High performance carbon-coated hollow Ni12P5 nanocrystals decorated on GNS as advanced anodes for lithium and sodium storage

Huinan Guo  
*Nankai University*

Chengcheng Chen  
*Nankai University*

Kai Chen  
*Nankai University*

Haichao Cai  
*Nankai University*

Xiaoya Chang  
*Nankai University*

*See next page for additional authors*
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Abstract
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Huinan Guo, Chengcheng Chen, Kai Chen, Haichao Cai, Xiaoya Chang, Song Liu, Weiqin Li, Yijing Wang, and Caiyun Wang

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Transition-metal phosphides have been considered as promising anode materials for rechargeable secondary batteries owing to their low cost and high capacity. However, the low electron conductivity and poor stability limit their further development. Herein, we have designed a template-free refluxing method for synthesizing the tailored carbon-coated hollow Ni$_{12}$P$_5$ nanocrystals in situ grown on reduced graphene oxide nanosheets (noted as Ni$_{12}$P$_5$@C/GNS). The hollow structure can accommodate the volume expansion and shorten the ion transfer path. GNS loading and carbon shell can efficiently prevent Ni$_{12}$P$_5$ from aggregating and improve the electronic conductivity. As an anode of Li-ion batteries (LIBs), hollow Ni$_{12}$P$_5$@C/GNS composite displays an excellent discharge specific capacity of 900 mAh g$^{-1}$ at a current density of 100 mA g$^{-1}$ after 100 cycles and outstanding rate capability. Meanwhile, it also shows a good Na storage capability with reversible capacity of 235 mAh g$^{-1}$ at 100 mA g$^{-1}$. Therefore, our work demonstrates this hollow Ni$_{12}$P$_5$@C/GNS composite will be a great potential material for Li/Na storage.

1. Introduction

Rechargeable secondary batteries are one of the most important energy storage technologies recently. LIBs have occupied dominant position of portable electronic devices for their high operating voltage, high energy density and limited self-discharging. Besides, sodium-ion batteries (SIBs) are expected to be broadly applied to large energy storage equipment owing to low cost and natural abundance of sodium sources. However, the commercial anode materials hardly show satisfied performances in both Li/Na storage systems, due to the worse sluggish kinetics of sodium than that of lithium. Therefore, it is urgent to search for advanced anode materials in terms of high capacity, long cycle life and superior rate capability for both LIBs and SIBs.

Transition-metal phosphides (TMPs) have been applied to several fields, especially in electrocatalysis and photocatalysis. Nevertheless, their applications as anode materials are not fully developed. Conversion type TMPs are rising anode materials for rechargeable secondary batteries in recent years. As anode materials for lithium/sodium ion batteries, based on the reversible reaction between phosphorus and Li/Na (Li$_3$P/Na$_3$P), TMPs display high specific capacity (up to ~1800 mAh g$^{-1}$), low operation potential, metallic feature as well as good thermal stability. However, the large volume expansion and poor electronic conductivity of TMPs seriously limit the kinetics property and cycling stability. Facing those similar defects, Lou's group prepared the tailored hollow structure aiming to buffer the substantial volume change and boost the electrochemical performances. This demonstrates the hollow structure can significantly accommodate pulverization by restricting volume expansion. Additionally, Mai and co-workers synthesized novel layered Li$_3$V$_2$(PO$_4$)$_3$ in which the LVP layers were uniformly alternated with reduced graphene oxide nanosheets to provide effective electron transport. The rate capacities of prepared Li$_3$V$_2$(PO$_4$)$_3$/rGO&C are better than most of the state-of-the-art reported results. Thus, it is interesting and worth designing the hollow nanocrystal, and combining with carbon materials to improve the electrochemical performances of Ni$_{12}$P$_5$.

