. 1 XRD pattern of Mn₃O₄ hollow microspheres.

shown in Fig. 2a displays that the crystals are formed in large scale with good dispersion and uniform diameter of about 500 nm. Moreover, observing from some broken microspheres shown in the partial enlarged detail in Fig. 2b, it is evident that these Mn₃O₄ microspheres are in hollow structure. Furthermore, the Mn₃O₄ microspheres were characterized with TEM to examine their fine structures. Fig. 2c shows that the Mn₃O₄ microspheres present a clear hollow structure with loose shell. A low magnification TEM image of the Mn₃O₄ hollow sphere shell is shown in Fig. 2d, which manifest that Mn₃O₄ nanocrystals were partially aggregated together. Measurement of about 100 nanocrystals from more HR-TEM images indicates that most nanocrystal sizes range from 7 to 15 nm, which is

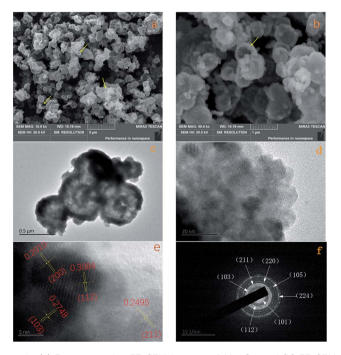


Fig. 2 (a) Representative FE-SEM images of Mn_3O_4 and (b) FE-SEM image of partial enlarged detail (c) TEM image of Mn_3O_4 . (d) TEM image of partial enlarged detail; (e) HR-TEM image of partial enlarged detail; (f) SAED image of Mn_3O_4 .

concordant with the value obtained from the XRD measurement. Also, there are some defects and vacancies on the surface of the sphere. Thus, it appears to be unsmooth. The vacancies and defects are beneficial to the Li-ion transport and redox reaction, which can improve the electrochemical performance of Mn₃O₄ as the lithium-ion battery material to some extent.¹¹ As high-resolution transmission electron microscopy (HR-TEM) image of the Mn₃O₄ nanoparticles is shown in Fig. 2e, the lattice fringes exhibited different crystal planes, which can be identified in (103), (112), (211) and (200) planes with matching dspacing of 0.2784, 0.3084, 0.2495 and 0.2910 nm, respectively. Fig. 2f is ring pattern of selected-area electron diffraction (SAED), the rings matched well with (101), (112), (103), (211), (220), (105), (224) planes, as reference to the JCPDS database (card of 27-0734) further demonstrate the tetragonal structure of Mn₃O₄ nanoparticles.

The specific surface area and pore size distribution of the synthesized Mn₃O₄ hollow microspheres were further investigated by N2 adsorption-desorption measurements. The adsorption isotherms profiles are displayed in Fig. 3. It shows that Mn₃O₄ hollow microspheres have a type isotherm of IV adsorption-desorption curve with a capillary condensation step and hysteresis loop which is characteristic of mesoporous materials. The BET surface area of the Mn₃O₄ sample is measured to be 87.1 m² g⁻¹ and a total pore volume of 0.2030 cm³ g⁻¹, much higher than the previous reports. For example, the nanotube Mn₃O₄ present BET surface area about 48.83 m² g⁻¹ and a total pore volume of $0.08 \text{ cm}^3 \text{ g}^{-1}$, the mesoporous Mn_3O_4 has a total pore volume of 0.150 cm³ g⁻¹. The pore-size distribution curve (the inset in Fig. 4), obtained by the Barrette-Joynere-Halenda (BJH) method, shows the pore size distribution located around at 5 nm. From the TEM photos in Fig. 2, such pores were formed mainly by the stacking of nanoparticles. The narrow pore size distribution and big volume will contribute much to the effective transport and storage of ion at the interface of electrode/electrolyte, which shows the tremendous potential applications in LIBs, sensors and electrochemical super capacitors.

