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# Research progress in transition metal chalcogenide based anodes for K-ion hybrid capacitor applications: a mini-review

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Metal ion capacitors have gained a lot of interest as a new kind of capacitor-battery hybrid energy storage system because of their high power density while maintaining energy density and a long lifetime. Potassium ion hybrid capacitors (PIHCs) have been suggested as possible alternatives to lithium-ion/sodium-ion capacitors because of the plentiful potassium supplies, and their lower standard electrode potential and low cost. However, due to the large radius of the potassium ion, PIHCs also face unsatisfactory reaction kinetics, low energy density, and short lifespan. Recently, transition metal chalcogenide (TMC)-based materials with distinctive structures and fascinating characteristics have been considered an emerging candidate for PIHCs, owing to their unique physical and chemical properties. This mini-review mainly focuses on the recent research progress on TMC-based materials for the PIHC applications summarized. Finally, the existing challenges and perspectives are given to improve further and construct advanced TMC-based electrode materials.

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## 1. Introduction

Lithium-ion batteries (LIBs) and supercapacitors (SCs) are currently successfully meeting the needs of commercial applications such as handheld devices and electric cars. They are outperforming other energy storage systems due to their ability to provide high energy and high capacity, respectively. The key



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drawbacks of LIBs are their low power density and poor cycling efficiency (within 4000 cycles).<sup>1</sup> An SC is a type of capacitor that can store a significant amount of energy per unit mass or volume, usually 10 to 100 times more than electrolytic capacitors. It is favoured over batteries because of its quicker and easier charging as well as faster charge delivery. SCs, on the other hand, have advantages in terms of power density, cycling lifespan (typically more than 100 000 cycles), and protection, but their energy density (only 1/10 that of LIBs) restricts them.<sup>2,3</sup> SCs cross the difference between batteries and traditional dielectric capacitors (cell voltage, specific power, and operating cost); the latter are considered to be ideal for rapid storage/release power systems,<sup>4</sup> with power distribution and absorption of 196 kW kg<sup>-1</sup> (10–100 times the energy density of electrolytic capacitors) in just a few seconds.<sup>5–7</sup> It is typically advantageous to provide a high energy density, and long cycle life for practical applications in automotive power storage systems and grid energy storage systems.<sup>8–19</sup>

A hybrid-ion capacitor (HIC) is a rapidly evolving technology that incorporates batteries' and SCs' best features to produce massive energy densities at high rates over a long period. According to previous studies, these HICs can supply energy between 60 and 200 W h kg<sup>-1</sup> (considering the mass of active materials), which is superior to a traditional SC, and their primary strength ranges between 200 and 20 000 W kg<sup>-1</sup>, which is considerably higher than that of batteries.<sup>20,21</sup> In comparison to Li (0.0017%), sodium (Na, 2.6%) and potassium (K, 2.1%) are plentiful in the earth's crust, rendering them promising substitutes to encourage batteries.<sup>22,23</sup> Furthermore, both K and Na belong to the same group after Li in the periodic table, displaying similar physicochemical properties. Consequently, research into Na<sup>+</sup>/K<sup>+</sup> storage technology is gaining momentum, paving the way to commercialize successful renewable energy storage systems.<sup>24</sup> K<sup>+</sup> storage devices are highlighted because they have a higher operating voltage and more excellent ionic conductivity in electrolytes than Na-ion storage devices. The K/K<sup>+</sup> redox pair, for example, has a potential of 2.93 V (vs. standard hydrogen electrode (SHE)), which is lower than that of Na/Na<sup>+</sup> (2.71) and quite similar to that of Li/Li<sup>+</sup> (3.3 V vs. SHE), maintaining the functional feasibility of high energy density and high operating potential of K<sup>+</sup> centered energy storage systems.<sup>21,25,26</sup> Furthermore, because of the weaker Lewis acidity, K<sup>+</sup> allows rapid transportation in the propylene carbonate electrolyte, resulting in a smaller Stokes radius (3.6). Simultaneously, Li<sup>+</sup> and Na<sup>+</sup> have Stokes radius values of 4.8 and 4.6, respectively.<sup>27–29</sup> With all these fascinating benefits, designing high-energy electrode materials (both anode and cathode) that guarantee overall energy values and good structural stability for long cycling performance is a significant challenge.

The comparatively plentiful reserves and low cost of potassium, potassium-ion hybrid capacitor (PIHCs) have recently emerged as a potential next-generation energy storage solution among the candidates.<sup>30,31</sup> The developmental history of PIHC is illustrated in Scheme 1a and b. Since 2015, there has been an increase in potassium ion batteries' research papers (Scheme 1c). Since potassium does not shape Al–K intermetallic

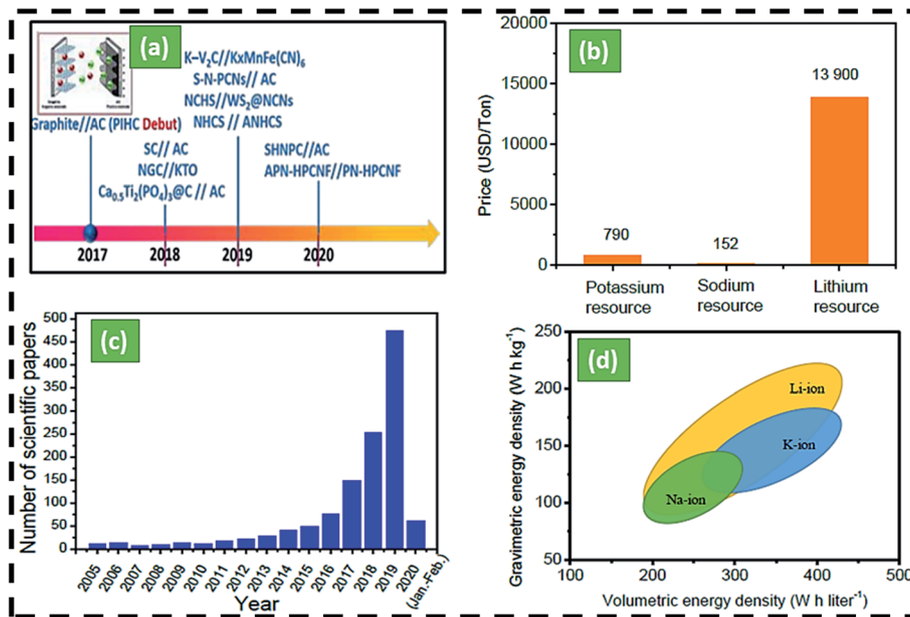
compounds thermodynamically, far less costly Al foil may be used as an anode current collector for KIBs. K has a lower standard electrode potential (–2.936 V vs. K<sup>+</sup>/K) than Na (–2.714 V vs. Na<sup>+</sup>/Na), which may contribute to a larger voltage window and better energy capacity than sodium-ion batteries (Scheme 1d). Due to the lower Lewis acidity of K<sup>+</sup>, it can shape smaller solvated ions (3.6) than Li<sup>+</sup> (4.8) and Na<sup>+</sup> (4.6). KIBs have a strong conductivity and a fast ion diffusion rate in the propylene carbonate (PC) solvent. Furthermore, unlike sodium-ion batteries, graphite can be used as an anode material for KIBs by creating the intercalation compound KC8, a theoretical power of 279 mA h g<sup>-1</sup>.<sup>31</sup> Another significant benefit of K ion batteries over sodium-ion batteries is that the K<sup>+</sup> intercalation capacity is 0.2 V vs. K<sup>+</sup>/K for other negative materials, resulting in fewer potassium metal plating and essentially preventing dendrite forming while charging in KIBs.<sup>24,28</sup> For hard carbon, the sodiation power in NIB is 0.05 V vs. Na<sup>+</sup>/Na, while in KIB, it is 0.2 V vs. K<sup>+</sup>/K. Since potassium ions have a high ionic radius (1.38), significant volume expansion of electrode materials, predominantly negative electrodes, during the charge and discharge phase results in a limited lifetime for most KIBs (usually 500 cycles) and a low power density. The technology for storing potassium ions as energy is also in its infancy.<sup>32–35</sup>

## 2. Transition metal chalcogenides nanomaterials for PIHCs

Massive volume shifts and pulverization occur in alloying metals, resulting in unexceptionally fast power decay within 200 cycles. The use of sylvite compounds in metalorganic composites reveals additional challenges to overcome, such as time-consuming planning, restricted performance, volatility, complicated K<sup>+</sup> ion insertion/extraction reactions, and poor reliability. Due to extreme side reactions between radicals and K<sup>+</sup> ions, insufficient active sites, and low electric/ionic conductivity, organic polymers typically have low initial coulombic efficiencies (CEs), restricted reversible capacities, and weak rate capabilities.