On the other hand, the synthesis methods of TMPs are usually defective. Generally speaking, there are two primary routes to prepare TMPs. High energy mechanical milling is simple, however, the products of this method are always without hollow morphology. Another way is a solid reaction using hypophosphites (NH$_4$H$_2$PO$_2$ and NaH$_2$PO$_2$) as the P source. This process releases very flammable and poisonous PH$_3$ gas. Hence, designing a facile and low toxic method for synthesis of TMPs is not only urgent, but still remains challenging.
Herein, we firstly report a novel hollow structure of carbon-coated Ni$_3$P$_3$ nanocrystals supported on reduced graphene oxide nanosheets via a simple refluxing method without any templates and surfactants, as high performance anodes for LIBs and SIBs. This pathway is facile and effective. During the synthesis, the phosphorus source is consumed in situ without hazardous gases escape. Meanwhile, the Ni$_3$P$_3$@C/GNS has endowed with three merits of targeted samples: (i) The hollow Ni$_3$P$_3$@C with average small size of 35 nm effectively shortens ions/electrons transport paths and facilitates the reaction kinetics; (ii) Hollow structure and carbon shell can buffer volume expansion wonderfully; (iii) The GNS prevent aggregation and provide an electric net between grains. As expected, when measured as an anode material for LIBs, the tailored Ni$_3$P$_3$@C/GNS electrode shows a high reversible capacity of 900 mAh g$^{-1}$ at a current density of 100 mA g$^{-1}$ after 100 cycles, and excellent rate capability of discharge special capacities of 702.2, 644.1, 566.5, 482.8, 423.2 and 905.9 mAh g$^{-1}$ at 100, 200, 500, 1000, 2000 and 100 mA g$^{-1}$, respectively. We also adopt this material as an anode material for the first time, which delivers excellent performance with reversible capacity of 235 mAh g$^{-1}$ at 100 mA g$^{-1}$. Therefore, Ni$_3$P$_3$@C/GNS composite is a promising anode material with high capacities and good rate capacities for LIBs and SIBs.

2. Experimental Section

2.1 Chemicals
Nickel (II) acetate tetrahydrate (AR) was purchased from Tianjin Guangfu Fine Chemical Research Institute and triphenylphosphine (TPP) was from Alfa Aesar. Oleylamine (OLA) was purchased from J&K. All reagents were used as received without further purification.

2.2 Synthesis of reduced graphene oxide
Graphene oxide was prepared following Hummers’ approach, then reduced by H$_2$ (5 %)/Ar gas.

2.3 Synthesis of Ni$_3$P$_3$/GNS
The Ni$_3$P$_3$ was synthesized using a facile refluxing condensation method. Briefly, 1 mmol Ni(AC)$_2$-4H$_2$O (0.5 g) was dispersed in 30 mmol OLA (8 g) in a 100 mL three-neck flask. The flask was heated by a heating panel magnetically stirred under Ar flow. The mixture solution was heated to 120 °C until all Ni(AC)$_2$-4H$_2$O was dissolved and the solution turned into green. Then 10 mmol TPP (2.6 g) and a certain amount of GNS was added into the solution. The growth solution was heating to 270 °C and kept at this temperature for 30 min. Then the solution was naturally cooled to ambient temperature. The black precipitate was isolated and washed three times with the mixture of hexane and ethanol by centrifugation. Finally, the products were dried at 80 °C under vacuum for further characterization.

2.4 Synthesis of Ni$_3$P$_3$@C/GNS
In a typical transformation process, the as-prepared Ni$_3$P$_3$/GNS was annealed at 450 °C in a H$_2$ (5 %)/Ar atmosphere for 30 min.

2.5 Characterization
The products were characterized by X-ray diffractometry using a Rigaku MiniFlexII diffractometer equipped with the Cu Ka radiation ($\lambda$=1.54178 Å) at a scan rate of 4° min$^{-1}$. Raman measurement was recorded on an Invia/Reflex Lasser Micro-Raman spectroscope (Horiba Jobin Yvon, Franch). The XPS spectra were collected by a VG Scientic ESCALAB 2201XL System equipped with a monochromatic Al Ka source. The weight percentage of carbon was characterized by elemental analysis (EA, Thermal Conductivity Detector, Elemental vario E cube). Inductively coupled plasma-atomic emission spectroscopy (ICP-AES) was performed on a PerkinElmer Optima 8300 instrument. Field-emission scanning electron microscopy (SEM) measurements were performed on a JOEL JSM-7500F (5 kV). Transmission electron microscopy (TEM) studies were conducted on a FEI Philips Tecnai electron microscope at an operating voltage of 200 kV.