Although it is easy to see the hollow sphere product is pure Mn₃O₄ with hausmannite crystal structure from the XRD analysis and has clean and neat crystal faces from HR-TEM photos, we further characterized the sample by X-ray photoelectron spectroscopy (XPS) and analysed with liquid-phase micro extraction high performance liquid chromatography (LPME-HPLC) analysis procedure, to investigate if there is chloride pyridine composition remnant on the surface or wrapped inside the Mn₃O₄ nanocrystals. Fig. 4a shows a typical over XPS spectra of the Mn₃O₄ hollow microspheres, suggesting that the hollow microspheres test samples are composed of Mn 2p, 3s, 3p, O 1s and 2s, respectively, as well as a weaker C 1s peak at 284.8 eV always come from environmental carbon pollution used for the XPS curve correction. It is clear that N and Cl not detected in the curve, which substantiates almost no existence of chloropyridine components in surface of hollow spheres after washing process. Fig. 4c depicts the XPS spectrum of Mn 2p, where two peaks located at 642.19 eV and 653.19 eV can be respectively attributed to Mn $2p_{3/2}$ and Mn $2p_{1/2}$ levels. In the Mn 3s spectra (Fig. 4d), two peaks at 83.40 eV and 89.04 eV with a separation of 5.64 eV

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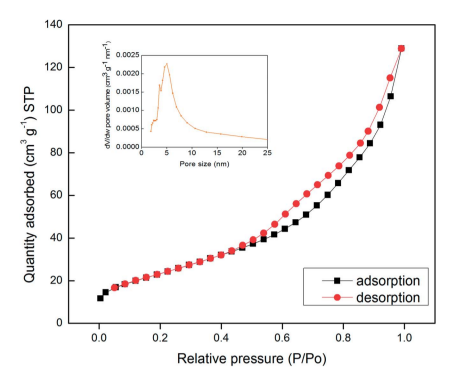


Fig. 3 N_2 adsorption isotherms and pore size distribution (insets) of Mn₃O₄ hollow microspheres

were observed, which agree well with that of reported for Mn₃O₄. 35,36 As for the O 1s spectra shown in Fig. 4b, the asymmetric peak was obviously superposed by two peaks a main at 529.89 eV and weaker 531.49 eV one, which could be attributed to lattice oxygen (O_{latt}) and adsorbed oxygen (O_{ads}), respectively.37 Adsorbed oxygen (Oads) can be associated with many kinds of species including low coordination situation, oxygencontaining surface contamination and surface chemisorbed oxygen. A higher ratio of O_{ads}/O_{latt} (in our work is 0.4) could indicate a larger amount of chemisorbed oxygen species,38 which could explain that the as-prepared Mn₃O₄ hollow microspheres have a good interfacial activity.

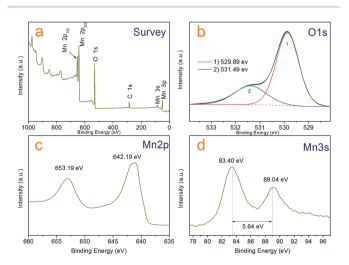


Fig. 4 XPS spectra of the Mn₃O₄ hollow microspheres (a) survey scan, (b) O 1s, (c) Mn 2p (d) Mn 3s spectra.

In the LPME-HPLC analysis procedure, 0.1 g as-synthesis Mn₃O₄ sample dissolved in hydrochloric acid, and then the transparent solution was transferred into a 10 mL volumetric flask and diluted with purified water. 0.5 mL toluene was added to extract the organic compounds for analysing. The result showed that the residue of TECP was lower than 2.0×10^{-7} g in every one gram Mn₃O₄ sample and other chloride pyridine compositions were not detected.

