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A review of phosphorus and phosphides as anode materials for advanced sodium-ion batteries

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Sodium-ion batteries (SIBs) are promising low-cost alternatives to lithium-ion batteries (LIBs) in energy storage applications because of the natural abundance of sodium as compared with lithium. However, the radius of Na⁺ ions is ~50% larger than that of Li⁺ ions, leading to challenging sodiation/desodiation at the anodes of SIBs and reduced performances. Therefore, the exploration of novel SIB anode materials to facilitate sodiation/ desodiation has become a leading area of research. Here, cost-effective phosphorus and metal/nonmetal phosphides, possessing high theoretical gravimetric capacities and volumetric specific capacities, have emerged as new and promising anode materials for SIBs. Based on this, this review will provide a comprehensive summary of the recent progress in the development of phosphorus and metal/nonmetal phosphide materials and corresponding composite SIB anodes in terms of material synthesis, characterization, sodium storage mechanism and performance validation in SIB apparatuses. In addition, challenges associated with the development of such advanced materials are summarized and analyzed. Finally, to facilitate the research and development of these emerging SIB anode materials, future research directions are proposed to overcome challenges toward the practical commercialization of sodium-ion batteries.

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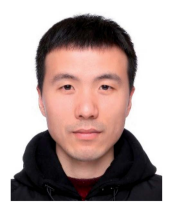
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1 Introduction

Global demand for advanced energy storage and conversion technologies, especially electrochemical energy technologies such as batteries, is rapidly growing and among various types of batteries, Li-ion batteries (LIB) have shown success in many applications.^{1,2} However, although LIBs can meet the tough demands of energy storage and conversion applications, the limited abundance of lithium (only 20 ppm) needs to be



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Review

considered in terms of sustainability.3 Therefore, the exploration of alternatives to replace lithium ion batteries is necessary. Fortunately, sodium has been identified as a suitable alternative to lithium as demonstrated by the practical feasibility of sodium-ion batteries (SIBs).⁴ Fig. 1a shows the comparison between lithium and sodium in terms of their atomic masses, electrode potentials, cationic radiuses, elemental abundances, and prices. It can be seen that sodium is the second-lightest and smallest alkali metal next to lithium and is one of the most abundant elements in Earth's crust with an average abundance of 28 400 ppm.⁵ And in battery applications, sodium can provide a suitable electrochemical redox potential of -2.71 V vs. the standard hydrogen electrode (SHE), which is only 0.3 V less negative than that of lithium. Furthermore, sodium is much cheaper than lithium. Due to these promising properties, SIBs are promising alternatives to LIBs, in particular for large-scale applications such as energy storage systems for power grids.⁵

However, the ionic radius of Na is larger than that of Li (1.02 Å for Na⁺ vs. 0.76 Å for Li⁺), which results in sluggish reaction kinetics and poor electrochemical performances in corresponding batteries. To address this, great efforts have been undertaken and in recent years, significant progress has been made for the cathodes of SIBs with a variety of materials being demonstrated as suitable cathodes for sodium ion storage.6-11 Fig. 2b summarizes the main difference between lithium-ion batteries (LIBs) and sodium-ion batteries (SIBs) in terms of their cost, power density, energy density and overcharge resistibility. In general, the exploration of low-cost anodes for both LIBs and SIBs to match their corresponding cathodes as well as the corresponding synthesis strategies remain challenging. Similar to LIBs, carbonaceous materials such as hard carbon,¹²⁻¹⁹ graphitic carbon,²⁰⁻²² and graphene²³⁻²⁸ along with novel materials such as metals,²⁹⁻³⁵ metal oxides,³⁶⁻⁴¹ metal sulfides,⁴²⁻⁴⁸ metal selenides,⁴⁹⁻⁵⁴ metal alloys55-64 and organic compounds65-69 have all been explored as

anode materials for SIBs. Most recently, phosphorus (P) and phosphides70-78 have also been extensively explored for Na storage. Among these P-based materials, red phosphorus (RP) was the first to attract attention from researchers due to favorable properties as anode materials in SIBs, including a highest theoretical specific capacity of ~ 2600 mA h g⁻¹ and a suitable insertion voltage of \sim 0.4 V vs. Na/Na⁺.⁷⁹ The major advantages of SIBs are summarized in Fig. 1b. Particularly, RP is abundant, easily available and nontoxic, being identified as the most promising anode candidate material for practical SIBs (Fig. 1c). However, RP also possesses drawbacks such as low electrical conductivity $(\sim 10^{-14} \text{ S cm}^{-1})^{80}$ and large volume expansion after full sodiation (300-500%),81-83 leading to shortened cycle lifespans and poor rate performances in sodium storage applications; in addition, the sodiated product Na₃P also shows a low electrical conductivity and the surface of Na₃P is highly reactive which may cause a severe electrolyte decomposition.⁸⁴ To overcome these issues, great efforts have been made and considerable progress has been achieved. Many composites such as amorphous RP/carbon,^{79,82} RP/carbon nanotubes,⁸⁵ RP/Ni-P⁸⁶ and hollow RP nanospheres⁸⁷ have been prepared to improve the Na storage performance of RP anodes, which could buffer the volume expansion of RP during cycles and/or enhance the conductivity of electrodes. In addition to RP, black phosphorus (BP) is also gaining prominence due to a similarly high theoretical specific capacity to RP (\sim 2600 mA h g⁻¹) and a higher electronic conductivity ($\sim 300 \text{ S} \text{ m}^{-1}$).⁸⁸ However, large volume expansion, sluggish reaction kinetics and rigorous synthesis conditions also hinder the application of BP in Na storage devices. Phosphorene is one or few layers of BP and has also attracted much attention as a promising anode material due to its unique 2D structure (similar to a graphene).89 Furthermore, metal phosphides, which usually possess higher electrical conductivities than phosphorus,77 can easily be synthesized into



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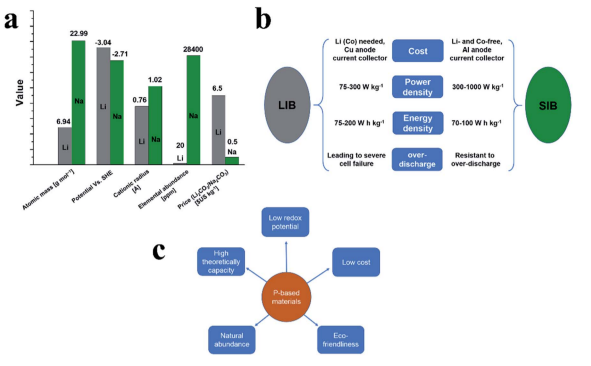


Fig. 1 (a) Major property differences between lithium and sodium; (b) main differences between lithium-ion batteries (LIBs) and sodium-ion batteries (SIBs); (c) main advantages of phosphorus/phosphides as anode materials for SIBs.

various nanostructures and are also promising anode materials for SIBs. Some unique metal phosphide nanostructures such as CoP₄ nanoarrays,⁹⁰ yolk–shell Sn₄P₃@C nanospheres⁹¹ and monodisperse Ni₂P nanocrystals⁹² have been prepared and showed excellent Na-ion storage performance as anode materials. Finally, non-metal phosphides and ternary phosphides as supplemental materials to metal phosphides have also experienced great progress for practical application in SIBs. Because of all these advancements, this review will comprehensively summarize the recent progress in the research and development of phosphorus and phosphide composites for SIB anodes. It covers the structural properties and sodium storage mechanisms of these composites as well as the design strategies. In addition, the advantages and limitations of current synthesis methods for the preparation of phosphorus/phosphide anodes are also discussed. And to facilitate future research efforts,

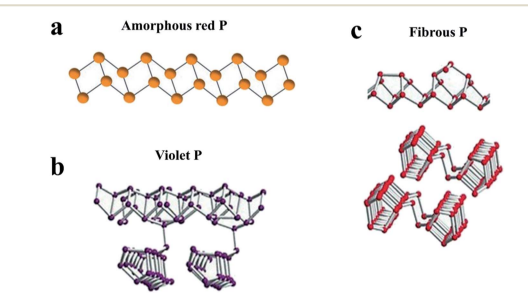


Fig. 2 Characteristic structural fragments of (a) amorphous RP; (b) violet P (Hittorf's P); (c) fibrous P. (a) Reproduced with permission.¹⁰⁰ Copyright 2019, WILEY-VCH. (b, c) Reproduced with permission.¹¹³ Copyright 2017, Wiley-VCH.

associated challenges are also presented and analyzed as well as possible research directions to address these challenges.

2 Phosphorus anodes for SIBs

In general, elemental phosphorus exists in three allotropes, namely, red, black and white phosphorus. White phosphorus is

flammable and toxic, and not suitable for SIBs, whereas RP and BP can react with sodium to form Na₃P and present high theoretical capacities of ~2600 mA h g⁻¹ as electrochemical sodium storage materials.⁹³ Compared with BP, which is difficult to synthesize, RP is more commercially available, and thus is regarded as the most promising candidate for SIB anode materials. In addition, phosphorene (one or several layers of BP)

Table 1 Summary of recent progress of phosphorus anodes in SIBs

Material	Synthetic method	Cycling performance	Rate capability	P content	Initial coulombic efficiency	Ref.
RP/CNT	Hand grinding	1283 mA h g^{-1} ,	_	70%	69%	121
RP/graphene	Ball milling	at 143 mA g ⁻¹ , 10th 1700 mA h g ⁻¹ , at 520 mA g ⁻¹ , 60th	520 mA h g^{-1} , 5200 mA g^{-1}	70%	83%	73
RP/CNT/PDA	Ball milling	470 mA h g^{-1} ,	$\begin{array}{c} 3200 \text{ mA g} \\ 1060 \text{ mA h g}^{-1}, \\ 2600 \text{ mA g}^{-1} \end{array}$	66.04%	$\sim 60\%$	122
RP/C	Ball milling/spray drying	at 5200 mA g^{-1} , 5000th 1408 mA h g_p^{-1} , at 0.1 A g^{-1} , 300th	$\begin{array}{c} 2000 \text{ mA g} \\ 986 \text{ mA h gp}^{-1}, \\ 2000 \text{ mA g}^{-1} \end{array}$	42.5%	86.2%	125
RP-graphene	Ball milling	2172 mA h g^{-1} , at 250 mA g^{-1} , 150th	$342 \text{ mA h g}^{-1},$ 2600 mA g ⁻¹	—	_	126
P-TiO ₂ -C	Ball milling	632 mA h g^{-1} ,	958.9 mA h g^{-1} , 1000 mA g^{-1}	60%	77%	130
RP/boron nitride/Gr	Ball milling	at 500 mA g^{-1} , 100th 947 mA h g^{-1} , at 50 mA g^{-1} , 100th	$776 \text{ mA h g}^{-1},$ 300 mA g $^{-1}$	42%	64%	179
Fe ₃ O ₄ /C/RP	Vaporization/condensation	1390 mA h g^{-1} , at 200 mA g^{-1} , 200th	$692 \text{ mA h g}^{-1},$ 2000 mA g $^{-1}$	25%	80.4%	129
RP/C	Vaporization/condensation	700 mA h g^{-1} , at 2 A g^{-1} , 920th	500 mA h g^{-1} , 10 A g ⁻¹	$\sim 60\%$	69.9%	134
P@RGO	Vaporization/condensation	914 mA h g^{-1} , at 1594 mA g^{-1} , 300th	519 mA h g^{-1} , 31.88A g^{-1}	61.4%	75.2%	132
RP/C	Vaporization/condensation	580 mA h g^{-1} , at 2500 mA g $^{-1}$, 800th	430 mA h g^{-1} , 8000 mA g^{-1}	${\sim}40\%$	69.8%	117
RP@RGO	Vaporization/condensation	1625 mA h g^{-1} , at 1000 mA g^{-1} , 200th	$679 \text{ mA h g}^{-1},$ 6000 mA g^{-1}	57.9%	73.2%	133
RP@YP	Vaporization/condensation	1064 mA h g_p^{-1} , at 420 mA g^{-1} , 100th 533 mA h g_p^{-1} , at 420 mA g^{-1} , 100th	601 mA h g^{-1} , 6780 mA g_p^{-1}	48%	68%	136
Hollow P sphere	Solvothermal	1500 mA h g ⁻¹ , at 1300 mA g ⁻¹ , 80th 970 mA h g ⁻¹ ,	278 mA h g ⁻¹ , 10.4 A g ⁻¹	99.9%	77.3%	87
Porous RP/GO	Boiling method	at 2600 mA g^{-1} , 600th 1250 mA h g^{-1} , at 173 mA g^{-1} , 150th	657 mA h g ⁻¹ , 3465 mA g ⁻¹	67.6%	78.5%	144
I-Doped hollow porous RP	Redox reaction Boiling process	1658 mA h g^{-1} , at 260 mA g^{-1} , 100th	$760 \text{ mA h g}^{-1},$ 5200 mA g^{-1}	$\sim 100\%$	75.6%	145
RPQDs/rGO	Hydrothermal	900 mA h g^{-1} , at 250 mA g^{-1} , 250th	193 mA h g^{-1} , 4000 mA g^{-1}	55.7%	55.5%	146
RP@C	Shear emulsifying electrospinning	821 mA h g^{-1} , at 1000 mA g^{-1} , 300th	$676 \text{ mA h g}^{-1},$ 5000 mA g $^{-1}$	51%	75%	127
NTO/C-BP	Ball milling	225 mA h g^{-1} , at 20 mA g $^{-1}$, 55th		20%	42%	177
BP-CNT	Ball milling	1092 mA h g^{-1} , at 520 mA g^{-1} , 200th	441 mA h g ⁻¹ , 11.7 A g ⁻¹	74.5%	87.7%	174
BP/G	Sonication Electrophoretic deposition	1297 mA h g^{-1} , at 100 mA g^{-1} , 100th	$\begin{array}{c} 11.7 \text{ Ag} \\ 623 \text{ mA h g}^{-1}, \\ 500 \text{ mA g}^{-1} \end{array}$	50%	75.6%	175
BP/C-MWNT	Ball milling	1700 mA h g^{-1} , at 1300 mA g^{-1} , 100th (for BP)	928 mA h g^{-1} , 3000 mA g^{-1}	70%	91.1%	173
Phosphorene-C	Liquid-phase exfoliation	2080 mA g^{-1} , 100 m g^{-1} , 10	$\begin{array}{c} \text{645 mA h g}^{-1}, \\ \text{26 A g}^{-1} \end{array}$	48.3%	80%	72
Phosphorene	Electrochemical exfoliation	\sim 1190 mA h g ⁻¹ , at 100 mA g ⁻¹ , 50th	591 mA h g^{-1} , 1500 mA g $^{-1}$	100%	${\sim}50\%$	188

Journal of Materials Chemistry A

has also attracted great interest for Na-ion storage because of its unique 2D structure, high electronic conductivity, and other advantages. Recently, some new allotropic modifications of P have been experimentally produced, such as blue phosphorus⁹⁴ and ring-shaped phosphorus within carbon nanotube nanoreactors;⁹⁵ nevertheless, these allotropes have not been experimentally applied in batteries yet.⁹⁶ So in this section, we will present recent advances in RP, BP and phosphorene anodes for SIBs. And Table 1 summarizes these recent developments.

2.1 Red phosphorus

RP is the most abundant and easily available allotrope of elemental phosphorus,97,98 and is chemically stable and nontoxic to the environment.99 Commercial RP exhibits an amorphous form or an intermediate state between amorphous and crystalline.79,100,101 RP has been widely used in our daily life and industry, such as safety matches,¹⁰² flame retardants¹⁰³ and chemical analysis.¹⁰⁴ Recently, the application of RP has extended to the field of semiconductors,105 rechargeable batteries⁸² and photocatalysts¹⁰⁶ with the progress of nanotechnology. And it is necessary to understand the structural information of RP for further research. Roth et al.¹⁰⁷ proposed five distinct allotropes of RP, four crystalline (Types II to V) and one amorphous (Type I). The exact structures of these four crystalline types can be determined by X-ray diffraction (XRD) or other methods. For example, Type IV (fibrous RP) and Type V (violet P or Hittorf's P) were confirmed to consist of tubes with a pentagonal cross section, which in turn are a regular sequence of three different building units (Fig. 2b and c).100,108-110,113 As for amorphous RP, the exact structure remained unclear until Zhang et al.¹⁰⁰ investigated some of its basic properties by several experimental methods, including scanning tunneling microscopy (STM), gel permeation chromatography (GPC) and single-molecule atomic-force microscopy (AFM). Based on the results, they proposed that amorphous RP should be a linear inorganic polymer with a broad molecular weight distribution (mainly at 31 000 \pm 20 000 Da) and with a zig-zag ladder chain structure (85 \pm 55 nm), as shown in Fig. 2a.¹⁰⁰ Compared with crystalline RP, amorphous RP seems more advantageous in accommodating the volumetric expansion during Li⁺/Na⁺ insertion-extraction cycles. In fact, most RP anodes for Li/Naion batteries in the literature are in the amorphous form, especially for Na-ion batteries.82,111 However, because RP anodes usually exhibit an amorphous form and most of the sodiated RP phases are also amorphous,82 they are difficult to investigate in terms of sodiation and desodiation mechanisms.82 Nevertheless, it is acknowledged that the sodiation of RP mainly occurs between 0.5 V and 0 V to form Na₃P, while the desodiation process involves a stepwise sodium extraction from Na₃P with possible intermediates such as Na₂P, NaP and NaP₇.^{76,79,87}

Although RP possesses great potential for practical application in SIBs due to high theoretical capacity, low redox potential (<0.4 V),⁸² natural abundance and eco-friendliness, major drawbacks such as low electronic conductivity ($\sim 10^{-14}$ S cm⁻¹),⁸⁰ large volume expansion during sodiation (300–500%)^{81,82,112} and low initial coulombic efficiency hinder its application. There are several approaches to overcome these drawbacks: one is to combine RP nanostructures with electronic conductive materials and supporting matrices including carbon-based materials, conductive polymers, metals, metal oxides or metal phosphides; another one is to synthesize novel or self-supported RP nanostructures, and particle surface modification may also be a potential method to address these issues.

2.1.1 Combination with carbon-based materials or conductive polymers. Various carbon-based materials possess high electronic conductivity and/or mechanical robustness, such as Super P® carbon, carbon nanotubes and graphene. These materials can be used to enhance the conductivity of RP based anodes and mitigate the expansion of RP during charge/ discharge cycles, leading to an enhanced Na storage performance. Recently, conductive polymers such as polyacrylonitrile (PAN) and poly-pyrrole (PPy) have also been used as host materials or a protective layer for RP.

As for the synthesis of RP anode composites, two conventional methods are mainly used, including ball milling (or other mechanical milling methods) and vaporization/condensation methods. Ball milling (high energy) is a mechanochemical process that is facile and cost-effective to produce a wide range of nano-powders from bulk raw materials. During a high energy and long-term ball milling process, chemical reactions and structural changes can occur and allow for the formation of new phases or complex composites.¹¹⁴ The vaporization-condensation method is also an effective approach to prepare RP/carbon composites.115 In a typical process, RP is first sublimated by heating and the gaseous P subsequently diffused into the pores or on the surface of the carbon matrix through capillary forces and pressure differences, and then the P is adsorbed and deposited onto the internal surface to form solid RP as the temperature drops down.81,116 Compared with mechanical strategies, the vaporization-condensation method can ensure the uniform dispersal or adsorption of RP into carbon matrices and facilitate intimate contact between RP and host materials.116 However, the RP content in the composite synthesized by this method was usually not high. Some other methods such as solvothermal and hydrothermal have also been applied to prepare RP nanostructure/nanocomposite, which will be fully discussed in latter sections. Among these various synthesis methods, the high energy ball milling method is the most widely used approach to prepare RP-based anodes for both LIBs and SIBs.

RP was used as the anode material in LIBs earlier than in SIBs,^{115,118,119} and researchers found that the lithium ion storage capacity of RP depended dramatically on its crystalline structure and morphology, and a better lithium ion storage performance could be obtained when the crystalline and large-size RP was transformed into amorphous RP nanoparticles.¹²⁰ Soon after these reports, Qian *et al.* and Kim *et al.*^{79,82} first introduced RP into SIBs, and respectively prepared an amorphous RP/carbon nanocomposite as an anode by simply ball milling commercially available amorphous RP and Super P (7 : 3). They found that the micro-sized commercially available RP could not react with Na reversibly in spite of a large initial specific capacity

Review

obtained, while the ball-milled RP nanoparticles show a much better Na-ion storage performance. During this long-term and high-energy ball milling process, bulk commercial phosphorus particles were crushed into amorphous nanoparticles under a strong mechanical force, dispersing into the conductive carbon matrix. Due to the short electron/ion transfer route and the amorphous form of RP, these P/C nanocomposites exhibited high reversible capacity and high rate capability, for example, a rate capability of 1540 mA h g^{-1} at a current density of 2.86 A g⁻¹ could be obtained.⁸² Interestingly, Li et al.¹²¹ demonstrated that commercial micro-sized RP could also deliver a high reversible capacity after being mixed with carbon nanotubes (CNTs) by simple hand milling. And the authors proposed that a good electronic conductivity is essential for RP to reversibly react with Na. This indicates that increasing the electronic conductivity of RP is an effective way to improve the Na storage performance of RP anodes in terms of either the rate capability and the specific capacity. As the combination of RP with CNTs shows noticeable success, predictably, ball milling of RP and carbon materials with high mechanical strength such as

CNTs or graphene may produce amazing results. Song *et al.*⁷³ found that after a ball milling process of RP and graphene stacks, the graphene stacks were mechanically exfoliated to nanosheets, and more importantly, these nanosheets were chemically bonded with the surfaces of RP particles. Benefiting from this robust and intimate contact, the graphene at the particle surfaces could help to maintain electrical contact and stabilize the solid electrolyte interphase upon the large volume change of RP during cycling, leading to an improved cycling stability.

However, these simply ball milled RP anodes often suffered from insufficient rate capability and long-term stability, mainly caused by the large and rough particle size for this method, as the larger particle size means a longer diffusion length for electrons/ions, leading to unsatisfactory rate performance. And it also causes severe pulverization of RP after repeated large volume changes between P and Na₃P, and as a result, lots of pulverized RP particles are electrically isolated in the anode,⁸³ and a continuous solid electrolyte interphase (SEI) growth occurs, leading to fast capacity decay. To address these issues,

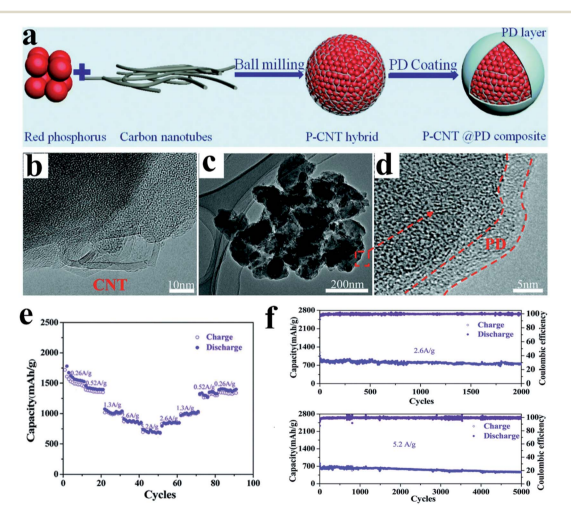


Fig. 3 (a) Schematic of the synthesis of the P-CNT@PD composite. (b) HRTEM image of the P-CNT hybrid. (c) TEM and (d) HRTEM images of the P-CNT@PD composite. (e) Rate performances at various current densities from 0.26 A g^{-1} to 5.2 A g^{-1} of the P-CNT@PD composite between 0.001 and 2.0 V (vs. Na⁺/Na). (f) Long-term cycling performances of the P-CNT@PD composite at 2.6 A g^{-1} and 5.2 A g^{-1} between 0.001 and 2.0 V (vs. Na⁺/Na). Reproduced with permission.¹²² Copyright 2018, The Royal Society of Chemistry.

combination with conductive polymers is an effective strategy. For example, Liu *et al.*¹²² recently designed a strategy to *in situ* coat ball milled P/C hybrids (phosphorus-carbon nanotube, P-CNT) with a polydopamine layer (Fig. 3a–d). They proposed that the coating layer could provide an elastic buffer to accommodate active material volume change and prevent their direct contact with electrolytes. As a result, the P-CNT@PD (PD, polydopamine) composite anode presented high rate capacities of 1060 mA h g⁻¹ in the second discharge and 730 mA h g⁻¹ after 2000 cycles at 2.6 A g⁻¹ and high cycling stabilities with 470 mA h g⁻¹ capacity retained after 5000 cycles at 5.2 A g⁻¹ (Fig. 3e and f), and this is the longest cycle lifespan that has ever been reported for RP-based materials.

In another exmaple,¹²³ a conductive polymer of sulfurized polyacrylonitrile (SPAN) was ball milled with RP, forming a chemically bonded RP/SPAN hybrid. The functional conductive matrix with robust P–S bonds linked to RP could accommodate the large volume change during cycles and enhance the conductivity of the anode. Consequently, this hybrid anode delivered a capacity of ~1300 mA h g⁻¹ at 520 mA g⁻¹ and good cycling performance (91% capacity retention after 100 cycles).

Song *et al.*¹²⁴ developed a novel cross-linked polymer binder and found that it could effectively maintain the structural stability of RP anodes during cycling. They first prepared a chemically bonded RP-carbon nanotube hybrid (P-CNT) as an active material through ball milling, and then, this cross-linked sodium carboxymethyl cellulose–citric acid was used as a binder to fabricate a robust RP electrode, in which citric acid functioned as a cross-linker to form a 3D network when drying the electrode at 150 °C for 2 h. As a result, a high reversible capacity of 2134 mA h g⁻¹ and good cycling stability (~91% capacity retention after 100 cycles) were achieved, with a high initial coulombic efficiency of 84.7%. Similarly, Xu *et al.*¹²⁵ used sodium alginate as a binder to synthesize an affordable

phosphorus/carbon (APC) anode and obtained an improved electrochemical performance. APC nano/microspheres were prepared by ball milling of RP, flake graphite, CNTs, Super P® carbon and sodium alginate (SA, a biomass binder), followed by a closed spray drying process (Fig. 4a). The researchers proposed that this binder-filled phosphorus/carbon structure could maintain the structural integrity after repetitive volume variation and enhance the overall electrical conductivity of the RP anode due to its porous structure and strong surface interactions (Fig. 4b). As a result, this anode delivered a high reversible capacity of 1408 mA h g_P^{-1} at 100 mA g^{-1} with a capacity retention of 82.6% for over 300 cycles and showed superior rate capabilities, demonstrating that this industrially adaptable process and the application of sodium alginate binders are promising methods for RP-based practical SIBs (Fig. 4c). However, the low RP content in this anode (\sim 42.5%) and the massive usage of carbon-based materials may cause issues with respect to gravimetric/volumetric capacity for practical application.