2.6 Electrochemical measurements.
Half cells were carried out in standard CR2032 coin-type model cells, with Li/Na foil as the counter and reference electrode. The working electrode was prepared by mixing Ni$_3$P$_3$@C/GNS powder, super P and polyvinylidene fluoride (PVDF) with a weight ratio of 7:2:1. Then the mixture was ground in a mortar with N-methyl-2-pyrrolidone (NMP) as solvent for making slurry and pasted on pure copper foil (99 %). The average mass loading of the active materials with the film was about 0.98 mg cm$^{-2}$. For lithium ion battery, the electrolyte was 1M LiPF$_6$ in ethylene carbonate (EC)/diethyl carbonate (DEC) (1:1 by volume ratio). The polypropylene film (Celgard 2400) was used as the separator. For sodium ion battery, the electrolyte was 0.1M NaN$_3$ in diethylene glycol dimethyl ether (DEG-DME) with addition of 5 % fluoroethylene carbonate (FEC). The glass fiber was used as the separator. The battery tests were assembled in an Ar-filled glovebox (H$_2$O, O$_2$< 1 ppm, Mikkouna). Cyclic performances were recorded on a CHI660E electrochemical station at a scan rate of 0.2 mV s$^{-1}$ from 0.1 to 3 V. Galvanostatic charging and discharging tests were performed using a LANDCT2001A instrument.

The electrochemical performances of full cells were also measured using a two electrode coin-type cell (CR2032). To prepare the positive electrode, a mixture of LiCoO$_2$, Super P and PVDF at a weight ratio of 7:2:1 were dispersed in NMP solvent and ground thoroughly into a slurry, then pasted on aluminum foil. After being dried at 80 °C, the aluminum foil with active materials was cut into small rounds for battery cathode. The negative electrode of Ni$_3$P$_3$@C/GNS was prepared by the same method. For assembling full cells, the mass ratio of cathode and anode was 16:1. The excess cathode was aimed to ensure the Ni$_3$P$_3$@C/GNS anode react thoroughly.

3. Results and Discussion

3.1 Structural and morphology characterization
Scheme 1. Schematic illustration showing the formation preparation of Ni$_{12}$P$_5$@C/GNS composite.

We employ GNS as an assistant agent to build hollow Ni$_{12}$P$_5$ nanocrystals through a facile template-free refluxing method (Scheme 1). The XRD patterns confirm the Ni$_{12}$P$_5$@C/GNS, pure Ni$_{12}$P$_5$@C and GNS are pure phase displayed in Fig. 1a and S1†. The products fully correspond to tetragonal unit cell (a=8.464 Å, c=5.070 Å, JCPDS card No. 22-1190) with the I4/m space group. In this structure (Fig. 1b), Ni atoms occupy 16i and 8h sites connected with two P atoms. While there are two kind of P atoms, which reside in the 2a sites connected with eight Ni and 8h sites connected with four Ni, respectively. SEM images (Fig. S2†) clearly show the sheet-like structure of GNS and nanocrystal structure of pure Ni$_{12}$P$_5$@C. Fig. S2b† displays the serious aggregation Ni$_{12}$P$_5$@C without GNS. Fig. 1c exhibits the SEM image of Ni$_{12}$P$_5$@C/GNS composite. Notably, Ni$_{12}$P$_5$@C nanocrystals with uniform size are homogeneously dispersed on the surface of GNS. In addition, compositional information is determined by energy-dispersive spectrum (EDS, Fig. S3†). The atomic ratio of Ni to P is measured to be 2.32:1, in accordance with that of stoichiometric Ni$_{12}$P$_5$. According to N$_2$ adsorption/desorption measurement and the Brunauer–Emmett–Teller (BET) analysis, the existence of mesoporous in the Ni$_{12}$P$_5$@C/GNS is observed and the specific surface area is estimated to be about 78.7 m$^2$ g$^{-1}$ (Fig. 1d), which is higher than pure Ni$_{12}$P$_5$@C of 41.8 m$^2$ g$^{-1}$ (Fig. S4†). Element analysis result shows that the carbon content of Ni$_{12}$P$_5$@C/GNS is 18.15 %. The content of Ni in the Ni$_{12}$P$_5$@C/GNS composite is estimated to be 66.3 wt.% by ICP instrument. Thus, the weight percentage of Ni$_{12}$P$_5$ in the hybrid is about 80.9 %. Raman spectrum is employed to investigate the conductivity of carbon of Ni$_{12}$P$_5$@C/GNS. As shown in Fig. S5†, two strong carbon peaks located at 1346 cm$^{-1}$ (D-band, sp$^3$-coordinated behavior)
Fig. 3 Electrochemical performances for lithium storage. (a) Cyclic voltammograms of Ni$_{12}$P$_5$@C/GNS electrode at 0.2 mV s$^{-1}$. (b) Charge/discharge profiles of Ni$_{12}$P$_5$@C/GNS electrode at 100 mA g$^{-1}$. (c) First charge/discharge profile of Ni$_{12}$P$_5$@C/GNS electrode at 100 mA g$^{-1}$. The marked points (from a to h) are collected to take XRD tests. (d) Ex situ XRD patterns of Ni$_{12}$P$_5$@C/GNS electrode at marked points. (e) XPS patterns of Ni 2p$_{3/2}$ in the 10th cycle, discharge to 0.005 V and charge to 3.0 V at a current density of 100 mA g$^{-1}$. (f-g) SAED patterns in the 10th cycle at a current density of 100 mA g$^{-1}$, discharge to 0.005 V and charge to 3.00 V.

