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#### Durable High-Rate Capability Na<sub>0.44</sub>MnO<sub>2</sub> Cathode Material for Sodium-ion Batteries

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#### ABSTRACT

Monocrystalline orthorhombic Na<sub>0.44</sub>MnO<sub>2</sub> nanoplate as a potential cathode material for sodium-ion batteries has been synthesized by a template-assisted sol-gel method. It exhibits high crystallinity, pure phase and homogeneous size distribution. During the synthesis, acidic and reductive conditions are applied to limit the production of unfavorable Birnessite phase in the precursor, and colloidal polystyrene is included to avoid morphology collapse during the

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gel formation and particle elongation in one direction. The decompositions of polystyrene and citric acid during high temperature firing offer a reductive carbothermal condition which can suppress the formation of unidimensional particles, and limit particle growth along the [001] direction. As a consequence, the material delivers 96 mAh g<sup>-1</sup> discharge capacity at 10 C (86% of 0.1 C capacity) and maintains 97.8% capacity after 100 cycles at 0.5 C. Such superior rate capability and cycling stability of this material are among the best to date, suggesting its great interest in practical applications.

Keywords: Sodium-ion batteries; Cathode; Nanoplate; Rate capability; Cycling stability

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#### 1. Introduction

The urgent demand for developing a sustainable and highly efficient energy storage system has led to a great deal of interest in lithium-ion batteries (LIBs) [1-3]. However, the geographically constrained Li mineral reserves combined with the increasing demand for Li commodity chemicals, linked to the foreseen growing implementation of large-format LIBs, will surely drive its price up. LIBs might become too expensive to be applied in the rapidly developing electric vehicles or in large stationary energy storage systems. In contrast to lithium, sodium is very abundant in nature and widely available in different forms [4]. It is also expected to show similar properties to lithium, according to their proximity in the periodic table. In addition, as sodium does not alloy with Al, it would allow substituting Cu

current collector at the anode for further gains in cost and weight. These advantages of sodium motivate the exploration of rechargeable sodium-ion batteries (NIBs) as a next generation energy storage system, which indeed has attracted a great research attention recently [5-7].

As a matter of fact, Na<sup>+</sup> ion (de-)intercalation and storage are still very challenging since the ionic radius of Na<sup>+</sup> (0.98 Å) is larger than that of Li<sup>+</sup> (0.69 Å) [8, 9], which makes the accommodation of Na<sup>+</sup> ions in a host material difficult and often leads to severe structural degradation [10]. Thus, developing active materials with improved rate performance and long term cycling stability for NIBs represents the main research challenge. Numerous active materials have been intensively studied as candidates for NIBs cathodes, including sodium transition metal (TM) pyrophosphates [11-13] and sodium transition metal phosphates (TM: Co, Mn, Fe, Ni, Cr, and multicomponent transition metals) [14-18]. These polyanion-based structures with stable host frameworks and strong P-O covalent bonds are thermally stable [11]. Nevertheless, most of these materials exhibit inferior electrochemical performance, especially a poor rate capability, originating from low electronic conductivity and reduced mobility at phase boundaries, which limits their practical capacity values. P2-type Na<sub>x</sub>TMO<sub>2</sub> are considered to be promising candidates; and these compounds are being introduced as cathode materials for NIBs due to their facile synthesis and structural stability [19-21]. Among them, Na<sub>0.44</sub>MnO<sub>2</sub> (Na<sub>4</sub>Mn<sub>9</sub>O<sub>18</sub>) presents interesting characteristics linked to its open structure with interconnected and large tunnels, which is able to sustain the multiple phase change that occurs during the  $Na^+$  (de)-insertion processes [22, 23]. Sauvage *et al.* demonstrated reversible insertion and extraction of Na<sup>+</sup> ions in pure Na<sub>0.44</sub>MnO<sub>2</sub> with an

initial capacity of c.a. 80 mAh g<sup>-1</sup> at 0.1 C [22]. However, after 50 cycles, only half of the original capacity was retained. More recently, research focused on downsizing particle, such as nanowires. The obtained  $Na_{0.44}MnO_2$  exhibited improved cycling performance combined with higher capacities [10] and less than 7% capacity fading during the first 30 cycles [10]. Although the cycling performance was improved to a certain extent, the rate capability, especially at high rate still needs to be enhanced.