To investigate whether the carbon defects of the carbon scaffold in a anode have an effect on the electrochemical performance, Li *et al.*¹²⁶ designed a carbon scaffold containing controlled single-walled carbon nano-horn (SWCNH) and single-walled carbon nanotube (SWCNT) compositions, and combined this carbon scaffold with RP by hand milling. The researcher chose these two kinds of carbon for the reason that SWCNHs exhibit a significant presence of sp³ hybridized carbon defects embedded into the sp² hybridized carbon matrix while SWCNTs are fully based on defect-free sp² hybridized carbon species, and thus carefully controlled defect density composites can be obtained. Here, they reported that the stable alloy region in which NaP and Na₅P₄ were formed between 0.40 and 0.15 V *versus* Na/Na⁺ was mostly independent of carbon composite matrix chemistry and that an unstable alloying region (0-0.15 V)

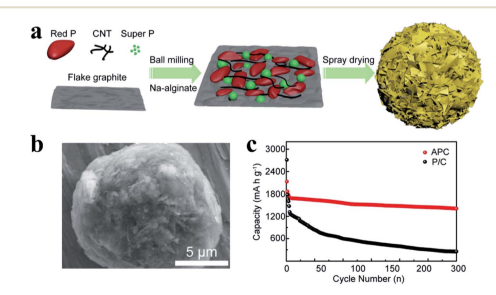


Fig. 4 (a) Schematic of the synthetic process of phosphorus/carbon (APC) nano/microspheres. (b) SEM image of the typical morphology of the APC nano/microspheres. (c) Cycling performance of the anode at a constant current density of 100 mA g^{-1} . P/C refers to the control material without SA binders. Reproduced with permission.¹²⁵ Copyright 2018, American Chemical Society.

where defects could facilitate the continuous formation of irreversible Na_3P products over the carbon surface. This work elucidates the mechanism why sp² hybridized carbon is better suited as a reversible host material for RP anodes, and suggests that future effort can leverage surface engineering routes to further improve performance in this unstable alloying regime.

Interestingly, besides this ball milling method, Liu et al.¹²⁷ broke RP particles into smaller clusters by using a high-shear emulsifying machine, and embedded these RP nanoparticles (~97 nm) homogeneously in porous nitrogen-doped carbon nanofibers (RP@C) by a electrospinning technique (Fig. 5a). They first ball milled commercial RP into RP particles, and then these RP particles were broken into smaller clusters under a high energy shear force during the emulsifying process. After standing, the RP with small sizes (\sim 97 nm) in the upper liquid were collected and were homogeneously embedded in porous nitrogen-doped carbon nanofibers via a electrospinning process and a heat treatment (Fig. 5b and c). Although the yield of these small RP particles may be low as they only exist in the upper liquid, this method is still interesting because of its uniform particle size, and the larger RP particles in the sediment can be recycled for further preparation. In addition, these RP embedded nanofibers could be fabricated as a binder- and current collector-free anode, which could not only increase the P content in the anode, but also reduce the fabrication cost for practical application. The anode showed a high reversible Na storage capacity of 1308 mA h g^{-1} at 200 mA g^{-1} , exceptionally

high rate capabilities of 637 mA h g⁻¹ at 5000 mA g⁻¹ and 343 mA h g⁻¹ at 10 000 mA g⁻¹ and long cycling stabilities of ~81% capacity retention over 1000 cycles (Fig. 5d and e). Moreover, researchers also assembled a soft package Na-ion full battery using RP@C as the anode and Na₃V₂(PO₄)₂F₃/C as the cathode and obtained an operational voltage of ~3.65 V and a SIB energy density of 161.8 W h kg⁻¹.

Aside from ball milling or other mechanical force methods, the vaporization-condensation method is also an effective approach to prepare RP/carbon composites.¹¹⁵ In a typical process of this method, RP is first sublimated by heating and subsequently diffused into the pores or surface of a carbon matrix through capillary forces and pressure differences, and then the gaseous P is adsorbed and deposited onto the internal surface.^{81,116} As compared with the mechanical strategy, the vaporization-condensation method can ensure the uniform dispersal or adsorption of RP into carbon matrices and can obtain nanosized RP.76,116 As a result, the P/C composite fabricated by this method usually shows better long-term stability than that by the ball milling method. For the first time, Marino et al.¹¹⁵ prepared a carbon-phosphorus composite using this method for application in LIBs. Subsequently, this method was widely applied in SIBs, for example, Zhu et al.85 prepared a RP/ single-walled carbon nanotube composite (P/SWCNT) as an anode, in which RP was uniformly distributed between tangled SWCNT bundles with an average particle size of \sim 5 μ m and a P content of \sim 40%. Benefiting from this mild and nondestructive

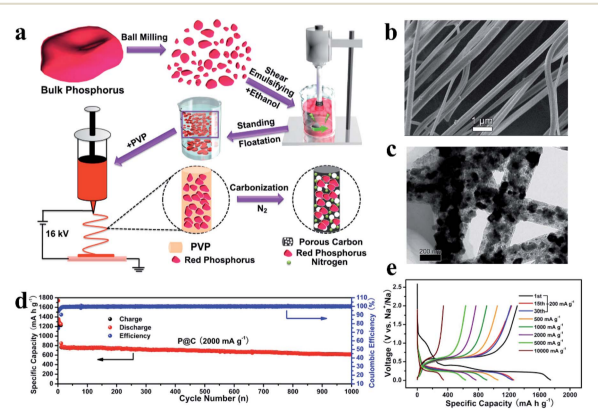


Fig. 5 (a) Schematic illustration of the preparation process for RP@C nanofibers. (b, c) SEM and TEM images of red P@C nanofibers. (d) Long-term cycling stability of the P@C electrode at a current density of 2000 mA g^{-1} . (e) Charge/discharge profiles obtained at various rates of the P@C electrode in the voltage range of 0.01–2 V vs. Na⁺/Na. Reproduced with permission.¹²⁷ Copyright 2017, Elsevier B.V.

preparation process, uniform mixing and intimate contact between RP and SWCNTs could be achieved, and the mechanically strong SWCNT network was preserved, which enhanced the conductivity of the composite, helped to maintain the integrality of the anode, and stabilized the solid electrolyte interphase, leading to a stable long-term cycling performance with 80% capacity retention after 2000 cycles. However, because the particle size was quite large ($\sim 5 \mu m$), the rate capability of the anode was not sufficient (\sim 300 mA h g⁻¹ at 2000 mA g⁻¹), and the high cost of SWCNTs is also a concern for practical application. In another study, ordered mesoporous carbon (commercial CMK-3) with a pore size of \sim 4 nm was also used as a carbon host material to load RP. In this RP/CMK-3 composite, the nanoscale of RP and the interconnected pores of the CMK-3 matrix provide many advantages for Na storage, such as accommodating the volume change of RP during cycling, reducing the diffusion length of Na^+/e^- and leading to an enhanced Na storage performance.128 However, the P content of this RP/CMK-3 composite was not high (\sim 31%), and the other RP/C composites prepared by this method in the literature also suffered from this same issue.117,129 The large amount of carbon in the composite is also a concern, which may lead to low volumetric capacity, low coulombic efficiency and other negative effects.130 Gao et al.131 developed a novel strategy and designed a unique structure: coating a 3D porous graphenehydrogel/RP precursor with a poly-pyrrole layer, and thus the P content in the composite could be effectively maintained during the following vaporization-condensation-redistribution process. This coated composite showed a P content of \sim 47.2%, which was more than twice that of the sample without a carbon layer (22.8%). In addition, the researchers proposed that the

unique nanostructure of this 3D porous carbon matrix with RP particles (10–20 nm) uniformly distributed and firmly sealed in it was an ideal design for an anode material in SIBs.¹³¹

Recently, Liu *et al.*¹³² developed a method to densely and uniformly deposit RP nanodots onto reduced graphene oxide sheets (P@RGO) with RP nanodot sizes ranging from tens to several hundred nanometers (Fig. 6a–c). Here, the researchers suggested that this unique size and structure could minimize sodium ion diffusion lengths and sodiation/desodiation stress whereas the RGO network could serve as an efficient pathway for electron transfer and accommodate the volume variations of phosphorus particles. As a result, the P@RGO flexible anode reportedly achieved specific charge capacities of 1165.4, 510.6 and 135.3 mA h g⁻¹ at 159.4, 31 878.9, and 47 818.3 mA g⁻¹ respectively along with a 914 mA h g⁻¹ capacity after 300 deep cycles at a current density of 1593.9 mA g⁻¹, which was much better than that of conventional RP anodes for SIBs (Fig. 6d–f).

Furthermore, Zhou *et al.*¹³³ developed a facile single-step flash-heat treatment to obtain reduced graphene oxide and deposit RP onto the rGO sheets simultaneously. This single-step method is energy-saving and is beneficial to preparation on a large scale. The resulting RP/rGO flexible film anode for SIBs produced an average capacity of 1625 mA h g^{-1} after 200 cycles at a charge/discharge current density of 1 A g^{-1} and also showed an excellent rate capability with average charge capacities of 1786, 1597, 1324 and 679 mA h g^{-1} at 1, 2, 4 and 6 A g^{-1} current densities respectively.

In addition to the application of these traditional carbon materials, the design and fabrication of novel carbon hosts with high electronic conductivity, high mechanical strength and porous morphology to load RP is an important topic for this

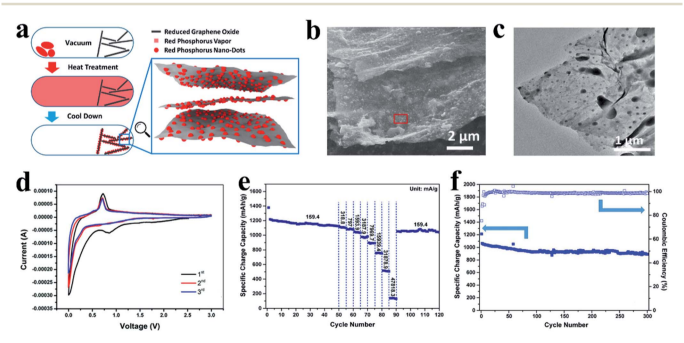


Fig. 6 (a) Schematic of the P@RGO synthesis. (b, c) SEM and TEM images of the P@RGO composite. (d) Cyclic voltammetry of the P@RGO anode at a scan rate of 0.1 mV s^{-1} between 0 and 3.0 V vs. Na/Na⁺. (e) Rate performance of the P@RGO anode. (f) Cycling performance of the P@RGO anode at a charge/discharge current density of 1593.9 mA g⁻¹. Reproduced with permission.¹³² Copyright 2017, American Chemical Society.

vaporization-condensation method to further improve the Na storage performance. Based on this, Sun et al.¹³⁴ used this approach to prepare composites of RP encapsulated into porous multichannel carbon nanofibers as a flexible anode material for SIBs and reported a rate capability of 500 mA h g^{-1} at 10 A g^{-1} and long cycle lifespans with 700 mA h g^{-1} capacity retention at 2 A g^{-1} after 920 cycles. The researchers suggested that the novel structure of the synthesized composite could reduce ion and electron diffusion barriers, accommodate volume variaduring sodiation/desodiation, improve electrode tions mechanical strength, and thus lead to excellent rate capabilities and cycling performances. Liu et al.117 designed a carbon nanotube-backboned mesoporous carbon (TBMC) material as a matrix for RP (Fig. 7a). In their preparation process, resorcinol-formaldehyde resin/SiO2 was first grown onto carbon nanotubes, with tetraethyl orthosilicate (TEOS) used as the SiO₂ precursor and resorcinol-formaldehyde (RF) resin as the carbon source, and then the resin was carbonized and the SiO₂ was removed to obtain TBMC. The researchers reported that the competitive growth of SiO₂ and RF resin during the synthesis process could lead to various SiO2 distributions and thus different final structures. Consequently, TBMC, hollow TBMC and deflated TBMC could be obtained based on high, medium and low dosages of RF respectively (Fig. 7a). These various TBMCs were subsequently infiltered with RP using the vaporization/condensation method to form P@TBMC composites. The researchers found that the TBMC with a high dosage of RF showed the best Na storage performance, providing a high reversible specific capacity of ~ 1000 mA h g⁻¹ at 0.05 A g⁻¹,

a superior rate performance of ~430 mA h g⁻¹ at 8 A g⁻¹ and a long cycle lifespan with no capacity decay for 800 cycles at 2.5 A g⁻¹ (Fig. 7b-d). They proposed that the multi-walled carbon nanotubes could facilitate electron transfer due to the high content of sp² carbon whereas the mesoporous carbon layers provide voids to load P and ensure adequate spacing to accommodate large volume variations of P during cycles, leading to an excellent Na storage performance.

Yao et al.135 also designed novel hollow porous carbon nanospheres (HPCNSs) as host structures for the vaporization/ condensation synthesis of P/C composites. Through the methods of molecular dynamic simulations and density functional theory calculations, they found that the morphology of polymeric P₄ and the P loading in the P/C composite were mainly dependent on the pore size and surface conditions of the carbon support. The micropores of 1-2 nm in diameter and oxygenated functional groups attached on the carbon surface were essential for achieving high P loadings and high structural stability. This HPCNS/amorphous RP composite with enhanced structural/functional features presented a low volume expansion of \sim 67.3% during anode cycles, much smaller than that of the theoretical value of commercial RP ($\approx 400\%$), and showed remarkable long-life cycling stabilities with a capacity retention of over 76% after 1000 cycles. This study signified the importance of the rational design of electrode materials to obtain cost-effective P/C composite anodes for SIBs. Furthermore, Li et al.76 reported that a composite of confined amorphous RP with particle sizes of less than 1 nm in MOF-derived N-doped microporous carbon also produced promising performances.

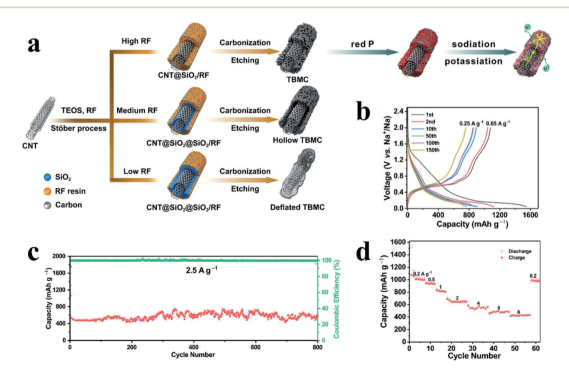


Fig. 7 (a) Schematic of the synthesis of nanotube-backboned mesoporous carbon (TBMC), hollow TBMC and deflated TBMC and the sodiation/ potassiation of the P@TBMC composite. (b) Representative galvanostatic discharge/charge voltage profiles of the P@TBMC electrode at 0.25 A g⁻¹ after the initial two-cycle activation at 0.05 A g⁻¹. (c) Long-term cycling performance of P@TBMC at 2.5 A g⁻¹ and (d) rate capability of P@TBMC. Reproduced with permission.¹¹⁷ Copyright 2018, Elsevier Ltd.

These studies indicate that the design of favorable carbon hosts for RP is an effective approach for improving the Na storage performance. In addition, facile preparation methods for these materials are also necessary for practical applications.

Compared with the ball milling method, the vaporization/ condensation method is a mild and nondestructive process; however, the phase-transformation process of P will inevitably generate white phosphorus, which is toxic and highly active.¹¹⁶ To address this, Yu et al.136 developed a modified vaporizationdeposition-conversion method in which a multi-step heating/ cooling process was used to avoid the generation of white phosphorus. By using this refined strategy, RP nanoparticles were adsorbed and confined within the nanopores of commercial microporous carbon YP-80F (P@YP) (Fig. 8a), ensuring high reversible capacity, excellent rate capability and long-term cycling stability with a superior capacity retention of 92% after 100 cycles and 46% after 1000 cycles (Fig. 8b-d). And as compared with P@YP, a P@CNT composite without confined P was deposited onto the surface of carbon nanotubes and presented rapid capacity decay with a capacity retention of only \sim 40.6% after 100 cycles. Here, the researchers suggested that the nanostructure of the P-C composite with P confined in the porous carbon is key to enhanced electrochemical performances and long-term cycling stabilities.136

2.1.2 Combination with other materials. Although RP/C composite approaches appear to be promising, the performance of the RP/C composites is found to be dependent on several factors such as fabrication methods.⁸¹ The large amount of carbon in the RP/C composite is also a concern, as the low density (\sim 2.16 g cm⁻³),¹³⁷ high specific area, poor air stability and low operating voltage (<0.1 V *vs.* Na/Na⁺)¹³⁸ of these carbonaceous materials usually result in low volumetric capacity, low

coulombic efficiency, autoignition, and possible sodium plating.^{86,130,138-141} The higher the carbon content, the more significant these problems are. In addition, the limited carbon coverage on RP and uncontrollability of carbon layer thickness may also hinder the electrochemical performance of RP/Cbased electrodes. In this regard, the search for noncarbonaceous materials that function similarly to those carbon materials is needed.¹³⁰

In fact, the combination of RP with other non-carbon materials, such as metals, metal oxides or metal phosphides also demonstrates improved Na storage performance. For example, Chin et al.142 mixed commercial RP with appropriate amounts of iron using high-energy ball milling to obtain a RP/ Fe composite. As a result, they found that the RP/Fe anode showed a higher sodium diffusion coefficient than pure RP anodes, and improved electrochemical performance of 1582, 1419, 884 and 676 mA h g^{-1} specific desodiation capacities at 0.2, 0.6, 6 and 10 A g^{-1} current densities respectively were obtained for the 9.1% Fe-containing RP composite. Although the cycle lifespan of this anode may not be long because of the pulverization of RP during charge/discharge processes, the addition of low cost iron is still a novel and effective approach to improve the electrochemical performance of RP. Based on the idea that the functional synergy of higher capacity of P with high-rate capability and high cyclability of Sb may obtain a compelling electrochemical performance, Walter et al.143 reported a P/Sb/Cu composite anode synthesized by simple mechanic mixing of monodisperse Sb nanocrystals with RP particles and Cu nanowires. Benefiting from the good electronic conductivity of Sb and Cu and the mechanical strength of Cu nanowires, this P/Sb/Cu anode showed much greater cycling stability than pure RP anodes, providing a capacity of above

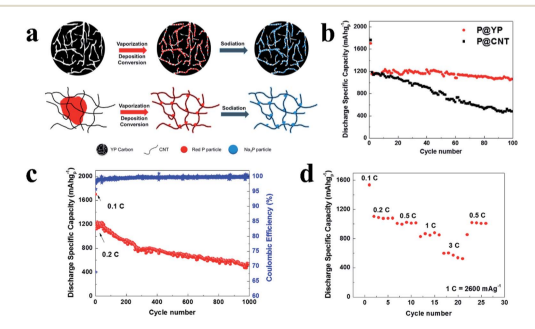


Fig. 8 (a) Schematics of the preparation and sodiation of P@YP and P@CNT composites. (b) Cycling performances of P@YP and P@CNT composites within 100 cycles. (c) Long term cycling performance of the P@YP composite. (d) Rate performance of the P@YP composite at C-rates between 0.1C and 3C ($1C = 2600 \text{ mA g}^{-1}$). Reproduced with permission.¹³⁶ Copyright 2017, Elsevier Ltd.

Review

1100 mA h g⁻¹ after 50 cycles at a current density of 125 mA g⁻¹, and an excellent rate capability of more than 900 mA h g⁻¹ at a high current density of 2000 mA g⁻¹. Compared with the addition of Fe within RP, the use of Sb and Cu in this study may increase the cost for practical application, as the P content in this P/Sb/Cu anode is low (40%). Nevertheless, the mechanical strength of Cu nanowires along with the matrix of Sb can ensure a longer cycle lifespan for this P/Sb/Cu anode.

Lan *et al.*¹³⁰ prepared a phosphorus–titania–carbon (P–TiO₂– C) composite as an anode for SIBs by ball milling of RP, titanium oxide and carbon powder. The researchers suggested that the P–TiO₂–C composites could alter local chemistries in both titanium oxide and phosphorus such as the doping of TiO₂ by P and/or C, as well as stabilization of P by the Ti–O–P and/or P–C chemical environment, and thus lead to an improved electronic conductivity and enhanced stability during repeated charge/ discharge cycles. The resulting composite anode delivered a high reversible capacity of 1147 mA h g⁻¹ at a current density of 100 mA g⁻¹ with good cyclability and a high rate performance with 81.5% of its original capacity being maintained as discharge rates increased from 100 to 1000 mA g⁻¹. In addition, a chain-like Fe₃O₄/C/RP (C means carbon) composite was also prepared by introducing a magnetic stimulus source during the vaporization/condensation process and also demonstrated high cycling performances and superior rate capabilities.¹²⁹ Interestingly, this assisted magnetic field could cause a high pressure driven force to uniformly deposit RP onto the Fe₃O₄/C surface, and consequently increase the P loading and enhance the contact between Fe₃O₄/C and RP. Although the content of RP in this anode is quite low (25%) and the RP is only loaded on the surface of the conductive carbon chain, the addition of RP can obviously increase the specific capacity and shows little negative effects on the cycling stability. This indicates that the combination of low-cost metal oxides and RP may be an effective approach to apply in practical SIBs; however, the low electronic conductive of metal oxides is still a concern and more effort should be dedicated to addressing this issue.

Liu *et al.*⁸⁶ designed RP@Ni–P core–shell nanostructures as anodes for SIBs through a two-step approach: electroless deposition of Ni on RP particles and then chemical dealloying of this RP@Ni compound, in which the Ni–P shell thicknesses could be controlled by regulating the dealloying time (Fig. 9a). Here, Liu *et al.*⁸⁶ suggested that *in situ* generated Ni₂P on RP particle surfaces could facilitate intimate contact between RP

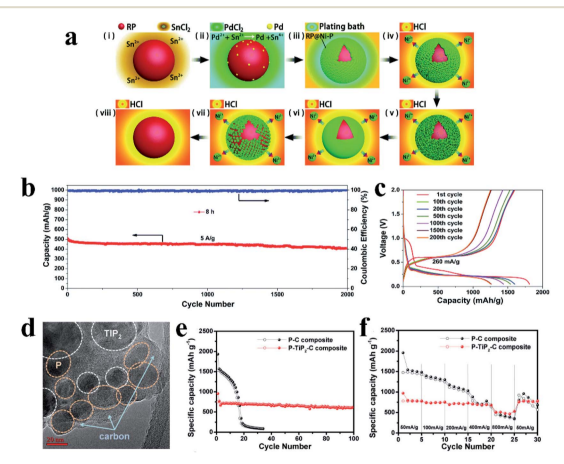


Fig. 9 (a) Schematics of the electroless deposition of Ni on RP nanoparticles (i–iii) and the evolution of the RP@Ni–P core@shell nanostructure through chemical dealloying (iv–viii). (b) Long-term cycling performance of the 8 h RP@Ni–P electrode at 5 A g^{-1} between 0.05 and 2.0 V. (c) Galvanostatic charge–discharge voltage profiles of the 8 h RP@Ni–P anode between 0.05 and 2.0 V at a current density of 260 mA g^{-1} . Reproduced with permission.⁸⁶ Copyright 2017, The Royal Society of Chemistry. (d) Low magnification TEM image of the P–TiP₂–C composite, (e) cycle performances and (f) rate capabilities of the P–C and P–TiP₂–C composite electrodes in sodium half cells. The specific capacity was calculated based on the total mass of the composite materials. Reproduced with permission.¹³⁹ Copyright 2016, American Chemical Society.

and the amorphous Ni-P outer shell which was mechanically strong and highly electron conductive, thus ensuring strong electrode structural integrity, stable solid electrolyte interphases and fast electronic transport. As a result, the RP@Ni-P composite with a dealloying time of 8 h displayed a super high capacity of 1256.2 mA h g^{-1} after 200 cycles at 260 mA g^{-1} , a superior rate capability of 491 mA h g^{-1} at 5200 mA g^{-1} and an long cycle lifespan with a capacity of 409.1 mA h g^{-1} retained after 2000 cycles at 5000 mA g^{-1} (Fig. 9b and c). When the RP particle size is below 200 nm, the outer protective shell is a key factor in obtaining long cycle stability, which can effectively prevent the pulverization of RP during cycles. In addition, the facile process of in situ growth and the low-cost of Ni make it a promising strategy for practical SIBs. In another study, Kim et al.¹³⁹ synthesized a P-TiP₂-C nanocomposite as an anode material for SIBs using a facile and simple one-step high energy mechanical milling process in which the resulting composite anode (Fig. 9d-f) displayed an initial desodiation capacity of 755 mA h g^{-1} with a coulombic efficiency of 80% and a good capacity retention of 80% after 100 cycles. Here, the researchers suggested that the greatly improved electrochemical performances could be ascribed to the unique nanostructure in which RP and crystalline TiP₂ were homogeneously embedded into a conductive carbon network. They also suggested that in this phosphorus-based composite, the conductive TiP₂ could provide structural stability as well as high conductivity during cycling, leading to better SIB cycling stabilities and rate capabilities as compared with nanocomposites without TiP₂.