In contrast to oxides and phosphides, transition metal chalcogenides (TMCs) have gained a lot of attention from researchers in recent years because of their broad theoretical specific power, limited volume shifts, and higher electrical conductivity, high surface area, and multiple oxidation states.<sup>38,39</sup> These properties enable them to store significant energy *via* an electrical double layer. The synergistic effects between TMDs and other materials in hybrid electrode designs and hybrid configurations (*e.g.*, PIHCs) meet the demand for high energy density requirements of modern electronic devices. In recent years, TMCs have been used as advanced hybrid electrochemical energy storage systems to solve the problems of power density and cycle life of LIBs, and low energy density of SCs,<sup>40,41</sup> and the schematic is shown in Fig. 2. These TMCs consist of a battery-type anode (electrochemical intercalation or conversion) and a capacitor-type cathode (physical adsorption) in an electrolyte containing metal ions. They include many of the features of LIBs and SCs.<sup>29</sup> TMC-based materials have





Scheme 1 (a) The history and development of PIHCs, (b) comparison of the price of Na/Li/K,<sup>1,9,22</sup> (c) number of publications according to the web of science with the topic of 'potassium-ion batteries' from 2005 to 2020, (d) volumetric and gravimetric density of Na/Li/K-ion batteries.<sup>36,37</sup>

a variety of special charge storage properties in PIHCs.<sup>42,43</sup> The ion diffusion duration is limited since TMC-based materials have a high particular surface area, resulting in multiple exposed active areas. Second, TMC-based materials have quick ion transport through 2D channels that can be easily expanded and contracted. Third, materials based on TMC have excellent electrical conductivity, increasing electrode dynamics and cycle stability. Fourth, intercalation or surface redox pseudocapacitance can be achieved using TMC-based materials. Finally, structures dependent on TMC have exceptional stability and mechanical ability. These remarkable properties suggest that metal chalcogenides-base materials can be used as cutting-edge active electrode materials in PIHCs.<sup>44–46</sup>

Due to the integration of two energy storage mechanisms in one device, TMCs show several preponderances, including (1) higher battery capacity and energy density than SCs, (2) higher power density than LIBs, (3) broad operating temperature range from  $-25$  to  $80$  °C, and (4) self-discharge properties lower than SCs.<sup>47,48</sup> Most metal chalcogenides have a two-dimensional (2D) layered structure and low reaction energy barriers for the handling of alkali metal ions. Because of the simplicity of topological expansion, several techniques may be used to build various morphologies of metal chalcogenides of the same crystal structure, extending the material's future applications. The TMCs, in particular, aren't only for K-ion batteries; they're "shining stars," with applications in K dual-ion batteries, KO<sub>2</sub> batteries, PIHCs, and other energy storage and conversion areas, as seen in Fig. 1b.<sup>49–52</sup> This study progress study addresses essential technological advancements and dominated challenges confronting existing TMC-based PIHCs. Meanwhile, some previous TMCs-based nanostructures and nano-composites that have been specially engineered or prepared are used to illustrate the critical alteration strategies. In the final

section of the review paper, the recent progress in TMC-based materials is highlighted.

In this mini-review, we summarize the most recent endeavors towards developing high-performance PIHCs using TMD-based electrodes. Finally, we share our insights into the present challenges and possible future research directions, hoping to provide some inspirations for the design and fabrication of high-performance metal chalcogenide-based anode materials for PIHCs.

### 3. Potassium-ion hybrid capacitors (PIHCs)

The first step toward high-performance K<sup>+</sup>-based energy storage technologies is to use host materials with superior potassium storage capacity and structural integrity to withstand repeated K<sup>+</sup> ion injection and extraction processes.<sup>53,54</sup> The goal mentioned above indicates that designing effective anode materials with the desirable features of high specific capacities, excellent cycling stabilities, and rapid diffusion/reaction kinetics is urgently needed to advance current PIHCs toward practical applications for PIHCs, the most representative K<sup>+</sup>-based rechargeable energy storage device.<sup>55–57</sup> PIHCs have been regarded as possible alternatives to lithium-ion/sodium-ion capacitors because of their plentiful potassium supplies, lower standard electrode potential, and low expense.<sup>29,42,58–60</sup> Fig. 1a illustrates how PIHCs correspond to other energy storage systems in energy density and power density diagrams.<sup>60</sup> Research on potassium ion hybrid capacitors has seldom been recorded because of the severe kinetics mismatch between the negative and positive electrodes. The study goes into the basics of how a potassium ion capacitor works and new advancements



in electrode formulations for dual-carbon and non-dual-carbon PIHCs. The electrolyte chemistry, binders, and electrode/electrolyte interface of PIHCs were also discussed. Finally, future problems and prospects for potassium ion capacitor production are discussed.

## 4. Energy storage mechanism in PIHCs

The developmental history of PIHCs is schematically presented in Fig. 1b. In 2017, Le Comte *et al.*<sup>42</sup> published a groundbreaking paper on a potassium ion hybrid capacitor made up of a graphite battery-style anode and an AC electric double-layer capacitor (EDLC) type positive electrode with an Al foil current collector for both electrodes in a 0.8 mol L<sup>-1</sup> KPF<sub>6</sub>/acetonitrile electrolyte. At the pouch cell size, superior electrochemical efficiency can be obtained, with cycling performance of up to 55 000 cycles at a high current density of 100C/100D (corresponding to a discharge period of 30 s). With a discharge rate of up to 300C in rigid 18 650 cells, corresponding to a discharge period of 12 s and an applied current of 21 A, the hybrid ion capacitor also has outstanding rate efficiency. The results reveal that the system has twice the energy density of a standard SC. The specific energy of 12 W h kg<sup>-1</sup> can be obtained under optimal conditions by combining all of the device's components, including all electrodes collector, binder, conductive carbon, electrolyte, and separator. Since the prototypes and their excellent electrochemical efficiency, PIHCs have been studied by a few classes. The distinction between dual-carbon and nondual-carbon PIHCs is when the anode is constructed of carbon content.

An ion insertion style negative electrode, a positive capacitor electrode with current collectors and an electrolyte (e.g., 0.8 m KPF<sub>6</sub>/EC : DEC) make up a potassium ion capacitor. Using the KPF<sub>6</sub> electrolyte as an example, during charging, K<sup>+</sup> from the electrolyte is injected into the negative electrode content. Simultaneously, the electrolyte's anion (PF<sub>6</sub><sup>-</sup>) is adsorbed on the positive electrode's back. A double layer of electricity is created. K<sup>+</sup> is liberated from the negative electrode content and returned to the electrolyte during discharge. Anions are emitted from the positive electrode surface when the electric double layer developed between the active carbon cathode and

electrolyte interface dissociates. Electrons transfer from the negative electrode to the positive electrode through the external circuit (Fig. 1b).<sup>42,61</sup> During the charge/discharge process, the battery-type anode retains a constant potential and is linked to the Faraday reaction, regulated by diffusion. The capacitor-type positive electrode exhibits a linear relationship between voltage and charge/discharge time during the charge/discharge phase and is similar to the non-Faraday reaction. During the charge/discharge operation, the battery-type anode retains a constant potential and is linked to the Faraday reaction, regulated by diffusion. According to eqn (1), the energy density of the PIHC is proportional to the sum of squared voltage and capacitance<sup>62</sup>

$$E = \frac{C_s \times \Delta V^2}{7.2} \quad (1)$$

where  $C$  is the specific capacitance of the electrode material and  $V$  is the working voltage window.

