Advanced Anode Materials of Potassium Ion Batteries: from Zero Dimension to Three Dimensions

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HIGHLIGHTS

• This review introduces the recent anode materials of potassium ion batteries classified into 0D, 1D, 2D, and 3D, mainly including carbon materials, metal-based chalcogenides and metal-based oxides, and alloying materials.

• The advantages, disadvantages, and optimized strategies of different dimensional anode materials are summarized.

• The relationship between different dimensional anode materials in potassium ion batteries and the corresponding electrochemical performances is outlined. And some strategies are proposed to deal with the current disadvantages of potassium ion batteries.

ABSTRACT Potassium ion batteries (PIBs) with the prominent advantages of sufficient reserves and economical cost are attractive candidates of new rechargeable batteries for large-grid electrochemical energy storage systems (EESs). However, there are still some obstacles like large size of K⁺ to commercial PIBs applications. Therefore, rational structural design based on appropriate materials is essential to obtain practical PIBs anode with K⁺ accommodated and fast diffused. Nanostructural design has been considered as one of the effective strategies to solve these issues owing to unique physicochemical properties. Accordingly, quite a few recent anode materials with different dimensions in PIBs have been reported, mainly involving in carbon materials, metal-based chalcogenides (MCs), metal-based oxides (MOs), and alloying materials. Among these anodes, nanostructural carbon materials with shorter ionic transfer path are beneficial for decreasing the resistances of transportation. Besides, MCs, MOs, and alloying materials with nanostructures can effectively alleviate their stress changes. Herein, these materials are classified into 0D, 1D, 2D, and 3D. Particularly, the relationship between different dimensional structures and the corresponding electrochemical performances has been outlined. Meanwhile, some strategies are proposed to deal with the current disadvantages. Hope that the readers are enlightened from this review to carry out further experiments better.

KEYWORDS Potassium ion batteries; Anode; Structure design; Nanomaterials; Dimensions

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1 Introduction

In 2004, it was the first time that Ali Eftekhari proposed the prototype of PIBs [1]. Nonetheless, the later stagnation about PIBs researches was ascribed to the safety issues about the K as well as technologies about other metal ion batteries becoming increasingly popular [2]. Owing to fossil fuels used increasingly as well as the severity of global warming, renewable and sustainable energy sources have been exploiting, including solar, wind, rain, geothermal, tide, and wave energies. Nevertheless, EESs must be considered to solve the intermittent issues of clean energy, which has a great effect on storing and delivering these energy resources [3, 4]. So far, lithium ion batteries (LIBs), as a representative energy storage technology, have been widely explored for portable devices, electrical vehicles, and large-grid EESs because of high energy density and stable cycling lifespan [5]. Unfortunately, there are some barriers for LIBs to develop sustainably, such as challenging lithium (Li) recovery, uneven distribution of Li, and increasing cost [6–8]. Accordingly, it is necessary to search for alternative rechargeable batteries’ technologies. Based on cost and resource considerations, scientists have put a lot of effort into developing a series of non-lithium ion batteries, including sodium ion batteries (SIBs) [9], PIBs [10], magnesium ion batteries (MIBs) [11], zinc ion batteries (ZIBs) [12], aluminum ion batteries (AlIBs) [13] and so on [14]. Since zinc, magnesium, and aluminum are less active than lithium, they could be used as anode materials for metal ion batteries [15–17]; especially, magnesium and aluminum anodes do not even form dendrites, so their corresponding ion batteries are able to meet the safe requirement [18, 19]. In addition, commercial LIBs mainly use graphite with a theoretical specific capacity of 372 mAh g⁻¹ as anode material, while the capacities of zinc (820 mAh g⁻¹) [20], magnesium (2205 mAh g⁻¹), and aluminum (2980 mAh g⁻¹) anodes are much higher than that of graphite [18]. Clearly, each multivalent cation can exchange more than one electron, suggesting that if a host material can store the same number of cations, the capacities of multivalent ion cells are several times that of monovalent ion cells. Therefore, these multivalent ion batteries may have higher energy densities. Furthermore, the ion radii of zinc ions (0.74 Å) [12], magnesium ions (0.72 Å) [21], and aluminum ions (0.535 Å) [21] are relatively small, having low trends to damage the structure of host material. However, although these multivalent metal ion batteries have various irreplaceable advantages, they also face some intractable problems. First of all, the surface charge density of these multivalent metal ions is relatively high, resulting in greater mutual repulsion between cations and greater interaction between cations and host materials, which is not conducive to high capacity and high rate performance for batteries [22]. Besides, the potentials of Zn/Zn²⁺ (−0.76 V vs. SHE) [23], Mg/Mg²⁺ (−2.37 V vs. SHE) [14], Al/Al³⁺ (−1.76 V vs. SHE) [13] are much higher than that of Li/Li⁺ (−3.04 V vs. SHE), indicating that it is difficult for these batteries to obtain high operating voltages. There are many other problems that hinder their development. For instance, the zinc anode is still plagued by dendrites problem [24] and the electrolyte used in MIB is not ideal [14, 25, 26]. Compared with MIBs, ZIBs, and AlIBs, both the working principle and the electrode materials and electrolytes used in SIBs and PIBs are similar to those of LIBs, for the reason that sodium, potassium, and lithium have similar physical and chemical properties [27]. It is expected that PIBs and SIBs have fewer obstacles for commercial applications.

PIBs aroused remarkable attentions again in 2015 and increasing researches have been publishing since then (the inset of Fig. 1). Firstly, PIBs possess similar rocking-chair operating principle compared with LIBs, which provides a favorable foundation for the studies of PIBs [28, 29]. Secondly, PIBs with lower price and sufficient resources are suitable for EESs [30]. Thirdly, the standard redox potential versus SHE of K⁺/K (−2.93 V) is not only even comparable to that of Li⁺/Li (−3.04 V) but also lower compared with that of Na⁺/Na (−2.71 V); thus, PIBs are beneficial to produce higher operating voltages [31, 32]. Meanwhile, the lowest potential versus SHE of K⁺/K (−2.88 V) is compared with that of Na⁺/Na (−2.56 V) and Li⁺/Li (−2.79 V) in some non-aqueous electrolytes like propylene carbonate solvent, which makes PIBs benefit from wider potential window to achieve high energy density [2, 33, 34]. Besides, Okoshi et al. indicated that weaker Lewis acidity and low K-ion desolvation energy brought about the smaller Stokes radius of solvated ions and the low interfacial reaction resistance, respectively, which made K electrolytes possess higher conductivity compared with Li as well as sodium (Na) electrolytes [35]. In addition, the commercialized graphite has been successfully explored for PIBs anode and the theoretical capacity is around 279 mAh g⁻¹, but it is not suitable
for SIBs [28, 29, 31]. Furthermore, the unwanted K-metal deposition on the surface of anode may be hindered due to the insertion potential versus K+/K of K-ion (0.2 V), which is beneficial for improving safety in terms of operation [32]. Moreover, compared with Li metal (180.54 °C) and Na metal (≈ 98 °C), K-metal lower melting point (63.38 °C) makes the dendritic K melting to provide a secure capability under an appropriate temperature [10, 36, 37]. It is worth mentioning that Cu foil can be replaced with Al foil as current collectors in anode electrode of PIBs without forming Al-K alloy, cutting down the batteries’ production expenses [29]. Given the above-mentioned advantages, PIBs are promising alternative batteries for LIBs. To date, the reported PIBs anode materials mainly contain carbon materials (e.g., carbon nanotubes (CNTs), graphene, and graphite), MCs (e.g., MoS2 and MoSe2), MOs (e.g., MoO2 and SnO2), and alloying materials (e.g., antimony (Sb), tin (Sn), bismuth (Bi), germanium (Ge), and phosphorous (P)) (Fig. 1).

Fig. 1 Cycle capacity of various PIBs anode materials reported recently. Sb2S3-SNG [38]; Co3O4-Fe2O3/C [39]; black phosphorus = BP [33]; SnP2/C [40]; CoS@G [41]; CNC [42]; PMC [43]; Sb@C [44]; P/C [45]; MoO2/rGO hollow sphere composites = MoO2/rGO [46]; Mn3O4@rGO [47]; SnP2/C [48]; SnO2 nanoparticles anchored on carbon foam = SnO2@CF [49]; P@TBMCS [50]; NCNF@CS [51]; alkali- ized Ti3C2 MXene nanoribbons = Ti3C2 [52]; chitin-derived NCNFs = C-NCNFs [53]; bamboo-like NCNTs = b-NCNTs [54]; MOF-derived NCNTs = M-NCNTs [55]; red P [56]; segment-like Sn nanorod encapsulated in hollow carbon tube = SnHCT [57]; K2Ti6O13 [58]; Bi@C [59]; Ti3CNTz [60]; MoS2@rGO [61]; MoS2@SnO2@C [62]; graphene-like VSe2 nanosheets = VSe2 [63]; Few-layered SnS2 nanosheets supported on rGO = SnS2/rGO [64]; FLNG [65]; PODG [66]; HPCS [76]; N/O co-doped mesoporous carbon octahedrons = NOHPHC [77]; rGO aerogel = rGO [78]; GNCs [79]; honeycomb hard carbon = HHC [80]. And the inset about number of articles obtained from Web of Science on PIBs anode materials and based on the corresponding Topic about potassium ion battery, potassium ion storage, potassium storage, and anode (accessed: July 1, 2020)
practical application of alkali ion batteries can be influenced because of the choice of suitable anode materials based on selection principle. Therefore, the selection principle of LIBs, SIBs, and PIBs should be considered about several aspects in detail: (1) The materials can be regarded as anode materials, depending on whether the material can react with alkali ions and the corresponding theoretical capacity; (2) The structural stability and durability should also be considered because the robust structures can better tolerate with the stress variation during long cycles; (3) The operability of the materials is also one of the considered factors, so the materials affected easily by environmental conditions like oxygen should be excluded; (4) The cost of materials also affects the practical application of batteries; (5) The recyclability of electrode materials should be paid more attention to make good use of resources and protect environment. Therefore, based on aforementioned selection principle, the anode materials can be better chosen. However, it does not mean that all anode electrodes used in LIBs can be successfully used in SIBs and PIBs, although LIBs, SIBs, and PIBs have similarities in the energy storage mechanism. For examples, the aforementioned graphite can be used as the commercial LIBs electrode materials but not suitable for Na+. This phenomenon is possibly attributed to two main reasons: (1) Ionic insertion can be affected because of different storage reactions between ions and the corresponding active materials and (2) Different ion radii are necessary to search for related different material accommodating their ionic size, such as Li⁺ (0.76 Å), Na⁺ (1.02 Å), and K⁺ (1.38 Å). So some anode materials may be conducive to hold Li⁺ but not in favor of containing large K⁺.