2.1.3 Morphology and/or size-controlled RP nanostructure. Although ball-milling and vaporization–condensation methods possess many advantages and are the most commonly used methods to prepare RP-based anodes, the uneven and large particle sizes of RP and the low P content in the ball-milled RP composite and the uncontrollable distribution and the low loading ratio of RP in the vaporization/condensation RP anode consequently cause poor rate performance, reduced long-term

stability, and decreased energy density. In addition, the toxic white P remnants produced by the vaporization/condensation method make it unfit for the popularization of SIBs.^{79,87,121,132,134} In view of this, synthesizing a morphology/size controlled RP nanostructure by novel preparation methods is an effective approach. For example, Zhou *et al.*⁸⁷ recently successfully fabricated hollow P nanospheres (HPNs) with porous shells by a solvothermal method (Fig. 10a). The synthesis progress could be summarized and expressed by the following equation:

$$10\text{NaN}_3 + 2\text{PCl}_5 \xrightarrow{\text{toluene}} 2\text{P} + 10\text{NaCl} + 15\text{N}_2$$
 (1)

Here, a gas-bubble-directed formation mechanism during the solvothermal process works to produce hollow P nanosphere with the produced P nuclei covering the surface of NaN₃ during the initial reaction stage. And with increasing gas-bubble sizes, P nuclei are continuously supplied at the interface between NaN₃ and the gas bubbles and gradually assemble around the surface of the gas bubbles. These bubbles eventually detach from the surface of NaN3 and form hollow phosphorus nanospheres. And with subsequent cooling and system pressure reduction, the gases confined by the P shells are released and thus help to form porous shells. Here, the researchers reported that different average diameters ranging from 150 nm to 1 µm could be obtained for these hollow phosphorus nanospheres through tuning the NaN3 concentration, and found that 300 nm hollow nanospheres (shell thickness: ~ 40 nm) exhibited optimal electrochemical performances as an anode for a SIB, delivering a high capacity of 1364.7 mA h g^{-1} at 0.2C (with 60% loading of P, 2274.5 mA h g^{-1} for RP, and $1C = 2.6 \text{ A } g^{-1}$) and long-term cycling performances.87

However, this pure phase RP nanostructure may suffer from insufficient electronic conductivity. Furthermore, the use of toxic and explosive NaN_3 in the preparation process is also a concern. In another study, iodine-doped RP nanoparticles with size ranging from 100 to 200 nm have also been

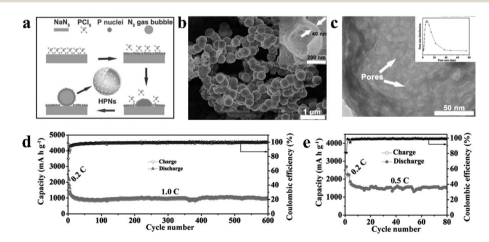


Fig. 10 (a) Schematic of the proposed formation mechanism of hollow phosphorus nanospheres (HPNs). (b) SEM image of the prepared HPNs; inset shows the SEM image of a broken nanosphere. (c) TEM image of the HPNs under higher magnification; inset shows the pore-size distribution of the HPNs. Cycling stability and coulombic efficiency of the HPNs at (d) 1.0C for 600 cycles and (e) at 0.5C for 80 cycles. $1C = 2.6 \text{ A g}^{-1}$. Reproduced with permission.⁸⁷ Copyright 2017, WILEY-VCH.

synthesized by a simple and large-scale solution phase method.¹⁴⁰ During the reaction process, the PI₃ precursor was reduced with ethylene glycol and simultaneously RP nanoparticles (RPNPs) were formed in the presence of CTAB under an ambient environment. The conductivity of these I-doped RPNPs was measured to be between 2.62×10^{-3} and 1.81×10^{-2} S m⁻¹, which is close to that of semiconductor germanium $(1.02 \times 10^{-2}$ S m⁻¹). Although the authors only investigated the Li-ion storage performance of these RPNPs, the unique morphology/nanostructure and the novel synthesis method are all progressive for the application of RP.

Based on this novel solution phase method, Liu *et al.*¹⁴⁴ synthesized nanoporous RP on reduced graphene oxide as a SIB anode (NPRP@RGO) by a boiling method. In brief, the RP@RGO precursor was first synthesized involving several steps, including the reduction of phosphorus triiodide (PI_3) to

produce RP, the uniform mixing of the as-prepared RP with GO in a solution, and a further reduction of RP@GO to RP@RGO. Then this reduced RP@RGO was subsequently subjected to centrifugation and the concentrated RP@RGO solution was heated in a vacuum drying chamber at 200 °C to allow the RP to assemble around gas bubbles. And with the growth of these bubbles, increasing amounts of disordered RP particles became dispersed onto the surface of the gas bubbles and the final NPRP@RGO was obtained after the gas bubbles were detached (Fig. 11a). Here, the researchers reported that the unique nanostructure of the nanoporous RP with sizes mainly in the range of 20-100 nm (Fig. 11b) was able to accommodate volume change and minimize ion diffusion lengths and that the highly conductive RGO network sheets could facilitate fast electron and ion transportation. As a result, this anode exhibited a high capacity of 1249.7 mA h g^{-1} after 150 cycles at 173.26 mA g^{-1} ,

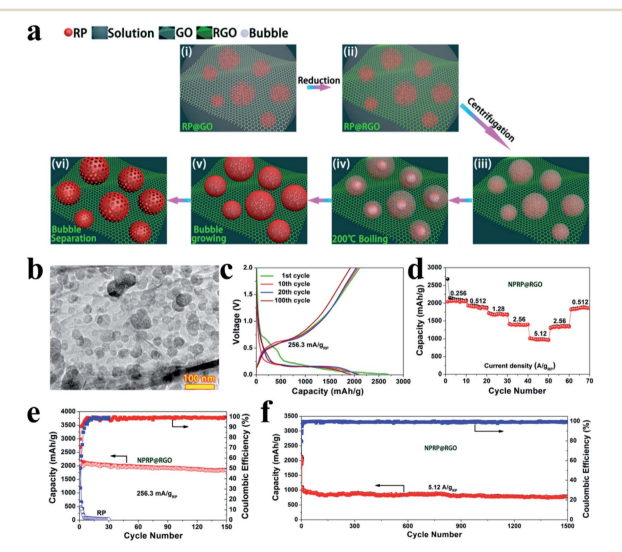


Fig. 11 (a) Schematic of the preparation of NPRP@RGO. (b) TEM image of NPRP@RGO. (c) Galvanostatic charge/discharge curves of the NPRP@RGO electrode at 256.3 mA g_{RP}^{-1} (173.26 mA $g_{composite}^{-1}$) within a voltage range of 0.01–2 V. (d) Rate capability of the NPRP@RGO electrode from 0.256 to 5.12 A g_{RP}^{-1} . (e) Cycling performances of NPRP@RGO and commercial RP electrodes at 256.3 mA g_{RP}^{-1} within the voltage range of 0.01–2 V. (f) Long-term cycling performance of the NPRP@RGO electrode at 5.12 A g_{RP}^{-1} (3465.28 mA $g_{composite}^{-1}$) between 0.01 and 2.0 V. Current density and specific capacity were calculated based on the mass of NPRP. Reproduced with permission.¹⁴⁴ Copyright 2018, American Chemical Society.

a superior rate capability of 656.9 mA h⁻¹ at 3465.28 mA g⁻¹ based on the anode composite and a long cycle lifespan of 1500 cycles with a capacity of 775.3 mA h g⁻¹ at 5.12 A g⁻¹ based on the RP mass (Fig. 11c and d).¹⁴⁴ In addition, in their another study,¹⁴⁵ iodine-doped hollow nanoporous RP (HNPRP) was prepared by a similar preparation method. When tested as an anode for SIBs,¹⁴⁰ this HNPRP exhibited a high capacity of 1658.2 mA h g⁻¹ after 100 cycles at 0.26 A g⁻¹, superior rate capability (759.6 mA h g⁻¹ at 5.2 A g⁻¹) and ultralong cycle-life (857.3 mA h g⁻¹ after 1000 cycles at 2.6 A g⁻¹).

Sun et al.¹⁰¹ further fabricated ultrafine RP particles with an average size of ~ 10 nm embedded in a 3D carbon framework through the carbothermic reduction of P₂O₅ and reported that these small sized phosphorus particles could accommodate large stress without cracking, decrease electron/ion diffusion lengths and connect strongly with the carbon framework. And as a result, their SIB anode provided a reversible specific capacity of 1027 mA h g⁻¹ and a high capacity retention of 88% over 160 cycles. RP quantum dots (RPQDs) were also prepared using a reliable hydrothermal method with commercial RP as the only phosphorus source.146 Although the yield of RPQDs in this synthesis process was low, this method was quite simple, and the size of these quantum dots was at the nano-level with an average size of 4.0 nm. And a hybrid of these dots with rGO (RPQDs/rGO) delivered an initial specific capacity of 1161 mA h g^{-1} and a low capacity deterioration rate of less than 0.12% per cycle after 250 cycles at 200 mA g^{-1} .

2.2 Black phosphorus and phosphorene

BP is another important allotrope of phosphorus and is most thermodynamically stable among these allotropes.88,147,148 Due to its layered structure and other properties, BP attracts much attention and shows great potential for many applications such as batteries,72,149 field-effect transistors,150 and photodetectors.149 BP possesses three crystalline modifications induced by different temperature and pressure,¹⁵¹ including orthorhombic, rhombohedral, and cubic, and also can present an amorphous form.^{152,153} Under normal temperature and pressure, BP usually shows an orthorhombic crystallinity with a layered structure, as shown in Fig. 12a and d.89 Here, phosphorus atoms are covalently bonded with three adjacent phosphorus atoms to form a puckered honeycomb structure in a single layer of BP,89,150 and these individual atomic layers are stacked together by van der Waals interactions, similar to graphite (Fig. 12a),^{148,150,154} giving a calculated density of \sim 2.69 g cm⁻³. The layer-to-layer spacing is about 5.3 Å, and there are no contacts shorter than 3.59 Å for atoms between successive layers.¹⁵⁴ BP shows good electronic conductivities (\sim 66–300 S m⁻¹ vs. \sim 10⁻¹² S m⁻¹ for RP)^{155,156} and has a suitable operating potential ($\approx 0.45 \text{ V} \nu s$. Na/Na⁺) for Na storage.¹⁵⁷ In addition, the interlayer channel size of BP is \sim 3.08 Å, which is suitable for the storage of both lithium (1.52 Å) and sodium (2.04 Å) ions in the channels (Fig. 12a).^{148,158} However, sluggish reaction kinetics and large volume changes (300-500%) during desodiation/sodiation hinder its application as a SIB anode.158,159 Furthermore, the lack of effective, low-cost

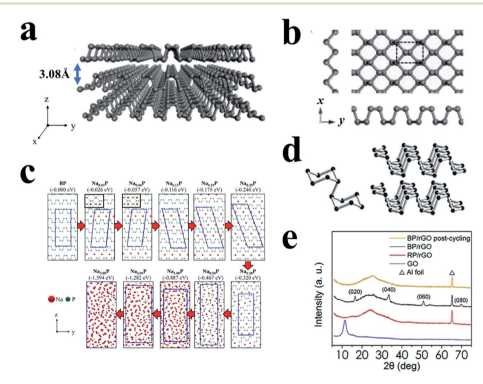


Fig. 12 (a) The puckered structure of a few-layer BP sheet. (b) Top view of monolayer BP. The *x* and *y* axes are along zigzag and armchair directions, respectively. Reproduced with permission.⁸⁹ Copyright 2015, Wiley-VCH. (c) Sodiation mechanism of black phosphorus. The numbers in parentheses indicate calculated formation energies of Na_xP structures. Reproduced with permission.¹⁶⁷ Copyright 2015, American Chemical Society. (d) Characteristic structural fragments of BP. Reproduced with permission.¹¹³ Copyright 2017, Wiley-VCH. (e) XRD patterns of the GO, RP/rGO precursor, as-prepared and post-cycling BP/rGO samples. Reproduced with permission.¹⁶² Copyright 2018, American Chemical Society.

View Article Online Journal of Materials Chemistry A

synthesis methods to produce BP on a large scale is a major concern for its practical application. Currently, the main strategies for preparing BP include high-energy mechanical milling,¹⁴⁷ mineralizer-assisted gas-phase transformation¹⁶⁰ and high-pressure (or with high-temperature) conversion from RP.^{161,162} All of these methods are critically dependent on the preparative conditions. With respect to this, some progress for novel methods to synthesize BP has been made recently.^{163–166}

Compared with RP, the electrochemical mechanism of BP to store Na is easier to unravel due to its crystalline structure. And a thorough understanding of the chemistry that occurs in sodiation and desodiation can elucidate the structure-performance relationships responsible for high capacities and help to devise strategies to mitigate the performance degradation.¹⁶⁸ Many theoretical calculations have been carried out to understand the process of sodiation for BP.167,169-171 Based on these investigations, sodiation of BP includes two steps: (1) an intercalation process occurring in the interlayer channels of BP (along the x-axial) until forming the composition of Na_{0.25}P (diffusion barrier of 0.18 eV); and (2) an alloying process to form a layered Na_xP structure and eventually an Na₃P phase. After the intercalation process (step 1) was completed, one Na atom tended to bind with four P atoms, resulting in sliding of the phosphorene layers (Fig. 12c). At Na levels beyond Na_{0.25}P, alloying with P could lead to the breaking of P-P bonds, in which the staggered P-P bonds were preferentially broken rather than the planar P-P bonds, generating P₂ dumbbells. As sodiation proceeded further, most of the P₂ dumbbells became isolated P atoms. Thus, after full sodiation to form the amorphous Na₃P phase, only low-coordinate P components such as isolated atoms (primarily) and dumbbells could be found.

However, the mechanism of the desodiation process was little reported in the literature. In addition, it seems that the crystalline BP was converted into an amorphous form during cycling, and this has been experimentally evidenced. In a study, Liu et al.¹⁶² found that the XRD characteristic peaks of BP disappeared after repeated charge and discharge (Fig. 12e), suggesting that the BP did not recover to its crystalline phase after desodiation, which was consistent with the results of Raman patterns, TEM and fast Fourier transform (FFT) images. If the BP was amorphized during cycling, it was difficult to explain why the cycling performance of anodes prepared from BP could continue to exceed that of anodes prepared from RP. To answer this question, researchers speculated that phosphorus (like C, H₂O, and silicate glasses) might be polyamorphous and that amorphous P formed from BP could retain both structural and electronic advantages over RP or amorphous products formed from RP, and BP-graphite/graphene composites might play a role in maintaining the layered structures of BP.72,162 In addition, it was proposed that during the charge-discharge process, the electrolyte molecules could intercalate into the layer spaces and then lead to exfoliation of the layered structures and thus BP-graphite/graphene composites or pillared BP might be good options to improve the stability of the layered structure of BP during sodiation.167,169-171

Recently, Marbella *et al.*¹⁶⁸ investigated phosphorus intermediates in BP anodes during the sodiation and desodiation process on a molecular level by a combined experimental and theoretical approach, involving techniques such as solid-state NMR (ssNMR), powder X-ray diffraction (XRD), and DFT and NMR calculations. Specifically, individual P binding sites within the amorphous Na_rP phases that were formed during cycling can be probed with 2D ³¹P phase-adjusted spinning sideband (PASS) ssNMR experiments, thus providing information on the geometry of chemical bonding environments and structures of molecular fragments. In their study, ex situ ³¹P magic-angle spinning (MAS) NMR spectra were acquired at different stages of (de)sodiation to probe the structural features of various Na_xP phases in BP anodes (Fig. 13a). During the first sodiation, ³¹P NMR shifts consistent with P helices/zig-zags were present as early as 0.60 V (Na_{0.52}P), suggesting that P-P cleavage began at this composition. The ex situ XRD results also indicated that an amorphous or highly disordered phase was present in this stage in addition to residual BP. P helices were the predominant structural motifs that were subsequently formed after this stage. In addition, P helices and zig-zags (which are likely truncated helices) might allow some of the original alternating planar/ staggered P-P bonds found in pristine BP to persist while also forming an energetically favorable NarP phase. As BP was further sodiated to 0.38 V, the amorphous Na_xP phase (780 mA h g^{-1} , approximate composition of NaP) showed ³¹P sites consistent with P helices and terminal P units from short P chains (four-membered P zig-zags), as shown in Fig. 13b-e. The corresponding ex situ XRD results also indicated that this phase was amorphous (a-Na_rP). In addition, the researchers proposed that dumbbell-containing structures previously predicted by others did not form in a measurable quantity in these SIBs during cycling. Once the potential dropped below 0.22 V, the formation of a lower energy crystalline Na₃P-P6₃cm structure occurred. And a plateau at 0.21 V (ranging from 1358-2340 mA h g⁻¹, corresponding x to 1.57–2.70) was observed and could be attributed to the formation of c-Na₃P-P6₃cm, which is the most stable crystal structure of crystalline Na₃P upon sodiation based on DFT calculations. During desodiation, the c-Na₃P–P6₃cm ³¹P resonance at –207 ppm persisted until >0.60 V (>1490 mA h g^{-1} , approximately Na_{1.28}P), as shown in Fig. 13a. At 0.8 V (1730 mA h g^{-1} , approximately NaP), P motifs of helices could again be observed in a-NaP, but with longer average Na-P interatomic distances (Fig. 13g and h). Increased structural disorder was also observed and might be due to a larger range of clusters that were formed from the isolated P ions in Na₃P, as opposed to the original P-P bonded BP phase. At the end of desodiation, the researchers found that BP was not re-formed, which might contribute to the poor capacity retention observed in the Na-P system. And the results of NMR also indicated that the Na ions were not completely removed during desodiation and/or were consumed in deleterious side reactions.

Compared with pure theoretical calculations, the use of experimental methods can make it more authentic and concrete for understanding the mechanism of Na storage, and more leveraging of advanced and suitable detecting instruments to study the behavior of active materials in SIBs is necessary.

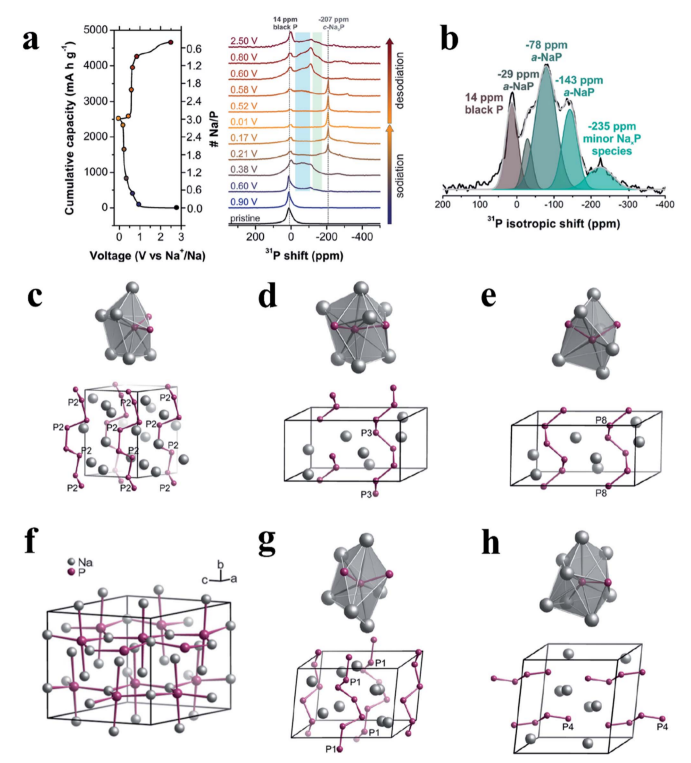


Fig. 13 (a) *Ex situ*³¹P MAS NMR spectra of BP recorded at various stages of sodiation/desodiation (right) and the corresponding electrochemistry (left). ³¹P chemical shift regions corresponding to P helical motifs and P near the end of chains are shaded in blue and green respectively. (b) Deconvolution of the ³¹P isotropic projection from 2D ³¹P phase-adjusted spinning sideband (PASS) experiments (black line) performed on a sample of approximate composition NaP, extracted from a SIB at 0.38 V (780 mA h g⁻¹) during sodiation. Five regions with centers-of-mass at 14 (BP), -29, -78, -143, and -235 ppm are observed. (c-e) The corresponding P sites (top) and extended P structures (bottom) of sideband patterns (calculated) for different isotropic chemical shifts, $\delta_{iso} = -29$ (c), -78 (d), and -143 ppm (e), extracted from a SIB at 0.38 V (780 mA h g⁻¹) during sodiation. (f) Crystal structures of Na₃P-P6₃*cm* (calculated). (g, h) The corresponding P sites (top) and extended P structures (bottom) of sideband patterns (calculated) for $\delta_{iso} = -29$ (g) and -78 (h), extracted from a SIB at 0.80 V (1730 mA h g⁻¹) during desodiation. The symbol of P_x, x = 2, 3, 8, 1, 4 (c-e, g and h), denotes individual ³¹P sites in certain structural models. Reproduced with permission.¹⁶⁸ Copyright 2018, American Chemical Society.

View Article Online Journal of Materials Chemistry A

To solve the issue of sluggish reaction kinetics, Dou et al.159 utilized nanoscale surface engineering to homogeneously deposit highly electron conductive poly(3,4ethylenedioxythiophene) (PEDOT) nanofibers onto special surface-modified BP nanosheets (Fig. 14a) and reported that this design could enhance charge transfer kinetics and super surface wettability with electrolytes, leading to enhancement in the reaction kinetics of sodium storage (Fig. 14b and c). In another study, Wang et al.¹⁷² used 4-nitrobenzenediazonium (4-NBD) to modify BP and further made it chemically bond with reduced graphene oxide to prepare a hybrid as an anode material. The researchers reported that due to enhanced interactions between the chemically functionalized BP and graphene, the cycling performance of the resulting SIB anode was significantly improved, exhibiting a specific capacity of 1472 mA h g^{-1} at 0.1 A g^{-1} in the 50th cycle and 650 mA h g^{-1} at 1 Ag^{-1} after 200 cycles. The results further indicated that these interactions could improve BP stability during long term cycling, reduce surface energies and increase BP thicknesses, all of which enabled the enlargement of channels between BP nanosheets and graphene, allowing more sodium ions to be stored for improved cycling performances. According to these studies, it can be seen that surface engineering is an effective way to improve the reaction kinetics of BP and thus can enhance the rate capability.

Similar to RP, direct combination with carbon-based materials is an effective approach to overcome the drawbacks of BP for Na storage. For example, Xu *et al.*¹⁷³ reported that a composite of ball milled BP/Ketjen-black/multiwalled carbon nanotubes (BPC) could exhibit high performances: a first-cycle reversible capacity of 2011.1 mA h g⁻¹, with an initial coulombic efficiency of 91.1% and a capacity of 1700 mA h g⁻¹ over 100 cycles at a current of 1.3 A g⁻¹. Similarly, Haghighat-Shishavan *et al.*¹⁷⁴ also prepared a composite of BP and multiwalled carbon nanotubes (BP-CNTs) through ball milling

process; in addition, they found that appropriate air exposure could successfully alter BP powder from a naturally hydrophobic form to a desired hydrophilic form, which is essential to the formation of stable bonds between BP and functionalized CNTs during ball milling. In this study, a binary polymeric binder of sodium carboxyl methyl cellulose-poly acrylic acid (NaCMC-PAA) was used to fabricate an anode; as a result, the resulting anode delivered a first discharge capacity of 2073 mA h g^{-1} at 0.2C for SIBs, with a high capacity retention of 75.3% after 200 cycles, and a rate capacity of 609 mA h g^{-1} at 2C was obtained. Here, the researchers suggested the strong bonds between BP and the hydroxyl groups of NaCMC and carboxylic acid groups of PAA played an important role for the improved Na storage performance of BP.¹⁷⁴ In another study, Li et al.¹⁷⁵ reported a BP/graphene (BP/G) 2D macroscopic heterostructure prepared by a co-exfoliation/electrophoretic deposition process. When used as a binder- and additive-free anode for SIBs, this composite demonstrated superior specific capacities of 2365 mA h g_P^{-1} , 1894 mA h g_P^{-1} and 1456 mA h g_P^{-1} at 100, 200 and 500 mA g_P^{-1} rates respectively and showed a stable cycling performance. This novel preparation method ensures the intimate contact of BP and graphene, and thus a binder- and additive-free anode can be obtained, not only increasing the specific capacity but also reducing the cost for practical application. Recently, a BP-graphene layered structure was synthesized directly from a RP-graphene precursor by a low-cost and scalable pressurization (8 GP) process at room temperature (Fig. 14d).¹⁶² These hybrids could also be directly used as anode materials without a conductive additive and binder for SIBs and showed excellent rate capabilities and cycling stabilities (\sim 1250 mA h g⁻¹ at 1 A g⁻¹ and \sim 640 mA h g⁻¹ at 40 A g⁻¹ after 600 cycles) (Fig. 14e). This study combines the synthesis of BP and the preparation of a BP-graphene hybrid anode in one step, offering excellent possibility for scalable production of composites for practical SIBs.

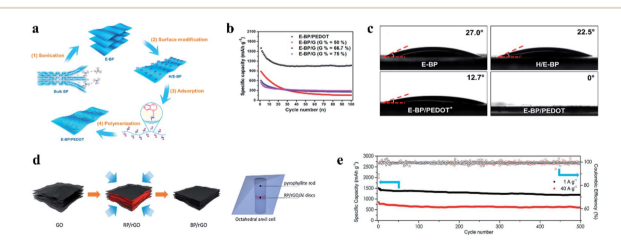


Fig. 14 (a) Schematic of the preparation of the E-BP/PEDOT architecture. (b) Cycling life of E-BP/PEDOT and E-BP/graphene hybrid electrodes with different ratios at 100 mA g^{-1} . (c) Characterization of wetting properties and optical images of contact angles between electrolytes and E-BP, H/E-BP, E-BP/PEDOT * (less PEDOT grown on BP nanosheets) and E-BP/PEDOT electrodes. (a–c) Reproduced with permission.¹⁵⁹ Copyright 2017, Elsevier Ltd. (d) Schematic description of the BP/rGO synthesis. (e) Cycling performance of the BP/rGO anodes at charge and discharge current densities of 1 and 40 A g^{-1} in black and red, respectively. (d, e) Reproduced with permission.¹⁶² Copyright 2018, American Chemical Society.