that Na<sub>0.44</sub>MnO<sub>2</sub> bi-dimensional structures Considering (nanoplates) extending perpendicularly to Na<sup>+</sup> most favorable insertion pathways would allow limiting surface defects, increasing the tap density and fast Na<sup>+</sup> (de)insertion (as compared to unidimensional particles), designing Na<sub>0.44</sub>MnO<sub>2</sub> material with this morphology is an efficient strategy to achieve enhanced rate capability and volumetric energy density. However, it is known that Birnessite nanoplates can form as an intermediate during the firing process of Na<sub>0.44</sub>MnO<sub>2</sub> [24], especially in alkaline and oxidative condition. They then split into Na<sub>0.44</sub>MnO<sub>2</sub> unidimensional particles upon high temperature annealing [25], while the obtaining of bi-dimensional Na<sub>0.44</sub>MnO<sub>2</sub> seems to be favored by direct high temperature sintering and the use of reductive condition, perhaps as a result of avoiding Birnessite intermediate structure. More recently, Xu et al. [26] synthesized Na<sub>0.44</sub>MnO<sub>2</sub> submicron slabs by a citric acid-based sol-gel synthesis, and obtained slightly extended (001) surface. However, the obtained slabs presented in wide range of widths and several grooves collapsing on their surface, reminiscent of Birnessite splitting mechanism.

Thus, in order to avoid extensive elongation in one direction and obtain homogenous morphology, a novel sol-gel method (NSG) with self-made in-house polystyrene (PS) is

applied to synthesize Na<sub>0.44</sub>MnO<sub>2</sub> material in this work. PS colloidal suspension is added to the precursor solution. It serves as a dispersion additive which can avoid the collapse of morphology during the gel formation, provides extra carbon source and offers a carbothermal reductive environment during firing. Orthorhombic Na<sub>0.44</sub>MnO<sub>2</sub> nanoplates are obtained, and its performance as cathode material for rechargeable NIBs is evaluated as well. In comparison, Na<sub>0.44</sub>MnO<sub>2</sub> material is also synthesized by the conventional sol-gel method (SG).

#### 2. Experimental

#### 2.1. Synthesis

Polystyrene (PS)-contained solution was synthesized by an emulsion polymerization method. Potassium persulfate (0.12 g) and sodium dodecyl sulfate (SDS 0.4 g) were dissolved in aqueous alcohol, and then drop 40ml of styrene monomer under Ar atmosphere and continuous rapid stirring. The colloidal suspension with dispersed PS was obtained by continuously stirring the mixture at 70 °C for 8 hours.

In syntheses of  $Na_{0.44}MnO_2$  active materials, stoichiometric amounts (calculated based on the 0.04 mol target material) of sodium nitrate (NaNO<sub>3</sub>, Sigma-Aldrich AR), manganese acetate (Mn(AC)<sub>2</sub>·4H<sub>2</sub>O, Sigma-Aldrich AR), as-prepared 10 ml PS-contained solution and 4 g citric acid (C<sub>6</sub>H<sub>8</sub>O<sub>7</sub>, Sigma-Aldrich AR) were dissolved in 100 ml distilled water. Here, additional to the chelating function, citric acid provided a reductive environment. After stirring for half hour, the mixture was heated to 120 °C until all the solvent evaporated. More citric acid was added in this process to maintain the pH value in the range of 4 to 5, which can contribute to the homogeneous distribution of the metal precursor. The xerogel was then