The aforementioned electrode materials have exhibited distinct electrochemical performances due to their unique features. Meanwhile, it is well known that the materials properties will be affected due to different dimensional structures. Zero-dimensional (0D) nanomaterials can promote ionic adsorption and mitigate stress variation owing to surface effect and small size effect. As 0D nanomaterials, SnO₂ nanoparticles anchored on carbon foam not only facilitated electrolyte penetration but also boosted K-ion transport, which realized the cycle capacity of 231.7 mAh g⁻¹ at 1 A g⁻¹ after 400 cycles as well as the rate capacity of 143.5 mAh g⁻¹ at 5 A g⁻¹ [49]. Furthermore, one-dimensional (1D) nanomaterials with high length-to-diameter ratio are beneficial to enhance the electronic and ionic transfer and provide high mechanical robustness. For example, owing to unique architecture, nitrogen (N)-doped carbon nanofibers (CNFs) as electrode materials in PIBs exhibited outstanding performances, delivering a capacity of 146 mAh g⁻¹ at 2 A g⁻¹ after 4000 cycles [81]. In addition, two-dimensional (2D) nanomaterials are also capable of improving ionic adsorption and facilitating ionic diffusion owing to high surface area as well as tunable interlayer spacing. As one of the typical 2D materials, few-layer bismuthene explored for PIBs anode materials could facilitate electrolyte infiltration, boost K-ion transfer, and buffer volumetric expansion in the course of charge and discharge process, so a capacity of more than 200 mAh g⁻¹ was delivered at 20 A g⁻¹ over 2500 cycles [82]. As for three-dimensional (3D) nanomaterials, with high mechanical strength and interconnected structures, they can effectively tolerate stress variation and make largely electrode contact with electrolyte as well as promote electronic conductivity. In 2019, 3D reduced graphene oxide (rGO) aerogel was fabricated to enhance K-ion transfer, improving rate performance (92 mAh g⁻¹ at 6.7 C) as well as cycle performance (267 mAh g⁻¹ at C/3 after 100 cycles) [78].

In this review, 0D, 1D, 2D, and 3D nanomaterials about the recent developed PIBs anode materials will be introduced (Fig. 2), mainly concentrating on carbon materials, MCs and MOs, and alloying materials. Among them, carbon materials with the advantage of low price are attributed to its abundant sources, which is beneficial for practical applications. In addition, the existing technology can be used to prepare carbon nanomaterials, making the synthesis process convenient. And then, the structures of carbon materials are relatively stable and durable, which helps to keep structural stability and improve the long life of the corresponding electrode during charging and discharging. As for MCs and MOs, they have high capacity and it is easy to synthesize. Besides, alloying materials not only possess high theoretical capacity, such as Sb (660 mAh g⁻¹) [83] and Bi (385 mAh g⁻¹) [83], but also can react with K-ion under low potentials (~0.1–0.8 V vs. K⁺/K⁺) [72, 84–88]. Additionally, similar to MCs and MOs, alloying materials can be prepared by some easy synthesis methods, so it is convenient to obtain the corresponding products [2]. According to these advantages, alloying materials can be considered as anode materials of PIBs. Furthermore, the structural design and the corresponding electrochemical performances of different dimensional anode materials will be summarized. To date, the capacities of 0D–3D carbon materials have varied from 200 to 600 mAh g⁻¹. Moreover, MCs and MOs...
with 0D–3D structures have delivered the capacities with 50–600 mAh g\(^{-1}\). In addition, the capacities of 0D–3D alloying materials varied from 300 to 1000 mAh g\(^{-1}\) have been studied [36]. Meanwhile, some practical strategies about solving some challenging issues will be proposed.

### 2 Zero-Dimensional Nanomaterials for PIBs

0D nanomaterials are defined as 1 nm to 100 nm in three dimensions. So far, 0D nanomaterials have various types, including quantum dots, nanoparticles, nanospheres, nanocages, core–shell structures, and so forth. Typically, these 0D nanomaterials have been widely studied in energy storage field due to structural features and properties including surface effect, small size effect, and so on; especially, the small size of 0D nanomaterials possesses large surface area, providing sufficient sites for ionic adsorption [94, 95]. Additionally, the stable structure plays a crucial role in buffering large volume expansion, like hollow 0D nanomaterials. Therefore, 0D nanomaterials are beneficial for anode materials of PIBs to keep contact with electrolyte and restrict volumetric variation. However, the electrochemical performances will be affected due to 0D nanomaterials with easier self-aggregation. Thus, it is better to be composited with other materials to inhibit aggregation for anode materials. Next, the relationship between 0D nanomaterials and the corresponding electrochemical performances will be elaborated in detail. In addition, the initial Coulombic efficiency (C.E.), rate performances, and cycle properties of recent reported 0D anode materials of PIBs are summarized in Table 1.

#### 2.1 Zero-Dimensional Carbon Materials

0D carbon materials with shorter ionic diffusion pathway are advantageous to reduce transport resistances. Accordingly, a few studies about nanoparticles have recently been reported so as to improve the electrochemical behavior. Carbon nanoparticles with N/P co-doping and expanded interlayer (NP-CNPs) were fabricated for the PIBs electrode materials [107]. The synergistic effect of nano-size and P/N-co-doping helped to obtain good electrochemical performances. Then, the NP-CNPs electrode had a capacity of 157 mAh g\(^{-1}\) at 5.0 A g\(^{-1}\) (Fig. 3c). In addition, a capacity of 190 mAh g\(^{-1}\) was obtained at 1.0 A g\(^{-1}\) after 4000 cycles (Fig. 3d). According to experimental test, the nanoparticles with the expanded interlayer and uniform ultrafine nanoparticles were exhibited (Fig. 3a, b). Obviously, the morphology of nanoparticles and its expanded interlayer could effectively improve the electrochemical performances, which not only provided shorter K-ion transfer path as well as improved electrical conductivity, but also enhanced adsorption capability toward K-ions.

Besides, novel hollow structures may be advantageous for accommodating the volumetric changes compared with solid nanoparticles. For example, graphite has been wildly utilized but still suffered from large volumetric variation during charge and discharge process as PIBs electrode. Although some progress has been achieved in terms of regulating structure including polynanocrystalline graphite, activated carbon, and expanded graphite, it is still challenging to ensure the robust stability of graphite during long cycling process [74, 108–110]. It is worthwhile that hollow nanocages have gradually been used for electrode materials. Cao et al. [42] synthesized graphitic carbon nanocage (CNC) as anode material for PIBs by taking

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**Fig. 2** Schematic diagram of various 0D–3D PIBs anode materials. Reproduced with permission from Ref. [42, 43, 68, 89]. Copyright 2018 (2019), John Wiley and Sons; from Ref. [66, 90, 91]. Copyright 2017 (2019, 2020), Royal Society of Chemistry; from Ref. [50]. Copyright 2018, Elsevier; from Ref. [69, 72, 92, 93]. Copyright 2017 (2018, 2019), American Chemical Society
advantage of Ketjen carbon black. From Fig. 3e, f, it can be clearly seen that the CNC possessed interlinked structural appearance, which benefited fast electronic transfer. As for the electrochemical performances, reversible capacities of the CNC electrode were 221.5 mAh g⁻¹ as well as 175 mAh g⁻¹ at 0.1 and 35 C, respectively (Fig. 3g). As expected, the electrochemical behavior of the obtained electrode was due to its unique structure features. In detail, the cage-like structure kept the stability of structure by reducing the anisotropy and avoiding interlayer slipping. Then, the cage-like hollow interior could retain the structural integrity by accommodating volume changes well in the course of charging and discharging. Similarly to soft carbon, hollow graphitized carbon nanocages (HGCNs) were also synthesized as effective PIBs electrode materials [111]. The authors exhibited that HGCN-1000 (1000 mean the carbonization temperature of 1000 °C) with highly graphitized carbon-cage, developed porosity, and hollow structure was endowed robust structural stability and good electrochemical performances. Therefore, the capacity retention of 95.9% for the HGCN-1000 electrode was achieved at 1 A g⁻¹ over 2000 cycles, which identified better cyclic stability.