Besides carbon materials to be combined with BP, researchers have also explored composites with other types of materials. For example, Meng et al.¹⁷⁶ prepared a composite of BP quantum dots (BPQDs) anchored onto Ti₃C₂ nanosheets (TNSs) as an anode material for SIBs by a assembling process with the assistance of sonication and long-term stirring. They reported that due to the addition of TNSs, the BPQDs were endowed with improved conductivity and reduced stress during cycling, allowing for a high initial discharge capacity of \sim 723 mA h g⁻¹ and outstanding long-term cycling stabilities up to 1000 cycles with a capacity retention of nearly 100%. In another study, Song et al.¹⁷⁷ employed a simple high-energy mechanical milling approach to fabricate a novel and stable sodium titanate/carbon-BP (Na2Ti3O7/C-BP) hybrid as a SIB anode material and found that the BP nanoparticles could interconnect with bare NTO through P-O-Ti bonds and/or P-C bonds. As a result, the NTO/C-BP hybrid delivered specific capacities of \sim 225 mA h g⁻¹ at 20 mA g⁻¹ after 55 cycles and \sim 183 mA h g⁻¹ after 100 cycles at 100 mA g⁻¹. The researchers also suggested that the formation of stable P-C bonds and P-O-Ti bonds was a promising strategy to fabricate other Ti-based and P-based electrode materials for practical application in SIBs. However, the BP content in this composite is quite low (20%) which results in a reduced special capacity, and the large specific surface area of these ball milled hybrids causes a low initial coulombic efficiency (42%), which is not suitable for practical applications. Xia's group178 synthesized a composite of sulfur-doped BP-TiO₂ (TiO₂-BP-S) as an anode material for SIBs using a feasible and large-scale high energy ball milling method and obtained a discharge specific capacity of up to 490 mA h g^{-1} after 100 cycles at 50 mA g⁻¹ and good cycling stabilities with 290 mA h g^{-1} capacity retention after 600 cycles at 500 mA g^{-1} . In another study, Ding et al.¹⁷⁹ also designed a strategy to produce BP composites in which they first ball milled amorphous RP with cubic boron nitride to obtain a BP/c-BN composite, and then wrapped this composite by nanoporous graphene networks via a sonication method. In this composite, the super hard c-BN could not only reduce the size of RP during ball milling, but also act as a mechanical skeleton to suppress and buffer the large volume expansion of BP during the sodiation process. As a result, this BP/c-BN/Gr composite anode demonstrated high cyclability with a stable specific capacity of 947 mA h g⁻¹ and a \sim 90% capacity retention after 100 cycles at a current density of 50 mA g^{-1} . This is the first time that a super hard material was introduced into P-based anodes for SIBs; however, the high electron resistance of such c-BN may cause insufficient rate capability, and finding materials that possess both high mechanical strength and high electronic conductivity to host BP is a promising strategy for practical SIBs.

Phosphorene is a new 2D material that contains one or few layers of BP (Fig. 12a and b)⁸⁹ and can be obtained from bulk BP through various methods^{180–183} or even from white phosphorus.¹⁶⁴ These 2D phosphorene materials possess unique properties such as enlarged specific surface areas, increased interlayer distances,⁷² and fast electron transfer and Na diffusivity (energy barrier of only 0.04 eV for monolayer BP),^{184,185} and have attracted significant attention and witnessed development

in SIB applications. For example, Sun *et al.*⁷² synthesized a hybrid with several layers of phosphorene sandwiched between graphene layers and obtained a specific capacity of 2440 mA h g⁻¹ (based on phosphorus) at a current density of 50 mA g⁻¹ and a 83% capacity retention after 100 cycles. The results of *in situ* TEM and *ex situ* XRD indicated a dual Na storage mechanism involving the intercalation of sodium ions along the *x*-axis of phosphorene and the formation of Na₃P, which is consistent with the theoretical calculations carried out by other researchers.¹⁶⁷ Furthermore, the graphene layers in this hybrid were found to serve as a mechanical backbone and an electrical highway to accommodate the anisotropic expansion of phosphorene layers along the *y*- and *z*-axial directions for stable cycling operations, leading to excellent stability.⁷²

The lack of large-scale and low-cost methods to prepare phosphorene hinders its practical applications; however, developments have been made after continuous efforts, such as mechanical exfoliation and liquid-phase peeling.180,182,183,186,187 Recently, Huang et al.¹⁸⁸ developed a electrochemical cationic intercalation method to prepare phosphorene from BP crystals and applied it as an anode material for SIBs (Fig. 15a). Here, tetraalkylammonium cations were used to intercalate BP, and FL-P-5 meant that a potential of -5 V was applied to exfoliate bulk BP; similarly, FL-P-10 and FL-P-15 correspond to potentials of -10 and -15, respectively. The researchers found that the number of BP layers (from 2 to 11 layers) could be manipulated by changing the applied potential and that the BP layers were free of surface functional groups (Fig. 15b-d). As a result, this phosphorene anode showed excellent performances for SIBs, delivering a capacity of 1968 mA h g^{-1} after 15 cycles and \sim 1190 mA h g⁻¹ after 50 cycles at a current density of 100 mA g^{-1} (Fig. 15f and g).

3 Metal phosphides

Metal phosphides (MPs) are also promising anode materials for SIBs due to their noticeably high gravimetric and volumetric specific capacities, appropriate redox potentials and higher electrical conductivities as compared with pure phosphorus.77,189 The higher performance of MPs as SIB anodes can mainly be attributed to the presence of metal atoms acting as electronic pathways that drastically enhance the electronic conductivity.¹⁹⁰ In addition, conductive metal particles can be formed in MP anodes during battery cycling which further increases the conductivity of electrodes.91,189 Furthermore, in some MPs, metal nanocrystals are embedded in phosphorus matrices during cycling and thus inhibit the agglomeration of metals and improve the cycling performance of MP anodes. However, there are also drawbacks of MP anodes, including large volume variations and poor diffusion kinetics during cycling which is similar to phosphorus. To overcome these disadvantages and allow MPs to become more practical for SIBs, extensive investigations have been conducted and various MP nanostructures have been prepared, such as yolk-shell Sn₄P₃@C nanospheres,⁹¹ CoP nanowires,¹⁹¹ monodisperse Ni₂P particles92 and hollow FeP@carbon nanocomposites,192 all of which possess unique nanostructures or conductive matrices

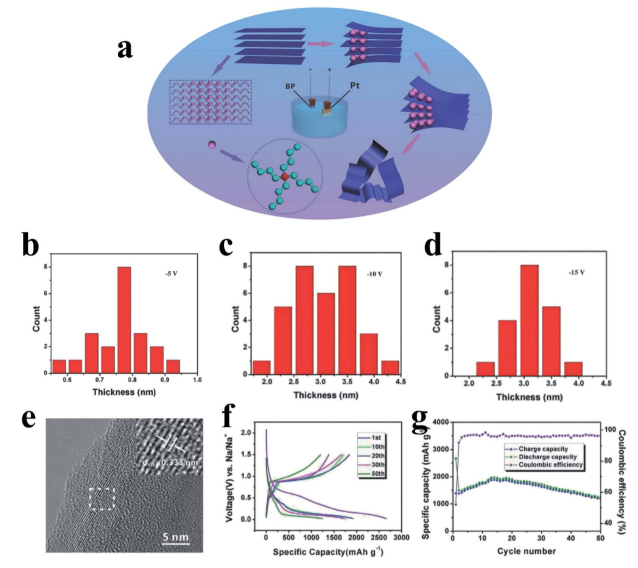


Fig. 15 (a) Schematic of the electrochemical exfoliation process of phosphorene. (b–d) The corresponding thickness distribution charts of FL-P-5. FL-P-10, and FL-P-15, respectively. (e) HRTEM image of FL-P-5. (f) Charge and discharge profiles of different cycles at 100 mA g^{-1} . (g) Cycling performances and coulombic efficiencies at 100 mA g^{-1} . Reproduced with permission.¹⁸⁸ Copyright 2017, WILEY-VCH.

that could accommodate volume expansion and offer more active sites for Na insertion/extraction and thus high electrochemical performances.

In general, the reaction mechanisms of MPs are more complex than those of pure phosphorus. Based on whether the metal elements in MPs can electrochemically react with sodium, MPs can be divided into metal-active phosphides and metal-inactive phosphides. For metal-active phosphides, both the metal element and P phase can react with Na, and in a typical reaction process, crystalline metal and P phases are formed after the first cycle through a conversion process and react with sodium during the subsequent cycles through a reversible alloying process respectively, as expressed as follows:^{74,77,91,193,194}

$$MP_x + (3x + y)Na^+ + (3x + y)e^- \rightarrow Na_yM + xNa_3P \qquad (2)$$

$$Na_{y}M \leftrightarrow M + yNa^{+} + ye^{-}$$
 (3)

$$Na_{3}P \leftrightarrow 3Na^{+} + P + 3e^{-}$$
(4)

Examples of metal-active phosphides include Sn_4P_3 (ref. 91 and 195) and GeP_5 .¹⁹³

For another kind of metal-active phosphide, crystalline metal and P phases can reform the original phase of MPs after desodiation, such as GeP₃ (ref. 196) and GeP,¹⁹⁷ and the process can be expressed as follows:

$$MP_x + (3x + y)Na^+ + (3x + y)e^- \leftrightarrow Na_yM + xNa_3P$$
 (5)

Similarly, metal-inactive phosphides also proceed through two main types of reactions: one allows for the reconversion to the original MP phase after desodiation whereas the other one cannot. In the former reaction, metallic and Na_xP phases are formed after sodiation and the metal and P atoms can re-bond to form MPs during desodiation. Examples of this type of material include CuP_2 ,¹⁸⁹ Cu_3P ,¹⁹⁸ NiP_3 (ref. 199) and CoP_3 (ref. 200) and the reaction process can be presented as eqn (5):

$$MP_x + 3xNa^+ + 3xe^- \leftrightarrow xNa_3P + M$$
(6)

As for the latter reaction, nanosized metallic nanocrystals and Na_xP phases are formed after metal–P bonds break during the first sodiation process. In the following desodiation process, a pure P phase is formed after Na-extraction and undergoes a reversible conversion reaction with Na in the subsequent cycles. Examples of this type of material include FeP^{201,202} and CoP^{203,204} and the reaction process can be expressed as follows:

$$MP_{x} + 3xNa^{+} + 3xe^{-} \rightarrow xNa_{3}P + M$$
(7)

$$Na_3P \leftrightarrow 3Na^+ + P + 3e^-$$
 (8)

Overall, details of Na-storage mechanisms for MPs have yet to be fully understood and more efforts and new advanced characterization methods are needed.

3.1 Tin phosphides

Tin phosphides such as Sn_4P_3 and SnP have attracted considerable attention because of their high theoretical volumetric specific capacity (6650 mA h cm⁻³ for Sn_4P_3 vs. 5710 mA h cm⁻³ for P)⁷⁷ and high electronic conductivity (30.7 S cm⁻¹ for Sn_4P_3 vs. $\sim 10^{-14}$ S cm⁻¹ for RP).^{77,80} Sn₄P₃ exhibits a rhombohedral lattice (JCPDS No. 71-2221) with a layered structure consisting of Sn layers and P layers packed alternately (Fig. 16a).^{205,206} Kim *et al.*⁷⁷ for the first time studied the electrochemical performance of Sn₄P₃ as a SIB anode obtained through ball milling and reported a high reversible capacity of 718 mA h g⁻¹ and

a high stability with negligible capacity fading over 100 cycles. In addition, the Sn₄P₃ sample in this study also showed an appropriately low redox potential of ~0.3 V ν s. Na/Na⁺, allowing the anode to achieve high energy densities in full cells. According to the reported literature,^{74,91,195} researchers suggested that the better cycle performance of Sn₄P₃ compared with Sn could be attributed to the confinement effect of Sn nanocrystallites in the amorphous phosphorus matrix during cycling. And the Na storage mechanism of Sn₄P₃ could be summarized based on their results,^{74,91,195} as expressed as follows:

$$Sn_4P_3 + 24Na^+ + 24e^- \rightarrow Na_{15}Sn_4 + 3Na_3P$$
 (9)

$$Na_{15}Sn_4 \leftrightarrow 4Sn + 15Na + 15e^{-}$$
(10)

$$Na_{3}P \leftrightarrow 3Na^{+} + P + 3e^{-}$$
(11)

According to the literature reported by Jung *et al.*,²⁰⁷ the sodiation/desodiation processes of Sn_4P_3 can be depicted as follows: in the initial stage of sodiation, the Na ions break the Sn–P bonds of Sn_4P_3 and react with P to form Na_xP due to the strong preference of Na for P over Sn; meanwhile, the Sn atoms are separated from P atoms and then join together to form local Sn particles. The Na_xP phase can surround the Sn particles because of its volume expansion with Na alloying, or Sn nanoparticles are dispersed in the amorphous-like P matrix. As sodiation continues, the Na_xP phase evolves into Na_3P , and the Sn particles evolve into Na_ySn by reacting with Na ions and eventually into $Na_{15}Sn_4$.⁷⁴ After full sodiation, Sn_4P_3 is changed to a morphology where $Na_{15}Sn_4$ particles are confined in a Na_3P matrix. After Na-extraction of these products, low crystallinity Sn nanoparticles are formed and highly dispersed in the

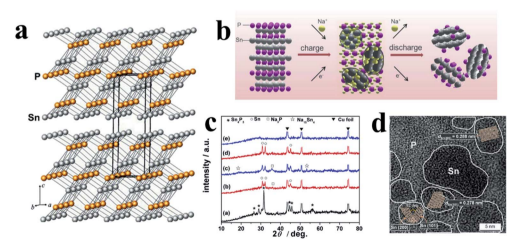


Fig. 16 (a) Optimized structure of hexagonal crystalline Sn_4P_3 ; orange and gray balls represent the Sn and P atoms, respectively. Reproduced with permission.²⁰⁶ Copyright 2006, Elsevier Inc. (b) Schematic illustration of the Na storage mechanism in the Sn_4P_3 electrode. Phase transformation of Sn_4P_3 by sodiation and desodiation (discharge) reactions. (c) *Ex situ* XRD patterns of the Sn_4P_3/C electrode in different charge and discharge states: (a) fresh electrode; (b) after the 1st charge to 0.25 V; (c) after completion of the 1st charge to 0.01 V; (d) after the 1st discharge to 0.5 V; and (e) full discharge to 2.0 V. Here the charge process represents sodiation. (c, d) Reproduced with permission.⁷⁴ Copyright 2014, American Chemical Society. (d) High magnification TEM image of Sn_4P_3 after the first desodiation (in an ionic liquid electrolyte). Reproduced with permission.²⁰⁸ Copyright 2017, American Chemical Society.

amorphous P host matrix. In the subsequent sodiation/ desodiation cycles, P and Sn undergo reversible alloving and de-alloying reactions, respectively. This is well consistent with the experimental results (Fig. 16c and d). In addition, this Na storage mechanism can well explain the excellent cycle performance of Sn_4P_3 . The core $(Na_{15}Sn_4)/matrix (Na_3P)$ morphology formed by initial sodiation has high structural integrity because the bulk modulus of Na₃P is 71% higher than that of Na₁₅Sn₄. Thus, Na₃P is much better able to withstand deformation than Na₁₅Sn₄, and effectively prevent the aggregation of the Sn particles in the next charge-discharge cycle, and this is largely responsible for the excellent cycle performance and high coulombic efficiency of Sn₄P₃.^{74,77,208} Furthermore, the Sn phase can change its morphology during sodiation-desodiation in response to P's volume changes; consequently, the void spaces could be filled by their morphological changes, preventing the electrical isolation of the active material layer.²⁰⁹ The structure of crystalline Sn₄P₃, the schematic illustration of the Na storage mechanism, and the experimental results are shown in Fig. 16a and b.

However, in the study carried out by Kim *et al.*,⁷⁷ the results of *ex situ* TEM and XRD analyses indicated that the conversion reaction of Sn_4P_3 to Na_xP and $Na_{15}Sn_4$ was reversible (Fig. 17). It is possible that the conversion reaction may be partially reversible, similar to the electrochemical behavior of Sn_4P_3 in LIBs.²¹⁰

In addition to the better cycle performance, Sn_4P_3 could also show excellent rate performance, due to not only its intrinsic high electronic conductivity, but also the Sn phase formed during the sodiation/desodiation processes, which could compensate for the poor electronic conductivity of the Na₃P phase.^{74,207,211} Moreover, Sn_4P_3 allows fast Na ion transport. The Na diffusivities in the intermediate product of amorphous

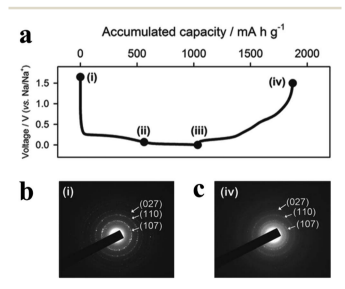


Fig. 17 (a) First galvanostatic charge and discharge profiles of the Sn_4P_3 electrode obtained without an FEC additive. The different points indicate (i) pristine, (ii) 0.07 V, (iii) 0 V and (iv) 1.5 V. (b, c) Electron diffraction (ED) patterns at the points (i) and (iv) in (a), respectively. Reproduced with permission.⁷⁷ Copyright 2014, WILEY-VCH.

 $Na_xSn_4P_3$ (x = 1.33-24) were found to be between 3.0×10^{-8} and 1.0×10^{-7} cm² s⁻¹, and for amorphous $Na_{15}Sn_4$ and Na_3P , they are 1.1×10^{-7} and 1.6×10^{-8} cm² s⁻¹, respectively.²⁰⁷ It was indicated that the Sn-rich environment around Na could effectively enhance the Na ion conductivity in Sn_4P_3 because Na ions could more easily break the weak Na–Sn bonds than the strong Na–P bonds during their transport.²⁰⁷ As a result, the fast electron/Na-ion transport could enable an excellent rate performance of Sn_4P_3 .

After Kim's pioneering study, a two phase nanocomposite of Sn_{4+x}P₃@Sn-P comprised of Sn_{4+x}P₃ crystals (size, ~30 nm) with an amorphous Sn-P coating layer (thickness, \sim 3 nm) has also been prepared by this ball milling method and showed an excellent Na storage performance.195 In another study, Wang et al.²¹² found that Sn agglomeration in pristine Sn₄P₃ particles during anode sodiation/desodiation was the major cause of performance fading. And they found that Sn agglomeration could be effectively suppressed through the introduction of TiC into Sn₄P₃ by a ball milling method. As a result, a significantly improved cycling stability has been obtained, delivering a capacity of 300 mA h g⁻¹ over 100 cycles at a current density of 100 mA g^{-1} (with 30 wt% TiC). It is noticeable that the performance of Sn₄P₃ in this study was different from that reported by Kim's group,77 which was probably caused by the different preparation conditions such as the molar ratio of precursors (Sn : P, 4 : 3 vs. 6 : 4) and the ball milling time (1 hour vs. 30) hours) and other reasons. Xu et al.213 also prepared a Sn₄P₃-P@graphene nanocomposite (Sn : P = 1 : 3) using ball milling and reported high rate capability retentions of >550 and 371 mA h g^{-1} at current densities of 1 and 2 A g^{-1} respectively over 1000 cycles as well as high rate capacities of >815, ~585 and \sim 315 mA h g⁻¹ at 0.1, 2 and 10 A g⁻¹ respectively. In this study, the ball milling process involved two steps: the mixing and milling of tin and RP powders under Ar atmosphere followed by the mixing and milling of the as-synthesized SnP₃ with graphene stacks under the same conditions to obtain the final Sn₄P₃-P@graphene nanocomposite.

However, similar to the preparation of RP, it is hard to obtain a desired small-size morphology or refined nanostructure for Sn₄P₃ via this mechanical approach. Nevertheless, many new nanostructured Sn₄P₃ samples have been reported by various preparation methods recently. For example, in situ phosphidation is often used to obtain a Sn₄P₃ nanostructure with Sn or SnO₂ as precursors. In most of the phosphidation processes for synthesizing metal phosphides (MPs), PH3 or gaseous RP is often used which can be generated from NaH_2PO_2 ($NaH_2PO_2 \cdot H_2O$) or solid RP under an inert atmosphere and high temperature conditions.90,202,214 Interestingly, researchers found that Sn could be phosphorized at low temperature by a solvothermal method. For instance, RP powder could be soluble in ethylenediamine during a solvothermal treatment, and then, the dissolved P species reacted with solid Sn and diffused towards the interior of the Sn particles to form Sn-P alloys.206,215 This mild preparation method avoids the toxic PH3 product or the white P by-product, during which lower temperature is needed. However, this solvothermal method could only be used to phosphorize the Sn precursor, and a traditional phosphorization approach with

 NaH_2PO_2 ($NaH_2PO_2 \cdot H_2O$) as the P source was often used for a SnO₂ precursor. For example, Liu *et al.*⁹¹ synthesized uniform yolk-shell Sn₄P₃@C nanospheres as anode materials for SIBs by a stepwise preparation approach, involving a hydrothermal method to obtain a SnO₂ template, a reduction process to obtain Sn, and a phosphorization process to transform Sn into Sn₄P₃ (Fig. 18a). During the reduction and phosphorization process, the hollow SnO₂ template transformed into one big nanoparticle and several small nanoparticles of Sn which were encapsulated in each carbon nanocage, and evenly formed a yolk-shell Sn₄P₃@C nanostructure. Due to the unique nanostructure, this Sn₄P₃@C anode exhibited a high reversible capacity of 790 mA h g^{-1} at 100 mA g^{-1} , superior rate capacities of 720, 651, 581, 505 and 421 mA h g^{-1} at 0.2C, 0.4C, 0.8C, 1.5C and 3C respectively (1C = 1000 mA g^{-1}) and stable cycling performances with a high capacity of 360 mA h g⁻¹ at 1.5C after 400 cycles. Similarly, Ma et al.²¹⁶ designed an integrated anode material of Sn₄P₃@C with a yolk-shell nanocube structure for SIBs by phosphorization of a Sn precursor and found that the voids and spaces between

Sn₄P₃ and carbon nanocubes could effectively buffer volume expansion during charge/discharge and that the highly conductive carbon material could promote fast electron transfer. As a result, this SIB anode exhibited a high discharge capacity of 701 mA h g^{-1} at 0.1 A g^{-1} after 50 cycles, remarkable rate capability of 508 mA h $\rm g^{-1}$ even at 2.0 A $\rm g^{-1}$ and highly stable cycling performances with 516 and 368 mA h g^{-1} capacity retention after 500 cycles at 1.0 and 2.0 A g^{-1} respectively. Furthermore, Choi et al.217 fabricated size-adjustable core-shell Sn₄P₃-C nanospheres using a carbonization/reduction and phosphorization process and reported that among the different sizes of Sn₄P₃-C nanospheres, the 140 nm Sn₄P₃-C nanosphere electrode demonstrated the highest reversible capacity and rate capability, with an ultra-long cycle stability with 420 mA h g^{-1} capacity retention after 2000 cycles at a current density of 2000 mA g^{-1} . In addition, researchers have also reported that the cycling life of Sn_4P_3/C could be largely extended through the reduction of particle size to the nanoscale which could reduce the pulverization of anodes.217

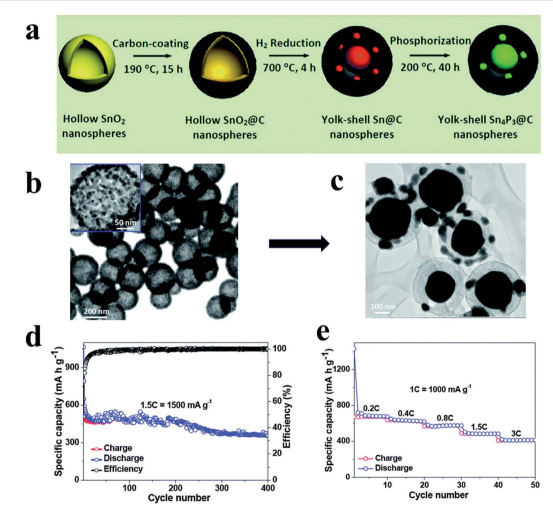


Fig. 18 (a) Schematic of the fabrication of uniform yolk-shell Sn_4P_3 @C nanosphere anodes. (b) TEM image of uniform SnO_2 hollow nanosphere templates; the inset shows a typical SnO_2 hollow nanosphere with a shell thickness of about ~25 nm. (c) TEM images of a uniform yolk-shell Sn_4P_3 @C nanosphere; (d) Long cycling performance of yolk-shell Sn_4P_3 @C nanospheres at 1.5C. (e) Rate capability at different rates (increased from 0.2C to 3C). $1C = 1000 \text{ mA g}^{-1}$. Reproduced with permission.⁹¹ Copyright 2015, The Royal Society of Chemistry.

These studies indicate that the combination of Sn_4P_3 with carbon materials is also an effective strategy to improve the performance of Na storage, and the refined nanostructures can give significant advantages for practical SIBs.

Shin et al.²¹³ also prepared Sn–P compounds (Sn₄P₃, Sn_{4–x}P₃ or Sn_4P_{3+x}) as anode materials for SIBs by this simple solvothermal method and found that the temperature and time period used for solvothermal treatment played a critical role in determining the phase and composition of the Sn-P compound and thus its electrochemical behavior. When solvothermal synthesis was conducted at lower or higher temperatures than a certain critical value (e.g., 200 °C), the resulting compound contained unreacted (residual) Sn or excess P species. The Sn₄P₃ compounds (Sn₄P₃ mixed with \sim 1.7% Sn) prepared with the optimized solvothermal parameters (T = 200 °C and t = 40 h) showed the best electrochemical Na storage performance. Yin et al.²¹⁸ designed a Sn₄P₃/reduced graphene oxide nanohybrid (Sn₄P₃/RGO) similarly by this low-temperature solution-based phosphorization reaction route. In the preparation process, NaBH₄ was used as the reducing agent to reduce Sn^{2+} and GO, with a Sn/RGO nanohybrid formed, and then the Sn/RGO nanostructures were used as the precursor and RP as the P source to synthesize Sn₄P₃/RGO by a solvothermal approach at 200 °C. Due to the hybrid structure and the high electronic conductivity of RGO, this nanohybrid SIB anode exhibited a high reversible capacity, an enhanced rate capability, and a superior long cycling life (362 mA h g^{-1} after 1500 cycles at a current density of 1.0 A g^{-1}). In another study, a novel morphology of pure phase and size-controllable Sn₄P₃ nanotops was developed through a novel facile solution chemical method using Sn and trioctylphosphine (TOP) as raw materials.²¹⁹ This Sn₄P₃ nanotop anode showed a good electrochemical

performance for SIBs, providing an initial capacity of 719.8 mA h g⁻¹ at a current density of 50 mA g⁻¹, an initial coulombic efficiency of \sim 72.7%, and good cyclability and stability (543 mA h g⁻¹ at 200 mA g⁻¹ for 80 cycles).