3.2 Reaction mechanism and formation processes of hollow microspheres

To study the reaction mechanism and formation process of Mn₃O₄ hollow microspheres, the source material and intermediates were tested by XRD and SEM. The intermediate samples

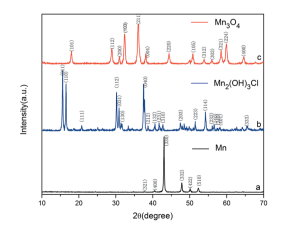


Fig. 5 XRD pattern of (a) Mn; (b) Mn₂(OH)₃Cl; (c) Mn₃O₄.

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were taken 4 hours after NH₄Ac solution were added dropwise at the temperature of 40 °C. Fig. 5a shows the XRD pattern of the metallic manganese source, which suggests that the source is pure Mn (JCPDS card of 32-0637, space group of $I\bar{4}3m$, a=8.912, b = 8.912, c = 8.912). Fig. 5b shows the XRD pattern of the intermediates, which indicates that the intermediate has a Mn₂(OH)₃Cl crystal structure (JCPDS card 25-1158 space group pnam, $a_0 = 6.490$, $b_0 = 9.52$, $c_0 = 7.12$). Fig. 5c shows the XRD pattern of final product Mn₃O₄. Based on the test results, the possible reaction mechanism of the first reaction step for obtaining TECP is described as follows.

$$2NH_4^+ + Mn \rightarrow Mn(NH_3)_2^{2+} + 2H$$
 (1)

$$2Cl^{\cdot} + Mn \rightarrow Mn^{2+} + 2Cl^{-}$$
 (3)

$$2Mn(NH_3)_2^{2+} + 3H_2O + Cl^- \rightarrow Mn_2(OH)_3Cl + 4NH_4^+$$
 (4)

$$2H' \rightarrow H_2 \uparrow$$
 (5)

In reaction (1), catalyst NH₄⁺ attaches on the surface of metallic Mn as a hydrogen carrier, where electron transferred from Mn (0) to NH₄⁺ and reactive hydrogen and Mn(NH₃)₂²⁺ generated. Then, Mn2+ could easily transfer from the surface of manganese powder into the solution in the form of Mn(NH₃)₂²⁺ complexes, which is very important for the exposure of intimal Mn. As in reaction (2) and (3), the reactive hydrogen replaces p-Cl from PCP selectively, which is a typical hydrodechlorination by SN₂ reaction process. At the meantime, substituted Cl gains another electron on the surface of metallic Mn and turned to Cl⁻¹. Unavoidably, reactive hydrogens can easily combine to H₂ as shown in reaction (5). Mn(NH₃)₂²⁺ will hydrolyse into

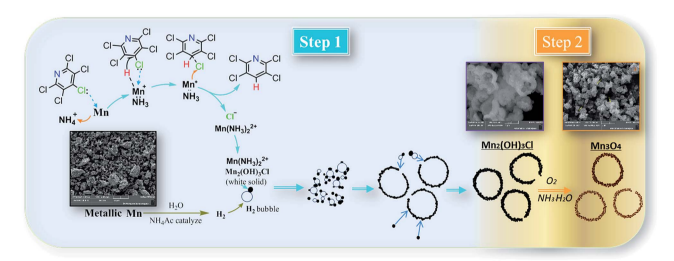
Mn₂(OH)₃Cl crystal as shown in reaction (4). Consequently, by using manganese powder as reducing agent, water as the hydrogen donor, NH₄Ac as catalyst, alcohol as solvent and in an eco-friendly temperature of 40 °C, the yield and selectivity of TECP achieved up to 99.2% and 99.5%, respectively. As the pH value adjusted to 8.0 with NH₃ H₂O solution, all the Mn²⁺ transformed into Mn(OH)2 white precipitation. When the air flowed in, its colour apparently turned to brown quickly, which is the specific phenomenon of Mn(OH)2 out of all the manganese compounds.39 Mn(OH)2 will then dehydrate and generate Mn_3O_4 . The reactions can be expressed as eqn (6)–(8).

