#### **REVIEW ARTICLE**



# High-Energy Room-Temperature Sodium–Sulfur and Sodium–Selenium Batteries for Sustainable Energy Storage

Zefu Huang<sup>1</sup> · Pauline Jaumaux<sup>1</sup> · Bing Sun<sup>1</sup> · Xin Guo<sup>1</sup> · Dong Zhou<sup>1</sup> · Devaraj Shanmukaraj<sup>2</sup> · Michel Armand<sup>2</sup> · Teofilo Rojo<sup>3</sup> · Guoxiu Wang<sup>1</sup>

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### Abstract

Rechargeable room-temperature sodium–sulfur (Na–S) and sodium–selenium (Na–Se) batteries are gaining extensive attention for potential large-scale energy storage applications owing to their low cost and high theoretical energy density. Optimization of electrode materials and investigation of mechanisms are essential to achieve high energy density and long-term cycling stability of Na–S(Se) batteries. Herein, we provide a comprehensive review of the recent progress in Na–S(Se) batteries. We elucidate the Na storage mechanisms and improvement strategies for battery performance. In particular, we discuss the advances in the development of battery components, including high-performance sulfur cathodes, optimized electrolytes, advanced Na metal anodes and modified separators. Combined with current research achievements, this review outlines remaining challenges and clear research directions for the future development of practical high-performance Na– S(Se) batteries.

**Keywords** Sodium–sulfur batteries  $\cdot$  Sodium–selenium batteries  $\cdot$  Sulfur cathodes  $\cdot$  Electrolyte engineering  $\cdot$  Solid-state electrolytes  $\cdot$  Sodium metal anodes

Zefu Huang and Pauline Jaumaux have contributed equally to this work.

Guoxiu Wang Guoxiu.wang@uts.edu.au

> Bing Sun bing.sun@uts.edu.au

Xin Guo xin.guo@uts.edu.au

Michel Armand marmand@cicenergigune.com

Teofilo Rojo teo.rojo@ehu.eus

- <sup>1</sup> Centre for Clean Energy Technology, School of Mathematical and Physical Sciences, Faculty of Science, University of Technology Sydney, Ultimo, NSW 2007, Australia
- <sup>2</sup> Centre for Cooperative Research on Alternative Energies (CIC energiGUNE), Basque Research and Technology Alliance (BRTA), Alava Technology Park, Albert Einstein 48, 01510 Vitoria-Gasteiz, Spain
- <sup>3</sup> Inorganic Chemistry Department, University of the Basque Country UPV/EHU, P.O. Box. 644, 48080 Bilbao, Spain

# **1** Introduction

Fast global warming, heavy environmental pollution, vast natural resource wastage, uncontrolled urbanization and severe climate change have become increasingly urgent challenges that need to be resolved in the next few decades. Extensive use of renewable energy from natural resources (e.g., solar, wind and hydro) and reduced usage of fossil fuels constitute a sustainable approach to relieve the pressure from these challenges. However, these new types of renewable energy sources are season/weather dependent and intermittent [1]. To ensure affordable and continuous energy supply to customers, the development of stationary electrical energy storage (SEES) systems is essential to boost the utilization of renewable energies [1, 2]. SEES is a technique that can convert and store the power from a power network and convert the power back to the network when needed. Several SEES technologies have been developed to match the energy storage demand, including electrochemical storage (e.g., batteries, supercapacitors), thermal storage (e.g., molten salt, ice storage) or mechanical storage (e.g., pumped hydroelectric systems, flywheels) [2–8]. Among them, rechargeable batteries have attracted extensive interest, especially

rechargeable lithium (Li)-ion batteries, which have dominated the market since their commercialization three decades ago, with applications ranging from small (e.g., smartphones, laptops) to large (e.g., electric vehicles, power walls) devices [9]. The fast-growing and higher demand energy storage market raises various concerns about (1) the limited raw material resources of lithium and cobalt (employed in cathode materials) or even nickel and copper and (2) the limited energy density of batteries based on graphite anodes and transition metal cathodes [9, 10]. Although employing Li metal anodes instead of graphite anodes could resolve the energy density limitation issue, the availability and cost of Li metal batteries remain problematic. Therefore, alternative battery technologies are being investigated, such as alkali metal (e.g., sodium (Na), potassium (K)) batteries and multivalent metal (e.g., zinc (Zn), magnesium (Mg), aluminum (Al), calcium (Ca)) batteries [11, 12].

Among all the alkali elements, Na has similar chemical characteristics to Li because it belongs to the same element group (group IA elements) and has attracted the most attention from researchers interested in studying beyond-lithiumion batteries. The redox electrode potential of the Na metal anode is -2.71 V (vs. the standard hydrogen electrode, SHE), which is only 0.35 V higher than that of the Li metal anode. In addition, Na metal anodes possess unique chemical and electrochemical properties, such as a high specific capacity of 1 165 mAh g<sup>-1</sup>, abundance in the Earth's crust, and low cost (sodium carbonate—155 US\$ t<sup>-1</sup> vs. lithium carbonate—17 000 US\$ t<sup>-1</sup>) [13–20]. Therefore, Na-based rechargeable batteries have recently attracted increasing attention.

To fulfill the low cost and high theoretical energy density requirements, room-temperature (RT) sodium-sulfur (selenium) (Na-S(Se)) batteries show the potential to be promising candidates for application in next-generation large-scale SEES systems. Sulfur (S) and selenium (Se) are promising cathode material candidates due to their high theoretical capacity (S—1 672 mAh  $g^{-1}$ , Se—678 mAh  $g^{-1}$ ), abundant storage on the Earth, and low prices [18–23]. Sulfur is priced very competitively in the current market at only approximately 69 US\$  $t^{-1}$  [18], which makes the cathode based on sulfur practical for application in large-scale energy storage systems. Na-S batteries have already been commercialized in high-temperature (HT) rechargeable batteries with molten Na as the anode, Na-ion-conductive ceramics as the electrolyte, and molten sulfur as the cathode. The high operating temperature of such batteries (above 300 °C) impedes their facile and safe application in large-scale energy storage systems [24–27]. Therefore, a surge of interest in RT Na metal batteries has occurred in the past decade, in which Na metal is directly employed as the anode. Additionally, selenium is also very attractive as a new cathode material due to its relatively high ionic conductivity  $(1 \times 10^{-3} \text{ S m}^{-1})$  compared to sulfur and competitive theoretical specific energy (644 Wh  $kg^{-1}$ ), which has resulted in the flourishing development of Na–Se batteries in recent years [28–30].

However, these battery technologies still face various challenges for practical applications. On the anode side, Na metal suffers from vigorous side reactions and dendrite growth, which cause a short cycle life and severe safety issues. On the cathode side, sluggish reaction kinetics and complex reaction mechanisms lead to poor cycling performance. Extensive research is ongoing to overcome these challenges, including developing high-efficiency Na metal anodes, employing electrolyte engineering strategies and formulating optimized cathodes.

In this review, we comprehensively summarize the recent progress in achieving high-energy-density RT Na-S and Na-Se batteries. First, several high-performance cathode materials, including porous carbon-based composites, 2D material-based composites, polymeric sulfur composites, and other sulfur-integrated composites, are discussed with respect to associated structure optimization tactics and underpinning mechanisms. Then, the recent progress in electrolytes for sulfur cathodes and Na metal anodes is separately summarized and discussed. In particular, this review also highlights the advances in developing gel polymer electrolytes (quasi-solid-state electrolytes) and solid-state electrolytes by enhancing their conductivity, mechanical strength, solid-solid interfacial contact with electrodes, and electrochemical and thermal stability, which can effectively improve the safety and electrochemical performance of Na-S(Se) batteries. Furthermore, we emphasize the development of advanced Na metal anodes from the points of view of nanostructure construction and advanced artificial protective layers that stabilize the solid electrolyte interphase (SEI). Additionally, several other strategies, such as separator modification and interlayer design that can simultaneously stabilize the cathode and anode, are reviewed, and the corresponding reaction mechanisms are simultaneously illustrated in each section. Finally, we provide future research directions to further improve the electrochemical performance of RT Na-S and Na-Se batteries.