Alternatively, a high temperature atmosphere phosphidation approach with SnO₂ as the precursor is another effective way to prepare Sn₄P₃ nanostructures. Compared with the solvothermal method, which often involves low temperature but a long process time (T = 200 °C and t = 40 h), the atmosphere phosphidation approach usually needs higher temperature but a shorter process time (~ 300 °C, less than a few hours) with NaH₂PO₂ (NaH₂PO₂·H₂O) as the P source.

As the initial coulombic efficiency (CE) of Sn₄P₃/C nanocomposites was found to be inefficient (40-50%)⁹¹ and it is challenging to achieve both long cycle life and high CE for high capacity tin phosphide anodes, Fan et al.²⁰⁵ reported an anode composed of pomegranate-structured Sn₄P₃@C spheres facile aerosol spray-pyrolysissynthesized using а phosphidation method. In brief for this process, the transparent precursor containing SnSO4 and sucrose was atomized into microsized droplets (0.1-2 µm) and then these droplets were pyrolyzed in a 600 °C furnace and transformed into SnO₂/C spheres (0.1-2 µm); finally, pomegranate-structured Sn₄P₃@C spheres were obtained after a phosphidation process with NaH₂PO₂ as the P source (Fig. 19a and c). It can be seen from the TEM images (Fig. 19b) that Sn_4P_3 nanoparticles with a size of 20-40 nm were encapsulated by thin carbon layers. The Sn₄P₃@C anode was able to provide a reversible capacity of \sim 800 mA h g⁻¹ with a record-high initial CE (>90%) and high cycle CE (~99.9%) for over 120 cycles at a current of 100 mA g^{-1} in an ether-based electrolyte (Fig. 19d). Here, the researchers suggested that to achieve both long cycling life and high CE,

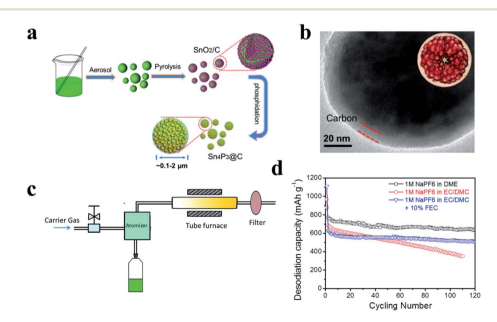


Fig. 19 (a) Schematic illustration of the fabrication process for $Sn_4P_3@C$ pomegranates. (b) TEM image of the as-synthesized pomegranatestructured $Sn_4P_3@C$ sphere; inset of (b) shows the ideal pomegranate structure. (c) Schematic illustration of the fabrication of the Sn_2/C spherical composite using the aerosol-spray-pyrolysis method. (d) Cycling performances of the pomegranate-structured $Sn_4P_3@C$ nanosphere at 100 mA using different electrolytes. Reproduced with permission.²⁰⁵ Copyright 2017, Elsevier Ltd.

ideal anode materials should possess microsized primary particles consisting of nanosized secondary high capacity active materials and nanopores in a carbon matrix, allowing for the formation of a SEI on only the primary particle surface and the maintenance of the electronically conductive nanoporous composite integrity during charge/discharge cycles. Based on this, the pomegranate-structured Sn_4P_3 @C spheres synthesized in this study were in accordance with the ideal anode they referred to.

Recently, Sn₄P₃ microspheres encapsulated in hollow carbon spheres $(Sn_4P_3(@C))$ were prepared by this phosphorization route using SnO₂@C as the precursor and NaH₂PO₂ as the P source. The void volume of this nanostructure and the elasticity of the protective carbon spherical shell could efficiently mitigate the volume change of active Sn₄P₃ microspheres (200 nm) during charge/discharge processes and form a stable solid electrolyte interface film on the surface of the carbon shell. As a result, the Sn₄P₃@C anode for SIBs showed a stable specific capacity of 420 mA h g^{-1} after 300 cycles at a current density of 0.2 A g^{-1} , as well as excellent rate capabilities of 253, 175, 111, 78, and 50 mA h g^{-1} at densities of 1, 2, 5, 10, and 20 A g^{-1} , respectively. More importantly, this anode also showed an ultralong cycling performance, delivering stable capacities of 205 and 103 mA h g^{-1} at large current densities of 2 and 5 A g^{-1} after 4000 cycles, respectively.

In addition to Sn₄P₃, tin monophosphide (SnP) also shows promise as anode materials for SIBs, possessing a high theoretical capacity of 1209 mA h g^{-1} due to its high P ratio. SnP exhibits a trigonal crystallographic phase and shows a layered structure consisting of Sn-P-P-Sn sandwiches that are stacked on top of each other.²²⁰ However, SnP has not yet been extensively explored in energy storage devices due to its challenging synthesis conditions. To address this, Liu et al. 190 synthesized spherical SnP nanocrystals (NCs) with a size of 34 \pm 5 nm through a liquid phase synthesis method. And a SIB based on the SnP NC anode with sodium(I) bis(fluorosulfonyl)imide (NaFSI) as the electrolyte exhibited a high reversible capacity of 600 mA h g^{-1} at 100 mA g $^{-1}$ and enhanced cycling stabilities for over 200 cycles. Here, the researchers suggested that the high cycling performance was associated with both the small size of the crystal domain and the particular composition and phase of SnP, which could prevent mechanical disintegration and major phase separation during charge/discharge.

Another tin phosphide material SnP₃, which has the highest P mass ratio among tin phosphides such as Sn₄P₃ and SnP, exhibits the highest theoretical capacity of 1616 mA h g^{-1.194} Fan *et al.*¹⁹⁴ prepared a composite of SnP₃/C by a simple ball milling method and demonstrated that SnP₃/C could reversibly react with Na through conversion and alloy reaction processes as SIB anodes. The corresponding anode could provide a high capacity of ~810 mA h g⁻¹ at a current density of 150 mA g⁻¹ without capacity fading after 150 cycles and was also able to retain 400 mA h g⁻¹ even at 2560 mA g⁻¹. Furthermore, detailed analysis using SEM, TEM and XPS revealed that the strong bonding interaction between Sn and P could successfully mitigate the pulverization and aggregation of Sn and P, allowing for the partial self-healing of the SnP₃/C composite structure and enhanced cycling stabilities.

3.2 Cobalt phosphides

Cobalt phosphides can also electrochemically react with Na.^{90,203} Although metallic Co is inactive with Na, researchers reported that it can act as a conductive matrix to buffer volume expansion and enhance electron transfer. CoP is the most commonly used cobalt phosphide for SIBs and usually exhibits an orthorhombic phase, as shown in Fig. 20a.^{221,222} Liu *et al.*²⁰³ synthesized nanosized CoP particles as an anode for SIBs through ball milling metal Co and commercial RP. The anode delivered a high initial specific capacity of 770 mA h g⁻¹ and high rate capabilities, demonstrating that CoP is a promising anode candidate for sodium ion storage. In addition, *ex situ* XPS and STEM were carried out to investigate the sodium storage mechanism of CoP in this study and revealed that the sodium storage of CoP was based on a reaction between P and Na and can be summarized as follows:

 $CoP + 3Na^{+} + 3e^{-} \rightarrow Co + Na_{3}P$ (first discharge) (12)

$$Na_3P \leftrightarrow 3Na^+ + P + 3e^-$$
 (subsequent cycles) (13)

It means that CoP decomposed into Co and P after the initial charge–discharge process and in the next cycles, element P reversibly reacts with Na to form Na₃P with intermediate phases (Na₂P, NaP and NaP₇), being consistent with other literature.²⁰⁴ Table 2 summarizes the recent progress of phosphides used as an anode for SIBs.

Similar to RP, CoPs obtained from ball milling methods often suffered from fast capacity fading during cycling and inferior rate performance due to the large and uneven particle size; however, complex and ultrafine nanostructures could be synthesized by methods such as hydrothermal reactions and solution phase processes for CoP, giving an effective approach to solve these issues. For example, Ge et al.204 developed a strategy to synthesize MOF derived core/shell structured CoP@C polyhedrons anchored onto 3D reduced graphene oxide (RGO) on nickel foam (NF) as a binder-free SIB anode. Here, the unique core/shell CoP@C polyhedral structure was obtained through an in situ low-temperature phosphidation process derived from ZIF-67 and was further combined with RGO/NF which could efficiently act as a binder and electrical conductor. As a result, the corresponding anode exhibited remarkable electrochemical performances and delivered a specific capacity of 473.1 mA h g^{-1} at a current density of 100 mA g^{-1} after 100 cycles. In another example, Kang *et al.*²²³ designed a nanostructure of small size CoP nanoparticles uniformly embedded in N-doped C nanosheets (CNSs) by a simple one-step calcination process with a Co-based MOF and RP as precursors. Due to the small CoP particles (~11.3 nm), the strong P-C bonds and the high conductivity of the CNSs, a resulting anode using this composite delivered a Na-storage capacity of 598 mA h g⁻¹ at 100 mA g⁻¹ (pure CoP, 831 mA h g⁻¹), a high rate capability of 174 mA h g⁻¹ at 20 A g⁻¹ and a long-term cyclability of 98.5% capacity retention after 900 cycles at 1 A g^{-1} . Furthermore, Zhang *et al.*¹⁹¹ introduced a conducting polymer (poly-pyrrole, PPy) coating layer to protect

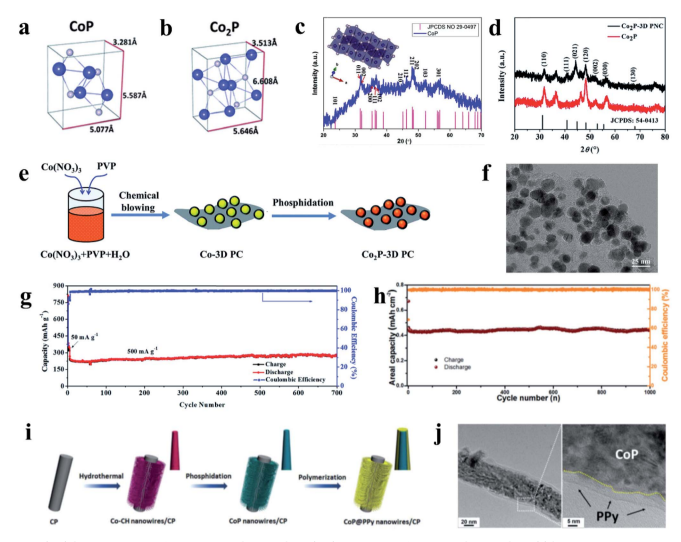


Fig. 20 (a, b) Crystal structures of orthorhombic CoP and Co₂P. (c, d) XRD pattern of crystalline CoP and Co₂P. (e) Schematic representation of the synthesis process of the Co₂P-3D PNC composite. (f) TEM image of the Co₂P-3D PNC composite. (g) Long-term cycling performance of the Co₂P-3D PNC composite at 500 mA g⁻¹. (h) Long-term cycling stability of the CoP@PPy NWs/CP electrode at 1.5 mA cm⁻². (i) Schematic diagram of the fabrication process of the CoP@PPy nanowires/CP electrode. (j) TEM images of CoP@PPy NWs. (a, b) Reproduced with permission.²²¹ Copyright 2011, The Royal Society of Chemistry. (c) Reproduced with permission.²⁰³ Copyright 2015, Elsevier B.V. (d–g) Reproduced with permission.²²⁴ Copyright 2018, The Royal Society of Chemistry. (h–j) Reproduced with permission.⁹¹ Copyright 2018, WILEY-VCH.

1D nanostructured CoP nanowires and fabricated a freestanding binder free SIB anode. Here, the synthesis process involved three steps, including the initial growth of 1D Cobased precursors on carbon paper using a hydrothermal method, a phosphidation process to obtain CoP nanowires and then a coating of the polymer PPy onto the surface of the CoP nanowires through polymerization and oxidation of pyrrole monomers (Fig. 20i and j). The researchers proposed that the obtained 1D core-shell CoP@PPy NWs/CP not only enhanced charge transfer and ion diffusion but also buffered volume expansion during cycling. As a result, these anodes exhibited superb electrochemical performance, delivering a high areal capacity of 0.521 mA h cm⁻² at 0.15 mA cm⁻² after 100 cycles and 0.443 mA h cm⁻² at 1.5 mA cm⁻² even after 1000 cycles (Fig. 20h), and a significant areal discharge capacity of 0.285 mA h cm⁻² even at a high current density of 3 mA cm⁻².

Other cobalt phosphides such as Co-rich phase Co₂P (Fig. 20b) and phosphorus rich phases CoP₃ and CoP₄ have also been reported as viable anode materials for SIBs. For example, Zhou et al.²²⁴ prepared a composite of Co₂P nanoparticle encapsulated 3D porous N-doped carbon nanosheet networks (3D-PNC) using a blowing method and an in situ phosphidation process (Fig. 20e). This unique Co₂P/3D-PNC nanostructure exhibited enhanced cycling stability and rate capability with a high initial discharge capacity of 827 mA h g^{-1} at 50 mA g^{-1} and 271 mA h g^{-1} after 700 cycles at 500 mA g^{-1} (Fig. 20g). Jin et al.²⁴⁰ utilized a cobalt-based metal organic framework (ZIF-67) as a self-template to obtain a nanostructure of Co₂P nanoparticles hybridized with N-doped carbon matrices (Co2P@N-C). For the preparation process, they first prepared a CoP@N-C nanostructure, and then combined this CoP@N-C with reduced graphene oxide by a simple hydrothermal method to form

Journal of Materials Chemistry A

Table 2 Summary of MPs and other phosphide anodes for SIBs

Materials	Synthetic method	Cycling performance	Rate capability	Initial coulombic efficiency	Ref.
Sn ₄ P ₃ powder	Ball milling	\sim 718 mA h g ⁻¹ , at 100 mA g ⁻¹ , 100th	_	~78%	77
Sn ₄ P ₃ @C	Hydrothermal/ phosphorization	360 mA h g^{-1} , at 1000 mA g^{-1} , 400th	360 mA h g ⁻¹ , 3000 mA g ⁻¹	_	91
Sn ₄ P ₃ /TiC	Ball milling	300 mA h g^{-1} , at 100 mA g^{-1} , 100th	_	85.5%	212
Sn ₄ P ₃ – P@graphene	Ball milling	607 mA h g^{-1} , at 1000 mA g^{-1} , 800th	371 mA h g ⁻¹ , 1000 mA g ⁻¹	73%	213
Sn_4P_3 nanotop	Solution chemistry method	543 mA h g^{-1} , at 200 mA g^{-1} , 80th	399 mA h g ⁻¹ , 1000 mA g ⁻¹	72.7%	219
Sn ₄ P ₃ –C sphere	Hydrothermal/ phosphorization	420 mA h g ⁻¹ , at 2000 mA g ⁻¹ , 2000th	260 mA h g ⁻¹ , 4000 mA g ⁻¹	60%	217
Sn ₄ P ₃ @C sphere	Aerosol spray-pyrolysis	700 mA h g ⁻¹ , at 100 mA g ⁻¹ , 120th	${\sim}500~\text{mA}~\text{h}~\text{g}^{-1}$, 800 mA g^{-1}	90.7%	205
Sn ₄ P ₃ @C yolk– shell	Liquid phase/sintering/ solvothermal	701 mA h g ⁻¹ , at 100 mA g ⁻¹ , 50th	508 mA h g ⁻¹ , 2000 mA g ⁻¹	63.7%	216
SnP ₃ @C	Ball milling	810 mA h g ⁻¹ , at 150 mA g ⁻¹ , 150th	${\sim}400~\text{mA}~\text{h}~\text{g}^{-1}$, 2560 mA g^{-1}	71.2%	194
SnP nanocrystal	Liquid phase synthesis	600 mA h g ⁻¹ , at 100 mA g ⁻¹ , 200th	396 mA h g ⁻¹ , 2500 mA g ⁻¹	_	190
СоР	Ball milling	315 mA h g ⁻¹ , at 100 mA g ⁻¹ , 25th	80 mA h g ⁻¹ , 2000 mA g ⁻¹	65.2%	203
CoP ₃ @C	Ball milling	144 mA h g ⁻¹ , at 100 mA g ⁻¹ , 320th	136.4 mA h g ⁻¹ , 250 mA g ⁻¹	87%	200
CoP@C polyhedron	Calcination/ phosphidation	473 mA h g ⁻¹ , at 100 mA g ⁻¹ , 100th	253.6 mA h g ⁻¹ , 800 mA g ⁻¹	47.3%	204
CoP/CNS	Calcination	386 mA h g^{-1} , at 1000 mA g $^{-1}$, 900th	174 mA h g ⁻¹ , 20 A g ⁻¹	64.6%	223
Co ₂ P-3D PNC	Blowing method/ phosphidation	271 mA h g^{-1} , at 500 mA g^{-1} , 700th	179 mA h g ⁻¹ , 3000 mA g ⁻¹	41%	224
CoP ₄ /CF	Hydrothermal/ phosphidation	851 mA h g^{-1} , at 300 mA g^{-1} , 300th	535 mA h g ⁻¹ , 4000 mA g ⁻¹	53%	90
FeP nanoparticle	Ball milling	321 mA h g^{-1} , at 50 mA g^{-1} , 60th	60 mA h g ⁻¹ , 500 mA g ⁻¹	60.2%	201
FeP nanoarray	Hydrothermal/ phosphidation	548 mA h g ⁻¹ , at 200 mA g ⁻¹ , 100th	184 mA h g ⁻¹ , 2000 mA g ⁻¹	55.7%	202
CNT@FeP@C	Solution phase	415 mA h g ⁻¹ , at 100 mA g ⁻¹ , 100th	268 mA h g ⁻¹ , 1500 mA g ⁻¹	59%	214
FeP ₄	Ball milling	1000 mA h g^{-1} , at 89 mA g^{-1} , 30th	900 mA h g ⁻¹ , 3578 mA g ⁻¹	84%	70
H–FeP@C@GR	Solvothermal/ phosphidation	446 mA h g ⁻¹ , at 100 mA g ⁻¹ , 100th	237 mA h g ⁻¹ , 1600 mA g ⁻¹	74%	192
NiP ₃	Ball milling	1079 mA h g^{-1} , at 176 mA g^{-1} , 15th	720 mA h g ⁻¹ , 1760 mA g ⁻¹	_	199
Yolk-shell Ni ₂ P	Hydrothermal/ phosphidation	181 mA h g^{-1} , at 200 mA g^{-1} , 100th	101 mA h g ⁻¹ , 2000 mA g ⁻¹	52%	225
NiP ₃ -CNT	Ball milling	668 mA h g^{-1} , at 200 mA g^{-1} , 120th	494 mA h g ⁻¹ , 3200 mA g ⁻¹	71.3%	226
Ni ₂ P/C	Solution phase/heat treatment	361 mA h g^{-1} , at 100 mA g^{-1} , 300th	100 mA h g ⁻¹ , 5000 mA g ⁻¹	76.7%	92
Ni ₂ P/graphene/ Ni ₂ P	Hydrothermal/ phosphidation	188 mA h g ⁻¹ , at 500 mA g ⁻¹ , 300th	152 mA h g ⁻¹ , 2000 mA g ⁻¹	69%	227
CuP_2/C	Ball milling	\sim 450 mA h g ⁻¹ , at 50 mA g ⁻¹ , 30th	178 mA h g ⁻¹ , 2000 mA g ⁻¹	$\sim 62\%$	189
Carbon sheet- CuP ₂	Solution phase/ phosphidation	410 mA h g ⁻¹ , at 80 mA g ⁻¹ , 200th	152 mA h g ⁻¹ , 2560 mA g ⁻¹	87%	228
CuP ₂ Cu ₃ P/C	Solution phase/	200th 221 mA h g ⁻¹ , at 100 mA g ⁻¹ , 100th	142 mA h g ⁻¹ , 2000 mA g ⁻¹	73%	198
GeP ₅ /C	annealing Ball milling	100th 1220 mA h g^{-1} , at 100 mA g^{-1} , 60th	900 mA h g ⁻¹ , 1500 mA g ⁻¹	92%	193
GeP ₅ /AB/p-rGO	Ball milling	400 mA h g ⁻¹ , at 500 mA g ⁻¹ ,	175 mA h g ⁻¹ , 5000 mA g ⁻¹	60%	229

Materials	Synthetic method	Cycling performance	Rate capability	Initial coulombic efficiency	Ref.
GeP _x	High temperature solvothermal	704 mA h g ⁻¹ , at 240 mA g ⁻¹ , 100th	130 mA h g ⁻¹ , 18 A g ⁻¹	82.6%	230
MoP nanorod/C	Solid phase reaction	398 mA h g ⁻¹ , at 240 mA g ⁻¹ , 100th	115.6 mA h g ⁻¹ , 1600 mA g ⁻¹	${\sim}40\%$	231
Zn_3P_2 nanowires	Chemical vapor deposition	${\sim}600~{\rm mA~h~g^{-1}},$ at 100 mA ${\rm g^{-1}},$ 85th	280 mA h g ⁻¹ , 5000 mA g ⁻¹	55%	232
InP	Ball milling	500 mA h g ⁻¹ , at 50 mA g ⁻¹ , 100th	_	—	209
SiP_2	Ball milling	572 mA h g ⁻¹ , at 100 mA g ⁻¹ , 15th	_	72%	233
Se_4P_4	Ball milling	804 mA h g ⁻¹ , at 50 mA g ⁻¹ , 60th	332 mA h g ⁻¹ , 3000 mA g ⁻¹	70.6%	234
P_2S_5/C	Ball milling	394 mA h g^{-1} , at 100 mA g^{-1} , 100th	114 mA h g ⁻¹ , 2000 mA g ⁻¹	—	235
NiPS ₃ nanosheet	Solution phase/ phosphidation	902 mA h g ⁻¹ , at 50 mA g ⁻¹ , 4th	316 mA h g ⁻¹ , 5000 mA g ⁻¹	70%	236
Cu ₄ SnP ₁₀ nanowires	High temperature solution phase	512 mA h g ⁻¹ , at 100 mA g ⁻¹ , 100th	412 mA h g ⁻¹ , 1000 mA g ⁻¹	63%	237
FeSi ₄ P ₄	Ball milling	180 mA h g^{-1} , at 100 mA g^{-1} , 100th	54 mA h g ⁻¹ , 4000 mA g ⁻¹	72%	238
SnPO/rGO	Hydrothermal	420 mA h g ⁻¹ , at 50 mA g ⁻¹ , 100th	200 mA h g ⁻¹ , 2000 mA g ⁻¹	53.2%	239
RP@Ni–P core@shell	Solution phase	1256 mA h g^{-1} , at 260 mA g^{-1} , 200th	491 mA h g ⁻¹ , 5200 mA g ⁻¹	88.2%	86

a Co₂P@N-C@rGO composite. The anode of this composite exhibited an excellent sodium storage performance with a high reversible capacity of 225 mA h g^{-1} at 50 mA g^{-1} after 100 cycles. In addition, they proposed that Co₂P underwent a Na storage mechanism similar to that of CoP, and decomposed into Co and P after the first discharge process, and then the P reversibly reacted with Na during the subsequent charge/discharge processes. In another study, Zhang et al.200 synthesized carbon-coated CoP₃ (CoP₃@C) nanocomposites as anodes for SIBs using a high energy mechanical milling method. Compared with plain CoP₃, this carbon-coated CoP₃ exhibited larger discharge capacities, enhanced cycling lifespans and better rate capabilities with a capacity of 212 mA h g^{-1} after 80 cycles at a current of 40 mA h g⁻¹. Sun et al.⁹⁰ fabricated a binder-free anode for SIBs by growing mesoporous CoP4 nanoarrays onto carbon felt by a hydrothermal and a phosphidation process and reported that this unique mesoporous nanoarray contained a secondary structure of ~10 nm ultrasmall primary particles. With the 3D carbon felt as a conductive network, this nanostructure demonstrated a capacity of 851 mA h g^{-1} after 300 cycles with an average coulombic efficiency of above 98%, a long cycling lifespan of up to 1000 cycles (>90% retention rate) and a high rate capability of 535 mA h g^{-1} at a current density of 4 A g^{-1} . This CoP₄/CF anode was also used in a full SIB with a $Na_3V_2(PO_4)_2F_3$ cathode and an average operating voltage of ≈ 3.0 V, a reversible capacity of 553 mA h g⁻¹ and a high energy density of \approx 280 W h kg⁻¹ were achieved.

In summary, these studies have shown the possible application of cobalt phosphides as anodes in SIBs, and refined Co–P nanostructures such as polyhedrons, nanowires, and nanoarrays often exhibit better Na storage performance than ball milled Co–P particles. However, the high cost of Co may hinder their practical application, especially for large-scale energy storage systems.

3.3 Iron phosphides

Iron phosphides such as FeP, FeP₂ and FeP₄ are also considered as promising anode materials for SIBs because of their high abundance, low-cost and environmentally friendly characteristics. Although Fe in FeP_x is electrochemically nonreactive towards Na, it can be used as a conductive matrix to relieve the volumetric expansion of P.²¹⁴ Because of this, great efforts have been devoted to the exploration of FeP_x as SIB anode materials. Among these various iron phosphides, FeP is most commonly used in SIBs and exhibits an orthorhombic phase (Fig. 21a)^{241,242} and possesses a theoretical capacity of 924 mA h g⁻¹. Li *et al.*²⁰¹ for the first time synthesized a low-cost FeP anode for SIBs with a high initial capacity of 764.7 mA h g⁻¹ through simple ball milling. The result of *ex situ* XRD and TEM indicated that the sodium storage mechanism of FeP was based on a conversion reaction, which can be expressed as follows:

$$FeP + 3Na^{+} + 3e^{-} \rightarrow Fe + Na_{3}P$$
(14)

$$Na_{3}P \leftrightarrow 3Na^{+} + P + 3e^{-}$$
(15)

The Na storage mechanism of FeP is similar to that of CoP: after the initial discharge process, FeP is converted into Co and

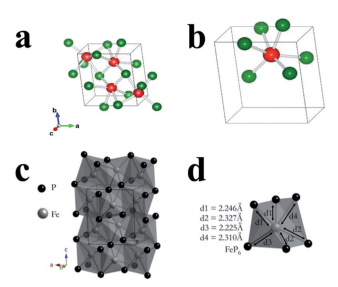


Fig. 21 (a, b) FeP crystal structure has an orthorhombic structure, and the iron atom in FeP has only one type of coordination: six. Red and orange atoms represent Fe and green atoms indicate P. Reproduced with permission.²⁴¹ Copyright 2018, Elsevier Ltd. (c) Crystal structure of orthorhombic FeP in another form. (d) FeP₆ polyhedron in orthorhombic FeP. The grey and black spheres represent Fe and P atoms, respectively. Reproduced with permission.²⁴² Copyright 2015, Elsevier B.V.