$$Mn_2(OH)_3Cl + NH_3 \cdot H_2O \rightarrow 2Mn(OH)_2 + NH_4Cl$$
 (6)

$$Mn(NH_3)_2^{2+} + 2H_2O \rightarrow Mn(OH)_2 + 2NH_4^{+}$$
 (7)

$$6Mn(OH)_2 + O_2 \rightarrow 2Mn_3O_4 + 6H_2O$$
 (8)

To explain formation processes of hollow microspheres clearly, a plausible conjecture involving bubble-templating process was described in Scheme 1, where the by produced H₂ plays the role of bubble templates. Actually, the gas bubbletemplating method has been proved to be a facile process to fabricate hollow microspheres of inorganic materials, such as Fe, 40 ZnO, 41 ZnSe, 42 CoOOH, 43 CaCO₃ (ref. 44) and $Cu_{2x}Se_{\nu}S_{1\nu}/c$ rGO.45 As illustrated in Scheme 1, a certain concentration of H2 was generated by reaction (5) and formed sub-microbubbles. At the same time, a large amount of Mn₂(OH)₃Cl nuclei was produced in solution through the relatively low nucleation process by reaction (4), leading to a relatively high local concentration of Mn₂(OH)₃Cl. Thus, it inevitably resulted in Mn₂(OH)₃Cl nuclei diffusing to the gas-liquid interfaces between H2 bubbles and liquid solution. Then these Mn2-(OH)₃Cl nuclei grow into small size nanocrystals with the dispersion effect of ethyl alcohol. Additionally, small size H₂ bubbles come into collisions in the dispersion system and coagulate to generate bigger H₂ bubbles, while the Mn₂(OH)₃Cl nanocrystals on the surface of bubbles assembled into hollow



Scheme 1 Possible mechanism for hydrodechlorination reaction process and formation processes of hollow microspheres.

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microspheres. Other $Mn_2(OH)_3Cl$ nanocrystals may also accumulate on the surface, forming the hollow sphere $Mn_2(OH)_3Cl$ with a thick shell, which is confirmed by the SEM photo of $Mn_2(OH)_3Cl$ with hollow structure as shown in Scheme 1. Driven by reaction (6)–(8), these nanocrystals generated smaller-volume Mn_3O_4 nanocrystals and left some space or pores between nanocrystals in the shell, forming as-prepared Mn_3O_4 hollow microspheres with macro- and mesoporous structures.

As a summary, two products 2,3,5,6-tetrachloropyridine and Mn₃O₄ hollow microspheres material were prepared in one pot via two simple steps with Mn, water, ethyl alcohol and air as inexpensive, non-toxic reductive agent, hydrogen source, solvent and oxidizing agent, respectively. With several advantages, including low temperature, low corrosion to equipment, good atom economy, organic-free template, low-cost, high conversion rate and selectivity, this hollow Mn₃O₄ microspheres preparation method was much better than those were previously reported. For example, Jian reported a method based on an in situ gas blowing mechanism in single aerosol droplets, in which Mn(NO₃)₂, sucrose and H₂O₂ used as raw material to synthesize hollow Mn₃O₄ microspheres.¹⁷ Yue used carbon nanospheres (CNSs) as a template and reagent to reduce KMnO4 into hollow MnO2 nanospheres, and then the hollow MnO2 nanospheres were reduced under an atmosphere of H₂/Ar into hollow Mn₃O₄ microspheres at 280 °C for 3 h.46 Sun reported a surface-layeradsorption and calcination (SLA-C) method for the fabrication of hollow Mn₃O₄ microspheres by templating carbonaceous microspheres. 47 Thus, it can be considered as a facile and green synthesis technology, as is summarized in reaction (9).