# 2 History, Reaction Mechanisms and Challenges of Na Metal Batteries

### 2.1 Roadmap from Na-Ion Batteries to Na Metal Batteries

The development of Na-ion batteries started in the 1970s side-by-side with that of lithium-ion (Li-ion) batteries [31–33], as shown in Fig. 1a. However, based on the discoveries by John Goodenough et al. during the 1970s to 1980s, Li-ion batteries were commercialized in the 1990s [34, 35].



Fig. 1 a Chronological review of the publications and the key points in the development of RT Na–S and Na–Se batteries. b Comparison of sodium, sulfur, and lithium in terms of the element abundance and current market prices of their raw materials, comparison of the ionic

conductivity between sulfur and selenium, and comparison of the theoretical specific energy and specific capacity among RT Na–S, Na–Se and Na-ion batteries.

Afterwards, Li-ion batteries dominated the market, which led to the research on Na-ion batteries almost stopping during that period. Subsequently, the cost of the raw materials for Li-ion batteries continued to increase, which forced the interest in Na-ion batteries to be renewed in the early 2010s [24]. However, state-of-the-art prototype Na-ion batteries can only deliver a specific energy density of approximately 150 Wh kg<sup>-1</sup>, which is a small fraction of their theoretical value [24]. This made researchers shift their focus toward high-energy Na metal batteries, such as RT Na–S and Na–Se batteries. The advantages of RT Na–S and Na–Se batteries can be identified as two parts. The first part is the raw material abundance and the cost of the batteries. As shown in Fig. 1b, sodium has an approximately 1 180 times higher content than lithium in the Earth's crust and a much lower price (approximately 110 times) than lithium, as does sulfur (approximately 18 times higher content and 246 times lower price than lithium). The second part is their electrochemical performance. RT Na–S batteries and Na–Se batteries have approximately 17 times and 8 times higher theoretical specific energy densities than Na-ion batteries, respectively. Moreover, Se has a higher conductivity than sulfur, and the theoretical volumetric capacity (3 253 mAh cm<sup>-3</sup>) is as high as that of sulfur (3 467 mAh cm<sup>-3</sup>) [36], which makes it promising for high-power and compact energy storage devices. These highlights of RT Na–S and Na–Se batteries give them the potential for further development in future stationary storage.

High-temperature sodium-sulfur (HT Na-S) batteries were first developed for electric vehicle (EV) applications

due to their high theoretical volumetric energy density. In 1968, Kummer et al. from Ford Motor Company first released the details of the HT Na-S battery system using a  $\beta$ "-alumina solid electrolyte [37]. According to their report, HT Na-S batteries need to operate at a temperature of approximately 300 to 350 °C, in which sodium metal, sulfur and the resulting polysulfides are all in molten states with high chemical reactivity. This causes severe safety concerns and limits the application of HT Na-S batteries for mobile devices and residential use. Furthermore, the discharge of HT Na-S batteries stops at the formation of Na<sub>2</sub>S<sub>3</sub>, which is the ultimate liquid-phase catholyte at 300 °C. This leads to the practical capacity decreasing from 1 672 mAh  $g^{-1}$  (complete conversion from  $\beta$ -S<sub>8</sub> to Na<sub>2</sub>S) to 557 mAh g<sup>-1</sup> (conversion from  $\beta$ -S<sub>8</sub> to Na<sub>2</sub>S<sub>3</sub>) [38]. In 2003, NGK Insulator Ltd. started the mass production of HT Na–S batteries [39]. After a few years of development, the production capacity of NGK Insulator Ltd. for HT Na-S batteries achieved 150 MW per year in 2010. However, a fire incident made them temporarily cease HT Na-S production in 2011 [40].

To decrease the operating temperature and improve the capacity of conventional HT Na–S batteries, Fielder et al. and Abraham et al. found that glymes could be suitable solvents for a catholyte that can dissolve both elemental sulfur and solid Na<sub>2</sub>S<sub>x</sub> (x < 3) at 150–300 °C [41]. This enabled the discovery of intermediate-temperature sodium–sulfur (IMT Na–S) battery systems [42]. Subsequently, studies on catholyte additives, Na alloy anodes, improvement of the  $\beta''$ -Al<sub>2</sub>O<sub>3</sub> purity, and solid inorganic electrolytes further enhanced the capacity and cycle life of IMT Na–S batteries [43]. However, IMT Na–S batteries still face similar safety problems owing to the employment of corrosive molten sodium, as well as a high maintenance cost.

RT Na–S batteries operating under ambient temperature (25 °C) were first proposed in 2006, suggesting the possibility of significantly reducing potential safety issues such as fire or explosion derived from highly reactive molten sodium and sulfur. Moreover, the maintenance cost of RT Na–S batteries is remarkably lower than that of HT Na–S and IMT Na–S batteries because they do not require stringent insulation cell packaging and extra heating device to maintain the high working temperature. More importantly, the theoretical energy density of RT Na–S batteries is up to 1 274 Wh kg<sup>-1</sup> owing to the complete transition from  $\beta$ -S<sub>8</sub> to Na<sub>2</sub>S, which is much higher than the theoretical energy density of HT Na–S batteries (760 Wh kg<sup>-1</sup>) [44]. All these advantages provide RT Na–S batteries with a promising future in the development of SEES systems.

As a chemical analogue of sulfur, Se has recently attracted growing research interest due to its much higher electronic conductivity  $(1 \times 10^{-3} \text{ S m}^{-1})$  than sulfur  $(5 \times 10^{-30} \text{ S m}^{-1})$  [45, 46]. Additionally, selenium also has an enhanced reaction activity with Na at RT. Meanwhile, Sodium–Selenium

(Na–Se) batteries have a comparable high volumetric capacity, which makes them attractive for energy storage with limited space. However, Na–Se batteries have been less investigated due to the intrinsic low theoretical gravimetric capacity of Se (678 mAh  $g^{-1}$ ) compared to S. In the following sections, we will systematically introduce the mechanisms, challenges, and up-to-date research progress for both RT Na–S and Na–Se batteries.

# 2.2 Electrochemical Mechanisms and Challenges of RT Na–S and Na–Se Batteries

The RT Na–S battery commonly consists of a sulfur cathode, a sodium metal anode, and a separator soaked with a Na-ionconducting electrolyte (Fig. 2). The working mechanism of the RT Na–S battery is similar to that of the Li–S battery system, which comprises a series of stepwise reactions starting from ring-opening of  $\beta$ -S<sub>8</sub>, followed by the formation of long-chain sodium polysulfides (Na<sub>2</sub>S<sub>x</sub>, 4 $\leq x \leq 8$ ) and then short-chain sodium polysulfide species (Na<sub>2</sub>S<sub>x</sub>, 1 $\leq x \leq 4$ ) [10, 26]. As shown in the inset of Fig. 2, a representative discharge curve of the RT Na–S battery can be identified as consisting of four stages.

- 1. In stage I, the solid elemental sulfur  $\beta$ -S<sub>8</sub> is reduced to a soluble long-chain Na polysulfide (Na<sub>2</sub>S<sub>8</sub>), corresponding to the plateau at ~2.2 V.
- 2. Stage II corresponds to the sloping curve in the voltage region between 2.2 and 1.65 V, in which  $Na_2S_8$  converts in solution to  $Na_2S_4$ .
- Stage III corresponds to the ~1.65 V plateau, where the liquid Na<sub>2</sub>S<sub>4</sub> reduces to insoluble short-chain Na polysulfides, i.e., Na<sub>2</sub>S<sub>3</sub>, Na<sub>2</sub>S<sub>2</sub>, and Na<sub>2</sub>S.
- Stage IV corresponds to the sloping curve in the voltage region between 1.65 and 1.2 V, in which a solid-to-solid transition from Na<sub>2</sub>S<sub>2</sub> to Na<sub>2</sub>S occurs [47].