P, and then element P reversibly reacts with Na to form Na_3P in the subsequent charge/discharge processes.

In another study, Wang et al.202 reported a SIB anode composed of FeP nanorod arrays on carbon cloth (FeP NAs/CC) (Fig. 22a), which delivered a high capacity of 829 mA h g^{-1} at 0.1 A g^{-1} and a high stability of 548 mA h g^{-1} at 0.2 A g^{-1} after 100 cycles with a capacity retention of 99.8%, and an excellent rate and cycle performance (Fig. 22b and c). The synthesis process was simple, involving growth of Fe₂O₃ nanorod arrays onto carbon cloth through a hydrothermal method followed by a phosphidation process to transform Fe₂O₃ into FeP (Fig. 22a). In another study, Han et al.²¹⁴ developed a synthetic strategy to construct carbon-coated FeP with an amorphous and mesoporous framework anchored onto carbon nanotubes as a SIB anode. The synthesis process involved a silica coating as a sacrificial internal spacer, which appeared to be key in the formation of a mesoporous framework. As a result, this FePbased anode demonstrated a reversible capacity of 415 mA h g^{-1} after 100 cycles at 100 mA g^{-1} and a long-life cycling capability of 295 mA h g^{-1} after 500 cycles at 500 mA g⁻¹. Furthermore, Wang et al.¹⁹² fabricated graphene encapsulated with a hollow FeP@carbon nanocomposite (H-FeP@C@GR) through a combination of solvothermal and hydrothermal methods with carbothermic reactions and phosphidation treatments (Fig. 22d) and reported that the synthesized H-FeP nanospheres were highly uniform with an average diameter of 300-400 nm and that the thickness of the carbon layer was ~ 4 nm (Fig. 22e). Here, the researchers also reported that such a 3D hierarchical architecture could exhibit high electrochemical performances where a reversible capacity of 446 mA h g^{-1} after 100 cycles and high rate capacities and stabilities were achieved (Fig. 22f and g), suggesting that the 3D conductive network of the GR skeleton and the carbon coating of FeP were key factors in the enhancement of Na storage performances, which could not only provide open pathways for electron/ion transport, but also prevent the continual reforming of a SEI on the surface of hollow FeP.

FeP₄ exhibits a monoclinic structure and also can be used as an anode for SIBs. Zhang *et al.*⁷⁰ synthesized iron phosphides of FeP₂ and FeP₄ as SIB anodes by a simple ball milling method and found that the FeP₄ anode demonstrated a highly reversible sodiation capacity of 1137 mA h g⁻¹ with a coulombic efficiency of 84.0% during the first cycle at 89 mA g⁻¹ and high stability with a capacity of ~1000 mA h g⁻¹ after 30 cycles along with good rate capabilities. However, the researchers also reported that FeP₂ showed no significant electrochemical reactivities as a SIB anode possibly because of its low operating potential (≤ 0 V vs. Na/Na⁺).²⁴³

Overall, iron phosphides can show great potential as anodes for practical SIBs due to their low-cost and the facile synthesis of these ultrafine Fe-P nanostructures. However, FeP composites often exhibit low initial Coulomb efficiency (55-75%) and insufficient cycle life-span, which is a concern for practical application. Although ball milled FeP₄ possesses a higher P content and has shown the highest gravimetric specific capacity as well as the highest initial coulombic efficiency (84%) among those Fe-P composites, the same issue of insufficient cycle stability hiders its application. In addition, its low electronic conductivity ($\sim 3 \times 10^{-3}$ S m⁻¹ for FeP₄ vs. $\sim 1.3 \times 10^{6}$ S m⁻¹ for FeP) is also a concern. It's presumed that FeP₄ may undergo a similar capacity fading mechanism to amorphous RP because of its high P content. Furthermore, as ball milled FeP₄ shows a large particle size $(1-10 \ \mu m)$, it is more likely to be pulverized, leading to insufficient cycle stability. With respect to this, more detailed and concrete Na storage mechanisms should be investigated, and the synthesis of novel ultrafine Fe-P nanostructures/composites especially for FeP4 is also necessary for their practical application.

3.4 Nickel phosphides

Nickel phosphides also have many advantages in Na storage, such as high electronic conductivity ($\sim 10^6$ S m⁻¹ for Ni₂P and Ni₃P)^{244,245} and low cost. In addition, various morphologies or refined nanostructures can be obtained for nickel phosphides. For example, Ni₂P nanoarrays,²²⁷ nanosheets,²⁴⁶ nanorods,²⁴⁷ nanocrystals,⁹² and Ni₃P films²⁴⁸ and particles²²⁶ were reported as anodes for Na-ion or Li-ion batteries. The crystal structures of cubic Ni₃P and hexagonal Ni₂P are shown in Fig. 23a. A nickel phosphide as a SIB anode material was first reported by Fullenwarth et al.¹⁹⁹ in 2014 when they demonstrated the great potential of NiP₃ in the Na storage field. In their study, two kinds of NiP₃ compounds were synthesized via high temperature treatment or ball milling methods, respectively, both with metal Ni and commercial RP as raw materials. And the Nastorage mechanism of NiP3 was also investigated, and proposed as follows:

Review

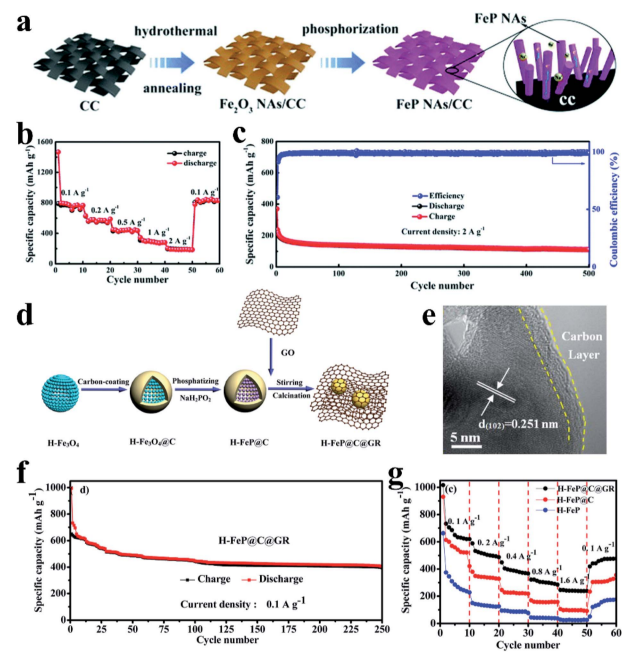


Fig. 22 (a) Schematic of the preparation of FeP NAs/CC. (b) Rate performance of the FeP NAs/CC electrode for SIBs. (c) Long-term cycling at 2 A g^{-1} . Reproduced with permission.²⁰² Copyright 2016, The Royal Society of Chemistry. (d) Schematic of the synthesis of H–FeP@C@GR nanocomposites. (e) HRTEM image of the H–FeP nanospheres. (f) Long-term cycling and (g) rate performance of a H–FeP@C@GR anode at a current density of 0.1 A g^{-1} . Reproduced with permission.¹⁹² Copyright 2017, American Chemical Society.

 $NiP_3 + 9Na^+ + 9e^- \rightarrow Ni + Na_3P$ (first discharge) (16)

$$Na_3P \leftrightarrow 3Na^+ + P + 3e^-$$
 (subsequent cycles) (17)

Here, cubic phase Ni_3P was directly transformed into Na_3P and nanosized metallic Ni particles in the discharge process, with no possible intermediate phases such as NaP and Na_3P_{11} evidenced. In the charge process, Na-extraction caused Na_3P transformation into amorphous P, and then the amorphous P underwent an alloying-dealloying process in the next cycles (amorphous Ni₃P or Ni–P compounds may also be part of the charged product). The as-prepared NiP₃ anode showed a relatively low redox potential of ~0.2 V *vs.* Na/Na⁺ and the NiP₃ synthesized by high temperature treatment exhibited a better Na storage performance than that synthesized from the ball milling method, delivering a discharge capacity of 1079 mA h g⁻¹ at 15th cycle.

Later, Kim *et al.*²²⁶ proposed an approach to synthesize NiP_3 particles to allow for chemical bonding with functionalized CNTs through a ball milling process (Fig. 24a). The

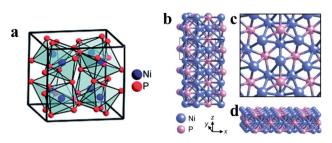


Fig. 23 (a) Crystal structure of cubic Ni₃P. Reproduced with permission.¹⁹⁹ Copyright 2014, The Royal Society of Chemistry. (b) Crystal structure of hexagonal Ni₂P: four unit cells stacked on top of one another, with a single unit cell outlined. (c) Top-down view of the Ni₂P (001) surface. (d) A two-dimensional slice of Ni₂P, showing the (001) surface on top. Reproduced with permission.²⁴⁹ Copyright 2013, American Chemical Society.

corresponding anode delivered a high initial reversible capacity of 853 mA h g^{-1} with more than 80% capacity retention after 120 cycles at 200 mA g^{-1} and a high rate capacity of 363.8 mA h g^{-1} after 200 cycles at 1600 mA g^{-1} (Fig. 24b). The results of physical characterization indicated that the P-C and P-O-C bonds between NiP3 and the functional groups on CNTs were generated during the ball milling process and played an important role in the enhancement of NiP3 anode performances in SIBs. Wu et al.225 also developed an assembly and selftemplate strategy for the synthesis of a 3D yolk-shell-like nanostructure in which Ni₂P nanoparticles with a size of \sim 24 nm were embedded into porous graphene networks. The overall fabrication process involved two main steps, an assembly step of NiNH4PO4 · H2O nanorods with graphene oxide (GO), and a following annealing step under a reducing atmosphere (H_2). Here, the researchers reported that the Ni₂P anode for SIBs delivered a reversible initial capacity of 181 mA h g^{-1} at 0.2 Ag^{-1} and maintained a capacity retention of 89% over 100 cycles. In addition, Dong et al.227 designed a sandwich-like nanoarchitecture hybrid of Ni₂P nanoarrays and nitrogendoped graphene (Ni₂P/NG/Ni₂P) as a SIB anode and reported that the as-prepared anode delivered a high capacity of 188 mA h g^{-1} (57% of initial capacity) at 0.5 A g^{-1} over 300 cycles. Shi et al.92 also designed a covalent heterostructure of monodisperse Ni₂P nanocrystals with sizes of 10-30 nm immobilized on N, P co-doped carbon nanosheets as a SIB anode (Fig. 24c-e) and reported that the anode showed enhanced cycling stabilities and high rate performances, maintaining 361 mA h g^{-1} at 100 mA g^{-1} after 300 cycles and 181 mA h g^{-1} at 500 mA g^{-1} after 1200 cycles (Fig. 24f and g). It is acknowledged that serious agglomeration during sodiation/ desodiation of transition metal phosphides (TMPs) leads to challenging kinetic issues and rapid capacity fading. Here, the researchers reported that an agglomeration free Na storage process was achieved and could be visualized using in situ TEM. In addition, the strong bonding between Ni₂P and NPC was evidenced through DFT calculations, which could stabilize Ni₂P nanocrystals and simultaneously enable synergistically enhanced charge storage by providing exponentially increased electronic states around the Fermi level.

Recently, a microporous structure of Ni₂P@C–N polyhedrons embedding in highly rough carbon fiber was designed as a selfsupporting anode material for SIBs.²⁵⁰ Due to the ultra-small size of Ni₂P particles (~5 nm) and the unique host nanostructure, the corresponding anode delivered a long-term cycling stability (196.8 mA h g⁻¹ at 1000 mA g⁻¹ over 1000 cycles with a capacity fading of 0.04% per cycle) and superior rate capability (197.1 mA h g⁻¹ at 2000 mA g⁻¹, and returned to 752.5 mA h g⁻¹ at 100 mA g⁻¹). Moreover, a flexible half-cell and a full cell based on this electrode could also give an excellent Na storage performance.

Guo *et al.*²⁵¹ synthesized tailored carbon-coated hollow $Ni_{12}P_5$ nanocrystals with an average size of 35 nm that were *in situ* grown on reduced graphene oxide nanosheets ($Ni_{12}P_5$ @C/GNS) using a template-free refluxing method. The corresponding composite anode showed good Na storage performances due to the finely designed dual carbon shell nanostructure and was found to be able to accommodate volume expansion, enhance ion/electron transfer, prevent $Ni_{12}P_5$ nanocrystal aggregation and thus deliver a reversible capacity of 235 mA h g⁻¹ at 100 mA g⁻¹.

To sum up, nickel phosphides such as NiP₃, Ni₂P, and Ni₁₂P₅ can show great potential as anodes for practical SIBs, among which NiP₃ possesses the highest P content as well as the highest theoretical gravimetric specific capacity. Although the pure NiP₃ anode can give a high specific capacity, the insufficient cycle stability hinders its application. In contrast, the NiP₃-CNT anode enables longer cycle lifespan, but a concern that the use of CNTs may increase the fabrication cost may be noticed. The high content of Ni in Ni₂P or Ni₁₂P₅ leads to smaller theoretical gravimetric specific capacity than that of NiP₃. Nevertheless, the high content of Ni may play an important role in achieving long cycle stability. For all these Ni-P anodes, the initial coulombic efficiency is not high (50-80%), and the specific capacities are also very low, far from the theoretical values (1591 mA h g^{-1} for NiP₃ and 1333 mA h g^{-1} for Ni₂P). With respect to this, more effort should be made, including design and synthesis of novel nanostructures/ composites and exploration of facile and large-scale preparation methods (especially for NiP₃, which has only been prepared by ball milling or high temperature methods).

3.5 Copper phosphides

Copper phosphides also possess various advantages in electrochemical Na storage applications. Among these copper phosphides, CuP_2 and Cu_3P are often used as anodes for Na-ion batteries. CuP_2 can exhibit a monoclinic structure while Cu_3P usually has a hexagonal structure. In the early stage of the study of these copper phosphides, ball milling is the most commonly used method to prepare anode materials. For example, Zhao *et al.*¹⁸⁹ synthesized a composite consisting of crystalline CuP_2 cores coated with carbon black nanoparticles (CuP_2/C) through two steps of ball milling processes, and reported that this structure could enable fast and reversible sodiation/ desodiation, allowing a capacity of more than 500 mA h g⁻¹ at 50 mA g⁻¹, a high rate capacity of 178 mA h g⁻¹ at 2000 mA g⁻¹



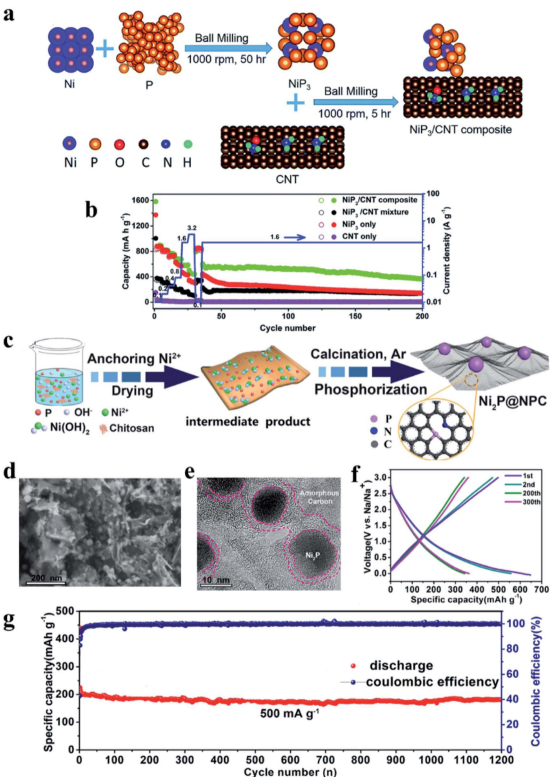


Fig. 24 (a) Schematic of the synthesis of the NiP₃/CNT composite. (b) Comparison of the cycling performances of CNT, NiP₃, NiP₃/CNT mixture and NiP₃/CNT composite electrodes. Reproduced with permission.²²⁶ Copyright 2018, The Royal Society of Chemistry. (c) Schematic of the synthesis of Ni₂P@NPC. (d) SEM and TEM images of Ni₂P@NPCS. (e) HR-TEM image of Ni₂P@NPC. (f) Galvanostatic charge/discharge profiles of the Ni₂P@NPC SIB anode for the 1st, 2nd, 200th and 300th cycles respectively. (g) Long-term cycling stability and coulombic efficiency of the Ni₂P@NPC composite at 500 mA g⁻¹. Reproduced with permission.⁹² Copyright 2018, Elsevier Ltd.

and decent short-term cycling stability. Then a chemically bonded CuP₂/C hybrid was also prepared by a one-step ball milling method,²⁵² and the strong P–O–C bond and the stable nanoscale conducting framework in this hybrid offered an enhanced Na storage performance. According to these studies and other literature,^{189,228,252} the electrochemical reaction mechanism of CuP₂ can be expressed as follows:

$$CuP_2 + 6Na^+ + 6e^- \leftrightarrow Cu + 2Na_3P$$
(18)

It means that crystalline CuP_2 can be recovered after one discharge/charge cycle and shows a high theoretical capacity of 1282 mA h g⁻¹ based on this conversion and alloy reaction.

Recently, other methods to prepare CuP_2 anodes such as the solution phase method coupled with a heat treatment have been developed.^{198,228} For example, Yu *et al.*²²⁸ designed a nanocomposite of cross-linked hollow carbon sheet encapsulated CuP_2 nanoparticles (CHCS– CuP_2) prepared in several steps including surfactant-assisted wet-chemical precipitation, carbon coating and *in situ* phosphorization (Fig. 25a). Here, the

researchers reported that the hollow structure and carbon shells could mitigate large volume change, prevent CuP₂ nanoparticle aggregation and significantly enhance electron/ion transport (Fig. 25b-d). In addition, the connectivity of the cross-linked nanocomposite could enhance structural stability and resist inner stress, allowing for longer lifespans during cycling. As a result, the corresponding SIB anode delivered a high reversible capacity of 451 mA h g^{-1} at a current density of 80 mA g^{-1} with a retention of 91% after 200 cycles, and an excellent rate performance was also obtained (Fig. 25e and f). Furthermore, a full cell made based on this anode also provided an excellent Na storage performance, showing great potential for practical SIBs. Similarly, Duan et al.²⁵³ reported a CuCl₂/chitosan monolith derived CuP₂/C composite where CuP₂ nanoparticles were uniformly embedded in the carbon matrix. The strong chemical bonding between electron rich groups in chitosan and the heavy metal ion (Cu²⁺) played a key role in the synthesis of this homogeneous monolithic composite. In addition, the chitosan derived carbon could prevent Cu and CuP2 particles from aggregation upon the subsequent thermal reduction and

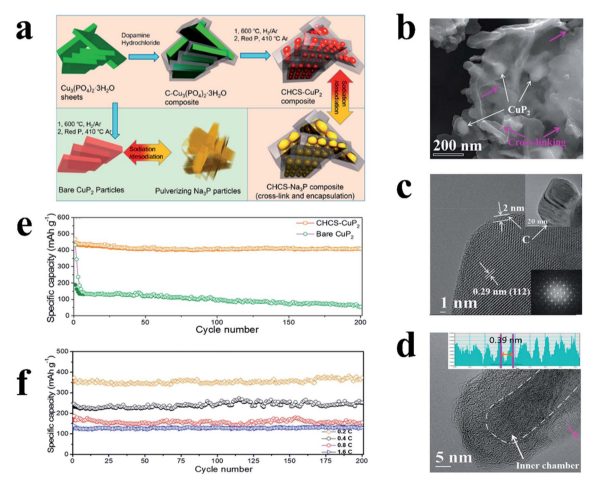


Fig. 25 (a) Schematic image of the preparation of the CHCS–CuP₂ composite and bare CuP₂ particles, as well as the reversible reactions with Na⁺ ions of these anodes. (b) SEM images of CHCS–CuP₂ composites. (c) HRTEM image of the CHCS–CuP₂ composite and its corresponding Fourier transformation in the inset, showing a thin carbon layer (~2 nm) and the (110) planes of CuP₂. (d) HRTEM image of a hollow carbon sheet with a wide interlayer distance (~0.39 nm), showing an apparent inner chamber allowing for volume changes of CuP₂ during cycling. (e) Cycling performance of bare CuP₂ and CHCS–CuP₂ composites at 80 mA g⁻¹. (f) Cycling performance of CHCS–CuP₂ composites at 0.2, 0.4, 0.8, and 1.6C (1C = 800 mA g⁻¹). Reproduced with permission.²²⁸ Copyright 2018, American Chemical Society.

phosphorization. Benefiting from the synergistic effect of the small particle size and conductive carbon matrix, the CuP₂/C composite anode for a SIB could deliver a high reversible capacity of 630 mA h g⁻¹ at a current density of 100 mA g⁻¹ and a capacity retention ratio of 91% after 200 cycles, while bare CuP₂ showed a rapid capacity decay within 50 cycles.

Cu₃P is also often used as an anode material for SIBs. Although Cu is inactive with Na⁺ and thus shows a relatively low theoretical capacity of 363 mA h g⁻¹ (similar to that of graphite), other advantages make it suitable as an anode for SIBs, such as air stability, high electronic conductivity, safe intercalation potential, and a high theoretical volumetric capacity of 4732 mA h cm⁻³ (830 mA h cm⁻³ for graphite).²⁵⁴ Fan *et al.*⁷⁸ reported an additive-free Cu₃P nanowire (CPNW) anode directly grown on a copper current collector *via* an *in situ* growth and a subsequent phosphidation process. Due to its structure features, the CPNW anode demonstrated highly stable cycling ability with an ~70% retention in capacity after 260 cycles. According to the literature, a reversible sodiation/desodiation of Cu₃P proceeds *via* a conversion mechanism, which is similar to that of CuP_2 , and can be expressed as follows:

$$Cu_3P + 3Na^+ + 3e^- \leftrightarrow 3Cu + Na_3P \tag{19}$$

After this study, many Cu₃P composites have been successfully prepared by various methods. Inspired by metal–organic framework (MOF) derived strategies to synthesize metal phosphides/carbon (MP@C) nanostructures, Kong *et al.*¹⁹⁸ designed a nanostructure of Cu₃P nanoparticles (~50 nm) coated with N, P-codoped carbon matrices using a metal– organophosphine framework (MOPF) derived strategy (Fig. 26a– d), where PTA represents 1,3,5-triaza-7-phospha-adamantane. This approach is relatively eco-friendly with no need for an additional P source, thus avoiding the release of harmful PH₃ which is inevitably produced in the phosphorization process for most MOF-derived MP@C nanostructures.^{204,255,256} Due to this unique nanostructure, the corresponding SIB anode of this composite exhibited a high initial reversible capacity of 332 mA h g⁻¹ at 50 mA g⁻¹, a rate capacity of 142.5 mA h g⁻¹ at



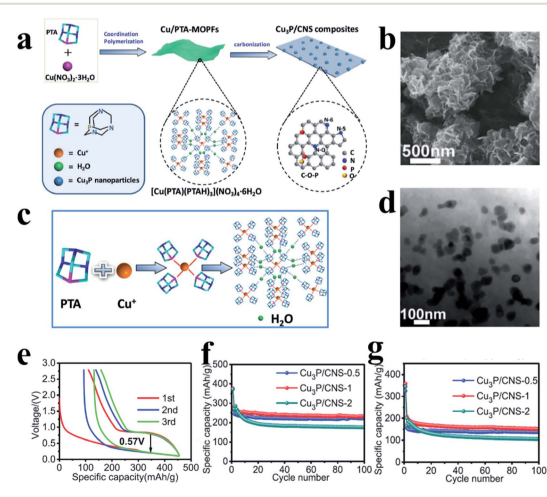


Fig. 26 (a) Schematic illustration of the evolution process from Cu/PTA-MOPFs to Cu_3P/CNS composites. (b) SEM images of Cu/PTA-MOPF-1 (molar ratio of PTA and $Cu(NO_3)_2$ is 1 : 1). (c) Schematic illustration of the coordination between PTA molecules and Cu ions in the precursors. (d) TEM images of the Cu_3P/CNS -1 composite. (e) The initial three charge/discharge curves of the Cu_3P/CNS -1 composite at 50 mA g⁻¹. (f, g) Cycling performances of three Cu_3P/CNS composites at 100 and 1000 mA g⁻¹, respectively (CNS-0.5, CNS-1, and CNS-2 represent the molar ratio of PTA and Cu(NO_3)_2 of 0.5, 1, and 2, respectively). Reproduced with permission.¹⁹⁸ Copyright 2018, WILEY-VCH.

2000 mA g⁻¹ and good cycling performances with a retention of 85% after 100 cycles (Fig. 26e–g). Zhu *et al.*²⁵⁷ reported a composite of 3D Cu₃P@C nanosheets synthesized by heating a mixture of copper foam and phosphorus-containing resin. During the heat treatment, an epitaxial phosphidation growth coupled with a carbon deposition process occurred on the surface of copper foam, and the 3D Cu₃P@C nanosheets were formed. In each individual nanosheet, the thin carbon shell could serve as an electron conductor and accommodate the volume change of the Cu₃P single-crystalline nanosheet. Due to its unique 3D nanoarchitecture, the Cu₃P@C anode delivered a high capacity retention of 286 mA h g⁻¹ after 300 cycles at 0.1 A g⁻¹ and 156 mA h g⁻¹ after 1000 cycles at 1 A g⁻¹.