collected and thermal treated at 800 °C for 10 hours under air to obtain the target material. For the convenience, the sample prepared under this procedure is named as NSG-Na<sub>0.44</sub>MnO<sub>2</sub>. To synthesize Na<sub>0.44</sub>MnO<sub>2</sub> by conventional sol-gel method, a solution was initially prepared by dissolving the stoichiometric amounts of sodium nitrate and manganese acetate in 100 ml distilled water. The mixed precursor was obtained by heating the prepared solution at 120 °C for 8 hours. The post thermal treatment was kept same as the NSG approach, and the sample is named as SG-Na<sub>0.44</sub>MnO<sub>2</sub>.

#### 2.2. Characterization

The crystal structure was characterized by X-ray diffraction (XRD) on a Bruker D8 Advance (Bruker) with Cu K<sub>a</sub> radiation at room temperature. The patterns were recorded in the  $2\theta$  range of 10-70°. Particle morphologies were evaluated using a field-emission scanning electron microscope Auriga) equipped with a focused ion beam (FIB) (FE-SEM, Zeiss). The detailed crystal structure was further analyzed by transmission electron microscopy (TEM, *Zeiss Libra 200 FE*) operating at 200 kV. Selected area electron diffraction (SAED) pattern was recorded by a Gatan CCD camera in a digital format. The electronic conductivity of prepared SG-Na<sub>0.44</sub>MnO<sub>2</sub> and NSG-Na<sub>0.44</sub>MnO<sub>2</sub> samples were investigated by an impedance spectroscopy with Novocontrol AN-alpha analyzer and POT/GAL 20/11 electrochemical test station. Each sample was pressed under the same pressure to obtain a pellet with 1.0 mm in thickness and 8 mm in diameter. Before the conductivity measurement, both sides of pellet were coated with 120 nm thick Au. During the investigation process, an alternating current (frequency range from 100 mHz to 10 MHz) with voltage of 1 V was utilized at room temperature.

#### 2.3. Electrochemical measurements

The slurry for preparing the cathode electrodes was obtained by mixing the active material, conductive carbon (Super C65, Imerys) and binder Poly (vinylidene fluoride) (PVdF, Kynar® FLEX 2801, Arkema Group) in a weight ratio of 80:10:10 in N-methyl-2-pyrrolidone (NMP, Sigma-Aldrich) using a ball milling method. The slurry was coated on an Al foil and dried at 80 °C overnight. After being punched into  $\emptyset$  12 mm disc, the electrodes were further dried for 12 hours under vacuum at 110 °C. The mass loading of prepared electrode is around 2 mg cm<sup>-2</sup>. The electrochemical performance was evaluated in 2032 coin cells, using sodium metal foil as counter electrode. Cathode and sodium metal were separated by glass fiber mats (Whatman GF/D) wetted with 1 M sodium perchlorate (NaCIO<sub>4</sub>, Sigma-Aldrich) in an ethylene carbonate (EC) / propylene carbonate (PC) equivolumic mixture. Galvanostatic cycling was carried out on MACCOR series 4000 battery testers in a voltage range of 2.0– 4.0 V at various current rates. All voltages reported in this work refer to the Na counter electrode. Cyclic Voltammetry measurements were conducted on a VSP electrochemical workstation (Bio-logic), applying a sweep rate of 0.02 mV s<sup>-1</sup>.