The power density can be calculated according to eqn (2).<sup>7,63</sup>

$$P = \frac{V^2}{4mR_s} \quad (2)$$

$V$  is the working voltage,  $m$  means the mass of electrodes, the equivalent series resistance determines  $R_s$ .

Besides, the mass ratio between battery-type and capacitor-type electrodes has a significant impact on the cycling life of PIHC. For hybrid capacitors, although one electrode acts as a battery electrode with different anode and cathode peaks, the cyclic voltammograms (CV) and galvanostatic charge/discharge curves of the entire device may also show more capacitor-like activity with noticeable deviation from ideal capacitive characteristics. In the case of a supercapacitor, the charge balance would be as follows:  $q_+ = q_-$ . Assume that the capacitance properties of a hybrid capacitor are ideal. In the electrode materials involving a different reaction mechanism, the applied current will split into the capacity of the individual electrodes. The charge balance between the two electrodes is necessary to achieve higher energy density in a hybrid capacitor configuration.<sup>64</sup>

$$Q_{\text{anode}} = Q_{\text{cathode}} \quad (3)$$

The total charge stored in each electrode is determined by the specific capacitance ( $C_{\text{electrode}}$ ), the mass of the electrode ( $m$ ), and the potential window ( $\Delta E$ ) of each electrode.

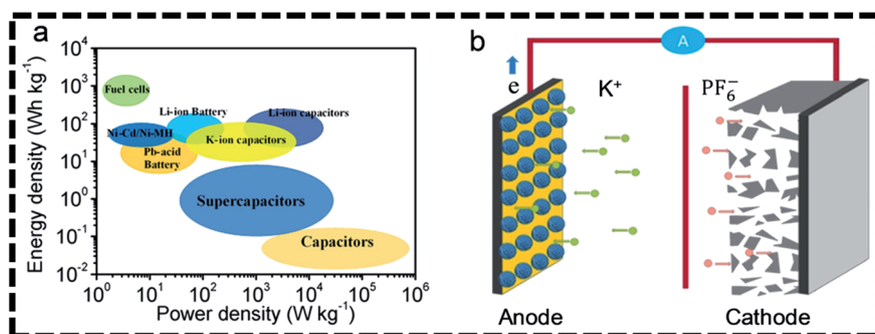


Fig. 1 (a) PIHCs and other energy storage systems have energy capacity and control density profiles (in the case of PIHCs, the values are centered on the Ragone plots in the literature), (b) working principle of a PIHCs.



$$Q_{\text{electrode}} = C_{\text{electrode}} \times m \times \Delta E \quad (4)$$

For asymmetrical PIHCs, the charge should be balanced.<sup>65</sup> Therefore, the mass ratio between two electrodes can be calculated by the following eqn (5)<sup>65</sup>

$$\frac{m_{\text{cathode}}}{m_{\text{anode}}} = \frac{C_{\text{anode}} \times \Delta E_{\text{anode}}}{C_{\text{cathode}} \times \Delta E_{\text{cathode}}} \quad (5)$$

where  $m$ ,  $C$  and  $\Delta E$  are the mass of active material, specific capacitance, and working voltage window for the cathode and anode, respectively.

## 5. Transition metal chalcogenides (TMCs) based composite electrodes for PIHCs

The basic structure and operating theory of “rocking-chair” PIBs are depicted schematically in Fig. 2a. Because of their excellent

physicochemical properties and strong theoretical ability, two-dimensional (2D) transition metal chalcogenides have drawn growing research attention for energy storage devices. TMCs have higher electrical conductivity, mechanical performance, and thermal stability than metal oxides, making them a promising candidate for HCs anode. Unfortunately, the conversion reaction's broad volume expansion induces rapid output degradation, stopping them from becoming suitable anodes for PIHCs. Furthermore, there are high hopes for maximizing the pseudo-capacitance benefits from the charge storage process to improve slow intercalation/deintercalation  $\text{Na}^+/\text{K}^+$  kinetics in transition metal dichalcogenides. Designing nanomaterials is thought to be a good way to shorten the electron and ion diffusion pathways while still exposing more active areas, resulting in better rate efficiency. Despite this, metal dichalcogenides' low intrinsic conductivity induces extreme polarization effects and slow electron utilization. In the charge/discharge phase for metal dichalcogenides, coupling with carbon materials will speed up electron transfer, thus reducing volume

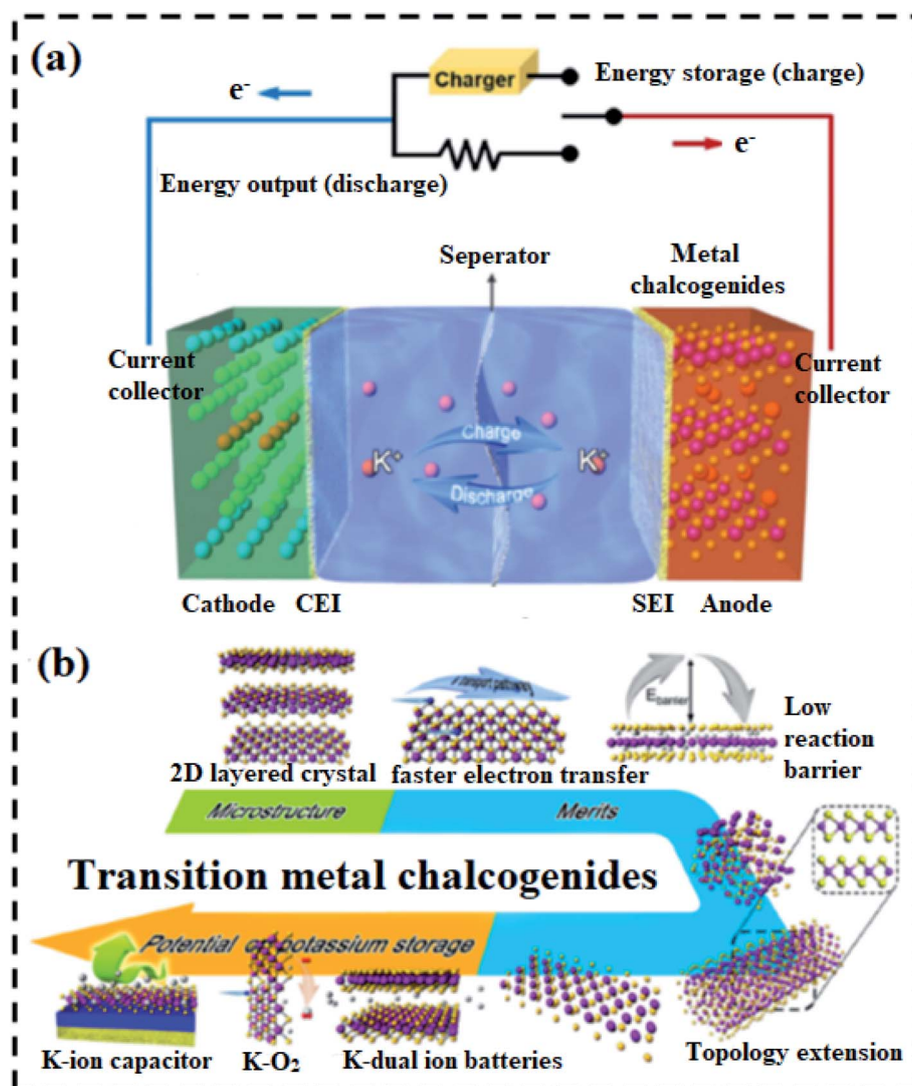


Fig. 2 (a) Schematic showing the setup and operation of a potassium-ion battery in the shape of a rocking chair, (b) a general illustration of the microstructure, benefits, and possible uses of metal chalcogenides in the area of potassium preservation.



transition. Metal chalcogenides are “shining stars,” with potential uses in K dual-ion batteries, K–O<sub>2</sub> batteries, PIHC, and other energy storage and conversion areas, as seen in Fig. 2b.