3.3 Electrochemical performance

The electrochemical properties of the as-synthesized Mn₃O₄ hollow microspheres were evaluated by cyclic voltammetry (CV)

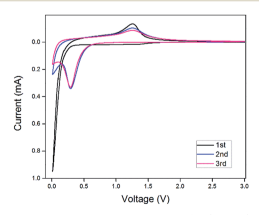
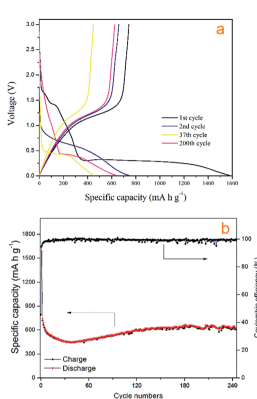


Fig. 6 Cyclic voltammetry (CV) curves of the 1^{st} 2^{nd} and 3^{rd} cycles for the Mn₃O₄ hollow microspheres at a scan rate of 0.40 mV s⁻¹ between 0.01 and 3.00 V.



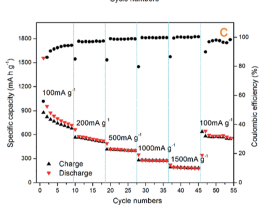


Fig. 7 (a) The $1^{\rm st}$, $2^{\rm nd}$, $37^{\rm th}$ and $200^{\rm th}$ voltage profiles of the Mn₃O₄ electrode between 0.01 and 3.00 V at a current density of 200 mA g⁻¹. (b) Cycle performance and coulombic efficiency *versus* cycle number of Mn₃O₄ hollow spheres at a current density of 200 mA g⁻¹. (c) Rate capabilities with increasing current density.

conducted in the range of 0.01 and 3.00 V at a scan rate of 0.40 mV s⁻¹, which is shown in Fig. 6. During the first cathodic process, there was a broad weak peak located at 1.10 V which disappeared afterwards. This peak can mainly be attributed to the formation of a solid electrolyte interphase (SEI) layer because of the decomposition of the electrolyte and formation of solid-state interface of the Mn₃O₄ hollow microspheres. The intensive peak which appeared in the low potential is ascribed to the reduction of Mn₃O₄ into Mn⁰ accompanied by the generation of Li₂O. In the following anodic process, it can be observed at around 1.3 V that Mn⁰ is oxidized to Mn₃O₄ and Li₂O is decomposed in the meantime. In the following cycles, the reduction peak shifted to a higher voltage about 0.3 V, which

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is primarily due to an irreversible structure change related to the lithium insertion. After the second cycle, the CV curves demonstrate good reversibility and stability of the Mn₃O₄ hollow microspheres.

Fig. 7a shows the cycling stability curve of the Mn₃O₄ hollow microspheres, examined by charging/discharging between 3.00 and 0.01 V at a current density of 200 mA g⁻¹ over 240 cycles. The electrode has a large initial discharge capacity of about 1577.8 mA h $\rm g^{-1}$. However, the coulombic efficiency for the first cycle was 46% with the first charge capacity of 720.48 mA h g^{-1} , then the discharge capacity gradually dropped to 448.2 mA h g⁻¹ at the 38th cycle, while the coulombic efficiency increased to 93% sharply at the second cycle, then gradually increased to and maintained at between 95-100% in the following several hundred cycles (Fig. 7b). The capacities of Mn₃O₄ exhibited a rising trend from the 39th cycle, and the electrode retains an average reversible capacity of 637.2 mA h $\rm g^{-1}$ during the 130 $^{\rm th}$ cycle to the 240th cycle. The Mn₃O₄ hollow microspheres electrode still retains a reversible capacity of 646.9 mA h g⁻¹ after total around 240th cycles. Such a rising trend of capacity between the 38th-130th is normally observed in transition metal oxides and is well-documented in literatures. 1,48-51 It is considered to be attributed to the reversible growth process of a polymeric gel-like film resulting from kinetically activated electrolyte degradation. In terms of discharge capacity and cycling stability, our electrode performs better than some previously reported Mn₃O₄ electrode.