#### 2.4. In-situ XRD measurement

*In-situ* XRD analysis on Na<sub>0.44</sub>MnO<sub>2</sub> cathode during Na ion insertion and extraction has been performed using a cell, as being reported in our previous work [27, 28]. The cell body was made of stainless steel covered internally by a Mylar foil for electrical insulation. The electrode slurry was cast directly on a glassy carbon SIGRADUR® window (180 µm, HTW Hochtemperatur Werkstoffe GmbH), which served both as current collector and "transparent window" for the X-ray beam. The coated glassy carbon was subsequently dried at 80 °C for 4

hours and at 40 °C under vacuum overnight. Metallic sodium foil served as counter electrode and Whatman GF/D glass fiber mats served as separators, drenched with 300 µl of electrolyte. The assembled cells were rested for 6 hours before the measurement. A blank measurement of pure PVdF was carried out to confirm that the binder does not show any reflection within the investigated  $2\theta$  range. Subsequently, the cell was charged and discharged by cyclic voltammetry applying a sweep rate of 0.02 mV s<sup>-1</sup> between 2.0 and 4.0 V starting from open circuit voltage (OVC) in the anodic direction (charge). In parallel, XRD analysis was performed, within the  $2\theta$  range of 10 - 90 °, with a step size of 0.01838 ° and a step time of 0.52 s, resulting in a complete individual measurement in 40 minutes, with a rest period of 140 s.

#### 3. Results and discussion

Figure 1a and 1b show the SEM images of the as prepared NSG-Na<sub>0,44</sub>MnO<sub>2</sub> material under different magnifications. The low magnification image (2kx, Figure 1a) shows a rather uniform size distribution of the particles (6 µm in length and 2µm in width), compared with SG-Na<sub>0,44</sub>MnO<sub>2</sub>, as can be seen in Figure 1g. Different from SG-Na<sub>0,44</sub>MnO<sub>2</sub> (Figure 1h), which shows more or less nanorod shape, the high magnification image (50kx, Figure1b) indicates that an individual particle of NSG-Na<sub>0,44</sub>MnO<sub>2</sub> is formed by several nanoplates bundled together. According to FIB image, showed in Figure 1c, NSG-Na<sub>0,44</sub>MnO<sub>2</sub> nanoplates have a specific lamellar with channels distributed between plates. The thin nanoplate (with the thickness of less than 100 nm) provides a large surface area and the channels improve the electrode/electrolyte contact. Both of them can facilitate the (de-) intercalation of Na<sup>+</sup>-ions, enhance the Coulombic efficiency of the material and reduce the electrode polarization. More

interestingly, an inner lamellar structure was also confirmed by TEM (Figure 1d). It can be observed that the slab-like shape particle consists of a series of adherent nanoflakes overlaying each other; these nanoflakes are in the wideness of 5-20 nm, forming very well lamellar structure. The result reveals that further crystallization happens during the heating process. However, no nanowire splits from the formed stable structure, which is different as literature reported [10, 25]. High resolution TEM (HRTEM) images of an individual particle (Figure 1e and 1f) show (200) and (002) planes of the crystalline phase of NSG-Na<sub>0.44</sub>MnO<sub>2</sub>, as being labeled in Figure 1f. The corresponding selected area electron diffraction (SAED) pattern recorded along [010] direction (insert of Figure 1f) also displays (200) and (002) planes of NSG-Na<sub>0.44</sub>MnO<sub>2</sub>. Thus, the crystal grows in parallel with (010) plane, resulting in the formation of nanoplates.

It has been reported that, if the ratio of Na to Mn in the starting material is slightly higher than 0.51, a mixture of orthorhombic Na<sub>0.44</sub>MnO<sub>2</sub> and hexagonal Na<sub>0.7</sub>MnO<sub>2</sub> could be obtained [29]. Therefore, in order to get the pure phase, the ratio of Na and Mn agents needs to be well controlled during synthesis. The XRD patterns, as showed in Figure 2a, confirm that the two materials obtained in this work are well consistent with the orthorhombic Na<sub>0.44</sub>MnO<sub>2</sub> phase (*Pbam* space group, JCPDS No. 27-0750). There is no trace of impurities viewed in the patterns, such as Mn<sub>2</sub>O<sub>3</sub>, which was detected in earlier reports [30, 31], or Na<sub>x</sub>MnO<sub>2</sub> with x value different from 0.44. The ICP-OES determined that the mole ratio of sodium and manganese of NSG-Na<sub>0.44</sub>MnO<sub>2</sub> sample was 0.435:1, within the 2% error of the expected 0.44:1. The crystallinity of NSG-Na<sub>0.44</sub>MnO<sub>2</sub> is much better than that of SG-Na<sub>0.44</sub>MnO<sub>2</sub> due to the relatively narrow and high diffraction peaks. The schematic illustration, in Figure 2b