and 1580 cm$^{-1}$ (G-band, sp$^2$-hybridized carbon) can apparently be observed. The Raman peak intensity ratio ($I_D/I_G=0.78$) is inversely proportional to the graphitization degree of materials, indicating a higher electric conductivity.

The detailed morphology and microstructure of pristine Ni$_{12}$P$_5$@C nanocrystals and prepared Ni$_{12}$P$_5$@C/GNS are further characterized by TEM. Low-resolution TEM images show the pure Ni$_{12}$P$_5$@C nanocrystals are aggregated with average diameter of ~19 nm (Fig. 2a, b and S6a†). The carbon layer thickness is ~3.14 nm. High-resolution TEM (HRTEM) image displays the characteristic lattice fringe spaces of 0.185 and 0.301 nm, which are consistent with the (312) and (220) plants of Ni$_{12}$P$_5$ nanocrystal, respectively, demonstrating a high degree of crystallinity (Fig. 2c). Low-magnification TEM images show the hollow structure of carbon-coated product and the particle size distribution histogram displays the ~35 nm size of Ni$_{12}$P$_5$@C/GNS (Fig. 2d, e and S6b†). Typical HRTEM image (Fig.
2f) reveals the Ni$_2$P$_5$@C anchored on GNS surface is fully hollow and the carbon shell is clear. The carbon layer thickness is ~2.64 nm and the thickness of Ni$_2$P$_5$ is ~6.34 nm. The characteristic lattice fringe spaces of 0.193, 0.235 and 0.217 nm are consistent with the (312), (330) and (321) plants of Ni$_2$P$_5$@C nanocrystal, respectively. Meanwhile, the angle between (312) and (330) lattice plane is 53.6° consistent with the calculation value. Additionally, the crystal type of pure Ni$_2$P$_5$@C and Ni$_2$P$_5$@C/GNS both are polycrystal (Fig. S7†). Notably, GNS effectively suppress aggregation of Ni$_2$P$_5$@C and adjust the growth direction of crystals to construct hollow architecture. The formation mechanism of hollow structure can be ascribed to Kirkendall Effect. It’s known that GNS are negatively charged. When dispersed in suspension, the surface functional groups of GNS greatly attract Ni$^{2+}$, which leads to the faster out-diffusion tendency of inner Ni$^{2+}$ and forms hollow.41-43 The carbon shell derives from the carbonation of OLA. Moreover, the element mapping analysis (Fig. 2g) further clearly confirms the uniform distribution of C, Ni and P elements in the Ni$_2$P$_5$@C/GNS.