Although RT Na-S batteries possess a high theoretical energy density depending on the above reactions, several issues must be solved before their practical application. First, the extremely low electronic conductivity of sulfur leads to sluggish electrochemical reaction kinetics and low utilization of active sulfur during discharging. Second, the tremendous volumetric change of sulfur during sodiation/ desodiation (~260% volume expansion upon full discharge) induces pulverization of the sulfur-based cathode, resulting in loss of active materials and fast capacity fading during cycling [26]. Third, the reaction kinetics of the transition from long-chain sodium polysulfide species to short-chain sodium polysulfides is slow, and vice versa, which results in low round-trip efficiency [48]. Notably, stage IV presents the highest polarization compared to the other stages, which can be ascribed to the slowest reaction kinetics caused by the



Fig. 2 Structure and challenges of RT Na-S batteries. Inset: reaction mechanisms of RT Na-S batteries

poor conductivity of solid Na<sub>2</sub>S<sub>2</sub> and Na<sub>2</sub>S. Fourth, the shuttle effect of the soluble sodium polysulfides ( $Na_2S_8$ ,  $Na_2S_6$ and  $Na_2S_4$ ) in conventional liquid electrolytes induces low Coulombic efficiency, high self-discharge and poor cycling performance. These long-chain sodium polysulfide intermediates can easily move across the traditional separators (e.g., polypropylene microporous membrane, glass fiber) to react with the sodium metal anode and form insoluble Na<sub>2</sub>S<sub>2</sub> and Na<sub>2</sub>S on the Na anode surface. The precipitates corrode the anode surface and cause the loss of both anode and cathode active materials [13]. Finally, sodium dendrite growth is a common and fundamental problem for all Na metal batteries [49]. These Na dendrites can cause continuous breaking and reconstruction of the SEI, which reduces the Coulombic efficiency of RT Na-S batteries. In addition, they can also penetrate the separator and lead to short-circuit of batteries and other associated safety issues [13].

The electrochemical reaction mechanism of Na–Se batteries is similar to that of RT Na–S batteries, with the formation of sodium polyselenides as intermediate products and sodium selenide (Na<sub>2</sub>Se) as the final discharge product. Likewise, the inherently low electronic conductivity and shuttling of high-order polyselenides (such as Na<sub>2</sub>Se<sub>4</sub>, Na<sub>2</sub>Se<sub>6</sub> and Na<sub>2</sub>Se<sub>8</sub>) lead to poor electrochemical performance. The high similarity of RT Na–S and Na–Se batteries resulted in researchers employing similar strategies to overcome the shortcomings and improve the performance, which allows us to merge these two topics and summarize their recent progress together in the third section.

To date, tremendous efforts have been devoted to overcoming the shortcomings of RT Na–S and Na–Se batteries mentioned above. The massive successful experience in improving the electrochemical performance of Li–S batteries has shed light on the routes for enhancing the performance of RT Na–S and Na–Se batteries. However, the larger radius of Na<sup>+</sup>, higher solubility of long-chain sodium polysulfides and polyselenides, and lower reactivity of Na<sub>2</sub>S and Na<sub>2</sub>Se compared to the Li counterparts cause more challenges in realizing large-scale practical applications.

# 3 Cathode Design for RT Na-S and Na-Se Batteries

Although S and Se are the active cathode materials in the corresponding Na metal batteries, neat S or Se powders deliver poor electrochemical performance due to the limitations summarized in the previous section. Rational cathode material design has been demonstrated to be one of the most effective strategies to increase the conductivity and activity of sulfur-based cathodes, buffer their volume expansion, and alleviate the dissolution of polysulfides. Currently, the adopted conventional strategies include infusing S or Se into a porous carbon architecture, anchoring sulfur onto

functional nanomaterials via physical and chemical adsorption, fabricating organosulfur composites, and integrating S or Se with other metallic (e.g., Mo) or nonmetallic (e.g., O) elements [44, 50]. In addition, developing catholytes consisting of soluble sodium polysulfides can increase the utilization of active sulfur and improve the electrochemical performance, which will be discussed in the electrolyte Sect. 4.1.3.

# 3.1 Porous Carbon-Based Composites

Porous carbon nanoarchitectures are the most popular sulfur and selenium hosts for RT Na–S and Na–Se batteries owing to their low cost and good conductivity. Beyond providing good electronic support for poorly conductive S and Se as well as the desirable final discharge products Na<sub>2</sub>S and Na<sub>2</sub>Se, the porous carbon structure can also offer space to accommodate the volume change and confine the intermediate polysulfides.

#### 3.1.1 Structure Optimization of Porous Carbon

According to the diameter of the pores, porous materials can be divided into three types: macroporous (pore size > 50 nm), mesoporous (pore size of 2–50 nm) and microporous (pore size < 2 nm) structures. The various porous structures have different impacts on the electrochemical performance of RT Na-S and Na-Se batteries. Macroporous and mesoporous architectures can enable a high sulfur loading and promote electrolyte impregnation. However, these pores are too large to effectively suppress the dissolution of sodium polysulfides or polyselenides into the electrolyte. For example, Wang et al. fabricated hierarchical interconnected mesoporous hollow carbon nanospheres (iMCHSs) as cathode hosts for RT Na-S batteries. S was embedded in the mesopores and encapsulated in the inner hollow spaces of carbon spheres, achieving a high sulfur loading of approximately 60 wt% (wt% means the weight percentage) (Fig. 3a) [51]. Although the RT Na-S batteries using S@iMCHS cathodes displayed a significantly improved electrochemical performance (~292 mAh g<sup>-1</sup> after 200 cycles at 100 mA g<sup>-1</sup>) compared to open carbon matrices, such as reduced graphene oxide or mesoporous carbon, self-discharge and capacity decay resulted from dissolution and shuttling of polysulfides, which has yet to be fully addressed.

In contrast, microporous carbon (MPC) materials are efficient in improving the cyclability of sulfur cathodes because the formation of soluble sodium polysulfides can be circumvented due to the micropore confinement. Theoretical calculations reveal that only small  $S_{2-4}$  molecules can stably exist in the micropores owing to space confinement (less than 0.7 nm) [52]. These micropores can also accommodate the formed final discharge product Na<sub>2</sub>S, which enables complete reduction of sulfur and leads to a high capacity. During charging, the in situ formed  $S^{2-}$  species convert to  $S_{2-4}$  molecules rather than  $S_8$  due to the confinement inside the micropores, leading to a good cycling performance (approximately 684 mAh  $g^{-1}$  at 1 C after 2 000 cycles) (Fig. 3b). Xin et al. first proposed and demonstrated this concept by employing a carbon nanotube-supported sulfurcontaining microporous carbon cathode (S/CNT@MPC) for RT Na-S batteries [38]. The electrochemical tests and theoretical simulation revealed a two-step reduction reaction from small sulfur molecules to Na<sub>2</sub>S<sub>2</sub> and Na<sub>2</sub>S occurring at the composite cathode (Fig. 3c). Benefiting from the high conductivity of the CNT core and physical confinement of MPC, the S/CNT@MPC cathode presented a high reversible capacity of 1 148 mAh  $g^{-1}$  and an excellent rate and cycling performance of 580 mAh  $g^{-1}$  at 2 C after 200 cycles. However, the sulfur loading ratio in the MPC host is severely restricted by the micropore volume, and the maximum S loading amount in previous reports was approximately 50 wt% [53]. In addition, the average output voltage of the S/ MPC cathode is low because of the absence of transformation of long-chain polysulfides at high voltage. Both issues remarkably decrease the energy density of RT Na-S batteries using the abovementioned materials. Thus, a hierarchical porous sulfur host that can combine the advantages of different pores and ameliorate their shortcomings will be desirable for practical application.