The charge/discharge profiles of the hollow Mn₃O₄ microspheres in the 1st, 2nd, 37th, and 200th cycles for the Mn₃O₄ hollow microspheres electrode cycled between 3.00 and 0.01 V at a current density of 200 mA g⁻¹ are shown in Fig. 6a. In the first discharge curve, it is observed a sloping voltage from 2.00 to 0.35 V is attributed to the generation of solid-electrolyte interphase (SEI) film and the reduction of Mn³⁺ to Mn²⁺. The well-defined voltage plateau at 0.35 V is related to the main reduction of Mn²⁺ to Mn⁰. In the charge curve, the Mn₃O₄ hollow microspheres electrode shows a gentle incline in the voltage range between 1.0 and 1.5 V, which is associated with the oxidation from Mn⁰ to Mn²⁺/Mn³⁺. In comply with the CV results, in the second cycle the lithiation plateau moves to a higher voltage of about 0.65 V and the sloping voltage rapidly decreased from 1.75 V to 1.00 V which are attributed to an irreversible capacity loss of 54%, resulted from the phase transformation in the first cycle. As the cycle times increase from 2nd cycle to 37th cycle, the lithiation plateau continue decreasing to a lower voltage of about 0.5 V, which is attributed to the continuing phase transformation occurred in the conversion process. The lithiation plateau of the 200th cycle hardly changed compared with the 37th cycle, but the voltage decrease from 2.00 V to 0.50 V is much slower, which contributes a lot to the increase of the reversible capacity, and it also confirms the dominant influence on Mn₃O₄/Li cell capacity from the reversible growth of polymeric gel-like film.

The perfect electrochemical performance of the synthesized Mn₃O₄ hollow microspheres could be associated with the big specific surface area and the big total pore volume of the macroand mesoporous. The porous sponge structure could improve

the efficient utilization of active material and allow lithium ions to transfer easily in and out. The unique structure is also able to accommodate the strain by volume change during conversion process and maintain the integrity in the reversible growth process of polymeric gel-like film.

We also investigated the rate capability of Mn₃O₄ hollow microspheres at increased stepwise current densities from 100 mA g⁻¹ to 1500 mA g⁻¹ between 0.01 and 3.00 V with each current density tested for 8 cycles (Fig. 7c). At current density of 100 mA g⁻¹, it delivered an initial capacity of about 955 mA h g^{-1} , then reduced to 718 mA h g^{-1} gradually. Even at 1500 mA g⁻¹ the Mn₃O₄ hollow microspheres can still deliver a specific capacity of nearly 200 mA h g⁻¹. When the test current density returns to 100 mA g^{-1} again, the capacity recovered to 645 mA hg⁻¹, which was close to the capacity at the initial rate of 100 mA g⁻¹, indicating a good reversibility of the Mn₃O₄ hollow microspheres. Thus, as-prepared Mn₃O₄ hollow microspheres showed good electrochemical properties and huge application potential in high capacity LIB anode materials.

Conclusions 4.

In summary, a new facile and green route for the preparation of Mn₃O₄ hollow microspheres and 2,3,5,6-tetrachloropyridine in one pot via two simple steps was developed using Mn, water, ethyl alcohol and air as inexpensive non-toxic reductive agent, hydrogen source, solvent and oxidizing agent, respectively. Mn₃O₄ hollow microspheres was made by H₂ gas bubbletemplating method presenting a high specific surface area $(87.1 \text{ m}^2 \text{ g}^{-1})$ and a big a total pore volume $(0.2030 \text{ cm}^3 \text{ g}^{-1})$. The Mn₃O₄ hollow microspheres electrode shows a good electrochemical performance with a reversible specific capacity of 646.9 mA h g⁻¹ at a current density of 200 mA g⁻¹ after 240 cycles. In addition, the yield and selectivity of 2,3,5,6-tetrachloropyridine achieved up to 99.2% and 99.5%, respectively. Above all, the whole process has several advantages, including low temperature, low corrosion to equipment, good atom economy, surfactant and/or solid free template, low-cost, high conversion rate and selectivity and short reaction process. The 2,3,5,6-tetrachloropyridine will significantly reduce the production costs of Mn₃O₄ product. The strategy of Mn₃O₄ preparation, substituting zinc with manganese can possibly be extended to other organic reductions using zinc dust as reducing agent.