Because of its high power density and low cost, PIHCs technology has many promises for large-scale energy devices. However, it's still constrained by the slow diffusion method and inadequate K<sup>+</sup> ion storage in the electrodes. Transition metal dichalcogenides with high power, a suitable voltage platform, and stability have emerged as a critical factor as a promising anode for PIHCs functional applications in recent years. In this regard, further research into suitable metal dichalcogenides with new approaches and structures that provide excellent pseudocapacitive conduct and apparent electrochemical kinetics is needed to realize realistic PIHCs devices with reasonable efficiency. As a superior anode material for PIHCs, Yi and colleagues created nitrogen-doped MoSe<sub>2</sub>/graphene (N-MoSe<sub>2</sub>/G) composites with good pseudocapacitive K<sup>+</sup> ion storage.<sup>66</sup> To make diatomite@N-MoSe<sub>2</sub>/G, diatomite templates were combined with graphite oxide, sodium molybdate, and Se powder, then hydrothermally processed and annealed. The 3D biomorphic N-MoSe<sub>2</sub>/G with large pore structures was obtained (Fig. 3a).<sup>66</sup> The SEM and TEM photos demonstrate that the diatomite-derived N-MoSe<sub>2</sub>/G composites maintain a significant number of spherical voids, resulting in a 3D porous N-MoSe<sub>2</sub>/G structure (Fig. 3b and c). The PIHC unit, which used N-MoSe<sub>2</sub>/G as an anode and bioderived activated carbon (AC) as

a cathode, provided a high energy production of 119 W h kg<sup>-1</sup> and a high power density of 7212 W kg<sup>-1</sup>, as well as a long lifetime of 3000 cycles at 1.0 A g<sup>-1</sup> (Fig. 3d). The unique hierarchical 3D nanoarchitecture contributes to improved electrode/electrolyte interfacial connectivity, fast ion/electron transport, reduced volume transition, and enriched active sites, resulting in a noticeable pseudocapacitive effect.

Ge J. *et al.*<sup>68</sup> generated ED-MoS<sub>2</sub>@CT hybrids by self-loading and anchoring well-distributed MoS<sub>2</sub> nanosheets on hollow tubular carbon skeletons (Fig. 3e). The self-loading active materials within hollow structures were accomplished, utilizing MoS<sub>2</sub>@CT composites extracted from well-dispersed anionic groups anchored on the inner surface of HTCSs. The intercalation of ethylene glycol (EG) and dopamine (DA) hydrochloride in a step-by-step manner regulates the number of layers and interlayer spacing of MoS<sub>2</sub>. This unique MoS<sub>2</sub>@CT with the accessible and interconnecting area widened layer spacing and spacious “roads” and “houses” for fast ion/electron diffusion, and the transition will effectively alleviate mechanical strain and provide spacious “roads” and “houses” for rapid ion/electron diffusion and transition. Consequently, the ED-MoS<sub>2</sub>@CT/PC PIHC system has a strong energy/power density and good cycling stability. N-doping carbon-coated FeSe<sub>2</sub> clusters were suggested as the anode for PIHCs by Ge and colleagues. FeSe<sub>2</sub>/N-C has a 3D structure with micro rods assembly and a diameter of about 100 nm and a carbon sheet

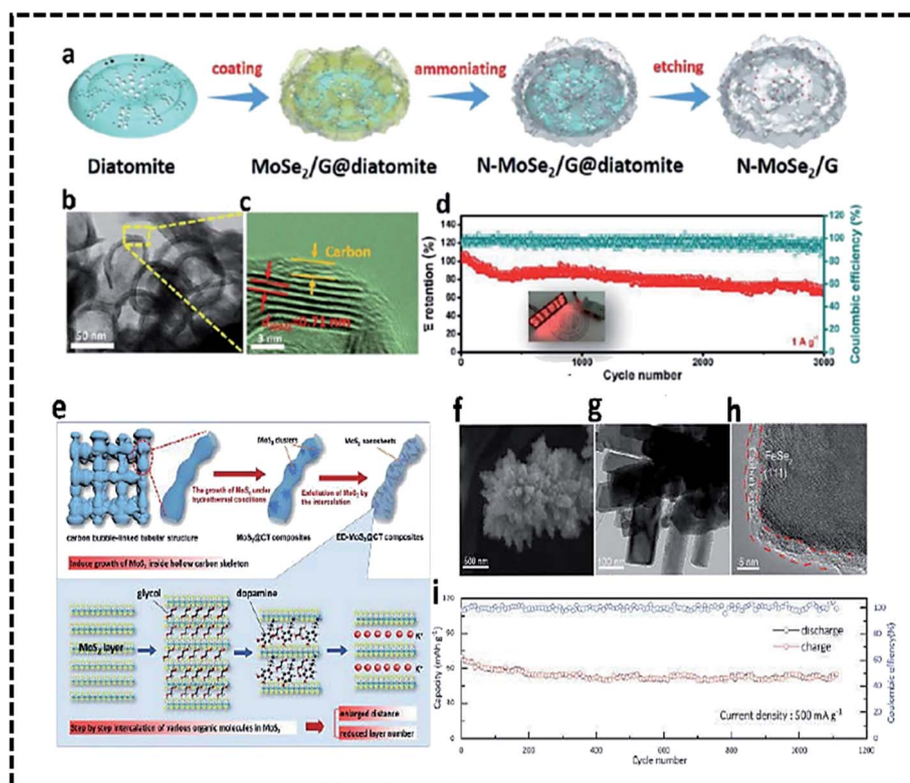


Fig. 3 Schematic diagram of the synthetic process (a) and TEM photos (b and c) of N-MoSe<sub>2</sub>/G as an anode for PIHCs. (d) The long-term cyclic output of the N-MoSe<sub>2</sub>/G/AC PIHC system, (e) schematic diagram of the ED-MoS<sub>2</sub>@CT synthesis method,<sup>67</sup> (f) FeSe<sub>2</sub>/N-C SEM graphic, (g) TEM image, and (h) HRTEM image.<sup>68</sup> (i) The cycling stability of the PIHC device.



with a thickness of about 3 nm, as seen in SEM and TEM pictures (Fig. 3f–h). With its unusual 3D nanostructures and carbon coating, the FeSe<sub>2</sub>/N–C effectively shortens the electron transport road, enhances electron conductivity, provides rich active sites, and reduces volume expansion. The PIHC achieved a high energy density of 230 W h kg<sup>-1</sup>, a strength density of 920 W kg<sup>-1</sup>, and excellent cycling endurance over 1100 cycles at 500 mA g<sup>-1</sup>, thanks to the above merits (Fig. 3i). Sheet-like MoSe<sub>2</sub>/carbon nanofibers produced by a simple electrospinning and selenization process were also investigated as anodes for PIHCs.<sup>69</sup> The composite has outstanding performance in PIHCs, including high capacity and stability, due to the structural stability of one-dimensional nanofibers and the extended interlayer spacing of MoSe<sub>2</sub>. A sheet-on-sheet nanostructure was developed by multilayer tungsten sulfide (WS<sub>2</sub>) nanosheets anchored on nitrogen-doped carbon nanosheets metal-chelate-assisted method followed by a sulfidation procedure. During sodium/potassium ion intercalation/extraction, the composite anodes with sheet-on-sheet architecture boost electron transport, ion migration, and volume expansion of WS<sub>2</sub>, and the interface between the two nanosheets will improve pseudocapacitive charge recovery. Furthermore, the nitrogen-doped carbon hollow nanospheres cathode, which was made up of vertically aligned ultrathin carbon nanosheets, had a wide specific area and pores for anion adsorption (Fig. 4a and b).<sup>70</sup> The nitrogen factor was uniformly distributed in the hollow carbon nanospheres according to EDX mapping (Fig. 4c). The fabricated PIHCs will produce a high energy density of 134.7 W h kg<sup>-1</sup> at a power density of 117.5 W kg<sup>-1</sup> due to unique characteristics and improved anode and cathode compatibility. The high energy of 103.4 W h kg<sup>-1</sup> can also be reached at a high

235 W kg<sup>-1</sup>. Meanwhile, the mounted PIHC has shown good cycling reliability over 500 cycles, with 78% power retention at 500 mA g<sup>-1</sup> (Fig. 4d). Ten processes of the WS<sub>2</sub> composite anode and carbon nanospheres cathode were presented at scan rates of 0.1 and 1 mV<sup>-1</sup>, respectively, prior to assembling the hybrid complete cell to reduce irreversible power.