#### 3.1.2 Defect Engineering of Porous Carbon

Although a rational porous carbon host architecture can alleviate dissolution of polysulfides, the weak physical interaction between nonpolar carbon and sulfur species is insufficient to address the shuttle effect. Doping heteroatoms (e.g., N, S) into the carbon material can increase its polarity, thereby suppressing the polysulfide shuttling owing to the enhanced chemical interaction between the polar host materials and sulfur species [54]. Seh et al. systematically investigated the interactions between various types of N-doped carbon and sulfur species and their influences on the performance of RT Na-S batteries [55]. As shown in Fig. 3d, density functional theory (DFT) calculations were performed to optimize the binding configurations and calculate the binding energies of N groups to different sulfur molecules (S4,  $S_3$  and  $S_2$ ) and sodium (poly)sulfides (Na<sub>2</sub>S<sub>4</sub>, Na<sub>2</sub>S<sub>3</sub>, Na<sub>2</sub>S<sub>2</sub>) and Na<sub>2</sub>S). The results in Fig. 3e show a stronger interaction between N-group-containing carbon and discharged sodium sulfide and polysulfides, while pristine carbon prefers to bind to sulfur in the charged state. The experimental results also demonstrate that the RT Na-S batteries with the highest proportion of pyridinic- and pyrrolic-type nitrogen groups deliver the best cycling performance.



Fig. 3 Transmission electron microscopy (TEM) images and corresponding schematics of **a** iMCHSs and **b** MPC. iMCHS picture: reprinted with permission from Ref. [51]. Copyright © 2016, American Chemical Society. MPC picture: reprinted with permission from Ref. [53]. Copyright © 2017, American Chemical Society. **c** Electrochemical reactions between S/MPC and Na<sup>+</sup> during discharge. Reprinted with permission from Ref. [38]. Copyright © 2014, John Wiley & Sons. **d** Atomic illustration of characteristic nitrogen groups, namely, pyridinic nitrogen (N-PD), pyrrolic nitrogen (N-PL), graphitic nitrogen (N-G), pyridine N-oxide (N-PDO), and pristine graphene (P-G), and optimized binding configurations of sodium sulfide

Dual heteroatom doping or increasing the ratio of heteroatoms can further enhance the chemical adsorption capability of the carbon host. For instance, the adsorption energies of N, S-codoped carbon to sodium sulfide/polysulfides are much more negative than those of carbon, implying stronger

with the corresponding N-doped carbon (where the large yellow ball represents Na, yellow ball represents S, brown ball represents C, blue ball represents N, red ball represents O, and pink ball represents H). **e** Binding energies of sodium sulfide, short-chain sodium polysulfides and sulfur allotropes to various N-doped carbons. Reprinted with permission from Ref. [55]. Copyright © 2021, American Chemical Society. **f** Atomic conformations and adsorption energies for Na<sub>2</sub>S, Na<sub>2</sub>S<sub>2</sub>, and Na<sub>2</sub>S<sub>4</sub> species on S, N-doped carbon (right) and pristine carbon (left). Reprinted with permission from Ref. [56]. Copyright © 2020, John Wiley & Sons

entrapment of these discharge products [56]. Qiang et al. synthesized hierarchical porous carbon with a high doping concentration of 40 atom% (atom% means the atomic percentage) (N, S) for long-cycle-life RT Na–S batteries [57]. Due to the ultrahigh heteroatom concentration, the functional carbon could effectively immobilize sodium polysulfides and prevent side reactions between polysulfides and the carbonate electrolyte. Consequently, the RT Na–S batteries using the as-fabricated cathode maintained a capacity of 378 mAh g<sup>-1</sup> at 230 mA g<sup>-1</sup> after 350 cycles (98% capacity retention) [57]. Dual-doped (N, S, or O doping) porous carbon materials have also been employed as selenium hosts for high-performance Na–Se batteries [28, 58]. DFT calculations demonstrated the strong chemical affinity of dualdoped carbon to sodium polyselenides, which resulted in enhanced cycling performance. Note that extensive heteroatom doping of the carbon host would inevitably sacrifice the electronic conductivity required for fast reaction kinetics. Therefore, an optimum doping concentration is desired to achieve high-performance RT Na–S and Na–Se batteries.

# 3.1.3 Metal Catalyst Decoration of Porous Carbon

Decorating the carbon matrix with polar components, such as metal oxides, metal chalcogenides and metal nanoparticles (NPs) or single atoms (SAs), can also directly improve the chemical adsorption capability of the host for polysulfides owing to the strong polar-polar interactions. However, the interactions are restricted to the polar surface of the sulfur host and Na polysulfides, which cannot efficiently prevent the dissolution of Na polysulfides into the electrolyte, particularly at a high sulfur loading. Due to the poor conductivity of sulfur and sodium polysulfides, the conversion reactions from S species to short-chain sodium polysulfides are sluggish. This means that the intermediate polysulfides formed at the early stage cannot be reduced to short-chain polysulfides in a timely manner, and vice versa, resulting in the dissolution of an increasing amount of long-chain polysulfides into the electrolyte [13]. The use of carbonsupported transition-metal-based catalytic hosts can boost the conversion reaction of long-chain polysulfides to insoluble short-chain polysulfides and thereby fundamentally suppress polysulfide shuttling. For example, Liu et al. delicately designed and synthesized a core-shell nanoarchitecture with multisulfiphilic sites for RT Na–S batteries [59]. The inner cores consist of ZnS and CoS2 embedded in N-doped carbon, which not only provide good electronic conductivity to sulfur but also possess strong adsorption of polysulfides. Meanwhile, CoS<sub>2</sub> and ZnS present good electrocatalytic capability, which speeds up sulfur conversion. Additionally, the outer MPC shell can further physically confine polysulfide dissolution and stabilize the cathode structure. Similarly, Xu's group reported a hollow, polar and catalytic bipyramid prism consisting of carbon CoS<sub>2</sub>/carbon as a cathode host for high-performance RT Na–S batteries (Fig. 4a) [60]. Due to the unique porous structure that can trap sodium polysulfides and accelerate the reduction reaction, the sulfur/  $CoS_2/C$  cathode achieves a high capacity of 675 mAh g<sup>-1</sup> after 800 cycles at 0.5 C, with an ultralow capacity decay rate of 0.012 6%.

Beyond transition-metal chalcogenides, bare transition metal and precious metal NPs/SAs are also employed to modify the carbon host [48, 61, 62]. These metal NPs/SAs are expected to enhance the sulfur conductivity, catalyze the sulfur conversion, and capture polysulfides via chemical bonding. For example, Mou et al. reported metal organic framework (MOF)-derived hierarchical porous nanocubes with self-grown CNTs and Co NPs embedded in a N-doped carbon matrix to efficiently immobilize sulfur species (Fig. 4b) [63]. The Co NPs and N-doped carbon derived from bimetallic zeolitic imidazolate frameworks (ZIFs) have strong polarity and electrocatalytic activity, which significantly promote sulfur redox reactions and suppress polysulfide shuttling.

Zhang et al. fabricated a sulfur host with atomic cobaltdecorated hollow carbon sphere-encapsulated S (S@Co<sub>n</sub>-HC) to enhance the sulfur activity and catalyze the reduction reaction from polysulfides to Na<sub>2</sub>S [64]. As shown in Fig. 4c, the shuttle effects were inhibited owing to the enhanced electrocatalysis of Co and the polar–polar interaction between Co clusters/SAs and polysulfides. As a result, the S@Co<sub>n</sub>-HC cathode presented a significantly enhanced rate and cycling performance compared to the S@HC electrode. Lai et al. developed a general method to synthesize a range of N-doped carbon-supported SAs, such as Fe, Ni, Mn, Ge, Ru, and Pt, and evaluated their electrocatalytic behavior in RT Na–S batteries [65]. Among these candidates, the Fe SA-based sulfur cathode exhibited the best performance for polysulfide conversion and immobilization.