(right), shows that the crystal structure of  $Na_{0.44}MnO_2$  includes two groups of manganese ion units: sheets of edge-sharing octahedra  $MnO_6$  which consists of all  $Mn^{4+}$  ions and half of the  $Mn^{3+}$ , and columns of square-pyramids  $MnO_5$  built by the rest  $Mn^{3+}$ . Two types of complex tunnels can be built from the different edge-linked chains.  $Na_1$  and  $Na_2$  sites are situated in large S-shaped tunnels, while  $Na_3$  site is located in smaller pentagonal tunnels [32, 33]. The main diffusion pathway for  $Na^+$  ions is along the c-axis ([001] direction), when the material is fully discharged, and all the sodium sites are occupied in  $Na_{0.66}MnO_2$  [22, 34]. The wide tunnel structure may accommodate the stresses associated with the structure change during electrochemical process and benefit the cycling performance. Therefore, the rate capability of NIBs employing  $Na_{0.44}MnO_2$  as cathode materials can be significantly improved by tuning the crystal habit.

Figure 3a illustrates the crystal growth of NSG-Na<sub>0.44</sub>MnO<sub>2</sub> material, which is concluded from TEM and XRD results. The *c* axis, which is also the crystal orientation [001], has been reported as the favorite direction of the crystal growth for Na<sub>0.44</sub>MnO<sub>2</sub> [35-37]. As the result, the nanorod shape for this material is preferred, and relatively lower amount of (001) planes on the particle surface is obtained. It is also proved by SG-Na<sub>0.44</sub>MnO<sub>2</sub> synthesized in this work, and the material crystalizes into rod-shape particles as showed in Figure 1h. Since it is also the direction of large S-shape channel for Na<sup>+</sup> (de)-insertion [26], the long distance of Na<sup>+</sup> diffusion pathway as well as the limited rate performance of the material unfortunately can be expected. Different from the nanorod, the crystal of nanoplate shape grows in parallel with (010) plane, which can not only limit the length of the crystal structure along *c* axis ([001] direction), but also increase the length along *a* axis ([100] direction). Thus, the size of

the particle surface perpendicular to c axis is enlarged and the better rate capability will be achieved. Meanwhile, as being showed in Figure 3b, the relatively thin nanoflake and tunnel among the overlapping nanoflakes could reduce the diffusion distance of Na<sup>+</sup> ions and benefit the electrolyte permeation. With these advantages, a great enhancement of electrochemical performance can be expected from the as prepared NSG-Na<sub>0.44</sub>MnO<sub>2</sub> nanoplates.

The electrochemical behaviors of the NSG-Na<sub>0.44</sub>MnO<sub>2</sub> were first investigated by cyclic voltammetry (CV) with a sweep rate of 0.02 mV s<sup>-1</sup> between 2.0 and 4.0 V. The resulting voltammograms are shown in Figure 4a. The first cycle exhibits at least six pairs of symmetrical redox peaks, which represent the different intercalation/deintercalation steps of Na<sup>+</sup> ions into/from NSG-Na<sub>0.44</sub>MnO<sub>2</sub>, presented by different colors of the first CV plot (Figure 4a). The redox peaks were attributed to consecutive two-phase reaction mechanism [10, 22, 38], but the highly reproduced shape and the very small voltage gap between each redox couple imply the fast diffusion of Na<sup>+</sup> ions during the electrochemical measurement. Nevertheless, a weak oxidation peak (when the voltage is higher than 3.6V) in the initial anodic scan has no corresponding reductive peak, which may be attributed to the electrolyte decompositions. Except this, the shape of the oxidation/reduction peaks after the first cycle remains unchanged, indicating an excellent reversibility of Na insertion into NSG-Na<sub>0.44</sub>MnO<sub>2</sub>.