Table 1 Comparison of the state-of-the-art performances of 0D anode materials in PIBs

| Materials                           | Initial C.E. (%) | Rate capacity (mAh g⁻¹) at the current density (mA g⁻¹) | Cycle capacity (mAh g⁻¹) at the current density (mA g⁻¹) (cycle number) | References |
|-------------------------------------|------------------|----------------------------------------------------------|------------------------------------------------------------------------|------------|
| Sb₂S₃-SNG                           | 69.7             | 340 at 1000                                              | 480.078 at 50 (100)                                                   | [38]       |
| Co₃O₄-Fe₂O₃/C                       | 54               | –                                                        | 220 at 50 (50)                                                        | [39]       |
| BP                                 | 60               | 300 at 2000                                              | 270 at 50 (50)                                                        | [33]       |
| Sn₃P₄/C                             | –                | 221.9 at 1000                                            | 307.2 at 50 (50)                                                     | [40]       |
| Sn@C                                | 75.8             | 127 at 2000                                              | 160 at 1000 (800)                                                    | [44]       |
| Sn@PC                               | 46.2             | 200 at 2000                                              | 90 at 500 (200)                                                      | [96]       |
| P/C                                 | 50.3             | 90 at 500                                                | 71.5 at 500 (500)                                                  | [45]       |
| ZNP/C                               | 58.5             | 46 at 2000                                               | 145 at 500 (300)                                                    | [97]       |
| Sn@NPMC                             | 50               | 161 at 1000                                              | 130 at 1000 (1500)                                                  | [98]       |
| MoO₂/rGO                            | 51.6             | 176.4 at 500                                             | 104.2 at 500 (500)                                                  | [46]       |
| Co₉Se QDs/C                         | 61.8             | 220 at 2000                                              | 228 at 1000 (500)                                                   | [99]       |
| S,N co-doped thin carbon            | –                | 64 at 4000                                               | 65 at 2000 (900)                                                   | [100]      |
| MnO₂@rGO                            | 66               | 95 at 10,000                                             | 635 at 500 (500)                                                    | [47]       |
| Fe₃O₄@NFLG                          | 46               | 176 at 5000                                              | 140 at 5000 (5000)                                                  | [101]      |
| Titanium oxyxnitride nanoparticles/carbon | –             | 72 at 1600                                               | 150 at 200 (1250)                                                   | [102]      |
| SnP/C                               | 58.8             | 221.8 at 1200                                            | 225 at 500 (80)                                                     | [48]       |
| SnO₂@3D PC                          | 13.99            | 144.6 at 2000                                            | 270.3 at 100 (200)                                                  | [103]      |
| VN-QDs/CM                           | 72.9             | 152 at 2000                                              | 215 at 500 (500)                                                    | [104]      |
| Co₉S₈/N-C@MoS₂                       | 89.1             | 50 at 1000                                               | 100 at 1000 (100)                                                   | [105]      |
| Carbon-coated mesoporous Co₉S₈ nanoparticles supported on rGO | 59.5             | 215.1 at 5000                                            | 210.8 at 1000 (1200)                                                | [106]      |
| SnO₂@CF                             | 44.43            | 143.5 at 5000                                            | 231.7 at 1000 (400)                                                 | [49]       |

2.2 Zero-Dimensional MCs and MOs

MCs and MOs have proved to be a kind of hopeful PIBs electrode materials, but severe volume changes and low electrical conductivity have hindered their development. It is well known that the minimization of materials is conducive to alleviating stress changes, so the preparation of 0D MCs
and MOs is an effective method. In order to solve low conductivity and agglomeration problems, MCs and MOs are usually combined with conductive materials.

Quantum dots with quantum confinement are practical architectures for boosting electronic and ionic transportation. A composite about cobalt sulfide compounded with graphene (CoS@G) was synthesized for PIBs (Fig. 4a) [41]. In this composite, graphene offering landing platform for CoS quantum dots not only contributed to restraining the agglomeration of CoS quantum dots, but also improved electronic conductivity. Then, numerous smaller CoS nanoclusters were uniformly dispersed onto the graphene nanosheets, consisting of the interlinked quantum dots (Fig. 4b, c). As illustrated in Fig. 4d, the

Fig. 3 NP-CNPs’ a schematic diagram, b SEM image, e as well as N–CNPs’ rate capacities in the range of 0.05–5.0 A g⁻¹, d long cycle capacities at 1.0 A g⁻¹ over 4000 cycles. Reproduced with permission from Ref. [107]. Copyright 2019, American Chemical Society. e SEM image of CNC. f Schematic illustration of CNC. g Discharge capability retention corresponding to rate performance (inset). Reproduced with permission from Ref. [42]. Copyright 2018, John Wiley and Sons
CoS@G-25 (25 means 25% graphene oxide) electrode delivered the discharge capacity of 310.8 mAh g⁻¹ at 500 mA g⁻¹ after 100 cycles. Moreover, the CoS@G-25 electrode’s capacities were retained 67.3% and 56.2% at 3 C and 4 C, respectively. Therefore, it is effective for novel shapes to improve the corresponding electrochemical performances. Coating is also an effective means to prevent agglomeration and buffer volume changes, such as building a core/shell structure. So Wang et al. [43] synthesized MoSe₂/C composites with pistachio-shuck-like morphology (PMC) as PIBs anodes to improve their performances. As shown in Fig. 4e, the packing density of the unique PMC was enhanced via plane-to-plane contact. It could expedite electronic transfer as well as K-ion diffusion and retain the structural stability during the process of potassium/depotassium. Additionally, the PMC exhibited pistachio-shuck-like morphology with the diameter of around 70–90 nm (Fig. 4f). In the electrochemical test, the PMC electrode delivered 226 mAh g⁻¹ at 1.0 A g⁻¹ after 1000 cycles (Fig. 4g). The PMC electrode’s electrochemical performances were due to the core of MoSe₂, the pistachio-shuck-like structure, and thin amorphous carbon
shell. These structural features could facilitate K-ion transfer and buffered the volume expansion.

Besides, titanium dioxides (TiO$_2$) have been utilized in LIBs and SIBs but seldom reported for PIBs. Therefore, in 2019, Fang et al. [121] synthesized a composite about TiO$_2$ nanoparticles anchored on rGO for PIBs. As for this composite, the small size TiO$_2$ could boost K-ion diffusion and the rGO could promote electronic transfer. So its electrochemical performances could be enhanced, which achieved the rate capacity of 107.1 mAh g$^{-1}$ at 1000 mA g$^{-1}$ and the cycling capacity of 88.4 mAh g$^{-1}$ at 1000 mA g$^{-1}$ after 1000 cycles. Nevertheless, its electrochemical behavior is necessary to be improved by further experiments.

2.3 Zero-Dimensional Alloying Materials

Alloying materials including Sb, Sn, Bi, Ge, as well as P, characterized by high capacity, have been studied for PIBs anode materials [84]. Volume expansion is one of their most obvious shortcomings, and the stress variation can be affected by small size structures. Therefore, it is urgent to design appropriate electrode materials and designing suitable nanostructures is a common modification method.

It was reported that nanoparticles have been widely fabricated as electrode materials to mitigate stress changes. As a typical example, engineering bulk materials to nanoparticles is beneficial to alleviate stress variation. Thus, the network of a carbon sphere enclosed Sb nanoparticles (Sb@CSN) as PIBs anode [90]. The schematic diagram of Sb@CSN and its TEM image are shown in Fig. 5a, b, which indicated the Sb nanoparticles were well encapsulated in the carbon sphere with uniform distribution. As for electrochemical performances, the Sb@CSN electrode presented high discharge capacity of 626 mAh g$^{-1}$ at 200 mA g$^{-1}$ after second cycle (Fig. 5c). In addition, the capacity of 504 mAh g$^{-1}$ was obtained at 200 mA g$^{-1}$ after 200 cycles (Fig. 5d). The Sb@CSN anode’s electrochemical behavior indicated the importance of small size nanoparticles and carbon sphere network, which could buffer volume expansion and promote the transportation of electrons. Similarly, Ge and co-workers [87] also constructed a composite composed of Sb nanoparticles and carbon materials. They encapsulated ultra-small Sb nanocrystals into CNFs for PIBs anode, which achieved 225 mAh g$^{-1}$ at 1 A g$^{-1}$ after 2000 cycles because ultra-small Sb nanocrystals and hollow nanochannels not only promoted K$^+$ fast diffusion but also buffered strain variation. Compared to carbon spheres, CNFs may have better connectivity and can better make the electrode stable. Additionally, the nanostructural design also has an impact on the electrochemical performances of red P. So tailoring different sizes of red P into 3D carbon could effectively overcome the above-mentioned disadvantages (Fig. 5e) [122]. In this composite, the small size of red P and 3D carbon nanosheet framework could reduce the degree of volume expansion; on the other hand, it also could boost the fast electron transportation during charge and discharge process. In addition, red P without agglomeration was displayed, which indicated that red P was uniformly dispersed in the 3D carbon nanosheet frameworks (Fig. 5f). Benefit from the structural design, the red P@CN composite anode delivered a charge capacity of 715.2 mAh g$^{-1}$ and a rate capacity of 323.7 mAh g$^{-1}$ at 2000 mA g$^{-1}$ (Fig. 5g, h). Based on nanoparticles with effective impact, different morphology of 0D alloying materials should be designed to enhance the electrochemical performances.