### 3.2 2D Material-Based Composites

Owing to the large active surface area, open channels for fast ion accessibility and good electronic conductivity, twodimensional (2D) nanomaterials such as graphene, MXenes, and layered MOFs have attracted intensive attention as cathode hosts for RT Na–S and Na–Se batteries [66–68]. The 2D materials are generally assembled into porous structures to increase the electronic contact with sulfur and sodium sulfide while simultaneously providing physical and chemical confinement to Na polysulfides and polyselenides [69]. For instance, our group reported a wrinkled S-doped MXene  $(S-Ti_3C_2T_r)$  as the sulfur host and achieved high areal sulfur loading up to  $4.5 \text{ mg cm}^{-2}$  for RT Na–S batteries [70]. S-Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> improved the electronic conductivity of the sulfur species and effectively restricted the diffusion of sodium polysulfides due to its high polarity. The RT Na-S batteries using S-Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub> as the cathode host offered an initial



**Fig. 4 a** Synthesis of bipyramid prism catalytic metal chalcogenide/ C@S composites. Reprinted with permission from Ref. [60]. Copyright © 2020, Nature Publishing Group. **b** Schematic illustration of the construction of S@CNTs/Co@NC. **c** TEM image of CNTs/ Co@NC. Reprinted with permission from Ref. [63]. Copyright ©

2021, John Wiley & Sons. **d** Schematic illustration of the synthesis of  $S@Co_n$ -HS. **e**, **f** High-angle annular dark field (HAADF) images of  $S@Co_n$ -HS showing the Co SAs and clusters dispersed on carbon spheres. Reprinted with permission from Ref. [64]. Copyright © 2018, Nature Publishing Group

capacity of 822 mAh  $g^{-1}$ , which remained at 577 mAh  $g^{-1}$  after 500 cycles at 2 C.

Ye et al. synthesized a 2D Ni-based MOF (Ni-MOF-2D) by exfoliating the bulk precursor and employed it as a sulfur host for RT Na–S batteries [71]. The 2D MOF exhibited strong adsorption and fast conversion kinetics for sodium polysulfides owing to their enhanced interaction. This can be ascribed to the dynamic electron states of Ni centers in the MOF, which cause electron transfer from the cathode to

polysulfides during discharge and boost the sodiation process, and vice versa (Fig. 5a). The high redox capability of Ni centers is associated with the charge redistribution and semiconducting property of the 2D structure (Fig. 5b), which was not found in the bulk counterpart. The resultant RT Na–S batteries presented a stable cycling performance with a capacity of 347 mAh g<sup>-1</sup> after 1 000 cycles (58% capacity retention). The electrocatalytic activity toward boosting the polysulfide redox reactions was also demonstrated in MoS<sub>2</sub>



Fig. 5 a Schematic illustration of sodium polysulfide confinement on Ni-MOF-2D. b High-resolution TEM (HRTEM) image of Ni-MOF-2D and corresponding atomic schematic. Reprinted with permission from Ref. [71]. Copyright © 2020, John Wiley & Sons. c Probable chemical structure of copolymer CS90 obtained by reaction of the Ca monomer with sulfur. d TEM image of the CS90-rGO composite.

nanosheets, which could be tuned by adjusting the sodiation depth [72].

Reprinted with permission from Ref. [75]. Copyright © 2017, American Chemical Society. e Illustration of the synthetic procedures for the composite cathode of  $Na_2S$  nanospheres embedded in a spongy carbon matrix. f TEM (top) and scanning electron microscopy (SEM) (bottom) images of the composite. Reprinted with permission from Ref. [79]. Copyright © 2019, John Wiley & Sons

# 3.3 Polymeric Sulfur Composites

Polymeric sulfur composites are advantageous in suppressing polysulfide shuttling via the formation of covalent bonds between sulfur and the polymer. Wang et al. reported the first polymeric sulfur composite, amorphous sulfur NPs embedded in sulfurized polyacrylonitrile (S-PAN), as the cathode for RT Na–S batteries [73]. The composite was prepared by mixing sublimed S with PAN at 300 °C. During this heat treatment, PAN cyclizes into a heterocyclic structure and is dehydrogenated by S, resulting in the formation of S-PAN. The RT Na-S with S-PAN cathode delivered an initial discharge capacity of ~655 mAh  $g^{-1}$ , which remained at 500 mAh g<sup>-1</sup> after 18 cycles at a current density of 0.1 mA cm<sup>-2</sup> (based on the mass of the composite). Hwang et al. increased the synthetic temperature to 450 °C to obtain a carbonizedpolyacrylonitrile and sulfur composite (S@c-PAN), which improved the electrochemical reaction kinetics due to the good conductivity of carbon and unique 1D porous architecture [74]. Consequently, the RT Na-S with S@c-PAN cathode retained 70% of the capacity after 500 cycles at 1 C. However, the cathode reported by Wang et al. only offered a sulfur ratio of approximately 45 wt%, and that reported by Hwang et al. provided an even lower sulfur loading of approximately 31 wt%, which is a concern in that it limits the energy density of polymeric sulfur composites.

Incorporating sulfur chains into the polymer backbones can prominently improve the sulfur content in polymeric sulfur composites and thus provide a higher capacity in RT Na-S battery systems. Ghosh et al. synthesized a sulfur-rich copolymer with reduced graphene oxide (CS90-rGO) as a cathode, and this type of polymeric material-based cathode offered a high sulfur loading of approximately 90 wt% [75]. CS90 was prepared through thermal ring-opening polymerization of cardanol benzoxazine, which in situ reacted with sulfur (Fig. 5c). The CS90 copolymer enhanced the reversibility of low-order polysulfides due to the plasticizing effects of small organosulfur units formed during deep discharge (the organosulfur units evenly dispersed in the insoluble polysulfides and converted them into a soluble phase). After coupling with high-conductivity rGO (Fig. 5d), the CS90-rGO cathode delivered an initial discharge capacity of 542 mAh  $g^{-1}$ , which remained at 335 mAh  $g^{-1}$  after 50 cycles at 200 mA  $g^{-1}$  [75]. Furthermore, small organosulfur molecules can also be applied as cathode materials to enhance the cycling stability of RT Na-S batteries. Wu et al. used the sulfur-carbon complex (SC-BDSA) prepared by using benzenedisulfonic acid (BDSA) and K<sub>2</sub>SO<sub>4</sub> as the raw materials to form a cathode that effectively suppressed the shuttle effect by confining the sulfur in the bridging bonds  $(C-S_x-C)$  [76]. The RT Na-S battery using the SC-BDSA cathode provided a high initial discharge capacity of 1 050 mAh g<sup>-1</sup>, which was maintained at 750 mAh g<sup>-1</sup> after 200 cycles at 250 mA  $g^{-1}$ .

### 3.4 Other Sulfur-Based Composites

In addition to using traditional sulfur as an active cathode material, sodium sulfide (Na<sub>2</sub>S) can also be employed as the starting cathode material for RT Na-S batteries. Na<sub>2</sub>S in the fully sodiated state shrinks when it is initially desodiated, generating space for subsequent volume expansion during cycling [77]. Therefore, the cathode structural stability is significantly improved compared to using elemental S. However, the insulating property of Na<sub>2</sub>S still requires the aforementioned functional materials as a conductive support and a polysulfide immobilizer. Fan et al. reported an all-solid-state RT Na-S battery by using cast-annealed Na<sub>2</sub>S-Na<sub>3</sub>PS<sub>4</sub>-CMK-3 mesoporous carbon as the cathode, which reduced the interfacial resistance and improved the electrochemical performance [78]. Furthermore, Wang et al. applied hollow Na<sub>2</sub>S nanospheres embedded in a polyvinyl pyrrolidone (PVP)-derived carbon matrix to substitute the sulfur cathode (Fig. 5e, f) [79]. The hollow nanoarchitecture of Na<sub>2</sub>S and its intimate contact with the spongy carbon matrix ensure fast Na<sup>+</sup> and electron transport, which improves the reaction kinetics and utilization of active materials. By pairing this material with a tin-based anode material, a Na metal-free RT Na-S battery was achieved with superior rate capability and cyclability.