Recently, Cu_2P_7 as an anode material for SIBs has also been reported and showed excellent electrochemical performance. For example, Li *et al.*²⁵⁸ successfully prepared a $Cu_2P_7/CuP_2/$ graphite composite by a facile ball milling method and used it as an anode material for SIBs. Benefiting from a synergistic effect of the high P concentration, the layered structure and heterointerfaces between the different phases, the corresponding anode could provide capacities of 1254 mA h g⁻¹ after 100 cycles at 200 mA g⁻¹ and 200 mA h g⁻¹ at 5 A g⁻¹ in a rate capability test. Similarly, a composite of Cu_2P_7 -BP-MWCNT (MWCNT, multiwalled carbon nanotube) was also prepared by a mechanical milling method.²⁵⁹ When used as an anode for Naion batteries, the composite delivered high comprehensive performances: initial coulombic efficiency up to 84%, 1170 mA h g⁻¹ after 200 cycles, and 580 mA h g⁻¹ at 5 A g⁻¹.

In summary, copper phosphides, including CuP₂, Cu₃P and Cu₂P₇, have also attracted much attention because of their lowcost, high electronic conductivity and other advantages. Although the ball milling method to prepare CuP₂ composites is facile, the insufficient cycle life-span of these anodes is a concern for practical application. In contrast, CuP₂/C nanoparticles prepared by the solution phase method (assisted by other processes) show better cycle stability. However, these CuP₂ anodes all suffer from insufficient gravimetric specific capacity and long-cycle lifespan, which need to be addressed in future research. Various Cu₃P nanostructures/composites such as nanowires, nanoparticles, and nanosheets have been prepared, and some high sodium storage performances are achieved in terms of rate capability and long-cycle stability. However, the low theoretical specific capacity may hinder their application, especially for electric vehicles or even large-scale energy storage systems. The Cu₂P₇ composite has been recently reported as an anode for SIBs and exhibited high specific capacities due to the high P content of Cu₂P₇. In addition, Cu₂P₇ promises high rate capability due to its layered structure and high electronic conductivity, which favor Na⁺/e⁻ transportation. All of this makes Cu₂P₇ a promising candidate as an anode material for practical SIBs.

3.6 Germanium phosphides

Germanium phosphides (GeP_x) such as GeP_5 and GeP_3 have recently been extensively investigated as anode materials for SIBs. GeP_5 possesses a large theoretical specific gravimetric capacity of 1888 mA h g⁻¹ due to the highest P atomic percentage among all the binary phosphides, and a high theoretical volumetric capacity of 6891 mA h cm⁻³ which is the best record in anodes for SIBs. In addition, rhombohedral GeP5 exhibits a layered structure similar to those of black P and graphite (Fig. 27a), with a high electrical conductivity of $\sim 10^6$ S m^{-1} , 4 orders of magnitude higher than that of black P and similar to that of graphite (XRD pattern is shown in Fig. 27b). All of this indicates that GeP5 is another promising candidate as an anode material for SIBs. Li et al.193 synthesized a GeP5/C composite by ball milling GeP5 and commercial graphite and reported a low potential of ~ 0.4 V vs. Na⁺/Na with a smooth charge/discharge profile. This GeP5/C anode showed an excellent Na storage performance, delivering a large reversible capacity of 1250 mA h g^{-1} with an initial coulombic efficiency of 93% and considerable cycling stabilities with 98% capacity retention after 60 cycles (Fig. 27d). Here, the synergetic effects between the sodiation of Ge and P reportedly leveled multistep plateaus smoothly and effectively reduced polarization between charge and discharge, leading to high coulombic efficiencies. Moreover, a full SIB cell using this anode coupled with a $Na_3V_2(PO_4)_3/C$ cathode provided a large capacity of 800 mA h g^{-1} with a high average output voltage of 2.65 V. The researchers also investigated the electrochemical Na storage mechanism of GeP₅, which can be described by equations as shown in the following:

 $GeP_5 + 16Na^+ + 16e^- \rightarrow 5Na_3P + NaGe$ (first discharge) (20)

$$Na_3P \leftrightarrow 3Na^+ + P + 3e^-$$
 (subsequent cycles) (21)

$$NaGe \leftrightarrow Ge + Na^{+} + 3e^{-}$$
(22)

It was confirmed that GeP₅ experienced conversion-alloying type sodium storage during an electrochemical reduction process. GeP₅ was converted into Na₃P and NaGe after the initial discharge process, and Na₃P and NaGe respectively underwent desodiation/sodiation processes during the next cycles. This mechanism could be evidenced by the TEM images of the sodiated and desodiated states of GeP₅ (Fig. 27c and d). Actually, in the theoretical capacity of GeP₅ (1888 mA h g⁻¹), Ge element can contribute 118 mA h g⁻¹ and 5 P atoms donate the remaining 1770 mA h g⁻¹.

GeP₅ could also be obtained through the ball milling of highpurity Ge powder and commercial RP, and the composite of this GeP₅ with carbon (GeP₅/C) also demonstrated good Na storage performances.²⁶⁰ Ning *et al.*²²⁹ also prepared this GeP₅ by ball milling Ge powder and RP, and further designed a dual-carbon conductive network enhanced GeP₅ composite by a ball milling method (GeP₅/AB/p-rGO; AB represented acetylene black, and prGO was partially reduced graphene oxide). They suggested that the dual carbon network not only provided more electron transport pathways but also relaxed the huge volume change during charge/discharge. As a result, this composite anode delivered a high reversible capacity of 597.5 and 175 mA h g⁻¹ at current densities of 0.1 and 5.0 A g⁻¹ respectively and high

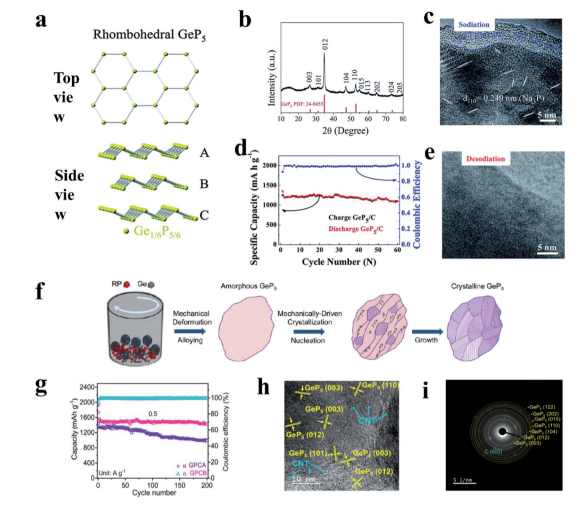


Fig. 27 (a) Crystal structure of rhombohedral GeP₅. Reproduced with permission.²⁶¹ Copyright 2015, The Royal Society of Chemistry. (b) XRD patterns of the as-synthesized GeP₅/C composite. (c, e) HRTEM of the GeP₅/C electrode in the full sodiation state (discharged to 0.005 V), and in the full desodiation state (charged to 3.0 V), respectively. (d) The cycle performance and corresponding coulombic efficiency of the assynthesized GeP₅/C composite. (b–e) Reproduced with permission.¹⁹³ Copyright 2017, The Royal Society of Chemistry. (f) Schematic explanation of mechanically induced crystallization that occurs during the milling of Ge and RP elemental powders. (g) Comparison of the cycling stability of GPCA and GPCB electrodes in the voltage range of 0.01–2.00 V at a current density of 0.50 A g⁻¹ for 200 cycles (GPCA and GPCB represent the amorphous and crystalline GeP₅-based composites, respectively). (h, i) HRTEM micrographs and SAED patterns of GPCB after the 50th cycle in the charged state. (f–i) Reproduced with permission.²⁶² Copyright 2019, American Chemical Society.

stability with 400 mA h g⁻¹ capacity retention at 0.5 A g⁻¹ after 50 cycles. Furthermore, Tseng *et al.*²³⁰ prepared mesoporous germanium phosphide (MGeP_x) microspheres as a SIB anode material through a high-temperature solvothermal method. These MGeP_x microspheres with diameters ranging from 0.5 to 1.5 µm were composed of 10 nm nanoparticles with a narrow pore size distribution of ~4 nm and possessed an amorphous structure with a weight ratio of P to Ge of 38.3 to 61.7. This unique porous nanostructure demonstrated high performance in sodium storage including high energy density, ultrahigh rate capability, and superior cyclability, showing a high reversible capacity of 704 mA h g⁻¹ after 100 cycles at 240 mA g⁻¹ and stable cycling performances of 278 mA g⁻¹ at 1200 mA g⁻¹ after 200 cycles.

Very recently, Cho *et al.*²⁶⁰ synthesized two kinds of GeP_5 , crystalline GeP_5 (the GeP_5 anodes reported by others were all of

this type) and amorphous GeP₅ (Fig. 27f); furthermore, the authors also prepared hybrids of GeP5 and multiwalled carbon nanotubes (GeP5/CNT). The researchers found that the crystalline GeP₅ exhibited better Na storage performance than the amorphous one and proposed that the crystalline GeP5 exhibited a higher electronic conductivity, more facility for the insertion of Na⁺, and smaller volume expansion after sodiation. As a result, this crystalline GeP5/CNT anode showed a high initial CE of 94.13% and a high capacity of 1430 mA h g^{-1} after 200 cycles at 0.5 A g^{-1} , corresponding to a capacity retention of 91.43% from the second cycle (Fig. 27g). Interestingly, the authors suggested that GeP5 could recover to its crystalline phase after desodiation, as evidenced by the TEM images (Fig. 27h and i), which was contradictory to other reported literature.193,260 Therefore, further investigation about this issue is needed.

Journal of Materials Chemistry A

Kim *et al.*²⁶³ introduced GeP₃ as a promising anode material for SIBs and carried out a comparative investigation of GeP₃ and GeP₅. In their study, GeP₃ was obtained by ball milling of Ge and RP, and then the GeP₃/C compound was obtained by further ball milling of GeP₃ and carbon (super P) for electrochemical measurements. The GeP₅/C composite was synthesized by a similarly procedure. It was confirmed that GeP₃ could give a rhombohedral crystal phase similar to GeP₅ (Fig. 28a), but with a larger cell volume and average bonding distance

compared with GeP₅. Based on theoretical calculations, they also found that the bonding strength of P–P was higher than that of P–Ge, and the elastic modulus of GeP₃ showed significantly softer features than that of GeP₅, indicating that the softer feature of Ge–P bonding could enable more effective accommodation for mechanical stress and deformation during charge/discharge cycles than P–P bonding. This insight could be experimentally confirmed: the synthesized GeP₃ compound maintained a high discharge capacity of 1274 mA h g⁻¹ after 150

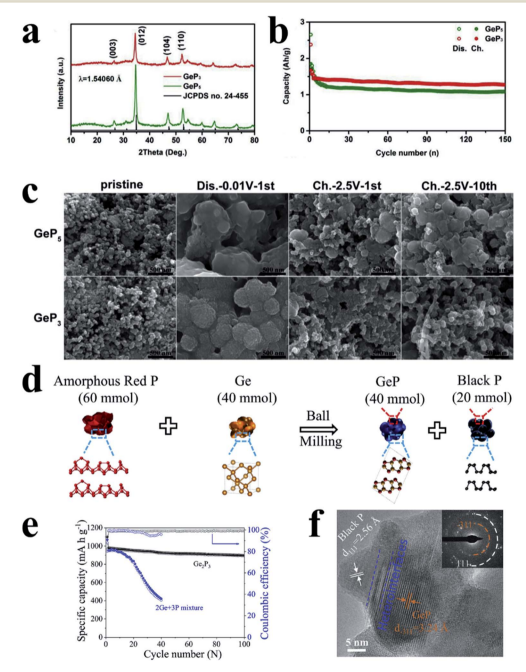


Fig. 28 (a) XRD patterns of the as-prepared GeP₃ and GeP₅ samples. (b) Cycling performance of GeP₃ and GeP₅. (c) A comparison of morphological changes for GeP₅ (up) and GeP₃ (down) compounds during the first 10 cycles. Reproduced with permission.²⁶³ Copyright 2018, Elsevier Ltd. (d) Illustrated preparation process of the layered GeP-black P composite (Ge₂P₃). (e) Cycle stability of the Ge₂P₃ composite. (f) High-resolution transmission electron microscopy (HR-TEM) image and the inset shows the selected area electron diffraction of the Ge₂P₃ composite. Reproduced with permission.²⁶⁴ Copyright 2019, Elsevier Ltd.

cycles, while the GeP₅ compound with more P–P bonding displayed an inferior cycling stability (Fig. 28b). Furthermore, the morphology and volume change of the GeP₅ electrode seemed more serious compared to those of the GeP₃ electrode during charge/discharge (Fig. 28c).

In another study, Nam *et al.*¹⁹⁶ used commercial Ge and synthesized black P as raw materials to synthesize crystalline GeP₃ through a high-power ball-milling (HPBM) method, and further prepared a GeP₃/carbon (GeP₃/C) nanocomposite by ball milling this GeP₃ with Super P. The GeP₃/C anode showed a highly reversible initial capacity of 984 mA h g⁻¹ with a high initial CE of 81.3%, a very stable capacity retention of 94.9% after 30 cycles, and a high rate capacity of 520 mA h g⁻¹ at 2C (Fig. 29e and f). Furthermore, the researchers investigated the Na storage mechanism of GeP₃ by using various techniques, such as extended X-ray absorption fine structure (EXAFS) (Fig. 29b and c), X-ray diffraction (XRD), high-resolution transmission electron microscopy (HR-TEM) (Fig. 29d), and differential capacity (dQ/dV) and cyclic voltammetry (CV). It was indicated that the reaction mechanism involved one-step conversion during Na-insertion and two-step recombination/ topotactic-transition during Na-extraction, as shown by the following equations:

Na-insertion:

$$GeP_3 \rightarrow NaGe + Na_3P$$
 (23)

Na-extraction:

$$NaGe + Na_{3}P \rightarrow Na_{x}GeP_{3} + (x \le 2.4) \rightarrow GeP_{3}$$
(24)

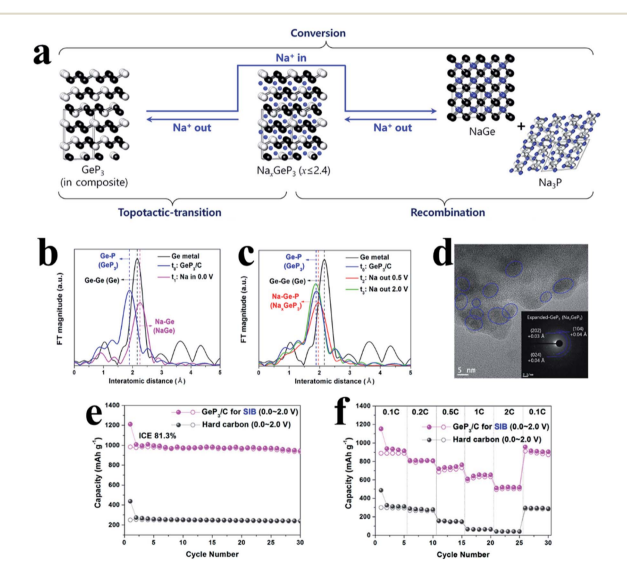


Fig. 29 (a) Schematic diagram of the electrochemically driven crystallographic phase-change mechanism of GeP₃ during Na-insertion/ extraction. (b, c) *Ex situ* EXAFS spectra of the GeP₃-based nanocomposite electrode during first Na insertion and first Na-extraction, respectively. (d) *Ex situ* HR-TEM image with the corresponding diffraction patterns in the Na-extracted state of 0.5 V for the GeP₃-based nanocomposite electrode. (e) Cycling performance of the GeP₃-based nanocomposite and hard carbon (cycling rate: 50 mA g⁻¹). (f) Rate capabilities of the GeP₃-based nanocomposite (1C = 950 mA h g⁻¹) and hard carbon (1C = 250 mA h g⁻¹) for the SIB. Reproduced with permission.¹⁹⁶ Copyright 2018, Elsevier B.V.

Total:

$$GeP_3 + 4Na^+ + 4e^- \leftrightarrow NaGe + Na_3P$$
 (25)

During Na-insertion, the layer structured GeP₃ can be converted to NaGe and Na₃P. During Na-extraction, however, Na₃P and NaGe are transformed into a Na_xGeP₃ ($x \le 2.4$) phase, which is then recovered to pristine GeP₃ with a topotactic-transition (Fig. 29a).

Recently, a layered GeP-black P composite (Ge₂P₃) as a binary-phase anode for SIBs was prepared by a simple ball milling method²⁶⁴ and also showed an excellent Na storage performance. During the milling process, element Ge and commercial RP were converted into GeP and BP (black P); in the meantime, these two products were hybridized to form a layered GeP-BP composite (Fig. 29d and f). The authors proposed that the numerous heterointerfaces between layered GeP and BP could increase the number of reaction sites and enhance the Na-ion and electron transport capability. Coupled with the intrinsic excellent Na storage capability of GeP and BP, this binary-phase anode showed a reversible capacity 970 mA h g^{-1} at 50 mA g^{-1} with an initial coulombic efficiency of 88%, and retained 890 mA h g^{-1} after 100 cycles (Fig. 29e). Moreover, it also exhibited a rate capacity of 275 mA h g^{-1} at 5 A g^{-1} , which could recover back nearly to its initial value when the current density was changed back to the initial value of 50 mA g^{-1} . This study shows a simple and cost-efficient strategy to prepare GeP and BP in one step, offering excellent possibility for application in practical SIBs. In another study, a GeP anode with a flexible layered structure was successfully synthesized by a simple mechanochemical method.¹⁹⁷ When tested as the anode for a SIB, the GeP anode delivered an initial discharge/ charge capacity of 900/840 mA h g^{-1} with an initial coulombic efficiency of up to 93% and a rate capacity of 360 mA h g^{-1} at a current density of 5 A g^{-1} . Similar to GeP₃, GeP underwent a reversible Na-storage process that involved intercalation, conversion, and alloying. The peculiar self-healing phenomenon was derived from its ultra-low formation energy (0.19 eV) for the reconstruction of the original layered structure, as confirmed by first-principles calculations. The low formation energy also enabled the formation of chemical bonding between layered GeP and graphite, thus achieving an excellent performance as an anode for SIBs.

Overall, germanium phosphides also attract much attention and show great potential as anode materials for SIBs. Despite the high cost of Ge, these Ge–P anodes may still be affordable for large scale application, especially GeP₅ which has a low molar ratio of Ge. Although high performances of these GeP₅ composite anodes have been achieved in terms of initial coulombic efficiency and specific capacity (especially for GeP₅/ CNT), up to now, all these GeP₅ composites are prepared by the ball milling method and show a large particle size. The large particle size might be a major reason for their insufficient longcycle stability. In addition, the poor rate capability is also a concern for practical application. The MGeP_x ($x \approx 1.6$) composite reported by Tseng *et al.*²³⁰ is the only case where a germanium phosphide anode was prepared by a solvothermal method. Due to the ultrafine nanostructure of this $MGeP_x$, a better rate capability could be obtained. Although the P content in GeP_3 is lower than that in GeP_5 , it was reported that a superior sodium storage performance could be achieved for GeP_3 compared to GeP_5 in terms of specific capacity and cycling stability. GeP and its composites also exhibit high performance for Na storage; however, the high molar ratio of Ge may increase the fabrication cost of these GeP anodes, and thus may hinder their practical application.

3.7 Other metal phosphides

Other binary metal phosphides have also been studied for SIB anodes. For example, Li et al.²³² synthesized a well-aligned Zn₃P₂ nanowire-assembly bundle/Ti foil integrated anode through a facile chemical vapor deposition method and reported that this anode showed high sodium storage performances, delivering a reversible capacity of \sim 600 mA h g⁻¹ at 100 mA g⁻¹ and a high rate capability of 280 mA h g⁻¹ at 5 A g⁻¹. Huang *et al.*²³¹ also successfully prepared molybdenum phosphide (MoP) nanorods wrapped within a thin carbon layer as a SIB anode material and reported that this MoP electrode exhibited high cycling performances with a discharge capacity of 398.4 mA h g^{-1} at a current density of 100 mA g^{-1} after 800 cycles and a remarkable rate capability of 104.5 mA g^{-1} at 1600 mA h g^{-1} after 10 000 cycles. These low-cost metals such as Zn and Mo give a competitive option for metal phosphides in fabricating large-scale SIB anodes. In another study, Usui et al.209 studied sodiation/desodiation reactions in indium phosphide (InP) anodes which were obtained by ball milling in ionic liquid electrolytes and found that InP could provide phase separation in the first sodiation to form elemental In and P, which could subsequently react with Na individually. And in the following desodiation process, Na-extraction of NaIn and Na₃P occurred and a nanostructure of crystalline In nanoparticles dispersed in an amorphous-like P matrix was formed. Based on this Na storage mechanism, InP can provide a theoretical capacity of 730 mA h g^{-1} (Fig. 30a–c). This mechanism was also observed in Sn₄P₃ anodes.²⁰⁸ These researchers also reported that the In phase in this InP anode could relax strain induced by P volume change and prevent electrical isolation of active material layers, and in addition, the high conductivity could compensate for the poor electronic conductivity of Na₃P phases created by P sodiation, allowing for a high capacity of 500 mA h g⁻¹ after 100 cycles at a current of 50 mA g⁻¹ (Fig. 30d).

4 Non-metal phosphides

Many non-metal phosphides also possess the ability to store Na ions, such as SiP_2 and Se_4P_4 .^{233,234} Duveau *et al.*²³³ prepared a SiP_2 material through simple ball milling of silicon and phosphorus powders at room temperature and demonstrated potential as a novel anode for SIBs in which the corresponding anode provided a sustained capacity of 572 mA h g⁻¹ after 15 cycles. Although the reversible capacity is much lower than the theoretical capacity of 1780 mA h g⁻¹ (considering that P can

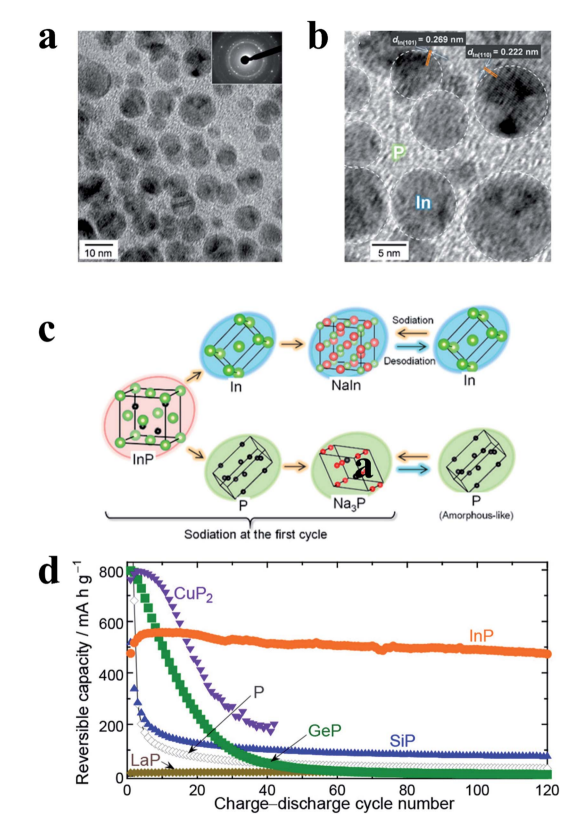


Fig. 30 (a) TEM image of InP after the first desodiation. The particle size was 9.3 ± 1.7 nm. (b) High-magnification TEM image showing crystalline In nanoparticles dispersed in an amorphous-like P matrix. (c) Phase changes of InP in charge/discharge reactions. (d) Anode performances of various phosphide electrodes cycled at a current density of 50 mA g⁻¹ in an ionic liquid electrolyte (NaFSA/Py13-FSA). For comparison, results are also shown for the electrode of P alone. Reproduced with permission.²⁰⁹ Copyright 2018, American Chemical Society.

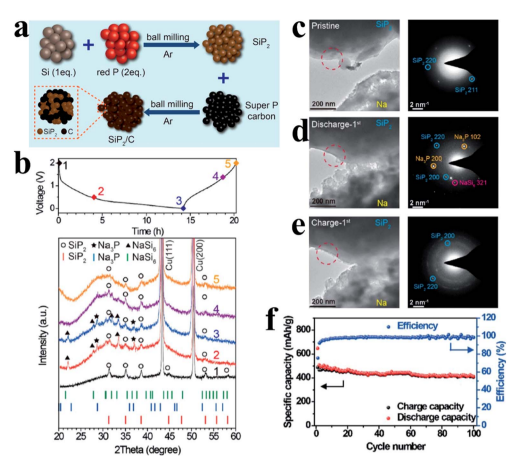


Fig. 31 (a) Schematic illustration of the synthesis process of the SiP₂/C composite. (b) *Ex situ* XRD of the SiP₂/C electrode in different discharge/ charge states in the first cycle. (c–e) *In situ* TEM investigation of a SiP₂–Na nanobattery. TEM images of electrodes and the corresponding SAED patterns of the SiP₂ electrode in selected regions acquired before battery working (c), when discharged at -3 V (d), and when further charged at +3 V (e). (f) Cycle performance of the SiP₂/C electrode acquired at a current density of 50 mA g⁻¹. Reproduced with permission.²⁶⁶ Copyright 2019, American Chemical Society.

react with 3 Na to form Na₃P and that Si has no electrochemical activity toward sodium),²⁶⁵ this pioneering study demonstrates the great potential of SiP₂ as a SIB anode material. Recently, Saddique *et al.*²⁶⁶ synthesized a SiP₂/C composite by a facile ball milling method (Fig. 31a). The well-constructed composite comprised of uniformly distributed SiP₂ nanocrystallites within a conductive carbon framework could greatly enhance the electrode conductivity and structural compatibility for repeated sodiation/desodiation conversions, demonstrating a high capacity of 410 mA h g⁻¹ at 50 mA g⁻¹ after 100 cycles (Fig. 31f), and a largely improved cyclability with 80% capacity retention for 500 cycles. Furthermore, the authors elucidated the reaction mechanism of SiP₂ in SIBs by various investigation methods and found that Si could also electrochemically react with Na, as expressed in the following equations:

$$SiP_2 \rightarrow Na_6Si + Na_3P$$
 (discharge) (26)

 $Na_6Si + Na_3P \rightarrow (SiP_2)_{crystalline/amorphous} + P (charge)$ (27)

This reaction mechanism was evidenced by the *ex situ* XRD and TEM images (Fig. 31b-e). During the first discharge

process, crystalline SiP₂ was gradually transformed into Na₆Si and Na₃P because of Na insertion. In the charge process, only parts of Na₆Si and Na₃P could reversibly form SiP₂, resulting in amorphization of part of SiP₂ after full charge. This could be evidenced by the XRD result where the SiP₂ peaks could not recover their initial intensities after full charge (Fig. 31b). Although Na storage performance is insufficient, this study demonstrates that SiP₂ can also reversibly react with Na. And in view of the low cost and environmental friendliness of both Si and P, these studies offer the possibility for scalable application of SiP₂ or other silicides in practical SIBs.