### 3.2 Electrochemical performances of LIBs

Electrochemical Li-storage performances of hollow Ni$_2$P$_5$@C/GNS are given in Fig. 3. As shown in Fig. 3a, cyclic voltammetry (CV) is adopted to verify the reversible Li$^+$ storage behavior of hollow Ni$_2$P$_5$@C/GNS. The CV curves display weak redox peaks and box-like shape. This is due to the sorption of lithium ions on the pores and functional groups of GNS surface, as well as the sorption on the edge of defective sites. Thus, the electrochemical behaviors are surface capacitive lithium storage behaviors.44-47 In the first cycle, the reduction peaks are located at around 1.5 and 0.7 V, which represent the irreversible structure change, lithiation and formation of solid electrolyte interphase (SEI), respectively. Besides, the oxidation peaks are observed at around 1.0, 1.5 and 2.4 V, which can be severally attributed to the gradual reduction of Li$_3$P. Subsequently, the curves keep stable in the following cycles implying its high reversibility. According to this mechanism, the reaction process could be summed as:

$$\text{Ni}_2\text{P}_5 + x\text{Li}^+ + xe^- \leftrightarrow m\text{Li}_3\text{P} + n\text{Ni} \quad (1)$$

Fig. 4 Electrochemical performances for lithium storage. (a) Cycling performances of Ni$_2$P$_5$@C/GNS with different carbon contents at 100 mA g$^{-1}$ (discharge capacities). (b) Rate performances of Ni$_2$P$_5$@C/GNS electrode with different carbon contents at different rate current densities (discharge capacities).

Fig. 5 Electrochemical performances for lithium storage. (a) Cycling performances of pure Ni$_2$P$_5$@C and Ni$_2$P$_5$@C/GNS at 100 mA g$^{-1}$ (discharge capacities). (b) Rate performances of pure Ni$_2$P$_5$@C and Ni$_2$P$_5$@C/GNS electrodes at different rate current densities (discharge capacities). (c) Long-term cycling performances and coulombic efficiency of the Ni$_2$P$_5$@C/GNS electrode at 2000 mA g$^{-1}$.
When considering the performance of a whole electrode, the proportion between Ni$_{12}$P$_5$@C and GNS can seriously influence the performances. Broadly speaking, with the increasing of GNS, the electroconductivity of Ni$_{12}$P$_5$@C/GNS gradually increases. However, the capacity will reach saturation point and not increase any more with the content of GNS increasing. Therefore, in order to explore the best ratio between the active material Ni$_{12}$P$_5$@C and GNS, we further prepare four Ni$_{12}$P$_5$@C/GNS composites with different carbon content (7.68, 11.13, 18.15 and 34.19 wt.%, respectively). The carbon contents are characterized by EA. Cycling tests and rate capacities are displayed in Fig. 4. It could be observed that the electrochemical performances of Ni$_{12}$P$_5$@C/GNS electrodes with carbon content above 18.15 wt.% are superior to that of 11.13 and 7.68 wt.%. The cycling performances of the Ni$_{12}$P$_5$@C/GNS with 7.68, 11.13, 18.15 and 34.19 wt.% carbon at 100 mA g$^{-1}$ are 517.7, 564.1, 900 and 923 mAh g$^{-1}$ after 100 cycles, respectively (Fig. 4a). Fig. 4b shows the rate capacities of Ni$_{12}$P$_5$@C/GNS with different carbon content. As expected, with the GNS content increasing, the capacities of Ni$_{12}$P$_5$@C/GNS tend to improve, because more GNS offer more ions and electrons transport paths to achieve good performances. However, the capacity of Ni$_{12}$P$_5$@C/GNS no longer infinitely grows with the further increasing of GNS amount. Considering the preparation of GNS is costly and time-consuming, the Ni$_{12}$P$_5$@C/GNS of 18.15 wt.% carbon is optimum.