Additionally, attention should also be paid to several emerging materials that have superior polysulfide capture ability. For instance, Sun's group first introduced a ferroelectric material (BaTiO<sub>3</sub>) with spontaneous polarization as a cathode additive to trap polysulfides [21]. Mitra et al. deposited a thin sheath of aluminum oxyhydroxide (AlOOH) onto the surface of a S/ carbon composite, and the AlOOH layer efficiently inhibited dissolution of polysulfides due to the structural encapsulation and chemical immobilization via Lewis acid–base interactions (AlOOH is a Lewis acid, and sodium polysulfides are considered to be Lewis bases) [80].

Overall, a great range of sulfur host materials has been reported thus far to tackle the challenges in developing RT Na-S batteries. A desirable cathode material should have the following features: (1) a hierarchical porous architecture to accommodate the volume change of S during discharge/ charge and physically confine polysulfide dissolution; (2) good conductivity to enable electron percolation to the active S and insoluble low-order sodium sulfides; (3) high polarity or chemical adsorption capability to immobilize polysulfides; and (4) excellent electrocatalytic behavior to boost the conversion reactions from long-chain polysulfides to Na<sub>2</sub>S, as well as the desodiation process from Na2S to S molecules. Note that thoroughly addressing all the concerns with one type of material is difficult, and one promising solution is to synergize the merits of different components in the composite cathode and the efforts on electrolytes and metal anodes [81]. The cathode design for Na-Se batteries follows very similar strategies,



◄Fig. 6 a Schematic illustration of the operation of RT Na–S batteries with (left) a 1 M (1 M = 1 mol  $L^{-1}$ ) NaTFSI in a PC electrolyte and (right) a 2 M NaTFSI in PC:FEC (with a volumetric ratio of 1:1) with a 10 mM InI<sub>3</sub> additive electrolyte. **b** Visual observation of Na polysulfide formation in different electrolytes (1#-1 M NaTFSI in PC; 2#-1 M NaTFSI in PC:FEC (with a volumetric ratio of 1:1); 3#-2 M NaTFSI in PC; 4#-2 M NaTFSI in PC:FEC (with a volumetric ratio of 1:1); 5#-2 M NaTFSI in PC:FEC (with a volumetric ratio of 1:1) with 10 mM  $InI_3$ ) with the aging time at 60 °C. Reprinted with permission from Ref. [44]. Copyright © 2018, Nature Publishing Group. c Schematic illustration of the in situ electrochemical treatment (in situ ET) method. The inserted TEM image shows the in situ ET cathode composite with an FEC-containing electrolyte. Reprinted with permission from Ref. [88]. Copyright © 2018, John Wiley & Sons. d Combustion tests of 2 M NaTFSI in EC:DEC (with a volumetric ratio of 1:1) and 2 M NaTFSI in TMP:FEC (with a volumetric ratio of 7:3) electrolytes. Reprinted with permission from Ref. [90]. Copyright © 2019, Elsevier

which will not be separately summarized due to the small number of publications in the field.

# 4 Electrolyte Engineering for RT Na-S and Na-Se Batteries

Although the research on electrolytes for RT Na-S and Na-Se batteries is still at its beginning, understanding the contribution of electrolytes to the electrochemical performance of batteries is essential. As an electron-insulator buffer between the anode and the cathode, the electrolyte is in contact with all the battery components. Ideally, electrolytes should be electrochemically inert to the electrodes, current collectors, and separators, serving only as a medium for ionic mobility. Practically, during the first cycle, the electrolyte decomposes at the surface of the electrodes and forms a passivation layer called the SEI, which prohibits continuous electrolyte decomposition during cycling. The electrolyte should be thermally and electrochemically stable to ensure battery operation over the wide temperature and voltage windows, dictated by the electrode couples. Finally, the cost and environmental impact should be minimized by selecting inexpensive, abundant and eco-friendly chemicals [82]. RT Na-S and Na-Se batteries, as mentioned earlier, still face many challenges, such as polysulfide/polyselenide dissolution and Na dendrite growth. Electrolyte optimization can contribute to overcoming these challenges. Many approaches have been explored to understand and improve the electrolyte composition, such as tuning the composition of liquid electrolytes (e.g., cosolvent, additives, ionic liquids), gelling electrolytes to form quasi-solid-state electrolytes, and developing solid-state electrolytes.

# 4.1 Liquid Electrolyte Optimization for Sulfur Cathodes

Generally, batteries employ liquid electrolytes owing to multiple advantages, such as high ionic conductivity, ease of synthesis, and availability of chemicals. RT Na–S and Na–Se liquid electrolytes consist of one or several Na salts dissolved in a solvent. Additives and cosolvents are frequently added to the electrolyte to enhance the performance of batteries. In the following section, we categorize the latest advancements in liquid electrolytes applied in RT Na–S and Na–Se batteries according to the nature of the solvent, such as carbonate-based electrolytes, ether-based electrolytes, catholytes, ionic liquids and inorganic liquid electrolytes.

#### 4.1.1 Carbonate-Based Electrolytes

Carbonate solvents, including linear carbonates (e.g., diethyl carbonate (DEC), dimethyl carbonate (DMC), ethyl methyl carbonate (EMC)) and cyclic carbonates (e.g., ethylene carbonate (EC), propylene carbonate (PC)), have been primarily employed in Na-based batteries owing to their high electrochemical stability and good solvation properties toward Na salts. Investigation of the interactions between Na<sup>+</sup> cations and carbonate solvents demonstrated that the sodium ion pair dissociation energy is smaller than that of the Li counterpart with the same solvent due to the weaker Lewis acidity of Na<sup>+</sup> [83]. Therefore, the knowledge acquired on Li SEI formation is not directly transferable to Na-based batteries [84]. Furthermore, a Na<sup>+</sup> ion possesses a larger ionic radius (<30%) and a larger coordination shell than a Li<sup>+</sup> ion. Thus, the electrolyte properties, such as viscosity and ionic conductivity, are also different between Li-based carbonate electrolytes and Na-based carbonate electrolytes [85]. Ponrouch et al. explored the properties of variable electrolyte composition with different Na salts (e.g., NaClO<sub>4</sub>, NaPF<sub>6</sub> and sodium bis(trifluoromethylsulfonyl)imide (NaTFSI)) dissolved in several common solvents (e.g., PC, EC, DMC, dimethoxymethane (DME), DEC, tetrahydrofuran (THF) and triglyme) or bisolvent mixtures (e.g., EC:DMC, EC:DME, EC:PC and EC:triglyme) [86]. They discovered that in terms of the electrolyte performance, i.e., ionic conductivity, viscosity and thermal stability, the choice of Na salt had a small influence in comparison with the solvent. Based on a 1 M  $(1 \text{ M}=1 \text{ mol } \text{L}^{-1}) \text{ NaClO}_4$  sodium salt concentration, the ionic conductivity of the as-prepared electrolytes followed the order of DMC < triglyme < EC:DEC < PC < EC:triglyme <EC:PC<EC:DMC<EC:DME, while the viscosity order was DMC < EC:DME < EC:DMC < EC:DEC < EC:triglyme <EC:PC < triglyme < PC. Furthermore, the EC:PC and EC:DMC mixture demonstrated an electrochemical stability window of  $\sim 5$  V, comparable to that of the single DEC

or PC solvent. This result illustrates the synergistic effect of solvents in enhancing the electrolyte properties.