In order to investigate the initial electrochemical reaction of NSG-Na<sub>0.44</sub>MnO<sub>2</sub>, *in-situ* XRD analysis coupled with electrochemical process has been performed using an *in-house* designed cell. The XRD patterns obtained during the first oxidation/reduction process of the Na/Na<sub>0.44</sub>MnO<sub>2</sub> cell in the restricted 20 degree region from 32-40° are displayed in Figure 4b.

The diffraction patterns are structurally close associated with the voltage plateaus from the beginning of Na extraction process, and each step is distinguished by different colors in relation to the initial curve of the CV measurement (Figure 4a). During the first charge process, most of the diffraction peaks of NSG-Na<sub>0.44</sub>MnO<sub>2</sub> shift to higher  $2\theta$  angle, indicating that the extraction of Na<sup>+</sup> ions from the structure would lead to the decrease of the lattice parameters. The peak intensity changes due to the two-phase reaction. During the following discharge process, the diffraction peaks shift back in a very symmetrical way. The peaks of the same voltage, for both charge and discharge, locate in the same  $2\theta$  position with little change in intensity. It is interesting to see that the peaks for the planes which are perpendicular, such as (0 10 0), and are nearly perpendicular, such as (1 11 0), to b axis, show obviously bigger shift than the others. This means that the expansion and contraction of the lattice during the Na extraction / insertion are along the b axis. All these appearances indicate a highly reversible process of the extraction / insertion of Na<sup>+</sup> ions. Based on the *in-situ* XRD results, the charge/discharge process does not lead to any new phases, and NSG-Na<sub>0.44</sub>MnO<sub>2</sub> cathode material has excellent reversibility, which proves again the conclusions from the CV measurement.

Considering the detrimental impact of the specific particle morphologies on the high rate discharge and capacity retention, we have investigated the rate and long-term cycling performances of the synthesized material. The discharge rate capability of the two  $Na_{0.44}MnO_2$  cathodes is shown in Figure 5a. The cells were cycled at various charge and discharge rates ranging from 0.1 to 10 C (1 C is defined as 120 mA g<sup>-1</sup>) in a voltage range of 2.0 - 4.0 V. It can be observed that NSG-Na<sub>0.44</sub>MnO<sub>2</sub> exhibits much better rate performance than