Guan's team looked at the FeS<sub>2</sub> potassium storage mechanism.<sup>71</sup> Fig. 5a–c indicates that when the discharge voltage decreases to 1.1 V, the peak amplitude of the FeS<sub>2</sub> phase is significantly reduced. In contrast, a slight peak corresponding to KFeS<sub>2</sub> appears, meaning that K<sup>+</sup> is intercalated into FeS<sub>2</sub> interlayers to form K<sub>x</sub>FeS<sub>2</sub>. A minor peak of the Fe process can be seen in the XRD pattern after the anode is discharged to 0.5 V. Because of the low crystallinity or the small particle size in discharge products, there were no prominent XRD peaks when the electrode was fully discharged. Peaks corresponding to K<sub>x</sub>FeS<sub>2</sub> and K<sub>2</sub>S phases were predominantly observed after the electrode was recharged to 2.8 V, suggesting that the final charge result is K<sub>x</sub>FeS<sub>2</sub>. Surprisingly, certain potassium chalcogenides have poor reaction kinetics and also loss chemical activity when de-alloyed. Furthermore, the TEM findings show that Fe and KFeS<sub>2</sub> are embedded in the discharge and charge materials, respectively, which agrees with the XRD results. Consequently, despite the reversibility of the deeper intercalation and conversion steps, the group concluded that the initial phase transformation between FeS<sub>2</sub> and K<sub>x</sub>FeS<sub>2</sub> is an irreversible potassium ion reaction, as seen in Fig. 5d. Guo's group<sup>72</sup> was the first to investigate Sb<sub>2</sub>S<sub>3</sub> as a PIB anode material. To understand the phase evolution of Sb<sub>2</sub>S<sub>3</sub> during discharge/charge, they used in operando synchrotrons XRD (=0.6888), as seen in Fig. 5e. The intercalation of K<sup>+</sup> extends the interlayer

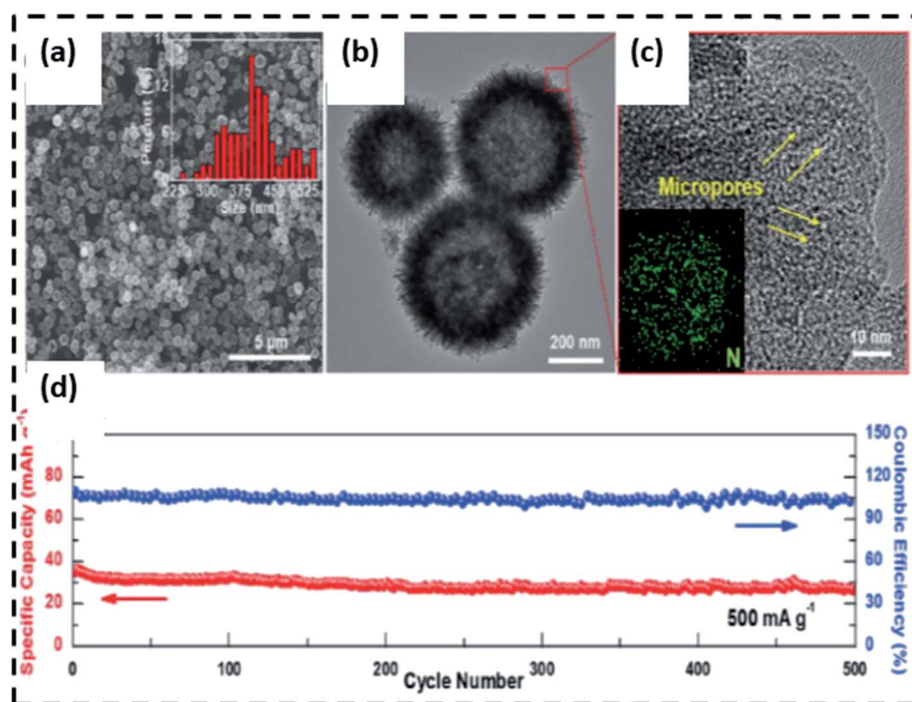


Fig. 4 (a–c) NCHS cathode SEM and TEM characterizations, (d) cycling efficiency of the assembled NCHS/WS<sub>2</sub>@NCNS PIHCs.<sup>70</sup>



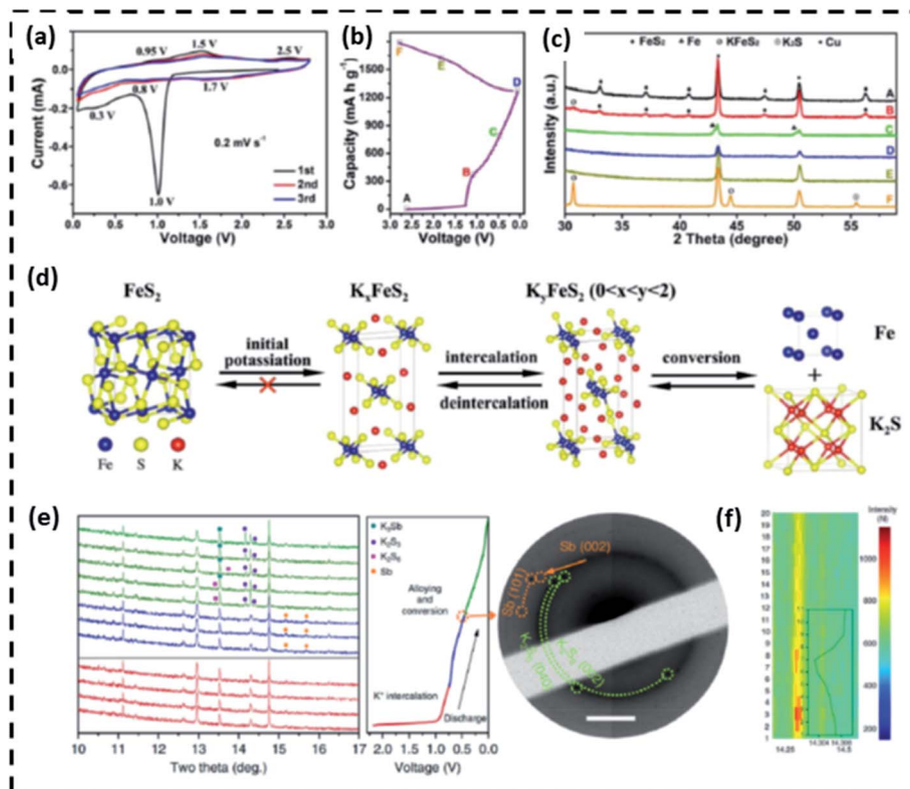
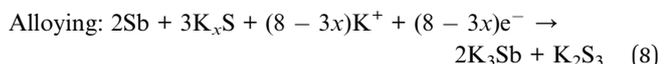
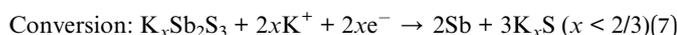
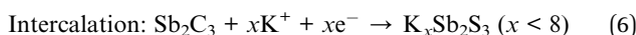


Fig. 5 At different states of pure  $\text{FeS}_2$  anode, (a) CV curves, (b) original discharge/charge curves, and (c) *ex situ* X-ray diffractometry (XRD) trends, (d) a diagram depicting the potassium storage reaction process,<sup>71</sup> (e) *in situ* synchrotron XRD patterns of  $\text{Sb}_2\text{S}_3$  upon K insertion (left) and *ex situ* selected area electron diffraction (SAED) pattern (right) at 0.5 V, (f) Image plots of the (212) reflection and corresponding fitted peak.<sup>72</sup>

gap of  $\text{Sb}_2\text{S}_3$ , as evidenced by the transition to smaller diffraction angles associated with  $\text{Sb}_2\text{S}_3$  above 0.7 V (Fig. 5f). Diffraction peaks attributed to Sb appeared when  $\text{Sb}_2\text{S}_3$  was discharged from 0.7 to 0.5 V, suggesting the existence of transfer reactions. Peaks attributed to  $\text{K}_2\text{S}_6$ ,  $\text{K}_2\text{S}_3$ , and  $\text{K}_3\text{Sb}$  were produced when the voltage reached 0.1 V, but they were weak and wide due to the nano crystallinity. *In situ* selected area electron diffraction (SAED) verified the existence of  $\text{K}_2\text{S}_6$  and Sb as intermediate products based on their polycrystalline diffraction rings, which is encouraging. Therefore, the potassiation mechanism for  $\text{Sb}_2\text{S}_3$  can be summarized by eqn (6)–(8).