3 One-Dimensional Nanomaterials for PIBs

1D nanomaterials are defined as high length-to-diameter aspect ratios, including nanotubes, nanorods, nanowires, nanofibers, and nanoribbons. Since CNTs were first discovered by Iijima [123], 1D nanomaterials with similar structures have been widely studied. The structural features of 1D nanomaterials not only include nanoscale and microscale joint effects, but also contain high aspect ratio and oriented growth direction [124]. Therefore, 1D nanomaterials are beneficial for fast transportation of electrons and ions and have an impact on tolerating the stress changes, which was considered as a class of most promising materials for high performances of energy storage systems [125]. Given these mentioned above, 1D nanomaterials can be utilized to boost the diffusion of electrons and ions as well as alleviate the stress variation as PIBs electrode materials. Significant progress has recently been achieved for anode materials with high performances by designing and fabricating 1D nanomaterials. Based on typical 1D nanomaterials, systematic discussion is carried out about the relationship between the related structures and their electrochemical performances. Furthermore, in order to comprehensively comprehend the electrochemical behavior of 1D nanomaterials, the initial
C.E., rate performances, and cycle properties of recent reported 1D anode materials of PIBs are outlined in Table 2.

### 3.1 One-Dimensional Carbon Materials

CNTs can be considered as a representative example among 1D carbon materials. It is well established that CNTs with remarkable mechanical strength and excellent electrical properties have been applied in PIBs field [136–139]. So far, CNTs not only can be used directly or doped with other elements, but also can be compositied with other materials to serve as electrode materials. As for composites, a CNT-backboned mesoporous carbon confining red P composite (P@TBMC) was formed as anode material for SIBs/PIBs (Fig. 6a) [50]. P encapsulated completely within the pores, and homogenous distributions in the composite were confirmed (Fig. 6b). In these composites, the main function of red P was to provide high K storage capacity. The multi-walled CNTs not only supported the structure as backbone, but also boosted the electrons transfer because of the high content of $sp^2$ carbon. Therefore, the P@TBMC-2.4 (2.4 means the resorcinol mass (g) in the recipe) anode exhibited a depotassiation capacity of 244 mAh g$^{-1}$ at 0.5 A g$^{-1}$ after 200 cycles, which displayed stable cyclability (Fig. 6c).

Generally, heteroatom-doped carbon materials have better K storage properties than non-doped ones, so
heteroatom-doped CNTs may perform better than ordinary CNTs. So a CoSe$_2$ strung by CNTs with N-doping (NCNF@CS) was prepared by two-step hydrothermal means as a PIBs anode (Fig. 6d) [51]. Then, the CNTs made every octahedral CoSe$_2$ particle arrange in sequence with zigzag void space among particles (Fig. 6e). Remarkably, the NCNF@CS-6 h (6 h means the reaction hours) anode reached the reversible capacity of 196 mAh g$^{-1}$ at 2.0 A g$^{-1}$ (Fig. 6f). Moreover, a capacity of 173 mAh g$^{-1}$ at 2.0 A g$^{-1}$ after over 600 cycles was exhibited (Fig. 6g). The electrochemical performances of NCNF@CS-6 h electrode were ascribed to the use of CNTs and octahedral CoSe$_2$ particles arranging in sequence, where conductive N-doped CNTs not only hindered the agglomeration and anchored the active materials as backbone but also effectively promoted the electronic transfer. Based on aforementioned work, CNTs can enhance the electrochemical behaviors by boosting electronic and ionic diffusion and keeping structural integrity. Therefore, novel composites about CNTs deserve to research for PIBs anode materials in the further experiments.

### 3.2 One-Dimensional MCs and MOs

To date, 1D MCs and MOs mainly include various K-Ti–O anode materials with common formula K$_2$Ti$_{6}O_{2n+1}$, which are advantageous for insertion/extraction of K-ion [140]. As for K$_2$Ti$_6$O$_{13}$, its crystal structure with open 3D framework and tunnel can facilitate K-ion transfer. So K$_2$Ti$_6$O$_{13}$ nanorods were synthesized as PIBs anode and the obtained electrode achieved stable long cycling performance, but its poor rate performance was exhibited [58]. Accordingly, carbon coating may be a practical way to further improve the rate capacities. Therefore, oriented nanorod-like K$_2$Ti$_6$O$_{13}$ bunches with a thin carbon layer (KTO/C) were fabricated as PIBs electrode materials (Fig. 7a) [76]. The size of nanorods of KTO/C-700 (700 means the heat treatment temperature

### Table 2 Comparison of state-of-the-art performances of 1D anode materials in PIBs

| Materials | Initial C.E. (%) | Rate capacity (mAh g$^{-1}$) at the current density (mA g$^{-1}$) | Cycle capacity (mAh g$^{-1}$) at the current density (mA g$^{-1}$) (cycle number) | References |
|-----------|-----------------|---------------------------------------------------------------|--------------------------------------------------------------------------------|------------|
| Ti$_3$C$_2$ | – | 60 at 300 | 42 at 200 (500) | [52] |
| Cryptomelane-type MnO$_2$/CNT hybrids | 40.07 | 127.2 at 1000 | 226.5 at 100 (500) | [126] |
| C-NCFNs | 37.8 | 84.7 at 5 C | 103.4 at 2 C (500) | [53] |
| b-NCNTs | 23.3 | 186 at 1000 | 204 at 500 (1000) | [54] |
| M-NCNTs | 24.45 | 102 at 2000 | 102 at 2000 (500) | [55] |
| Highly N-doped CNFs | 49 | 101 at 20,000 | 146 at 2000 (4000) | [81] |
| Cup-stacked NCNT mats | 14.2 | 75 at 1000 | 236 at 20 (100) | [127] |
| Red P | 68.26 | 71 at 3000 | 300 at 1000 (60) | [56] |
| Sb/HCT | 70 | 211.5 at 5000 | 300.1 at 2000 (120) | [57] |
| 3D amorphous carbon encapsulated CoS/CNTs on CoS-coated CNTs | 57.6 | 133.1 at 6400 | $\approx$130 at 3200 (600) | [128] |
| Carbon-encapsulated CoP nanoparticles embedded in CNTs supported on CNFs | 53.2 | $\approx$292 at 3200 | 247 at 800 (1000) | [129] |
| Sub-micro-carbon fiber@CNTs | – | 108 at 5 C | more than 193 at 1 C (300) | [130] |
| Pyrrrolic/pyridinic-N-doped necklace-like hollow carbon | – | 204.8 at 2000 | 161.3 at 1000 (1600) | [131] |
| Bi-nanorod networks confined in N, S co-doped carbon matrix | 65 | 289 at 6000 | 285 at 5000 (1000) | [132] |
| rGO/CNT hybrid papers | – | 110 at 100 | 148 at 50 (200) | [133] |
| Porous Mn–Fe-Se composite adhered/inserted with interlaced CNTs | 33.43 | 83 at 800 | 141 at 50 (70) | [134] |
| Hollow NCNFs anchored hierarchical FeP nanosheets | 57 | 103 at 800 | 210 at 100 (1000) | [135] |
of 700 °C) was well maintained compared with that of bare KTO (Fig. 7b). As for rate capacities, the KTO/C-700 electrode delivered 65.1 mAh g\(^{-1}\) at 500 mA g\(^{-1}\) (Fig. 7c). The electrochemical performances of the KTO/C-700 electrode were due to K-ion diffusion boosted by crystal orientations of KTO and electron conductivity improved by carbon layer. Besides, in 2016, K\(_2\)Ti\(_4\)O\(_9\) was fabricated as PIBs anode materials for the first time, but delivered low capacities and poor cycling performance [141]. It is necessary to control the morphology of K\(_2\)Ti\(_4\)O\(_9\) well so as to improve the electrochemical behavior. Therefore, K\(_2\)Ti\(_4\)O\(_9\) nanoribbons derived from MXene have been studied for PIBs anode materials due to unique structural features. Wu's group [92] reported that ultrathin nanoribbons of potassium titanate
Fig. 7  a Schematic diagram of KTO/C.  b SEM image of KTO/C-700.  c Rate capacities of the obtained hybrids electrode. Reproduced with permission from Ref. [140]. Copyright 2020, Royal Society of Chemistry.  d M-KTO nanoribbons’ schematic illustration.  e M-KTO’s HRTEM image with the corresponding SAED patterns (inset).  f M-KTO’s rate performance.  g Long cycle capacities as well as C.E. of the obtained sample at 200 mA g⁻¹. Reproduced with permission from Ref. [92]. Copyright 2017, American Chemical Society
(M-KTO, K$_2$Ti$_4$O$_9$) were synthesized as anode materials for PIBs. The morphology of nanoribbons and the interlayer space with 0.93 nm were exhibited, which benefited K-ion insertion/extraction (Fig. 7d, e). As for electrochemical performances, the rate capacity of 81 mA h g$^{-1}$ was attained at 300 mA g$^{-1}$ as well as the retention of 51% (of the second charge capacity) was exhibited at 200 mA g$^{-1}$ after over 900 cycles (Fig. 7f, g). The M-KTO electrode’s electrochemical performances were ascribed to suitable interlayer spacing, narrow widths, ultrathin thickness, as well as open macroporous architectures. The convincing example verified that nanoribbons are beneficial to enhance electrochemical behavior due to well-designed structures. However, because only single material is used, the K storage capacity is not high enough, and it may need to be compounded with other materials in the subsequent research to further improve the electrochemical performances.