SG-Na<sub>0.44</sub>MnO<sub>2</sub>, especially at high rate. Specially, NSG-Na<sub>0.44</sub>MnO<sub>2</sub> delivers a discharge capacity of 96 mAh g<sup>-1</sup> at current density 10 C, which is actually 86 % of the capacity (112 mAh g<sup>-1</sup>) at 0.1 C. Strikingly, a capacity of about 112 mAh g<sup>-1</sup>, same as the initial capacity, can be recovered when the current density returns to 0.1 C after the high rates test. These results suggest that NSG-Na<sub>0.44</sub>MnO<sub>2</sub> electrode can fully afford rapid charge and discharge. Generally, Na<sup>+</sup> ions insertion into and extraction from oxide hosts are considered with poor kinetics, compared with their Li-ion analogues due to the much larger size of Na<sup>+</sup> ions than Li<sup>+</sup> ions and 1D diffusion pathway of Na<sup>+</sup> ions for the orthorhombic Na<sub>0.44</sub>MnO<sub>2</sub>, but this is not the case of the material synthesized in this work. The good rate capability of the NSG-Na<sub>0.44</sub>MnO<sub>2</sub> electrode was also confirmed by electronic conductivity measurement. The bulk electronic conductivity of NSG-  $Na_{0.44}MnO_2$  sample is  $7.7 \times 10^{-5}$  S cm<sup>-1</sup> which has improved around one or two orders of magnitude comparing to the result of SG-Na<sub>0.44</sub>MnO<sub>2</sub> sample  $(5 \times 10^{-6} \text{ S cm}^{-1})$ . In summary, the relatively high rate capability is related to the pure phase, stabilized structure, improved electronic conductivity, well controlled morphology with limited crystal growth along the [001] direction and less polarization of the Na<sub>044</sub>MnO<sub>2</sub> material obtained by the NSG method, as indicated by structure, morphology characterizations, conductivity measurement and CV analysis. The cycling stability of the two Na<sub>0.44</sub>MnO<sub>2</sub> cathodes are tested under the current rate of 0.5 C, and the results are showed in Figure 5b. Superior to SG-Na<sub>0.44</sub>MnO<sub>2</sub>, which maintains 78.9 % initial capacity after 100 cycles, NSG-Na<sub>0.44</sub>MnO<sub>2</sub> presents excellent capacity retention, with no more than 2.2 % capacity loss after 100 cycles (from 108 mAh g<sup>-1</sup> in the first cycle decreases to 105 mAh g<sup>-1</sup> in the 100<sup>th</sup> cycle), almost unnoticeable for the plot. The slight 3 mAh g<sup>-1</sup> capacity decrease may

be ascribed to the electrolyte degradation, or the strain caused by Jahn-Teller distortion and the structure degradation upon cycling [38]. Further study is needed to fully understand these subtle yet critical phenomena. Nevertheless, it is obvious that the sodium (de)-intercalation in NSG-Na<sub>0.44</sub>MnO<sub>2</sub> shows much better reversibility than SG-Na<sub>0.44</sub>MnO<sub>2</sub>. The better crystallinity and homogenous particle morphology could be responsible for that.

The initial different discharge profiles of the NSG-Na<sub>0.44</sub>MnO<sub>2</sub> electrodes were shown in Figure 5c. The data were obtained by testing the electrodes at the same charge current density of 0.2 C, but discharged at 0.2, 2 and 5 C, respectively. It delivers reversible discharge capacities of 112, 111 and 110 mAh g<sup>-1</sup>, respectively. Agree with the CV plots, six voltage plateaus can be viewed. With increasing the discharge current density, the material still retains high capacities, although the discharge voltage profiles slightly shift to lower voltage. This behavior can affect the energy density of the NIBs using Na<sub>0.44</sub>MnO<sub>2</sub> as active material. However, no obvious voltage decay is found when the current density changes from 2 to 5 C. Compared the rate capability of NSG-Na<sub>0.44</sub>MnO<sub>2</sub> (based on Figure 5a) with the literature reported Na<sub>0.44</sub>MnO<sub>2</sub> materials (Figure 5d) synthesized by solid state reactions (SS), sol-gel routes (SG), polymer-pyrolysis (PP) methods [10, 22, 25, 29, 39-42], our results are evidently encouraging. To the best of our knowledge, this material exhibits by far the best performance in the case of high rate capability and excellent cycling stability for Na<sup>+</sup>-ions storage. The high discharge capacity and power density, significant high rate capability and the low cost of the Na<sub>0.44</sub>MnO<sub>2</sub> nanoplate make it practically applicable in large power devices and EVs.

#### 4. Conclusions

In summary, the pure monocrystal Na<sub>0.44</sub>MnO<sub>2</sub> nanoplate was synthesized by the template-assisted sol-gel method and applied as cathodes for sodium ion batteries. The material shows high crystallinity and homogeneous nanoplate morphology. The lamellar structure Na<sub>0.44</sub>MnO<sub>2</sub> material, owning to well distributed channels and controlled morphology with limited crystal growth along [001] direction, can provide a mechanically stable construction corresponded with short diffusion path for Na<sup>+</sup> ions insertion/extraction. Electrochemical performance revealed the obtained Na<sub>0.44</sub>MnO<sub>2</sub> nanoplates exhibited excellent reversibility, outstanding high rate capability (96 mAh g<sup>-1</sup> at 10 C) and remarkable cycling stability (less than 2.2 % loss in 100 cycles at 0.5 C), which are much better than other previously reported Na<sub>0.44</sub>MnO<sub>2</sub> nanoplates represent a promising cathode material for high power NIBs.