Electrochemical lithium storage performances of Ni$_{12}$P$_5$@C/GNS are given in Fig. 5. Fig. 5a displays the cycling performances of the pure Ni$_{12}$P$_5$@C and Ni$_{12}$P$_5$@C/GNS at a current density of 100 mA g$^{-1}$. The discharge specific capacity of Ni$_{12}$P$_5$@C/GNS remains at 900 mAh g$^{-1}$ after 100 cycles much better than that of Ni$_{12}$P$_5$@C (420 mAh g$^{-1}$). The TEM images of Ni$_{12}$P$_5$@C/GNS composite after 100 cycles are shown in Fig. 5b, which shows Ni$_{12}$P$_5$@C/GNS can maintain original appearance and no significant aggregation is observed. The batteries show a gradual gain in capacity. According to previous references, the major reasons can be summarized as the follows: (1) The slow increased capacities are driven by the activation effects. (2) The stabilization process of the SEI layer will result in a gradually increasing capacities. (3) Decreasing polarization and resistances are directly connected to the increase of capacities. Then rate capacities are examined by discharging/charging at various current densities (Fig. 5b). Ni$_{12}$P$_5$@C/GNS delivers discharge specific capacities of 702.2, 644.1, 566.5, 482.8, 423.2 and 905.9 mAh g$^{-1}$ at 100, 200, 500, 1000, 2000 and 100 mA g$^{-1}$, respectively, which are much higher than that of pure Ni$_{12}$P$_5$@C. Moreover, Ni$_{12}$P$_5$@C/GNS can be cycled with high stability at higher current density of 500 mA g$^{-1}$ (Fig. S9†) with discharge capacity keeping 308.9 mAh g$^{-1}$ after 250 cycles and 2 A g$^{-1}$ (Fig. 5c) with the discharge capacity reaching up to 237.3 mAh g$^{-1}$ after 200 cycles. It firmly verifies their excellent cycling stability and long service life. Table 1 shows the electrochemical performances of the present work for LIBs anode are higher than most of the related reported in seven years. 50-59

Fig. 5 Electrochemical performances for lithium storage. (a) Nyquist plots of the pure Ni$_{12}$P$_5$@C and Ni$_{12}$P$_5$@C/GNS half cells. (b) Real parts of the impedance (Z”) versus the reciprocal square root of the lower angular frequency (ω) of the pure Ni$_{12}$P$_5$@C and Ni$_{12}$P$_5$@C/GNS. (c) Schematic diagram of electron/ ion transportation in the special Ni$_{12}$P$_5$@C/GNS composite vs. pure Ni$_{12}$P$_5$@C nanocrystals.

Electrochemical impedance spectroscopy (Fig. 6a) demonstrates that Ni$_{12}$P$_5$@C/GNS shows a lower charge transfer resistance (279.3 Ω, fitted by Zview) than pure Ni$_{12}$P$_5$@C (1021.0 Ω). The ion diffusion capabilities of electrodes are estimated according to the slope of the lines between Z’ and ω$^{-1/2}$ (α, the value of which represents Warburg factor, Fig. 6b). Because α$^2$ is in inverse ratio to the diffusion coefficient of Na$^+$ ions (D$_{Na^+}$), it indicates that the D$_{Na^+}$ of Ni$_{12}$P$_5$@C/GNS is higher than that of pure Ni$_{12}$P$_5$@C. These results declare the cathode and Ni$_{12}$P$_5$@C/GNS anode at 100 mA g$^{-1}$ (inset is digital picture of a full cell that lights LEDs). Ni$_{12}$P$_5$@C/GNS electrode performs superior electronic/ionic conductivity, which could be reasonably attributed to two merits (Fig. 6c). (1) Hollow nanocrystal structure shortens ion transport path and provides ample room to accommodate the volume expansion, so that the cycling performance is improved drastically. (2) Carbon shell and adding of GNS