### 3.3 One-Dimensional Alloying Materials

Up to now, few studies about 1D alloying materials have involved in nanotubes, nanowires, nanofibers, and nanoribbons in the PIBs fields. Therefore, nanorod-like alloying materials with shorter ionic transfer path will be introduced by typical examples [57, 59, 91, 132, 142].

Nanorod-like Bi has been composites with carbon materials as anode materials for PIBs. For instance, Bi nanorods coated with mesoporous carbon were fabricated [142]. The unique structure with the core of Bi nanorods and outside carbon mesoporous shell had an effect on boosting the electronic and ionic diffusion as well as buffering the volumetric variation. Thus, the storage capacity reached 425 mA h g$^{-1}$ at 0.2 A g$^{-1}$. Compared with mesoporous carbon coatings, CNTs may encapsulate and confine Bi nanorods better due to their hollow tubular structures and good mechanical properties. Li et al. [91] synthesized Bi nanorods confined by hollow N-doped CNTs (Bi@N–CT) as anode materials of PIBs. The Bi nanorod structure was encapsulated in hollow structure with the carbon coating layer thickness of about 40 nm (Fig. 8a, b). In the electrochemical test, the capacity was 297 mA h g$^{-1}$ at 20 C (Fig. 8c). Additionally, the obtained electrode reached the capacity of 266 mA h g$^{-1}$ at 10 C after over 1000 cycles (Fig. 8d). These electrochemical performances were largely ascribed to the nanostructural design as well as the perfect coordination of Bi nanorods and CNTs. In addition, the storage mechanism could be understood well by in situ XRD, which indicated the high reversibility of structure (Fig. 8e). Coincidentally, Bi nanorods were encapsulated in N-doped CNTs (Bi@C nanorods) for PIBs anode [59]. The N-doped CNTs were originated from the carbonization of polydopamine, different from the above example using polypyrrole. The morphology of Bi nanorods, hollow structural robustness, and carbon coating layer conductive network were beneficial for the obtained anode to deliver the capacity of 179.1 mA h g$^{-1}$ at 0.5 A g$^{-1}$ after over 300 cycles. Therefore, Bi nanorods compounded with hollow structure carbon materials are advantageous to restrict volume changes and keep long cycle stability.

### 4 Two-Dimensional Nanomaterials for PIBs

2D nanomaterials have attracted increasing attentions since the discovery of graphene in 2004, and they have been extensively studied for electrochemical energy storage because of unique structural features as well as physicochemical properties [143–151]. On the one hand, the large surface area of 2D nanomaterials is conducive to ionic adsorption, which is beneficial to improve capacitance. On the other hand, 2D nanomaterials with high conductivity and tunable interlayer spacing can boost electronic transfer and benefit ionic intercalation, respectively [152]. Based on these advantages, 2D materials have become favored materials for researchers. Therefore, the corresponding 2D materials will be introduced typically. And then, the initial C.E., rate performances, and cycle properties of recent reported 2D anode materials of PIBs are summarized in Table 3.

#### 4.1 Two-Dimensional Carbon Materials

As a typical example of 2D carbon materials, graphene has been used for PIBs because of its high surface area, extraordinary mechanical strength, as well as high electrical conductivity [167–169]. In 2015, it was the first time that the K-ion intercalation of rGO film was studied by Luo and co-workers [170]. Although the reversible capacity was 222 mA h g$^{-1}$, rGO with poor rate capability was due to its inferior electronic conductivity. Therefore, graphene need to be modified and modifying its structure may be one of the effective strategies to improve its electrochemical performances. Firstly, single heteroatom
Fig. 8  a Bi@N–CT’s schematic diagram. b Bi@N–CT’s TEM image. c Rate performance of the obtained electrode at different C-rates (1 C is 385 mAh g\textsuperscript{-1}). d Long-term cycling performance of Bi@N–CT at 10 C. e Contour plot of the obtained electrode’s in situ XRD results in the course of discharge/charge process about the first two cycles with discharge/charge curves. Reproduced with permission from Ref. [91]. Copyright 2020, Royal Society of Chemistry
doping has been used to regulate the structure of graphene. Few-layered graphene with N-doping was reported as PIBs anode materials, which realized the charge capacity of over 350 mAh g\(^{-1}\) at 50 mA g\(^{-1}\) and the cycle capacity of more than 210 mAh g\(^{-1}\) at 100 mA g\(^{-1}\) after 100 cycles because the obtained electrode could provide abundant sites for ion storage and boost ionic transfer [171].

Based on an improved method, similar few-layer N-doped graphene (FLNG) was prepared as electrode materials in PIBs [65]. Its storage mechanism was made up of two parts including trapping K-ion on the surface and into the defect sites (Fig. 9a, b). The FLNG could supply more K-ion storage active sites and enhance electronic as well as ionic diffusion because of few-layer structure, N-doping impact and high surface area, so the long cycle capacity of 150 mAh g\(^{-1}\) was realized at 500 mA g\(^{-1}\) after 500 cycles. Besides, in order to regulate the structure of graphene better, co-doping with two different heteroatoms has been utilized. Ma et al. [66] fabricated graphene with P and O co-doping (PODG) as PIBs electrode materials (Fig. 9c). The ultrathin film of PODG with only a few layers was displayed in the TEM image (Fig. 9d). As for PODG, graphene and elements with P and O not only boosted K-ion diffusion by the expanding interlayer spacing, but also promoted K-ion adsorption because of large surface area as well as sufficient defects. Therefore, the PODG electrode delivered 165 mAh g\(^{-1}\) at 2000 mA g\(^{-1}\) (Fig. 9e). Moreover, PODG electrode delivered about 385, 235, and 160 mAh g\(^{-1}\) at 500, 1000, as well as 2000 mA g\(^{-1}\) after 600 cycles, respectively (Fig. 9f). Accordingly, expanding interlayer spacing and abundant active sites can be achieved by dual doping, so Luan and co-workers [172] further synthesized the multilayer graphene with N and P dual-doping (NPG) in PIBs electrode materials field. As for its electrochemical behavior, the rate capacity of 194 mAh g\(^{-1}\) was achieved at 1000 mA g\(^{-1}\). In addition, the cycle capacity of 242 mAh g\(^{-1}\) was obtained at 500 mA g\(^{-1}\) after 500 cycles. These electrochemical

| Materials                                      | Initial C.E. (%) | Rate capacity (mAh g\(^{-1}\) at the current density (mA g\(^{-1}\)) | Cycle capacity (mAh g\(^{-1}\)) at the current density (mA g\(^{-1}\)) at the current density (mA g\(^{-1}\)) | Cycle capacity (mAh g\(^{-1}\)) at the current density (mA g\(^{-1}\)) | References |
|-----------------------------------------------|------------------|-------------------------------------------------|--------------------------------------------------------------------------------|--------------------------------------------------------------------------------|------------|
| Nanocrystalline SnS\(_2\) coated onto rGO     | –                | 120 at 2000                                     | 280 at 25 (25)                                                                 |                                                                                   | [153]      |
| Ti\(_3\)CNTz                                  | 28.4             | 32 at 500                                        | 75 at 20 (100)                                                                 |                                                                                   | [60]       |
| Mo\(_5\)S\(_2\)@rGO                          | –                | 178 at 500                                       | 381 at 100 (100)                                                                |                                                                                   | [61]       |
| Mo\(_5\)S\(_2\)@SnO\(_2\)@C                  | 73               | 86 at 800                                        | 250 at 100 (20)                                                                 |                                                                                   | [62]       |
| Mo\(_5\)S/C                                   | –                | 164 at 2000                                      | 180 at 500 (240)                                                                |                                                                                   | [154]      |
| Envelope-like N-doped carbon nanosheets       | ≈20              | 168 at 2000                                      | 151 at 1000 (1000)                                                              |                                                                                   | [155]      |
| VSe\(_2\)                                     | 69.1             | 172 at 2000                                      | ≈87.3% of the capacity retention at 2000 (500)                                  |                                                                                   | [63]       |
| FSCC                                          | 62.7             | 139 at 1000                                      | 226 at 100 (100)                                                                |                                                                                   | [156]      |
| SnS\(_2\)@graphene                            | 40.5             | 290 at 2000                                      | 559 at 100 (50)                                                                 |                                                                                   | [157]      |
| CuNanoplates                                  | 50.8             | 163 at 2000                                      | 206 at 1000 (100)                                                               |                                                                                   | [158]      |
| SnS\(_2\)@rGO                                 | 51.2             | 247 at 1000                                      | 205 at 1000 (300)                                                               |                                                                                   | [64]       |
| Amorphous carbon/graphitic carbon nanoribbons  | 15.7             | 120 at 5000                                      | 192 at 1000 (5200)                                                              |                                                                                   | [159]      |
| SnS\(_2\)@C@rGO                              | 53.0             | 287.8 at 500                                     | 170.9 at 500 (500)                                                              |                                                                                   | [160]      |
| HeTiO\(_2\)C micro-tubes                      | 49.1             | 97.3 at 2000                                     | 132.8 at 500 (1200)                                                             |                                                                                   | [161]      |
| MnCO\(_3\) nanorods@rGO                      | –                | 98 at 2000                                       | 701 at 200 (500)                                                                |                                                                                   | [162]      |
| N and P co-doped vertical graphene/ carbon cloth | 53.47           | 156.1 at 2000                                   | 142.4 at 1000 (1000)                                                            |                                                                                   | [163]      |
| rGO@p-FeS\(_2\)@C composite                   | –                | 298 at 2000                                      | 322 at 1000 (30)                                                                |                                                                                   | [164]      |
| SnP\(_{0.94}\) nanoribbons/graphene oxide composite | 42             | 57 at 1000                                       | 106 at 200 (100)                                                                |                                                                                   | [165]      |
| Activated crumbled graphene                    | ≈39              | 210 at 2000                                      | 245 at 500 (2800)                                                               |                                                                                   | [166]      |
performances were mainly due to the NPG with sufficient active sites for ion storage, expanding interlayer spacing, and enhanced electrochemical conductivity. Given the aforementioned examples, graphene can improve electrochemical performances of PIBs anode by heteroatom doping (e.g., N, P, and O). However, the obstructions of graphene in practical applications are attributed to low initial coulomb efficiency, potential plateau lack, and the large voltage hysteresis [173]. Therefore, some strategies should be undertaken in the further experiments.