Wang et al. prepared a "cocktail optimized" electrolyte for RT Na-S batteries by introducing an InI<sub>3</sub> additive into a concentrated 2 M NaTFSI in PC:fluoroethylene carbonate (FEC) electrolyte [44]. As schematically illustrated in Fig. 6a, on the cathode side, the dissolution of polysulfides is inhibited by the synergistic effects of the high-concentration Na salt and FEC-rich solvent. In addition, during the charging process, I<sup>-</sup> from the InI<sub>3</sub> additive is oxidized to I<sub>3</sub><sup>-</sup> and subsequently reacts with the irreversible Na<sub>2</sub>S end-product to form Na polysulfides, thereby avoiding Na<sub>2</sub>S deposition on the cathode. On the anode side, increasing salt concentration contributes to decreasing free solvent molecules in the electrolyte. Therefore, the decomposition of the electrolyte at the surface of the electrodes is diminished. Figure 6b visually reveals the absence of polysulfides in the "optimized cocktail" electrolyte (#5) through the conservation of the yellow color from InI<sub>3</sub> after aging for 36 h. However, FEC has also been observed to have a positive effect on inhibiting polysulfide dissolution.

The two electrolyte samples (#2 and #4) containing FEC remained mostly transparent. This observation has also been proven by Zhao et al., who employed a 1 M NaClO<sub>4</sub> in EC:PC with a 5 wt% FEC electrolyte [87]. They discovered that the FEC additive had a positive impact on the cycling performance of the full Na-S battery by creating a strong SEI and allowing an easier transformation of S into Na<sub>2</sub>S. The analogous RT Na-Se batteries demonstrated a similar mechanism. By introducing 3 wt% FEC additive into the electrolyte, the anodic corrosion due to polyselenides was diminished [22]. FEC not only protects the Na anode by creating a strong SEI layer but also prevents polysulfide/polyselenide dissolution in the electrolyte. Lee et al. protected the cathode electrode by exploring an "in situ electrochemical treatment" (in situ ET) method to form an SEI layer on the sulfur-carbon cathode [88]. The in situ ET consisted of a galvanostatic and potentiostatic discharge followed by a galvanostatic and potentiostatic charge of the RT Na-S batteries. Figure 6c shows the mechanism of the in situ formation of a protective layer on the cathode electrode. When 8 wt% FEC is introduced into a 1.5 M NaClO<sub>4</sub> in PC electrolyte, the pretreated cathode displays an SEI layer of approximately 20 nm (SEM images in Fig. 6c). During the in situ ET discharge, the SEI is formed on the cathode by reduction of the FEC-based electrolyte and is maintained after the in situ ET charge process. The in situ ET method and FEC additive benefit the full cell, with a capacity increase of about 120% (from 270 to 590 mAh  $g^{-1}$  at the 50th cycle) and a high Coulombic efficiency of 99.5% over 200 cycles at 2 C, confirming the efficiency of this new in situ ET method.

Carbonate-based electrolytes, although widely employed in batteries, are highly flammable. Therefore, nonflammable

or fire-retardant additives have been introduced into carbonate-based electrolytes to reduce the ignition risk. Phosphatebased species demonstrate good fire-retardant properties owing to their hydrogen radical scavenging ability [11]. Trimethyl phosphate (TMP) was introduced into a 1.0 M  $NaClO_4$  in the EC/PC electrolyte to address the safety concerns of carbonate-based electrolytes in RT Na-S batteries [89]. When 15 wt% TMP was introduced into the electrolyte, the latter exhibited nonflammability, good thermal stability and electrochemical compatibility, allowing RT Na-S batteries to achieve stable capacities of 441 and 177 mAh  $g^{-1}$  for 200 cycles at rates of 1 C and 5 C, respectively. However, with 25 wt% TMP added to the electrolyte, the specific capacity of the batteries significantly declined due to an unstable SEI. An excessive amount of TMP leads to poorer cycling performance due to the lack of compatibility of phosphate compounds with electrodes [89]. Wu et al. prepared a nonflammable electrolyte based on 2 M NaTFSI dissolved in a mixture of TMP/EC (with a volumetric ratio of 7:3) solvents [90]. As shown in Fig. 6d, the electrolyte containing TMP can extinguish the fire, whereas a common carbonate-based electrolyte, such as 2 M NaTFSI in EC/ DEC, burns until all the combustible is consumed. The RT Na-S battery with the electrolyte of 2 M NaTFSI in TMP/ FEC delivered a reversible capacity of 788 mAh g<sup>-1</sup> after 300 cycles at 1 C, with a negligible capacity decay below 0.04% per cycle.

#### 4.1.2 Ether-Based Electrolytes

Ether-based electrolytes were overshadowed by the more successful carbonate-based electrolytes in Li-ion batteries for a long time due to their lack of electrochemical stability at high voltage. In the past decade, ether solvents, particularly the glyme family (e.g., monoglyme, diglyme, triglyme, tetraglyme), have been crucial in the development of electrolytes for Li-S batteries owing to their better compatibility with metal anodes (i.e., dendrite growth prevention) and sulfur composite cathodes [91, 92]. Moreover, long-chain ether solvents such as tetraethylene glycol dimethyl ether (TEGDME) are less flammable than carbonate solvents, reducing safety concerns. For instance, an ether-based electrolyte composed of 1 M NaTFSI salt dissolved in TEGDME resisted direct flame exposure for 30 s. The nonflammable electrolyte possesses a high ionic conductivity of  $3 \times 10^{-3}$  S  $cm^{-1}$  at RT, a high Na-ion transference number ( $t^+ = 0.72$ ), and an electrochemical stability window suitable for RT Na-S battery applications [93].

Inspired by the discoveries generated in Li–S batteries, researchers developed ether-based electrolytes for RT Na–S batteries, where ether compounds also contributed to enhancing the stability of sulfur composite cathodes and Na metal anodes [94, 95]. Ether species contain C–O–C groups, while carbonates possess O-C(=O)-O groups; thus, the electrolyte solvation properties will differ, as will the Na-S chemistry mechanism. Furthermore, the sulfur content in the composite cathode should be over 70% to reach a satisfactory energy density [96]. At such high sulfur loading, sulfur molecules inevitably exist on the surface of the composite cathode; thus, side reactions between sulfur molecules and the electrolyte occur to a greater extent, which results in a low specific capacity and capacity decay. When a common composite cathode is used (CNT/S, ~70 wt% S), an irreversible solutionphase reaction has been demonstrated to occur in carbonate solvents. However, in ether-based electrolytes, sulfur reduction involves several conversion reaction processes with the formation of polysulfide intermediates, which allows higher electrochemical reversibility [97, 98]. Liu et al. also observed that a high-sulfur-loading cathode performed better in ether-based than carbonate-based electrolytes (Fig. 7a) [99]. In the TEGDME electrolyte, when elemental sulfur is deposited on the surface of the composite cathode (i.e., the cathode with high sulfur loading), a "solid-liquid" reaction occurs in which elemental sulfur reacts with Na to form liquid polysulfides (Na<sub>2</sub>S<sub>\*</sub>). In comparison, in carbonate-based electrolytes, side reactions will occur between the surficial sulfur and the electrolytes, leading to low specific capacity and Coulombic efficiency. When elemental sulfur is confined inside the pores of the composite cathode (i.e., the cathode with low sulfur loading), a "solid-solid" reaction, in which elemental sulfur is directly transformed into Na<sub>2</sub>S, occurs in both etherbased and carbonate-based electrolytes (Fig. 7b). However, the soluble liquid polysulfides formed in ether electrolytes contribute to the shuttle effect, which leads to side reactions with the Na metal anode. To address this issue, additives can be introduced into the electrolytes to prevent the dissolution or shuttling of the liquid polysulfides.