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Notes and references

1 Q. Hao, J. Wang and C. Xu, J. Mater. Chem. A, 2014, 2, 87-93.

Paper

Sources, 2013, 244, 56-62.

2 I. Nam, N. D. Kim, G.-P. Kim, J. Park and J. Yi, J. Power

- 3 J. Wang, N. Du, H. Wu, H. Zhang, J. Yu and D. Yang, *J. Power Sources*, 2013, **222**, 32–37.
- 4 J. Gao, M. A. Lowe and H. D. Abruna, *Chem. Mater.*, 2011, 23, 3223–3227.
- 5 G. Wang, B. Huang, Z. Lou, Z. Wang, X. Qin, X. Zhang and Y. Dai, *Appl. Catal.*, *B*, 2016, **180**, 6–12.
- 6 Y. Li, J. Qu, F. Gao, S. Lv, L. Shi, C. He and J. Sun, *Appl. Catal.*, *B*, 2015, **162**, 268–274.
- 7 W. B. S. Machini and M. F. S. Teixeira, *Electroanalysis*, 2014, 26, 2301.
- 8 L. C. Dong, Y. B. Zhong, S. Zhe, T. Y. Zheng and H. Wang, *RSC Adv.*, 2016, **6**, 21037–21042.
- 9 I. Ryu, G. Kim, D. Park and S. Yim, J. Power Sources, 2015, 297, 98-104.
- 10 X. W. Lou, L. A. Archer and Z. Yang, Adv. Mater., 2008, 20, 3987–4019.
- 11 X. Lai, J. E. Halpert and D. Wang, *Energy Environ. Sci.*, 2012, 5, 5604–5618.
- 12 Z. Wang, L. Zhou and X. W. Lou, *Adv. Mater.*, 2012, 24, 1903–1911.
- 13 Z. Bai, N. Fan, Z. Ju, C. Guo, Y. Qian, B. Tang and S. Xiong, J. Mater. Chem. A, 2013, 1, 10985–10990.
- 14 D. Pasero, N. Reeves and A. R. West, *J. Power Sources*, 2005, **141**, 156–158.
- 15 C. Wang, L. Yin, D. Xiang and Y. Qi, ACS Appl. Mater. Interfaces, 2012, 4, 1636–1642.
- 16 D. P. Dubal and R. Holze, RSC Adv., 2012, 2, 12096–12100.
- 17 G. Jian, Y. Xu, L.-C. Lai, C. Wang and M. R. Zachariah, *J. Mater. Chem. A*, 2014, 2, 4627–4632.
- 18 N. Lavoie, P. R. L. Malenfant, F. M. Courtel, Y. Abu-Lebdeh and I. J. Davidson, *J. Power Sources*, 2012, 213, 249–254.
- 19 S. Liu, Y. Wang, J. Jiang and Z. Jin, *Green Chem.*, 2009, 11, 1397-1400.
- 20 J. Kroemer, C. Kirkpatrick, B. Maricle, R. Gawrych, M. D. Mosher and D. Kaufman, *Tetrahedron Lett.*, 2006, 47, 6339–6341.
- 21 S. Suresh, Open Waste Manage. J., 2009, 2, 6-16.
- 22 H. Chen and S. Liu, Chem. Eng. Sci., 2015, 138, 510-515.
- 23 G. N. Jovanovic, J. E. Atwater, P. Znidarsic-Plazl and I. Plazl, *Chem. Eng. J.*, 2015, 274, 50–60.
- 24 Y.-h. Shih, M.-Y. Chen and Y.-F. Su, *Appl. Catal.*, *B*, 2011, **105**, 24–29.
- 25 C.-B. Wang and W.-x. Zhang, Environ. Sci. Technol., 1997, 31, 2154–2156.