4.2 Two-Dimensional MCs and MOs

Among 2D MCs and MOs, 2D transition metal chalcogenides (2D TMCs), including MoS₂, MoSe₂, VS₂, VSe₂, V₅S₈, etc., have been studied in PIBs anode materials for the reason that they have sufficient active sites, short ionic transfer pathways, and low intercalation barriers [174]. Taking MoSe₂ as an example, it possesses sandwich-like lamellar structure, but low intrinsic conductivity, compounding with carbon materials, could be considered to enhance its
Fig. 10  a TEM image of MoSe$_2$/N–C.  b HRTEM image of MoSe$_2$/N–C.  c Rate capability of MoSe$_2$/N–C from 100 to 2000 mA g$^{-1}$.  d Long Cycle capacities of MoSe$_2$/N–C, MoSe$_2$, as well as C at 100 mA g$^{-1}$. Reproduced with permission from Ref. [67]. Copyright 2018, John Wiley and Sons.  e Schematic diagram of expanding process of EF-TNS.  f HRTEM image of pristine Ta$_2$NiSe$_5$.  g HRTEM image of intercalated Ta$_2$NiSe$_5$.  h Stable cycle capacities of EF-TNS anode at 500 mA g$^{-1}$. Reproduced with permission from Ref. [68]. Copyright 2019, John Wiley and Sons.
electrochemical performances. So MoSe$_2$/N-doped carbon (MoSe$_2$/N–C) composite was synthesized as anode materials in PIBs [67]. Spherical structure was composed of a great number of nanosheets, and the interplanar spacing with 0.678 nm was beneficial to K$^+$ insertion/extraction (Fig. 10a, b). In addition, the coated carbon layer encapsulating the MoSe$_2$ nanosheets effectively improved conductivity and obstructed the aggregation of the nanosheets. As for electrochemical performances, the MoSe$_2$/N-C electrode remained 178 mAh g$^{-1}$ at 2000 mA g$^{-1}$ as well as maintained 258.2 mAh g$^{-1}$ after 300 cycles at 100 mA g$^{-1}$ (Fig. 10c, d). Besides, quite a few binary 2D TMCs have also been reported to improve the electrochemical performances of PIBs anode, but a few studies about ternary 2D TMCs have been published. Gradually, ternary 2D TMCs have aroused increasing interesting in PIBs anode due to their unique structure and properties [68, 175, 176]. Then, expanded few-layered ternary Ta$_2$NiSe$_5$ (EF-TNS) flakes were reported [68]. The Ta$_2$NiSe$_5$ with expanded interlayer was realized by Mg$^{2+}$/NO$_3^-$ ion assisted intercalation (Fig. 10e). First of all, ternary Ta$_2$NiSe$_5$ was beneficial to increase the capacity. Secondly, the obtained EF-TNS had a few-layered structure, which was in favor of buffering the volume variation. Thirdly, the interlayer distance of EF-TNS was expanded from 0.6 to 1.1 nm after intercalation, which facilitated K-ion diffusion and benefited for K$^+$ ions intercalating along zigzag pathways (Fig. 10f, g). These three advantages made it have good K storage performances and become promising anode materials. As a consequence, the EF-TNS electrode delivered the capacity of 116 mAh g$^{-1}$ at 500 mA g$^{-1}$ after 1100 cycles (Fig. 10h).

In order to further search for appropriate anode materials, 2D metal-based oxides have been studied in PIBs [177]. Sb$_2$O$_3$ flakes anchored onto rGO were synthesized by Li and co-workers [178], which delivered the long cycle capacity of 201 mAh g$^{-1}$ at 500 mA g$^{-1}$ after 3300 cycles. As for the reason of the remarkable electrochemical behavior, it was found that the Sb$_2$O$_3$ flakes and rGO could not only make electrode and electrolyte close, but also enhance conductivity and buffer the volume expansion. In addition, further strategies should be taken so as to relieve the stress variation of Sb$_2$O$_3$ better, such as designing rational structures and synthesizing bimetallic compounds; especially, bimetallic compounds with the better electrochemical behavior are ascribed to the synergistic effects, more active sites, and interfacial effects, compared with the single-metal counterparts [179–182]. Accordingly, Wang et al. [183] fabricated Sb$_2$MoO$_6$ nanoplates compositied with rGO as anode materials of PIBs. In these composites, with the help of Mo element, the conductivity could be improved and the volume changes of Sb could be relieved in the course of charge and discharge process. Therefore, the electrochemical performances would be enhanced and the capacity of 247 mAh g$^{-1}$ was delivered at 500 mA g$^{-1}$ after 100 cycles.

4.3 Two-Dimensional Alloying Materials

Phosphorene can be regarded as a typical 2D alloying materials in PIBs. In 2014, monolayer and few-layer phosphorene were exfoliated from black phosphorus by scotch-tape-based micro-cleavage method [184, 185]. However, monolayer phosphorene could not be utilized directly as electrode materials, which was ascribed to be easily oxidized when it exposed to air [186, 187]. Therefore, He et al. [188] indirectly verified that the monolayer phosphorene was beneficial to improve performances of GeSe electrode by first-principles calculation. Moreover, phosphorene with sulfur doping could be considered as the PIBs anode materials by using ab initio density functional theory [189]. Furthermore, few-layer phosphorene has started to be directly utilized for PIBs because of high carrier mobility and superior mechanical flexibility [190]. And then, Nikhil Koratkar’s group [69] fabricated few-layer phosphorene composited with rGO (FLP/rGO) as anode materials (Fig. 11a). The TEM image of FLP/rGO indicated FLP encapsulated with rGO and the existing evidence of both FLP and rGO was provided by the SAED pattern (Fig. 11b). In these composites, FLP and rGO were beneficial for enhancing electrical conductivity as well as buffering volumetric variation in the course of alloying reaction. The FLP/rGO (1:3) electrode delivered different capacities at different current density, especially ~ 400 and ~ 230 mAh g$^{-1}$ at ~ 0.6 C as well as ~ 1.2 C, respectively. As for cycle performance, FLP/rGO (1:3) delivered the capacity of ~ 230 mAh g$^{-1}$ at ~ 0.5 C after 300 cycles (Fig. 11c, d). Furthermore, in terms of alloying mechanism,
the author confirmed that $\text{K}_4\text{P}_3$, bringing about high capacity was the type of alloy and used DFT calculation to give a further understanding about the formation of $\text{K}_4\text{P}_3$ (Fig. 11e). Further experiments can focus on phosphorene compounded with other materials or doped with heteroatom to enhance the electrochemical performances.

**Fig. 11** a Schematic diagram and b TEM image (inset shows SAED pattern) of FLP/rGO. c Rate capability of FLP/rGO (1:3) at various C-rates between ~0.1 and ~1.2 C. d Cycling performance of FLP/rGO (1:3) at ~0.5 C. e Various alloys’ formation energies calculated by DFT. Reproduced with permission from Ref. [69]. Copyright 2019, American Chemical Society
Three-Dimensional Nanomaterials for PIBs

3D nanomaterials are essential to the application of batteries, due to unique features such as high specific areas, interlinked porous channels, high conductivity and outstanding structural mechanical stability [191, 192]. 3D nanomaterials could overcome the insufficiency of 1D and 2D nanomaterials caused by evident aggregation [193]. In addition, some 3D nanomaterials can be directly used as free-standing electrodes, simplifying the preparation process of battery. To date, 3D nanomaterials in PIBs electrode materials field mainly include 3D carbon nanomaterials,