Fig. 6 Electrochemical performances for lithium storage. (a) Charge/discharge profiles and (b) Cycling performance of the full cell coupled by LiCoO$_2$. 6
effectively prevent the agglomeration between nanocrystals and improve the electronic conductivity of Ni$_{12}$P$_5$@C accelerating the transmission of electron, which greatly elevate the rate capacities. Thus, the designed Ni$_{12}$P$_5$@C/GNS exhibits excellent cycling performance and outstanding rate capability, indicating its overwhelming superiority in applications. We also further test the full battery cycling performances with prepared Ni$_{12}$P$_5$@C/GNS as anode and commercial LiCoO$_2$ as cathode. Fig. 7a displays the galvanostatic charge/discharge curves of the full LIBs at 100 mA g$^{-1}$. The reversible specific capacity (based on the anode material) can reach 350 mAh g$^{-1}$, and the average working voltage is about 2.3 V. What’s more, the curves overlap well in the process of 50 to 100 cycles, which reveals the high reversibility and stability of the LiCoO$_2$/Ni$_{12}$P$_5$@C/GNS full cell. Fig. 7b shows the stable capacity of 315.1 mAh g$^{-1}$ can be achieved at the current density of 100 mA g$^{-1}$ after 50 cycles. To demonstrate its potential application, the full cell is used to light up red LED (Fig. 7b, inset).

### 3.3 Electrochemical performances of SIBs

As mentioned before, the Ni$_{12}$P$_5$@C/GNS electrode will also show promising electrochemical performances for sodium-ion storage in half cells. Fig. 8a shows the charge and discharge capacities are 212.7 and 234.9 mAh g$^{-1}$ of 5 th cycle at 100 mA g$^{-1}$, respectively. Remarkably, the Ni$_{12}$P$_5$@C/GNS electrode shows a stable cycling performance (Fig. 8b). After 500 cycles, the discharge capacity remains 164.8 mAh g$^{-1}$. Fig. 8c exhibits the rate performances of the Ni$_{12}$P$_5$@C/GNS electrode. The reversible capacities are 171.6, 162.4, 144.4, 126.8, 110 and 105.6 mAh g$^{-1}$ for current densities of 50, 100, 200, 500, 1000 and 2000 mA g$^{-1}$, respectively. More importantly, the capacity can restore 199.2 mAh g$^{-1}$ when the current density returns to 100 mA g$^{-1}$, implying their high current adaptation. Since Na$^+$ (1.02 Å) is heavier and larger than that of Li$^+$ (0.76 Å), leading to the inferior reaction dynamics of Na$^+$. This significantly influences the electrochemical performances of Ni$_{12}$P$_5$@C/GNS electrode in Na$^+$ storage resulting in the lower capacity and

#### Table 1 Comparison of cycling performances of some TMPs materials for LIBs anodes.

<table>
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<tr>
<th>Material description</th>
<th>Preparation strategy</th>
<th>Current density (mA g$^{-1}$)</th>
<th>Cycle number</th>
<th>Discharge capacity (mAh g$^{-1}$)</th>
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poor stability. Therefore, further studies should be developed to improve its sodium storage properties with some novel strategies.

Conclusions

In summary, the hollow Ni$_3$P$_2$@C/GNS composite is firstly prepared by a simple refluxing method. The Ni$_3$P$_2$@C nanocrystals with particularly hollow structure and thin carbon shell are anchored on the GNS. This architecture combines the advantages of hollow structure and high electric GNS microstructure revealing excellent electrochemical performances at LIBs and SIBs in terms of cycle stability and rate capability. As an anode for LIBs, it displays an excellent special capacity of 900 mAh g$^{-1}$ at a current density of 100 mA g$^{-1}$ after 100 cycles, and high rate capability of special capabilities of 702.2, 644.1, 566.5, 482.8, 423.2 and 905.9 mAh g$^{-1}$ at 100, 200, 500, 1000, 2000 and 100 mA g$^{-1}$, respectively. As for Na-ion storage, this hollow Ni$_3$P$_2$@C/GNS composite also maintains a reversible capacity of approximate 235 mAh g$^{-1}$ at 100 mA g$^{-1}$. These promising features make hollow Ni$_3$P$_2$@C/GNS ideal candidate for the next generation of battery material and can be extended to electrochemical applications in various fields, such as fuel cells, super capacitors, electrochemical sensors, and so forth.

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Notes and references


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