Apart from acknowledging undecomposed  $\text{K}_3\text{Sb}$  and  $\text{K}_2\text{S}_3$  in recharged materials, the phase evolution during charging was not addressed in the literature. Furthermore, Guo *et al.* suggested that the rapid decay in potential is due to the continuous pulverization of  $\text{Sb}_2\text{S}_3$  particles induced by the massive volume shift (approximately 300%) during cycling.<sup>72</sup> Furthermore, Xia's team looked into the potassiation/depotassiation method.<sup>71</sup> *In*

*situ* XRD, TEM, and SAED measurements were used to research the actions of  $\text{SnS}_2$ . The XRD findings reveal that the conversion reaction of  $\text{SnS}_2$  starts above 1.0 V, as shown by the existence of diffraction peaks for  $\text{SnS}$ ,  $\text{K}_2\text{S}_5$ , and  $\text{K}_2\text{S}$ . Following that, further conversion and alloying reactions take place, with the prevailing power input happening between 0.01 and 0.5 V. The  $\text{KSn}$  and  $\text{K}_2\text{S}$  (primary) and  $-\text{Sn}$  and  $\text{K}_x\text{Sn}_{2.3}$  (secondary) were found to be the final discharge products (minor). Despite the electrode being recharged to 2.0 V, only  $\text{SnS}$  coupled with  $\text{K}_2\text{S}$  (rather than  $\text{SnS}_2$ ) was detected. The morphology, microstructure, components, and interfacial bonding of metal-chalcogenide-based PIHCs anodes are intimately dependent on their morphology, microstructure, and components, and phase transition is very likely to be incompletely reversible, as shown by the preceding works.

$\text{MnSe}$  is a strongly conductive substance that often affects ion transport and storage.  $\text{MnSe}$  promotes a stable base structure, which reduces diffusive capacitive behavior and improves  $\text{NiCo}_2\text{O}_4$  surface charge transfer. The output of manganese selenide ( $\text{MnSe}$ ) and manganese selenide ( $\text{MnSe}$ )/nickel cobalt oxide ( $\text{NiCo}_2\text{O}_4$ ) as electrode materials for SCs was investigated using an electrodeposition technique. The formation and configuration of the  $\text{MnSe}$  and  $\text{MnSe}(20)/\text{NiCo}_2\text{O}_4$  functioning electrodes are exposed by physical characterization techniques such as X-ray diffraction, field emission scanning electron microscopy, X-ray photoelectron spectroscopy, and



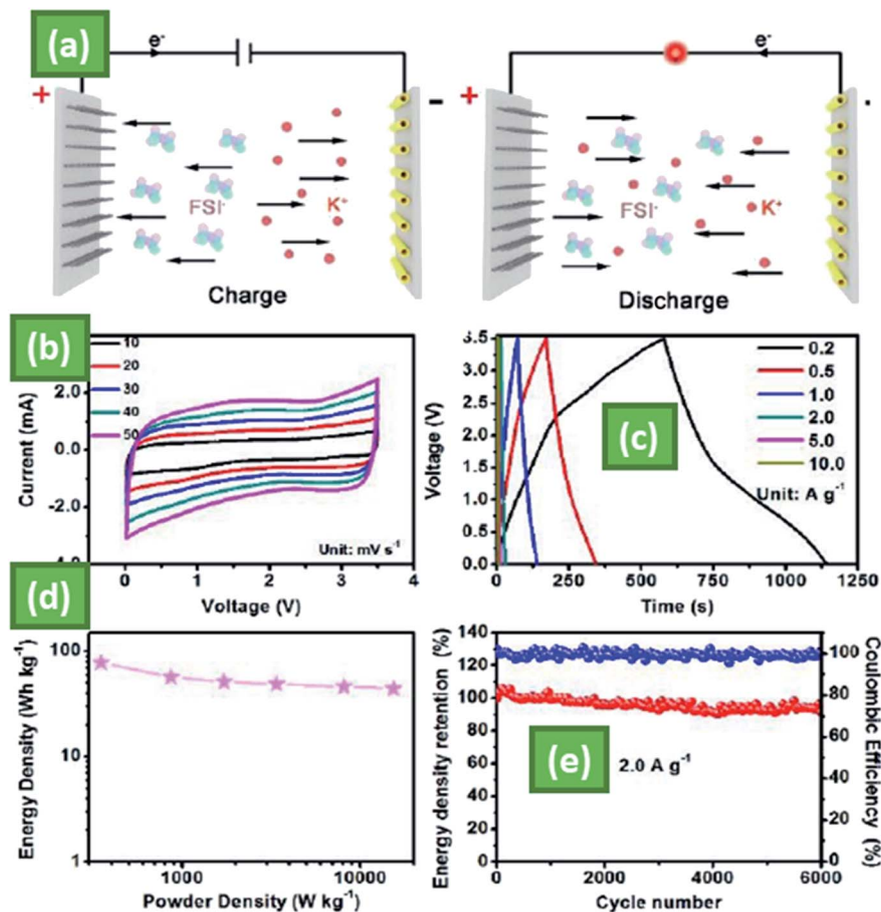


Fig. 6 The following are the results of the PIHC device using a MoS<sub>2/3</sub>Se<sub>4/3</sub>/C-HNT anode and an industrial AC cathode: (a) a schematic diagram of the device's operating principle, (b) CV curves at the scan rate ranging from 10 to 50 mV s<sup>-1</sup> the device's output, (c) galvanostatic charge and discharge curves; (d) Ragone plot, (e) cycling stability at the current density of 2 A g<sup>-1</sup>.

transmission electron microscopy. The electrochemical activity of electrodeposited MnSe and MnSe(20)/NiCo<sub>2</sub>O<sub>4</sub> was observed and analyzed for the first time. At a current density of 0.1 A g<sup>-1</sup>, the electrodeposited MnSe and MnSe(20)/NiCo<sub>2</sub>O<sub>4</sub> single electrodes had specific capacitances of 184.92 F g<sup>-1</sup> and 450.75 F g<sup>-1</sup>, respectively. Furthermore, the MnSe(20)/NiCo<sub>2</sub>O<sub>4</sub> asymmetric cell showed a particular capacitance of 45.78 F g<sup>-1</sup> at 0.1 A g<sup>-1</sup> and 12.46 W h kg<sup>-1</sup> at a current density of 0.1 A g<sup>-1</sup>. The MnSe(20)/NiCo<sub>2</sub>O<sub>4</sub> asymmetric cell demonstrated excellent long-term cycling stability over 5000 cycles, with an 86 percent capacitive retention.<sup>46</sup> The PIHC system centered on MoS<sub>2/3</sub>Se<sub>4/3</sub>/C-HNT anode and industrial AC cathode (denoted as MoS<sub>2/3</sub>Se<sub>4/3</sub>/C-HNT/AC) is further constructed to determine its possible applications, as shown schematically in Fig. 6a. During the charging phase, K<sup>+</sup> is intercalated into MoS<sub>2/3</sub>Se<sub>4/3</sub>/C-HNT anode, and FSI is absorbed onto the surface of the AC cathode before being extracted/desorbed and releasing the electron. In the voltage range of 0.01–3.5 V, Fig. 6b displays successive CV curves under variable scan speeds from 10 to 50 mV s<sup>-1</sup>, where minor differences from the ideal rectangular form can be shown, indicating the coexistence of faradaic and non-faradaic reactions. Because of the differing energy-storage kinetics of the anode and cathode in the hybrid capacitor, the

galvanostatic charge and discharge curves at different current densities (Fig. 6c) do not show a linear relationship like a capacitor or voltage plateaus like a battery, which is compatible with the CV performance. The Ragone plot of the PIHC can be determined from the charge/discharge profiles using the combined mass of the cathode and anode with the IR decrease omitted, as seen in Fig. 6d.