Table 4 Comparison of the state-of-the-art performances of 3D anode materials in PIBs

| Materials                                      | Initial C.E. (%) | Rate capacity (mAh g⁻¹) at the current density (mA g⁻¹) | Cycle capacity (mAh g⁻¹) at the current density (mA g⁻¹) (cycle number) | References |
|-----------------------------------------------|------------------|----------------------------------------------------------|------------------------------------------------------------------------|------------|
| N- and O-rich CNF                              | 35               | 70 at 10 C                                               | 160 at 1 C (300)                                                        | [198]      |
| Cobalt(II) terephthalate-based layered MOF     | 60.65            | 131 at 1000                                             | 188 at 1000 (600)                                                      | [199]      |
| Tire-derived carbons                           | 37.1             | 60 at 2 C                                               | 155 at C/2 (200)                                                       | [200]      |
| Hard wood-based hard carbon                    | 56               | 135 at 100                                              | –                                                                      | [201]      |
| AC                                            | –                | 30 at 1000                                              | 100.3 at 200 (100)                                                     | [74]       |
| Porous CNF paper                              | 24.1             | 140 at 5000                                             | 211 at 200 (1200)                                                      | [202]      |
| S/O co-doped porous hard carbon microspheres  | 61.7             | 158 at 1000                                             | 108.4 at 1000 (2000 cycles)                                           | [203]      |
| Co₃[Co(CN)₆]₂                                   | 45.5             | 112 at 2000                                             | 297.5 at 100 (200)                                                    | [75]       |
| Skimmed cotton-derived hard carbon             | 73               | 165.2 at 4000                                           | 240 at 200 (150)                                                       | [204]      |
| HPCS                                          | 45.3             | 150 at 500                                              | 276.4 at 500 (100)                                                    | [76]       |
| SnO₂-graphene-CNFS                             | 44.13            | 114.81 at 1000                                          | 202.06 at 1000 (100)                                                  | [205]      |
| MoS₂/N-Doped C                                | –                | 131 at 2000                                             | 151 at 5000 (1000)                                                    | [206]      |
| Hierarchically N-doped porous carbon           | 43.1             | 185 at 10,000                                           | 144.4 at 5000 (1000)                                                  | [89]       |
| HNTO/CS                                       | –                | 50 at 1000                                              | 88.9 at 1000 (1555)                                                   | [93]       |
| High pyridine NPC                             | –                | 186.2 at 2000                                           | 231.6 at 5000 (2000)                                                  | [207]      |
| Zero-strain potassium fluoromanganate hollow nanocubes | 65 | 78 at 1000 | 110 at 4000 (10,000) | [208] |
| N-doped hierarchically porous carbon           | 30.28            | 193.1 at 500                                            | 121.3 at 5000 (1000)                                                  | [209]      |
| KTi₂(PO₄)₆C nanocomposites                    | 35.3             | 131.1 at 1000                                           | 69.7 at 1000 (1000)                                                   | [210]      |
| NOHPC                                         | 25               | 118 at 3000                                             | 230.6 at 500 (100)                                                    | [77]       |
| SnP₃ in N-doped carbon fibers                  | 64.17            | 169.6 at 2000                                           | 160.7 at 5000 after 1000                                              | [211]      |
| Bi@3DGFs                                      | 51.1             | 113 at 10,000                                           | 164 at 1000 (400)                                                     | [212]      |
| N-doped biomorphic carbon                      | 55.1             | 102.6 at 2000                                           | 119.9 at 1000 (1000)                                                  | [213]      |
| Nanosheets-assembled CuSe Crystal Pillar      | 92.4             | 280 at 5000                                             | 337 at 100 (40)                                                       | [214]      |
| rGO aerogel                                    | 44               | 92 at 6.7 C                                             | 125 at 1.6 C (500)                                                    | [78]       |
| KVPO₄F                                        | –                | 65 at 2000                                              | 133 at 100 (100)                                                      | [215]      |
| MXene@Sb                                      | 57.29            | 270.81 at 500                                           | capacity retention of 79.1435% at 500 (500 cycles)                    | [216]      |
| Red P@N-PHCNFs                                 | –                | 342 at 5000                                             | 282 at 5000 (800 cycles)                                               | [217]      |
| NOHPC                                         | –                | 110 at 1000                                             | 80 at 2000 (3000)                                                     | [218]      |
| Yolk shell FeP@C nanoboxes                    | 47               | 37 at 2000                                              | 205 at 1000 (300)                                                     | [219]      |
| Multicore shell Bi@N-doped carbon nanospheres | 43               | 152 at 100,000                                          | 203 at 10,000 (1000)                                                  | [220]      |
| Graphitic nanocarbons                          | –                | 56.6 at 5000                                            | 189 at 200 (200)                                                      | [79]       |
| HHC                                           | 46.88            | 42 at 3200                                              | 67.6 at 500 (100)                                                     | [80]       |
| N-doped soft carbon frameworks built of well-interconnected nanocapsules | 30.9 | 151 at 5000 | 165 at 1000 (500) | [221] |
3D MCs and MOs, and 3D alloying materials [194–197]. At the following section, the relationship between these materials and their electrochemical performances will be discussed systematically. Meanwhile, the initial C.E., rate performances, and cycle properties of recent reported 3D anode materials of PIBs are summarized in Table 4.

5.1 Three-Dimensional Carbon Materials

3D carbon materials mainly include 3D interconnected structures, which can promote electronic and ionic transport as well as improve mechanical stability [222–224]. Among 3D interconnected networks, 3D carbon nanofiber frameworks are typical materials and were used in PIBs anode materials field in 2018. Li et al. [225] reported bacterial-derived and compressible carbon nanofiber foam (CNFF) with hierarchical pores as electrode materials in PIBs (Fig. 12a). The diameters of fibers were between 10 and 30 nm shown in the SEM image (Fig. 12b). According to the authors’ report, two kinds of pores were found in this CNFF. The first was numerous nanopores originating from nanofibers’ surface; the other was hierarchical pores between fiber and fiber. Those
pores could benefit K-ions absorption in the course of charge and discharge process (Fig. 12c, d). In the electrochemical test, the CNFF electrode achieved a reversible capacity of 158 mAh g$^{-1}$ at 1000 mA g$^{-1}$ after 2000 cycles. Furthermore, after a period of time, the same battery could also run 1500 and 1000 cycles at 2000 and 5000 mA g$^{-1}$, respectively (Fig. 12e), showing superior cycle stability. The splendid electrochemical behavior of the CNFF anode was ascribed to the hierarchical pores of the structure, the 3D carbon foam, and the quasi-amorphous carbon, which not only enhanced ionic adsorption and diffusion but also relieved the volume variation. Additionally, 3D porous structures have also been used in PIBs. Bin et al. [226] fabricated interconnected carbon architecture with hollow structure and neuron-like morphology (HINCA) as anode materials. As for HINCA, interconnected tetrapod backbones with tubular structure and hollow structural spherical joint were exhibited and better comprehend by cartoon (Fig. 12f-i). The aforementioned structural characteristics could facilitate electronic conductivity and possess flexible mechanical robustness. Then, the HINCA-type electrode delivered the capacity of 340 mAh g$^{-1}$ as well as showed only slight capacity decay of ~0.05% per cycle over 500 cycles at 1 C (Fig. 12j, k). The HINCA-type electrode’s electrochemical performances were due to carbon structure with hollow feature, which kept the stability of structure during potassiation/depotassiation process and promoted the ions and electrons transport. With similar structures, hollow neuronal carbon skeleton (HNCS) was fabricated, which exhibited interlinked hollow architecture with high content of pyridinic N [227]. The interlinked hollow framework and pyridinic N could enhance ionic transfer and adsorption as well as tolerate the stress variation to improve corresponding performances. As for electrochemical performances, the electrode delivered the rate capacity of 110 mAh g$^{-1}$ at 1000 mA g$^{-1}$ and 134 mAh g$^{-1}$ at 0.5 A g$^{-1}$ after 500 cycles. Overall, 3D carbon materials are beneficial to boost ionic and electronic transport and keep high mechanical robustness, resulting in improved electrochemical performances.

5.2 Three-Dimensional MCs and MOs

3D MCs and MOs have aroused increasing attentions in PIBs electrode materials. For example, in 2017, it was the first time that Ren et al. [70] used MoS$_2$ particles as electrode materials in PIBs. The author indicated that the micro-sized MoS$_2$ with layer structure was beneficial to stably form K-ion intercalation compound (K$_{0.4}$MoS$_2$) during potassiation to 0.5 V. It is demonstrated that MoS$_2$ may be considered as another choice of PIBs anode materials. Besides, increasing 3D metal-based oxides have been tried for PIBs anode. In 2018, Li and co-workers [228] firstly introduced orthorhombic niobium pentoxide (T-Nb$_2$O$_5$) into PIBs anode field. The urchin-like interlinked hierarchical structure of T-Nb$_2$O$_5$ assembled by nanowires was exhibited (Fig. 13a), which could boost K-ion diffusion. Therefore, the capacity of 104 mAh g$^{-1}$ was delivered at 0.4 A g$^{-1}$ in terms of rate performance (Fig. 13b, c). In addition, interconnected K$_2$Ti$_6$O$_13$ nanowires framework was fabricated as electrode materials in PIBs [71]. The K$_2$Ti$_6$O$_13$ nanowires with a diameter of around 5.5 nm (name as TBTN) were displayed (Fig. 13d). In this framework, ionic diffusion and mechanical robustness could be enhanced due to 3D interlinked architectures. Thus, the TBTN electrode delivered the rate capacities of 11 mAh g$^{-1}$ at 10 C (Fig. 13e). Besides, the cycle capacity of around 120 mAh g$^{-1}$ was exhibited at 0.2 C after 20th cycle (Fig. 13f). From the above analysis, it can be concluded that good structural design helps to improve the performances of 3D MCs and MOs, but these properties are not satisfactory enough. Judging from the characteristics of these materials, the reason may put down to their poor conductivity. Therefore, it is necessary for 3D MCs and MOs to combine other modification methods to obtain satisfactory electrochemical performances.