- 26 A.-Y. Guan, C.-L. Liu, X.-F. Sun, Y. Xie and M.-A. Wang, *Bioorg. Med. Chem.*, 2016, 24, 342–353.
- 27 R. H. Rigterink, US. Pat., 3, 244, 586, 1966.
- 28 Y. Matsuzaki, WO. Pat., 2015012245A1, 2015.
- 29 Z. Xiaomao, D. Huijuan and L. wei, CN. Pat., 102718700A, 2012.
- 30 J. C. Yang, M. Li, Q. Wu, C. L. Liu and X. H. Chang, *Bioorg. Med. Chem.*, 2015, 24, 383–390.
- 31 F. H. Murphy, US. Pat. 470 3123 A, 1987.
- 32 O. Zaberca, A. Gillorin, B. Durand and J. Y. Chane-Ching, *J. Mater. Chem.*, 2011, 21, 6483–6486.
- 33 Y. Zhao, Z. Zhang and H. Dang, J. Phys. Chem. B, 2003, 107, 7574–7576.
- 34 F. Yang, M. Zhao, Q. Sun and Y. Qiao, RSC Adv., 2015, 5, 9843–9847.
- 35 G. Zhao, J. Li, W. Zhu, X. Ma, Y. Guo, Z. Liu and Y. Yang, New J. Chem., 2016, 40, 10108–10115.
- 36 J. W. Lee, A. S. Hall, J.-D. Kim and T. E. Mallouk, *Chem. Mater.*, 2012, 24, 1158–1164.
- 37 J. Li, X. Liang, S. Xu and J. Hao, *Appl. Catal.*, *B*, 2009, **90**, 307–312.
- 38 Y. Wang, A.-P. Jia, M.-F. Luo and J.-Q. Lu, *Appl. Catal.*, *B*, 2015, **165**, 477–486.
- 39 M. Fang, X. Tan, M. Liu, S. Kang, X. Hu and L. Zhang, *CrystEngComm*, 2011, **13**, 4915–4920.
- 40 G. X. Tong, J. G. Guan, Z. D. Xiao, F. Z. Mou, W. Wang and G. Q. Yan, *Chem. Mater.*, 2008, **20**, 3535–3539.
- 41 C. Yan and D. Xue, J. Alloys Compd., 2007, 431, 241-245.
- 42 Q. Peng, Y. Dong and Y. Li, *Angew. Chem., Int. Ed.*, 2003, 42, 3027–3030.
- 43 J. Yang and T. Sasaki, Chem. Mater., 2008, 20, 2049-2056.
- 44 G. Hadiko, Y. S. Han, M. Fuji and M. Takahashi, *Mater. Lett.*, 2005, **59**, 2519–2522.
- 45 W. L. Li, H. Y. Zou, J. Lan, Q. Wang, Y. F. Li and C. Z. Huang, *RSC Adv.*, 2015, 5, 91206–91212.
- 46 J. Yue, X. Gu, L. Chen, N. Wang, X. Jiang, H. Xu, J. Yang and Y. Qian, *J. Mater. Chem. A*, 2014, **2**, 17421–17426.
- 47 X. Sun, J. Liu and Y. Li, Chem.-Eur. J., 2006, 12, 2039-2047.
- 48 S. Laruelle, S. Grugeon, P. Poizot, M. Dolle, L. Dupont and J. M. Tarascon, *J. Electrochem. Soc.*, 2002, **149**, A627–A634.
- 49 G. Zhou, D.-W. Wang, F. Li, L. Zhang, N. Li, Z.-S. Wu, L. Wen, G. Q. Lu and H.-M. Cheng, *Chem. Mater.*, 2010, 22, 5306– 5313.
- 50 S. Grugeon, S. Laruelle, L. Dupont and J. M. Tarascon, *Solid State Sci.*, 2003, 5, 895–904.
- 51 J.-S. Do and C.-H. Weng, J. Power Sources, 2005, 146, 482–486.