At a power density of 348.1 W kg<sup>-1</sup>, an impressive energy density of 77.4 W h kg<sup>-1</sup> was achieved. It can have an excellent energy density of 44.1 W h kg<sup>-1</sup> even at an ultrahigh power output of 15 414.0 W kg<sup>-1</sup>, which may be useful to practical applications requiring high power density. Importantly, the hybrid capacitor system will maintain 91.3% of its energy density after 6000 cycles with a current density of 2.0 A g<sup>-1</sup> (Fig. 6e) and a coulombic performance of nearly 100%, showing the great ability of as-synthesized MoS<sub>2/3</sub>Se<sub>4/3</sub>/C-HNTs as anode candidates for advanced K<sup>+</sup> dependent energy storage applications.<sup>73</sup>

## 6. Summary and outlook

According to previous research, TMC-based composites have a lot of capacity for potassium preservation. The reaction



kinetics of TMC, which are influenced by their elements, microstructure, particle size, and hybridization, determine the depth of potassiation and depotassiation. Limiting discharge width, nanostructure engineering, containment by carbon nanophases, and ternary alloying are four standard modification techniques that can be used to overcome TMC's inherent structural instability and low electric/ionic conductivity, resulting in significant improvements in electrochemical efficiency. However, these modifications tactics pose several concerns that should be considered. Finally, we believe that future research focusing on TMC for potassium storage should pay more attention to the following points.

- Build heterojunctions dependent on metal chalcogenides. This orientation tends to be unique to PIHCs dependent on metal chalcogenides. Surface/interface control, on the other side, can be an important way of guiding modifications to achieve high efficiency.

- During the synthesis of TMC-based compounds, high-temperature vulcanization occurs, thus requires high-tech equipment. This is inconvenient in terms of manufacturing and implementation. The mass development and low-cost processing of high-quality TMC-based materials must be solved before TMC-based materials can be used in PIHCs.

- Increase the amount of electrochemical activity that is regulated by capacitance. Even at high rates, most recorded TMC-based PIHCs anodes have a small capacitance-based power contribution (80%). If this aspect of power contribution can be encouraged, capacity retention can be improved at high rates. This will be particularly useful in consumer markets where quick charging is expected.

- To better understand the underlying potassium storage mechanisms, further *in situ* characterization is required. It is hoped that a better understanding of the process would help create new TMC-based nanostructures and provide guidance for potential material modification.

- Develop innovative potassium-based hybrid energy storage systems that incorporate the benefits of multiple energy storage technologies. This modern hybrid system is expected to make greater use of modified metal chalcogenides' superior potassium storage capacity, resulting in higher energy densities.

Current TMC for PIHCs and other potassium-based energy storage systems is expected to be overcome by advances in design, preparation, and characterization techniques, making potassium-based energy storage solutions a feasible option for daily use.

## Author contributions

Dr Muhammad Sajjad: investigation, formal analysis, data curation, conceptualization, validation, writing-original draft, writing-review, and editing, Dr Fang Cheng: investigation, Prof. Wen Lu: supervision.

## Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have influenced the work reported in this paper.

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## References

- 1 X. Zou, P. Xiong, J. Zhao, J. Hu, Z. Liu and Y. Xu, *Phys. Chem. Chem. Phys.*, 2017, **19**, 26495–26506.
- 2 M. Yan, H. Chen, Y. Yu, H. Zhao, C. F. Li, Z. Y. Hu, P. Wu, L. Chen, H. Wang and D. Peng, *Adv. Energy Mater.*, 2018, **8**, 1801066.
- 3 M. Yan, W.-P. Wang, Y.-X. Yin, L.-J. Wan and Y.-G. Guo, *EnergyChem*, 2019, **1**, 100002.
- 4 J. Huang, B. G. Sumpter and V. Meunier, *Chem.–Eur. J.*, 2008, **14**, 6614–6626.
- 5 M. Hashemi, M. S. Rahmanifar, M. F. El-Kady, A. Noori, M. F. Mousavi and R. B. Kaner, *Nano Energy*, 2018, **44**, 489–498.
- 6 A. González, E. Goikolea, J. A. Barrena and R. Mysyk, *Renewable Sustainable Energy Rev.*, 2016, **58**, 1189–1206.
- 7 M. Sajjad, *J. Inorg. Organomet. Polym. Mater.*, 2021, 1–19.
- 8 A. S. Arico, P. Bruce, B. Scrosati, J.-M. Tarascon and W. Van Schalkwijk, *Nature Materials*, 2011, 148–159.
- 9 B. Dunn, H. Kamath and J.-M. Tarascon, *Science*, 2011, **334**, 928–935.
- 10 J. Kim and A. Manthiram, *Nature*, 1997, **390**, 265–267.
- 11 P. Poizot, S. Laruelle, S. Grugeon, L. Dupont and J. Tarascon, *Nature*, 2000, **407**, 496–499.
- 12 S. J. Lee, *Nat. Nanotechnol.*, 2010, **5**, 531.
- 13 D. Zhao, R. Zhao, S. Dong, X. Miao, Z. Zhang, C. Wang and L. Yin, *Energy Environ. Sci.*, 2019, **12**, 2422–2432.
- 14 T. Liu, J. P. Vivek, E. W. Zhao, J. Lei, N. Garcia-Araez and C. P. Grey, *Chem. Rev.*, 2020, **120**, 6558–6625.
- 15 K. J. Griffith, K. M. Wiaderek, G. Cibir, L. E. Marbella and C. P. Grey, *Nature*, 2018, **559**, 556–563.
- 16 S. Chu, Y. Cui and N. Liu, *Nat. Mater.*, 2017, **16**, 16–22.
- 17 A. González, E. Goikolea, J. A. Barrena and R. Mysyk, *Renewable Sustainable Energy Rev.*, 2016, **58**, 1189–1206.
- 18 A. Jagadale, X. Zhou, R. Xiong, D. P. Dubal, J. Xu and S. Yang, *Energy Storage Materials*, 2019, **19**, 314–329.
- 19 M. Ma, Y. Yao, Y. Wu and Y. Yu, *Advanced Fiber Materials*, 2020, 1–24.
- 20 A. Eftekhari, Z. Jian and X. Ji, *ACS Appl. Mater. Interfaces*, 2017, **9**, 4404–4419.
- 21 X. Wu, D. P. Leonard and X. Ji, *Chem. Mater.*, 2017, **29**, 5031–5042.
- 22 T. Hosaka, K. Kubota, A. S. Hameed and S. Komaba, *Chem. Rev.*, 2020, **120**, 6358–6466.
- 23 D. Li, Q. Sun, Y. Zhang, L. Chen, Z. Wang, Z. Liang, P. Si and L. Ci, *ChemSusChem*, 2019, **12**, 2689–2700.
- 24 B. Ji, F. Zhang, X. Song and Y. Tang, *Adv. Mater.*, 2017, **29**, 1700519.
- 25 L. Fan, R. Ma, Q. Zhang, X. Jia and B. Lu, *Angew. Chem., Int. Ed.*, 2019, **58**, 10500–10505.