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#### **Figure captions**

**Fig. 1** SEM micrographs at the magnifications of (a) 2kx, (b) 50kx and (c) FIB-SEM micrograph for the prepared NSG-Na<sub>0.44</sub>MnO<sub>2</sub> material. (d) TEM image of an individual NSG-Na<sub>0.44</sub>MnO<sub>2</sub> particle. (e) HRTEM image and (f) SAED pattern of a monocrystalline Na<sub>0.44</sub>MnO<sub>2</sub> nanoplate. SEM micrographs of the SG-Na<sub>0.44</sub>MnO<sub>2</sub> material under the magnifications of (g) 2kx and (h) 50kx.

**Fig. 2** XRD patterns of SG-Na<sub>0.44</sub>MnO<sub>2</sub> and NSG-Na<sub>0.44</sub>MnO<sub>2</sub> samples (a) and corresponding structure schematic illustration (b) of the orthorhombic  $Na_{0.44}MnO_2$  in the *Pbam* space group (view perpendicular to the ab plane).

**Fig.3** Schematic illustration for crystal growth of nanorod and nanoplate (a); the structural formation of nanoplate particles (b).

**Fig. 4** Cyclic voltammogramms of (a) NSG-Na<sub>0.44</sub>MnO<sub>2</sub> electrode between 2.0 and 4.0 V at a potential sweep rate of 0.1 mV s<sup>-1</sup>. *In situ* evolution of the XRD pattern (b) recorded at 0.02 mV s<sup>-1</sup> voltage sweep rate of NSG-Na<sub>0.44</sub>MnO<sub>2</sub> between 2.0 and 4.0 V.

**Fig. 5** (a) Rate capabilities of the as-prepared SG-Na<sub>0.44</sub>MnO<sub>2</sub> and NSG-Na<sub>0.44</sub>MnO<sub>2</sub> nanoplates. (b) Cycling behaviors of the SG-Na<sub>0.44</sub>MnO<sub>2</sub> and NSG-Na<sub>0.44</sub>MnO<sub>2</sub> nanoplates at 0.5 C for 100 cycles. (c) The discharge profiles of NSG-Na<sub>0.44</sub>MnO<sub>2</sub> at 0.2, 2 and 5 C (The charge current rate is fixed at 0.2 C). (d) Comparison in the rate performance of the NSG-Na<sub>0.44</sub>MnO<sub>2</sub> with the literature reports for the same materials prepared by solid state reactions (SS) [22, 41], sol-gel routes (SG) [29, 42] and polymer-pyrolysis (PP) method [10].



Fig. 1



Fig. 2



Fig. 3



Fig. 4





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#### Highlights

- A template assisted sol-gel method was used to prepare Na<sub>0.44</sub>MnO<sub>2</sub> nanoplates for sodium ion batteries.
- The materials showed high purity, homogeneous size distribution and excellent electrochemical performance, especially the rate capability.
- The as-prepared nanoplates morphology can limit the crystal growth along [001] direction,

can provide a mechanically stable construction corresponded with short diffusion path for

Na<sup>+</sup> ions insertion/extraction.

Graphical abstract

The  $Na_{0.44}MnO_2$  materials were achieved with a template assisted sol-gel method. The prepared materials showed high crystallinity, pure phase and homogeneous size distribution. The materials exhibited superior electrochemical performance in both rate capability and cycling stability due to the limited crystal growth along [001] direction.