 Se_4P_4 is another attractive anode material for SIBs because selenium is active towards Na and possesses good electrical conductivity and Se_4P_4 provides a theoretical capacity of 1217 mA h g⁻¹ in reactions with 20 Na ions during sodiation.^{234,267} For example, Lu *et al.*²³⁴ synthesized an amorphous Se_4P_4 composite through a facile mechanical milling technique and reported that this Se_4P_4 anode exhibited a high reversible capacity of 1048 mA h g⁻¹ at 50 mA g⁻¹, good cycling stabilities of 804 mA h g⁻¹ after 60 cycles and outstanding rate capabilities of 724 mA h g⁻¹ at 500 mA g⁻¹ and 332 mA h g⁻¹ at 3000 mA g⁻¹ respectively, which were superior to those of both P and Se. Furthermore, *ex situ* XRD, TEM and *in situ* Raman analysis revealed that the reaction mechanism during the charge/ discharge of Se_4P_4 was as follows:

$$Se_4P_4 + 20Na^+ + 20e^- \leftrightarrow 4Na_3P + Na_2Se$$
(28)

 Se_4P_4 was converted into Na_2Se and Na_3P after full discharge and subsequently reformed to amorphous Se_4P_4 in the subsequent charging process. Furthermore, a full cell with a Se_4P_4 anode and a Na₃(VO_{0.5})₂(PO₄)₂F₂/C cathode delivered a discharge capacity of 985 mA h g⁻¹ (based on the mass of Se₄P₄) and an average output voltage of ~2.5 V. Cao *et al.*²⁶⁸ also successfully synthesized P₄Se₃ microparticles (higher P content as compared with Se₄P₄) through simple heating of selenium and amorphous RP at low temperatures (Fig. 32a and b) and reported that these orthorhombic-phase microparticles as a SIB anode could exhibit a high reversible capacity of 654 mA h g⁻¹ at

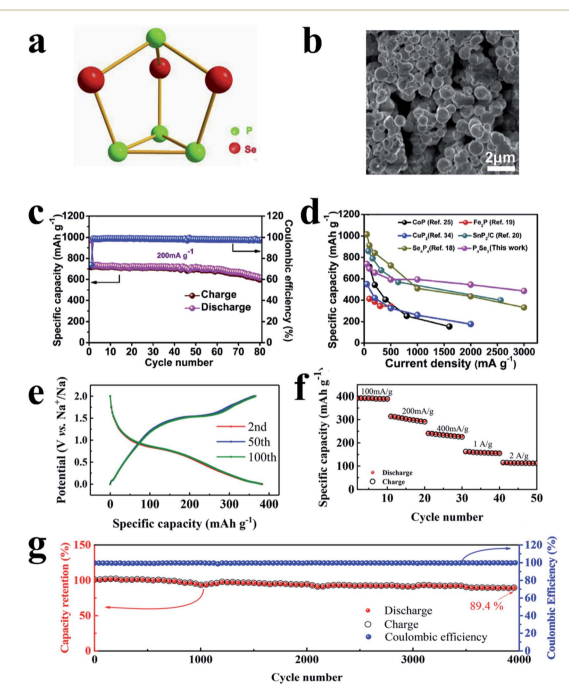


Fig. 32 (a) Illustration of the cage structure of P_4Se_3 . (b) SEM image of P_4Se_3 . (c) Cycling stability tested at 200 mA g⁻¹ and coulombic efficiency of P_4Se_3 anodes. (d) Rate capability comparisons of reported studies and this study. Reproduced with permission.²⁶⁸ Copyright 2018, Elsevier B.V. (e and f) Electrochemical performances of the P_2S_5/C composite after 70 hours of ball milling. (e) Typical potential profiles of the second, fifth and 100th cycles. (f) Rate capability of the P_2S_5/C composite. (g) Performance and coulombic efficiency at 2000 mA g⁻¹. Reproduced with permission.²³⁵ Copyright 2017, American Chemical Society.

Review

200 mA g⁻¹ after 70 cycles and high rate capabilities with a reversible capacity of 486 mA h g⁻¹ at 3 A g⁻¹ (Fig. 32c and d). Here, the researchers attributed this high electrochemical performance to the formation of more conductive Na₂Se and Se ($\sim 10^{-6}$ and $\sim 10^{-5}$ S cm⁻¹) during cycling and the dominance of capacitive behavior, all of which could accelerate kinetic behaviors.

In another example, Li *et al.*²³⁵ prepared a P_2S_5/C composite using a simple and low-cost high energy ball milling method and investigated its reaction mechanisms with Na. Here, the researchers found that during the ball milling process, crystalline P_2S_5 transformed into amorphous P_2S_5 to form an amorphous P_2S_5/C composite that could exhibit a safe average potential of 0.82 V, and the reaction mechanism of the first discharge process may be based on conversion and alloying processes as follows:

Conversion:

$$10Na^{+} + 10e^{-} + P_2S_5 \rightarrow 5Na_2S + 2P$$
 (29)

Alloying:

$$6Na^{+} + 6e^{-} + 2P \rightarrow 5Na_{3}P$$
 (30)

As a result, an anode of this P_2S_5/C material delivered a high reversible capacity of 396 mA h g⁻¹ at 100 mA g⁻¹, a high capacity retention of 99.5% after 100 cycles at 100 mA g⁻¹ and a moderate capacity of 114 mA h g⁻¹ at 2 A g⁻¹ at a high rate (Fig. 32e and f). In addition, a high capacity retention of 89.4% was achieved even after the amorphous P_2S_5/C was cycled over 4000 times at 2 A g⁻¹ (Fig. 32g). However, the initial coulombic efficiency (ICE) of this anode is not high (<40%), which is insufficient for practical SIBs. Even so, the low cost and facile P_2S_5 still shows significant potential for sodium storage if the issue of low ICE can be well addressed.

Theoretical calculations and predictions are also important in the search for new materials for SIB anode applications. For example, Zhao *et al.*²⁶⁹ identified a hitherto unknown P₃C monolayer with a puckered honeycomb structure using firstprinciples swarm-intelligence structural calculations in which they predicted that the theoretical capacity of P₃C as a SIB anode could reach 1022 mA h g⁻¹ and that the acceptable barrier energy and lower open-circuit voltage could ensure good rate capability and safety in practical applications. Here, the researchers attributed the high performances to the unique arrangement of two types of hexagon rings that could produce highly cohesive energy and thermodynamic stability, making P₃C monolayers a promising 2D material for SIB applications.

5 Ternary phosphides

Ternary phosphides are a class of novel phosphide materials in which the synergistic effects of different elements may enhance electron/ion conductivity and mechanical stability, and provide richer redox reactions, or bring about other beneficial effects on SIB anode performances.²⁷⁰ For example, Liang *et al.*²³⁶ prepared 2D MPS₃ (M = Fe, Co, Ni) nanosheets using an efficient solid-

state method in which the MPS₃ single-crystal nanosheets were exposed at the (00l) facet and possessed an average lateral size of \approx 200 nm with an average thickness of \approx 18 nm. And due to the layered structure, good crystallinity, large surface area, excellent electrical conductivity and high pseudocapacitance from surface reactions, the MPS₃ nanosheets provided superior performances as anode materials in SIBs; for example, NiPS₃ nanosheets could deliver a high initial coulombic efficiency of 70% and a reversible capacity of 910 and 316 mA h g^{-1} at current densities of 0.05 and 5 Å g^{-1} , respectively, and a capacity retention of 88% over 200 cycles at 2.0 A g^{-1} . In addition, the interlayer spacing of these MPX₃ nanosheets could be expanded by intercalating guest propylamine molecules, leading to an improved Na storage performance (1090 and 536 mA h g^{-1} at 0.05 and 5.0 A g^{-1} respectively for intercalated NiPS₃ nanosheets). These superior performances as well as the facile preparation method and the low cost of Fe, Ni, P and S indicate that these 2D MPS₃ (M = Fe, Co, Ni) nanosheet composites are promising candidates for practical SIBs. In another example, Marino et al.²⁷¹ synthesized a ternary Ni₂SnP composite as an anode material for SIBs using a simple ball milling method and reported that Ni₂SnP only exhibited a reversible specific charge capacity of \sim 200 mA h g⁻¹ in SIBs, which was much lower than that of the corresponding Li-ion battery. The researchers found that in SIBs, only phosphorus was active in the electrochemical reaction, unlike in the case of binary Sn-based phosphides such as Sn_3P_4 (ref. 91) in which Sn was also active towards Na. Here, the researchers suggested that this reaction mechanism could explain the poor cyclability of anode materials for SIBs and that in particular, the low conductivity of P could result in the formation of only small proportions of Na₃P. The insufficient sodium storage performance of the Ni2SnP composite may hinder its application in SIBs; however, this study gives a suggestion that the total or partial recombination of the pristine phase is crucial to the electrochemical performance of ternary electrode materials. Lan et al.²³⁷ prepared pure phase ternary phosphide-copper tin phosphide (Cu₄SnP₁₀) nanowires by a solution-liquid-solid growth method and investigated the performance of these composites as SIB anodes. The researchers found that Cu incorporation could effectively alleviate Sn aggregation in anodes during charge/discharge cycles, as Sn aggregation was considered to be a major reason for the reduced cycling performances in Sn-P compounds. And a composite of Cu₄SnP₁₀ nanowires combined with multiwalled carbon nanotubes delivered a stable capacity of 512 mA h g^{-1} after 100 cycles at a current density of 100 mA g^{-1} and a rate capacity of 412 mA h g⁻¹ at 1 A g⁻¹. Furthermore, Liu et al.²³⁸ synthesized a ball milled FeSi₄P₄ composite as a SIB anode material and reported a reversible capacity of 180 mA h g⁻¹ at 100 mA g^{-1} with a capacity retention of 99% after 100 cycles. Although the specific capacity is low, the high short-term cycle stability is quite surprising.

In another study, Wang *et al.*²⁷⁰ prepared a composite of holey-structured 2D Ni_{1.5}Co_{0.5}P_x nanosheets as a SIB anode material through a simple solution method and a subsequent phosphorization process. Then these as-prepared 2D holey-structured Ni_{1.5}Co_{0.5}P_x nanosheets were subsequently

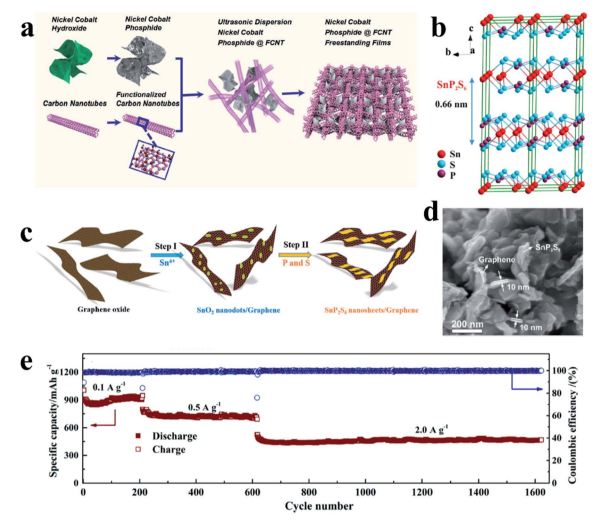


Fig. 33 (a) Schematic diagram of the synthesis of NCP@FCNT-FS free-standing films. Reproduced with permission.²⁷⁰ Copyright 2018, WILEY-VCH. (b) Crystal model showing the asymmetric-layered structure of SnP_2S_6 . (c) Schematic illustration of the *in situ* growth of ultrathin SnP_2S_6 nanosheets on graphene. (d) SEM image of the SPS/G hybrid. (e) Long-term cycling performance at current densities of 0.1, 0.5, and 2.0 A g⁻¹. Reproduced with permission. Copyright 2018, American Chemical Society.

hybridized with surface-functionalized carbon nanotubes (CNTs) to form a lightweight, free-standing and binder-free integrated flexible electrode (Fig. 33a). The researchers reported that the synergistic effects of Co and Ni atoms could enhance charge transfer and promote the release of more capacity as compared with single metal phosphides. In addition, they also suggested that the connected CNT network could affect the formation of conversion reaction products by forming a film structure rather than big-sized microspheres, which could effectively alleviate volumetric changes. As a result, this flexible anode achieved a reversible capacity of 496.4 mA h g^{-1} at 0.5C and a good rate capacity of 276.1 mA h g^{-1} at 8C. However the high cost of CNTs may increase the fabrication cost of this composite and thus hinder its practical application especially for large-scale energy storage systems. Nevertheless, the flexibility of this anode makes it very suitable for wearable products. In addition, this free-standing and binder-free integrated electrode is also beneficial to increasing the volumetric specific capacity and reducing the fabrication cost. Cheng *et al.*²³⁹ designed a strategy to *in situ* coat a tin–phosphorus– oxygen (SnPO) composite with reduced graphene oxide (rGO) and reported that the coating protection could not only increase electrical conductivity, but also buffer volume expansion and suppress the pulverization of electrode materials during cycling. As a result, the SnPO/rGO composite delivered an initial discharge/charge capacity of 843 mA h g⁻¹/450 mA h g⁻¹ at a current density of 50 mA g⁻¹ and retained a reversible discharge capacity of 200 mA h g⁻¹ at a current density of 2000 mA g⁻¹.

Recently, a composite of 2D ternary tin thiophosphate (SnP_2S_6) nanosheets (~10 nm thickness) grown on graphene (denoted as the SPS/G hybrid) was also demonstrated as an intriguing anode for SIBs (Fig. 33b–d).²⁷² Benefiting from the 2D hetero-structural nanojunction and compositional advantages, the SPS/G hybrid showed good electrical conductivity and mechanical resilience to realize rapid ion and electron transfer as well as accommodation of volume change during the

Table 3 Advantages and disadvantages of different preparation methods for RP anodes

Preparation methods	Advantages	Disadvantages	Reference
Ball milling	Simple/facile Large-scale	Uneven/big particle size ($\sim \mu m$)	82 and 121
Vaporization-condensation	Uniform/small size Intimate contact with host materials	Low P loading ratio Uncontrollable distribution	76, 85 and 131
	Well-controlled nanostructure	White P remnants	
Novel methods (solution method and solvothermal method	Controllable size	Simplex morphology (particle or sphere)	87, 127, 144 and 145
Carbothermic reduction)	Refined nanostructure	Toxic reagent involved	

electrochemical process. As a result, the SPS/G hybrid anode exhibited a high reversible capacity (1230 mA h g^{-1} at 50 mA g^{-1}), superior rate capability (200 mA h g^{-1} at 15 A g^{-1}), and an exceptional capacity retention of 76% after 1000 cycles at 2.0 A g^{-1} (Fig. 33e). A full cell based on this anode also exhibited an excellent performance, affording a capacity of 470 mA h g^{-1} at 30 mA g^{-1} (on the basis of anode weight) and a good cycling capacity of 360 mA h g^{-1} at 150 mA g^{-1} after 50 cycles corresponding to a retention of 90.6%. Due to the superior sodium storage performance, the low-cost of Sn, P and S, and the facile and scalable solid-state preparation route, this SPS/G hybrid can have great potential for practical SIBs.

The combination of different kinds of metal phosphides is an interesting and effective strategy to improve performances as SIB anodes. For example, Li et al.75 synthesized a unique coreshell porous FeP@CoP phosphide microcube interconnected with reduced graphene oxide (RGO) nanosheets (RGO@-CoP@FeP) through a solution method combined with a lowtemperature phosphorization process. The researchers reported that the FeP@CoP core-shell structure could provide enough cushion space for volume change and shorten Na⁺ diffusion paths. In addition, the interconnected RGO nanosheets and carbon layers wrapped on the FeP core cubes could build a conductive highway to enhance charge transfer kinetics, leading to a reversible capacity of 551.4 mA h g^{-1} in the first cycle and a capacity of 456.2 mA h g^{-1} maintained in the 200th cycle. In this study, the combination strategy enables facile preparation of this sophisticated core-shell structure, and thus provides an improved sodium storage performance. Zhang et al.273 ball milled a mixture of elementary Sn, Sb, P, and C, and obtained a Sn₅SbP₃/C powder composite which consisted of micrometric agglomerates of active Sn₄P₃, SnSb and Sn nanoparticles. Here, the researchers reported that the carbon in the composites not only acted as a conducting matrix, but also enhanced the ball milling efficiencies and extended electrode cycle lifespans. In addition, each of the active Sn₄P₃, SnSb and Sn phases in the composite could function mutually as a buffer for one another. And as a result, this ternary composite SIB anode delivered a capacity of 352 mA h g^{-1} at a current density of 2 A g^{-1} , which was notably higher than that of binary Sn₄P₃/C and SnSb/C composites produced under the same conditions. In another study,²⁷⁴ a self-supporting three-dimensional (3D) composite of Cu₃P and Co₂P interconnected by N-doped C fibers $(Cu_3P-Co_2P/N-C)$ was prepared by a simple

electrospinning method. The synergistic effects between Cu₃P, Co₂P, and N-doped carbon could increase the electrical conductivity and active sites, ensuring more ion storage. The Cu₃P-Co₂P/N-C anode for SIBs delivered high initial discharge/ charge capacities of 429.5/321.2 mA h g^{-1} accompanied by a high ICE of 74.8%, as well as a stable capacity of around 166.4 mA h g^{-1} after 50 cycles. As the Na storage performance of this anode was inferior to its Li storage performance, it seemed to be more suitable for application in LIBs rather than in SIBs. Nevertheless, this novel nanostructure as well as the corresponding preparation method can still give inspiration for future design of SIB anode materials. Table 3 summarizes the advantages and disadvantages of different preparation methods for RP anodes. Table 4 summarizes some basic properties of RP, BP and metal phosphides, as well as the advantages and disadvantages of these P-based materials used as anodes for SIBs.

6 Summary, challenges and future research directions

6.1 Summary

Sodium-ion batteries are promising alternatives to lithium-ion batteries and have attracted considerable attention due to the abundant availability of sodium and similar electrochemical properties to lithium as well as their great potential to meet the increasing demand for large-scale energy storage systems. Recently, phosphorus-based materials such as red phosphorus and metal/nonmetal phosphides have been considered as the most promising candidates as anodes for SIBs and attracted much research effort. Therefore, to facilitate the further research and development of these materials, this review has comprehensively discussed the status of phosphorus-based materials as anodes for sodium-ion batteries, including red/ black phosphorus, phosphorene, metal/nonmetal-phosphides and ternary phosphides, in which the material design, synthesis strategy, structural/morphology characterization, sodium storage mechanism and electrochemical performance were analyzed. In general, the combination of red phosphorus with carbon or other materials is an effective approach to enhance the conductivity of the anode and to accommodate large volume change during charge/discharge cycles, thus leading to improved specific capacities, rate performances and

				theoretical				
Materials	Crystal structure	Electronic conductivity	Reaction potential (vs. Na/Na ⁺)	gravimetric capacity (mA h g ⁻¹)	Highest initial CE ever reported	Advantages	Disadvantages	Reference
RP (amorphous)	Polymer	$\sim 10^{-12}~\mathrm{S~m^{-1}}$	~0.4 V	2596	86.2%	Natural abundance, eco- friendliness High theoretical	Low electronic conductivity Low initial CE	82 and 125
BP	Orthorhombic	$\sim 3 imes 10^2 \ \mathrm{S} \ \mathrm{m}^{-1}$	\sim 0.45 V	2596	87.7%	gravimetric capacity High electronic conductivity	Difficult to prepare	157 and 174
${ m Sn}_4{ m P}_3$	Hexagonal	$\sim 3 imes 10^3 \ \mathrm{S \ m^{-1}}$	~0.3 V	1132	90.7%	Good maintenance of the structure during cycles High electronic conductivity Good maintenance of the	Low initial CE	77 and 217
						structure High theoretical volumetric capacity Facile synthesis Fast Na diffusion Low cost of Sn and P		
SnP	Trigonal			1209				190
SnP ₃	Rhombohedral	I	~0.3 V	1616	72%	High theoretical volumetric Fast capacity decay capacity High P content, reversibility of the conversion reaction	Fast capacity decay	194
CoP Co ₂ P	Orthorhombic Orthorhombic		~0.2 V	894 540	77.1% 41%	11	High cost of Co Low initial CE Low P content High cost of Co	204
CoP_3	Cubic	\sim 1.8 $ imes$ 10 ³ S m ⁻¹	<0.4 V	1588	87%	High P content	D	200, 228 and 275
FeP	Orthorhombic	$\sim\!\!1.3 imes10^{6}~{ m S~m^{-1}}$	$\sim 0.4~{ m V}$	924	74%	Low cost		201 and 276
FeP_4	Monoclinic	$\sim 3 imes 10^{-3} \mathrm{~S~m}^{-1}$	\sim 0.25 V	1790	84%	High P content, low cost	Low electronic conductivity	70, 192 and 277
NiP ₃	Cubic	$\sim 2 imes 10^{6} \ \mathrm{S} \ \mathrm{m}^{-1}$	\sim 0.2 V	1591	71.3%	High P content, low cost High electronic conductivity		226 and 245
NiP_2	Hexagonal	$\sim 3 \times 10^6 \text{ S m}^{-1}$	~0.1 V	1333	76.7%	High electronic conductivity High P content Low cost	1	92, 245 and 250
CuP,	Monoclinic	${\sim}10^3~{ m S}~{ m m}^{-1}$	$\sim 0.5 \text{ V}$	1282	87%	Low cost	I	2.78

Table 4 Structure phase, electronic conductivity, and electrochemical properties of RP, BP and metal phosphides, and the advantages and disadvantages of these P-based materials as anodes

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Materials	Crystal structure	Electronic conductivity	Reaction potential ($\nu s. Na/Na^+$)	gravimetric capacity (mA h g^{-1})	Highest initial CE ever reported Advantages	Advantages	Disadvantages	Reference
Cu ₃ P	Hexagonal	${\sim}2 imes10^{6}~{ m S~m^{-1}}$	~0.4 V	363	80.6%	Low cost	Low theoretical gravimetric capacity Low P content	78, 257 and 279
GeP ₅	Rhombohedral	${\sim}10^{6}~{ m S}~{ m m}^{-1}$	~0.4 V	1888	94.1%	High initial CE High P content High electronic conductivity	High cost of Ge	193 and 280
GeP ₃ GeP	Rhombohedral Monoclinic	$-5 imes 10^3~{ m S~m^{-1}}$	\sim 0.25	1619 1033	81.3% 93%	High cycle stability High initial CE Low P content	Low initial CE High Ge ratio (high cost)	196 and 263 197 and 281
^{<i>a</i>} The advantage table.	s/disadvantages of all	these P-based anode 1	materials such as a suita	ıble operating potential	, large volume chang	^a The advantages/disadvantages of all these P-based anode materials such as a suitable operating potential, large volume change, and insufficient cycle stability during cycles are not listed in this table.	ability during cycles are n	ot listed in th

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durability. Various red phosphorus nanostructures/composites can be obtained through conventional ball milling or vaporization/condensation techniques and new strategies such as solution-based methods with well-controlled sizes/ morphologies are emerging. Layer-structured black phosphorus and phosphorene also show great potential as anode materials as 2D materials, and many nanostructures/ composites of BP/phosphorene materials have been explored for sodium storage and a comprehensive Na storage mechanism of BP was presented. Metal phosphides show higher electronic conductivities than red phosphorus and are more available than BP/phosphorene. Many nanostructures/ composites of metal phosphides, including Fe-P, Co-P, Ni-P, Sn-P, Cu-P and Ge-P, have been successfully synthesized and demonstrate excellent rate capabilities and cycle stabilities. Non-metal phosphides and ternary phosphides including metal-metal phosphides and metal-nonmetal phosphides have also been explored as anodes of sodium-ion batteries with some promising results obtained.

6.2 Challenges and future research directions

With the rapid advancement of different nanotechnologies, great progress has been achieved in the development of phosphorus-based composites as anode materials for sodiumion batteries. However, several challenges must be overcome before realizing their practical application, such as (1) fast degradation of sodium-ion batteries caused by large volume change of anode materials during charge and discharge; (2) low rate capability, low initial coulombic efficiency (60–85%), and reversible capacity caused by the low electronic conductivity and/or poor diffusion kinetics of sodium ions in phosphorus-based materials.

To overcome these challenges, several possible research directions are proposed:

(1) Further exploration of new phosphorus-composited materials as well as corresponding synthetic methods to obtain novel nanostructures/composites to accommodate large volume change during charge/discharge cycles. Here, the optimization of the structure/morphology/size and the introduction of counter materials with high conductivity and mechanical strengthen are promising approaches.

(2) Further improvements to the rate performance of phosphorus-based anode materials by solving the issues of poor diffusion kinetics, slow reaction kinetics and low electronic conductivity through surface engineering, compositing strategies, doping and other methods/strategies. And further improving the initial coulombic efficiency through rational composition/structure/morphology design and particle surface modification.

(3) Increasing the P content in P-composite anodes to further improve their gravimetric/volumetric specific capacities. Enhancing the phosphorus content in the composite (>70%) or constructing a self-supported anode (binder-/conductive additive-free anode) are all effective strategies to achieve this goal. In addition, fabricating P-based anodes for flexible SIBs is also interesting and meaningful; searching for large-scale, high

Table 4 (Contd.

yield and low-cost synthesis methods to prepare P-based anode materials is also important for practical applications.

(4) Combining experimental characterization with theoretical studies to obtain fundamental understandings of the relationship between the sodium storage performance and composition/structure as well as sodium storage mechanisms. This will allow for the prediction of new anode materials and guide the optimization of the design, synthesis and performance of phosphorus-composited materials for sodium-ion batteries.

(5) Further exploration of the degradation mechanisms of sodium-ion battery anode materials during charge and discharge through both theoretical and experimental approaches along with the development of mitigation strategies to enhance stability and durability.

(6) Further optimization of the design and fabrication of sodium-ion batteries using matching phosphorus-composited anodes, cathodes and electrolytes with respect to battery performance towards large-scale application in energy storage systems.

Conflicts of interest

There are no conflicts of interest to declare.

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