- 26 K. Share, A. P. Cohn, R. Carter, B. Rogers and C. L. Pint, *ACS Nano*, 2016, **10**, 9738–9744.
- 27 R. Rajagopalan, Y. Tang, X. Ji, C. Jia and H. Wang, *Adv. Funct. Mater.*, 2020, **30**, 1909486.
- 28 X. Zhang, D. Yang, X. Rui, Y. Yu and S. Huang, *Curr. Opin. Electrochem.*, 2019, **18**, 24–30.
- 29 S. Komaba, T. Hasegawa, M. Dahbi and K. Kubota, *Electrochem. Commun.*, 2015, **60**, 172–175.
- 30 L. Jiang, Y. Lu, C. Zhao, L. Liu, J. Zhang, Q. Zhang, X. Shen, J. Zhao, X. Yu and H. Li, *Nat. Energy*, 2019, **4**, 495–503.
- 31 J. C. Pramudita, D. Sehwat, D. Goonetilleke and N. Sharma, *Adv. Energy Mater.*, 2017, **7**, 1602911.
- 32 C. D. Wessells, S. V. Peddada, R. A. Huggins and Y. Cui, *Nano Lett.*, 2011, **11**, 5421–5425.
- 33 H. Huang, R. Xu, Y. Feng, S. Zeng, Y. Jiang, H. Wang, W. Luo and Y. Yu, *Adv. Mater.*, 2020, **32**, 1904320.
- 34 K. Song, C. Liu, L. Mi, S. Chou, W. Chen and C. Shen, *Small*, 2019, 1903194.
- 35 L. Zhou, Z. Cao, W. Wahyudi, J. Zhang, J.-Y. Hwang, Y. Cheng, L. Wang, L. Cavallo, T. Anthopoulos and Y.-K. Sun, *ACS Energy Lett.*, 2020, **5**, 766–776.
- 36 W. Zhang, Y. Liu and Z. Guo, *Sci. Adv.*, 2019, **5**, eaav7412.
- 37 M. Liu, L. Chang, Z. Le, J. Jiang, J. Li, H. Wang, C. Zhao, T. Xu, P. Nie and L. Wang, *ChemSusChem*, 2020, **13**, 5837–5862.
- 38 S. Chong, L. Sun, C. Shu, S. Guo, Y. Liu, W. A. Wang and H. K. Liu, *Nano Energy*, 2019, **63**, 103868.
- 39 C. Chen, Y. Yang, X. Tang, R. Qiu, S. Wang, G. Cao and M. Zhang, *Small*, 2019, **15**, 1804740.
- 40 V. Aravindan, J. Gnanaraj, Y.-S. Lee and S. Madhavi, *Chem. Rev.*, 2014, **114**, 11619–11635.
- 41 Y. Zhang, J. Jiang, Y. An, L. Wu, H. Dou, J. Zhang, Y. Zhang, S. Wu, X. Zhang and M. Dong, *ChemSusChem*, 2020, 651–658.
- 42 A. Le Comte, Y. Reynier, C. Vincens, C. Leys and P. Azaïs, *J. Power Sources*, 2017, **363**, 34–43.
- 43 L. Zhou, M. Zhang, Y. Wang, Y. Zhu, L. Fu, X. Liu, Y. Wu and W. Huang, *Electrochim. Acta*, 2017, **232**, 106–113.
- 44 Z. Hai and S. Zhuiykov, *Adv. Mater. Interfaces*, 2018, **5**, 1701385.
- 45 Y.-W. Lee, B.-S. Kim, J. Hong, H. Choi, H.-S. Jang, B. Hou, S. Pak, J. Lee, S.-H. Lee and S. M. Morris, *Nano Energy*, 2017, **37**, 15–23.
- 46 V. Raman, D. Chinnadurai, R. Rajmohan, V. T. Chebrolu, V. Rajangam and H.-J. Kim, *New J. Chem.*, 2019, **43**, 12630–12640.
- 47 L. Jin, C. Shen, A. Shellikeri, Q. Wu, J. Zheng, P. Andrei, J.-G. Zhang and J. P. Zheng, *Energy Environ. Sci.*, 2020, **13**, 2341–2362.
- 48 H. Wang, C. Zhu, D. Chao, Q. Yan and H. J. Fan, *Adv. Mater.*, 2017, **29**, 1702093.
- 49 A. Vasileff, Y. Zheng and S. Z. Qiao, *Adv. Energy Mater.*, 2017, **7**, 1700759.
- 50 C. Sun, Q. Dong, J. Yang, Z. Dai, J. Lin, P. Chen, W. Huang and X. Dong, *Nano Res.*, 2016, **9**, 2234–2243.
- 51 L. Yang, W. Zhou, J. Lu, D. Hou, Y. Ke, G. Li, Z. Tang, X. Kang and S. Chen, *Nano Energy*, 2016, **22**, 490–498.
- 52 Y. Qiao, P. Yuan, Y. Hu, J. Zhang, S. Mu, J. Zhou, H. Li, H. Xia, J. He and Q. Xu, *Adv. Mater.*, 2018, **30**, 1804504.
- 53 Y. Wu, H. B. Huang, Y. Feng, Z. S. Wu and Y. Yu, *Adv. Mater.*, 2019, **31**, 1901414.
- 54 S. Yao, J. Cui, J. Huang, J. Q. Huang, W. G. Chong, L. Qin, Y. W. Mai and J. K. Kim, *Adv. Energy Mater.*, 2018, **8**, 1702267.
- 55 Y. Qian, S. Jiang, Y. Li, Z. Yi, J. Zhou, T. Li, Y. Han, Y. Wang, J. Tian and N. Lin, *Adv. Energy Mater.*, 2019, **9**, 1901676.
- 56 D. Qiu, J. Guan, M. Li, C. Kang, J. Wei, Y. Li, Z. Xie, F. Wang and R. Yang, *Adv. Funct. Mater.*, 2019, **29**, 1903496.
- 57 Z. Li, Y. Dong, J. Feng, T. Xu, H. Ren, C. Gao, Y. Li, M. Cheng, W. Wu and M. Wu, *ACS Nano*, 2019, **13**, 9227–9236.
- 58 Z. Xu, M. Wu, Z. Chen, C. Chen, J. Yang, T. Feng, E. Paek and D. Mitlin, *Adv. Sci.*, 2019, **6**, 1802272.
- 59 J. Chen, B. Yang, B. Liu, J. Lang and X. Yan, *Curr. Opin. Electrochem.*, 2019, **18**, 1–8.
- 60 B. Yang, J. Chen, L. Liu, P. Ma, B. Liu, J. Lang, Y. Tang and X. Yan, *Energy Storage Materials*, 2019, **23**, 522–529.
- 61 Y. Luo, L. Liu, K. Lei, J. Shi, G. Xu, F. Li and J. Chen, *Chem. Sci.*, 2019, **10**, 2048–2052.
- 62 M. Sajjad, R. Tao, K. Kang, S. Luo and L. Qiu, *ACS Appl. Energy Mater.*, 2021, **4**, 828–838.
- 63 H. Yuan, L. Kong, T. Li and Q. Zhang, *Chin. Chem. Lett.*, 2017, **28**, 2180–2194.
- 64 M. Sajjad, Y. Khan and W. Lu, *J. Energy Storage*, 2021, **35**, 102336.
- 65 M. Sajjad and Y. Khan, *CrystEngComm*, 2021, **23**(15), 2869–2879.
- 66 Y. Yi, Z. Sun, C. Li, Z. Tian, C. Lu, Y. Shao, J. Li, J. Sun and Z. Liu, *Adv. Funct. Mater.*, 2020, **30**, 1903878.
- 67 Y. Cui, W. Liu, W. Feng, Y. Zhang, Y. Du, S. Liu, H. Wang, M. Chen and J. Zhou, *Adv. Funct. Mater.*, 2020, **30**, 1908755.
- 68 J. Ge, B. Wang, J. Wang, Q. Zhang and B. Lu, *Adv. Energy Mater.*, 2020, **10**, 1903277.
- 69 Q. Shen, P. Jiang, H. He, C. Chen, Y. Liu and M. Zhang, *Nanoscale*, 2019, **11**, 13511–13520.
- 70 Y. Li, Y. Yang, P. Zhou, T. Gao, Z. Xu, S. Lin, H. Chen, J. Zhou and S. Guo, *Matter*, 2019, **1**, 893–910.
- 71 Y. Zhao, J. Zhu, S. J. H. Ong, Q. Yao, X. Shi, K. Hou, Z. J. Xu and L. Guan, *Adv. Energy Mater.*, 2018, **8**, 1802565.
- 72 Y. Liu, Z. Tai, J. Zhang, W. K. Pang, Q. Zhang, H. Feng, K. Konstantinov, Z. Guo and H. K. Liu, *Nat. Commun.*, 2018, **9**, 1–10.
- 73 J. Gao, G. Wang, Y. Liu, J. Li, B. Peng, S. Jiao, S. Zeng and G. Zhang, *J. Mater. Chem. A*, 2020, **8**, 13946–13954.