5.3 Three-Dimensional Alloying Materials

3D alloying materials are beneficial to contact largely with electrolyte and buffer the volume expansion, especially 3D porous structures. Pristine Bi block was used as PIBs anode and would gradually transform into 3D porous networks after 100 cycles during charge and discharge process (Fig. 14a, b) [229]. The formed porous networks not only enhanced ionic transfer but also restricted the stress variation in the course of potassium/depotassium process. Therefore, the capacity retention realized 86.9% after 300 cycles at 2 C (Fig. 14c). This method provides new ideas for constructing porous alloying materials. So nanoporous Sb (NP-Sb) was synthesized as PIBs anode materials (Fig. 14d) [72]. The pores of NP-Sb-20 (20 means the percentage of Sb atom) were uniformly distributed in the continuous porous
structure (Fig. 14e). Benefiting from the 3D porous architecture, the NP-Sb-20 could effectively tolerate stress variation and promote ions transportation. As for rate performance, the bulk Sb only delivered 30 mAh g⁻¹ at 500 mA g⁻¹, while the NP-Sb-20 delivered 265 mAh g⁻¹ (Fig. 14f). Obviously, the performances improvement was due to the nanoporous structure of Sb. Besides, in 2019, nanoporous Ge was fabricated for the first time, which delivered the capacity of around 120 mAh g⁻¹ at 20 mA g⁻¹ over 400 cycles [73]. Based on nanoporous Ge with the numerous pores as well as nanoscale ligaments, the stable cycling performance was ascribed to shorter diffusion distance of K-ion as well as the adequate space for volumetric variation. Overall, 3D porous structures are of benefit to alloying materials and further efforts should be made to improve the electrochemical behavior.

Additionally, given the discussion about multi-dimensional structures, related nanomaterials possess attractive advantages and unfavorable disadvantages, so optimized strategies should be taken to improve their electrochemical performances. Accordingly, the corresponding advantages, disadvantages, and optimized strategies are outlined in Table 5.

6 Summary and Outlook

As aforementioned, PIBs have been investigated due to better advantages compared with LIBs and SIBs, such as abundant resources, lower price, and smaller stokes radius of solvated ions. However, there are still some obstacles to PIBs for commercial applications, such as large volumetric variation induced by large size of K-ion during charge and discharge process, weak ionic diffusivity in solid phase, inferior kinetics of K⁺ reaction, the existence of side and irreversible reactions, the imperfect energy store mechanism compared with LIBs, the production of dendrite and related safety problems. Accordingly, nanostructural design has been considered as one of the effective strategies to enhance corresponding performances.
In summary, 0D–3D nanomaterials about PIBs anode materials have been summarized and involved in the relationship with the corresponding electrochemical performances, mainly concerning carbon materials, MCs and MOs, and alloying materials. 0D nanomaterials have been utilized in PIBs anode materials due to nano-size and large surface area, which can boost ionic transportation and alleviate the stress changes. In addition, 1D nanomaterials with high length-to-diameter aspect ratios possess high mechanical robustness as well as shorter electronic and ionic transport path. Moreover, 2D nanomaterials have also possessed large surface area to enhance ionic adsorption and diffusion in PIBs anode materials field. Finally, the interconnected structure about 3D nanomaterials can make largely electrode and electrolyte contact, which can facilitate ionic transfer.

Given aforementioned information, 0D–3D nanomaterials possess different structural and morphological features, corresponding to different electrochemical performances. Therefore, in order to better understand the different effect of multi-dimensional nanomaterials, it is necessary to make a comparison about their electrochemical performances. Furthermore, the corresponding comparision would be discussed by taking 0D–3D carbon materials as examples. And then, typical carbon nanomaterials will be introduced, including carbon nanocage, CNTs, graphene, and graphite. Among them, 0D carbon nanocage possesses large surface area and unique cage-like structure, which makes it have well electrochemical performances (e.g., 195 mAh g⁻¹ at 0.2 C after 100 cycles for CNC) [42]. Additionally, as representative 1D carbon nanomaterial, compared with 0D nanocage, CNTs with high aspect ratio are conducive to enhance the mechanical strength of electrode materials, so the structure of related electrode can be maintained stable and durable during charging and discharging, improving the long cycling life (e.g., 244 mAh g⁻¹ at 0.5 A g⁻¹ after 200 cycles for P@TBMC-2.4) [50]. As for graphene, with large landing platform, they are beneficial for adsorbing K-ion and are different from 0D as well as 1D carbon materials, which can evidently improve the capacity by adsorption mechanism (e.g., 385 mAh g⁻¹ at 500 mA g⁻¹ after 600 cycles for graphene) [213].

![Fig. 14](image-url)  
**Fig. 14** SEM image of a pristine Bi and b Bi after 100th cycle. c Cycle performance of Bi electrode (1 C = 384.7 mAh g⁻¹). Reproduced with permission from Ref. [229]. Copyright 2018, John Wiley and Sons. d Schematic illustration of NP-Sb. e TEM image of NP-Sb-20. d Rate performance of two samples in the range of 50–500 mA g⁻¹. Reproduced with permission from Ref. [72]. Copyright 2018, American Chemical Society
cycles for PODG) [66]. Moreover, graphite can be used to inhibit the aggregation as a comparison of 0D–2D nanomaterials, which can keep structure stable and retain the stable capacity (e.g., 174 mAh g\(^{-1}\) at 200 mA g\(^{-1}\) after 500 cycles for expanded graphite) [109]. According to the comparison and discussion of 0D–3D carbon materials, the difference of electrochemical performances of multi-dimensional nanomaterials can be well understood. Additionally, the synthetic methods are important to prepare nanostructures with excellent electrochemical performances. Therefore, the main preparation methods about their advantages and disadvantages are summarized in Table 6. From the results of comparison, Ball milling may be considered as the practical strategies to obtain nanomaterials in the industries, which is attributed to its facile operability, low cost, and large scale.

Although significant progress has been achieved due to nanostructure design with different dimensions, more efforts should be made for PIBs anode materials to improve electrochemical performances as followed:

1. Novel architecture should be proposed. For example, carbon dots have been directly explored for LIBs and SIBs, while carbon dots only as raw materials were used to fabricate PIBs anode materials. Thus, further experiments may focus on novel structure design.
2. Novel synthetic strategies about different dimensional structures like 3D printing method can be used in PIBs field. Therefore, it is necessary to create methods with simple process and low cost.
3. Different dimensional structures can be assembled to fabricate various materials with unique morphology to sufficiently achieve the properties of every component.
4. Using new materials to design different dimensional structures is beneficial for increasing the variety of anodes as well as trying to improve the electrochemical performances.
5. The relationship between different dimensional structural electrodes and various electrolytes should be studied. To date, a few studies have involved in the impact of electrolytes on corresponding performances.
6. Some energy store mechanisms of PIBs still keep unclear, so more efforts should be undertaken.

All in all, it is one of the key preconditions for commercial PIBs anode materials to utilize simple preparation methods and simplified processes in different dimensional materials fabrication. Meanwhile, high-performance, low-cost, and good-stability materials will be considered as the desire

| Materials | Advantages | Disadvantages | Optimized strategies |
|-----------|------------|---------------|---------------------|
| 0D nanomaterials | Sufficient sites for ionic adsorption; buffering large volume expansion; shorter ionic diffusion pathway | Easy aggregation; low tap density | Combining with carbon materials; compounding with micro-sized materials |
| 1D nanomaterials | High mechanical robustness; facilitating electronic and ionic transfer | Nonadjustable specific surface area and porosity properties | Coating; second phases used to decoration; forming the interconnected network |
| 2D nanomaterials | Large surface area; enhanced open-edge morphology; high conductivity; tunable interlayer spacing; relieving the stress changes | Side reactions; easy agglomeration | Surface coating; compounding with other materials; doping with other heteroatoms |
| 3D nanomaterials | Inhibition of agglomeration; free-standing; high mechanical stability; tolerating the stress variation; facile ionic access | Numerous macropores affecting the capacitive-controlled contribution; low initial Coulombic efficiency | Combining with low-dimensional materials; surface engineering |
choice in the practical application. Therefore, this review is devoted to provide new insights for further research.

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Table 6 Summary of advantages and disadvantage of various synthesis methods for multi-dimensional structures

| Synthesis methods                  | Advantages                                      | Disadvantages                                      | References                                                                 |
|-----------------------------------|-------------------------------------------------|---------------------------------------------------|----------------------------------------------------------------------------|
| Ball milling method               | Facile; highly effective; economic; scalable    | Products with uneven and large size; generation of noise | [33, 40, 45, 48, 250]                                                      |
| Hydrothermal/solvothermal method  | Facile morphological control; controllable size; high purity | High cost; simple morphology; difficult controlled processes; existing safety problems | [38, 46, 61, 62, 99, 106, 157, 199]                                         |
| Solution method                   | Well-controlled size; uniform dispersion        | Related toxic reagent                              | [47, 63, 95, 165, 214, 251, 252]                                            |
| Vaporization-condensation method  | Uniform and small size; controllable nanostructure | Difficult controlled distribution; low production efficiency | [50, 122, 217, 253]                                                        |
| Calcination(carbothermal reduction)| Well operability; generation of new phases      | Large energy consumption; generation of toxic gases | [44, 53–55, 57, 80, 81, 89, 96–98, 100–102, 104, 105, 128, 129, 131, 132, 135, 198, 202] |
| Electrodeposition method          | Uniform size; controllable                      | Related toxic reagent; large energy consumption    | [49]                                                                       |
| Solid–liquid reaction method      | Simple; facile operability                      | Related dangerous solution and toxic reagent       | [52, 60, 103, 254–256]                                                      |
| Chemical vapor deposition method  | Precise controlled products; high purity        | Required relatively high deposition temperatures; involving some dangerous precursors; large energy consumption | [127, 257]                                                                 |
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