Similar to additives in carbonate electrolytes, FEC and NaNO<sub>3</sub> additives demonstrate a positive effect in TEGDMEbased electrolytes (Fig. 7c, d). FEC effectively protects the Na anode by creating a F-rich SEI layer on the anode, while the NaNO<sub>3</sub> additive limits polysulfide dissolution and contributes to the formation of a Na-O-containing SEI on the Na anode, which further inhibits Na<sub>2</sub>S deposition on the anode [99]. Another approach consists of introducing soluble polysulfidophosphate complexes (Na<sub>2</sub>S/P<sub>2</sub>S<sub>5</sub>, with a molar ratio of 1:1) into ether-based electrolytes, which enables higher Coulombic efficiency owing to stronger phosphate-containing anode protection [100].  $P_2S_5$  species have been introduced as an additive into Li-S and RT Na-S batteries owing to their ability to facilitate the dissolution of the insoluble lithium Li<sub>2</sub>S or sodium Na<sub>2</sub>S by forming Li<sub>2</sub>S/P<sub>2</sub>S<sub>5</sub> or Na<sub>2</sub>S/P<sub>2</sub>S<sub>5</sub> complexes, respectively [101]. Although Na<sub>2</sub>S and  $P_2S_5$  are separately insoluble in diglyme, the complex

becomes soluble when they are mixed in a specific ratio, as shown in Fig. 7e. Raman spectroscopy was employed to determine the effect of the  $Na_2S-P_2S_5$  complex on the surface of the Na anode (Fig. 7f). When the Na anode was submerged overnight in a solution of  $Na_2S-P_2S_5$ , a chemical passivation layer was observed on the surface of the anode, which further protected the anode. According to Wang et al., the reaction between the Na–P–S complex and Na forms an amorphous Na–P–S protective layer, which may consist of  $Na_3PS_4$ ,  $Na_2PS_4$ , and  $Na_4PS_6$  species [79].

### 4.1.3 Catholytes

Due to the inevitable polysulfide dissolution in ether-based electrolytes and the low utilization of sulfur in the cathode material, several research groups have focused on developing catholytes for RT Na-S batteries. Catholytes consist of an electrolyte containing the active material, namely, a "liquid-phase cathode", which enables a homogeneous distribution of sulfur in the form of liquid polysulfides in the positive electrode (e.g., a carbon cloth current collector), thus improving the sulfur utilization. Yu et al. investigated the origin of capacity fading in RT Na-S batteries [102]. By employing a Na<sub>2</sub>S<sub>6</sub> in TEGDME catholyte, the first discharge reaction step corresponding to the transformation of elemental sulfur into long-chain sodium polysulfides is suppressed. As a result, the electrochemical reaction mechanism in the lower-voltage-plateau region corresponding to the transformation of long-chain polysulfides to short-chain polysulfides can be more easily examined. The transformation between elemental sulfur and long-chain polysulfides  $(Na_2S_n, 4 \le n \le 8)$  has been demonstrated to be highly reversible, whereas the transformation from Na2S4 into short-chain polysulfides (Na<sub>2</sub>S<sub>n</sub>,  $1 \le n \le 4$ ) remains mostly irreversible mainly due to the deposition of the end-product on the anode, resulting in loss of active material [103].

To avoid side reactions between polysulfides and anodes, catholytes are generally employed with ion-selective membranes such as Na-β"-aluminate or Nafion-like polymers as separators [104, 105]. However, such ion-selective membranes are expensive. Kumar et al. employed a Na<sub>2</sub>S<sub>6</sub> catholyte with a carbon cloth current collector decorated with  $MnO_2$  nanoarrays [106]. This method enabled the reduction of shuttling of polysulfides owing to polysulfide adsorption on MnO2. The NalNa2S6 catholytelMnO2-carbon cloth battery displayed an initial energy density of 946 Wh kg<sup>-1</sup> and an energy density after 500 cycles of 728 Wh kg<sup>-1</sup>. More recently, the same group employed an activated carbon cloth current collector combined with a Na<sub>2</sub>S<sub>6</sub> in TEGDME catholyte and a Na metal anode [107]. They claimed that the activated carbon cloth serves as a catalyst to stabilize freeradical species by enhancing the kinetics of conversion of long-chain polysulfides into small-chain polysulfides  $(Na_2S_2)$ 



**Fig. 7 a** Cycling performance of the cathode with a high sulfur loading at 0.1 A g<sup>-1</sup> based on the mass of sulfur. **b** Schematic illustrations of the mechanisms in ether and carbonate electrolytes for the cathode with a high sulfur loading (sulfur on the surface) and the cathode with a low sulfur loading (sulfur in the pores). **c** Cycling performance of the cathode with a low sulfur loading in a 1.0 M NaClO<sub>4</sub> and 0.2 M NaNO<sub>3</sub> in a TEGDME electrolyte with or without a 5 wt% FEC additive at 1.0 A g<sup>-1</sup>. **d** Cycling performance of the cathode with a high sulfur loading in TEGDME with or without a NaNO<sub>3</sub> additive based on the mass of sulfur. Reprinted with permission from Ref. [99]. Copyright © 2021, Springer. **e** Photo of undissolved Na<sub>2</sub>S and P<sub>2</sub>S<sub>5</sub> as well as the Na<sub>2</sub>S–P<sub>2</sub>S<sub>5</sub> complex. **f** Raman analysis of pre-

passivated Na (surface), the Na<sub>2</sub>S–P<sub>2</sub>S<sub>5</sub> complex, Na<sub>2</sub>S powder, P<sub>2</sub>S<sub>5</sub> powder, and the blank solvent. Reprinted with permission from Ref. [79]. Copyright © 2018, John Wiley & Sons. **g** Proposed impact of activated carbon cloth (ACC) on the reaction mechanism during discharge of Na<sub>2</sub>S<sub>6</sub>. **h** Reaction mechanism describing the catalytic effect of ACC in the Na<sub>2</sub>S<sub>6</sub>@ACC cathode. **i** Second cycle discharge profiles of the Na<sub>2</sub>S<sub>6</sub> catholyte-impregnated ACC cathode (blue) and Na<sub>2</sub>S<sub>6</sub> catholyte-impregnated pristine carbon cloth cathode (red) at 0.2 C. **j** Long-term cycling of the Na<sub>2</sub>S<sub>6</sub> catholyte-impregnated ACC cathode at 0.5 C. Reprinted with permission from Ref. [107]. Copyright © 2020, American Chemical Society

through a two-step mechanism (Fig. 7g, h). This was further confirmed by the appearance of three plateaus on the discharge curve profile of the NalNa<sub>2</sub>S<sub>6</sub> catholytelactivated carbon cloth battery, whereas the discharge curve profile of the NalNa<sub>2</sub>S<sub>6</sub> catholytelpristine carbon cloth battery displayed the usual two plateaus corresponding to the two reaction steps of the regular Na–S chemistry (Fig. 7i). The catholyte combined with the effective catalysis of the activated carbon cloth permitted achievement of long cycling stability with low specific capacity fading (Fig. 7j).

Ether compounds are preferentially used as solvents to synthesize catholytes owing to the facile solubility of longchain polysulfides. However, Medenbach et al. innovated this approach by employing tetramethylurea as a solvent with a mixture of 0.5 M NaOTf and 1 mM Na<sub>2</sub>S<sub>5</sub> sodium salts and a  $P_2S_5$  additive [104]. This novel catholyte could achieve a specific capacity of approximately 800 mAh  $g_8^{-1}$ over 30 cycles at 0.1 C, without any notable fading. Only a limited number of research articles have been published about catholyte electrolytes for Na-S or Na-Se batteries. Additional investigations on the catholyte composition, such as studying other solvents, cosolvents and additives, could significantly improve the battery cycling performance by achieving better protection of the Na anode to suppress side reactions between short-chain polysulfide species and the Na anode.

#### 4.1.4 Ionic Liquids

Ionic liquids (ILs) are molten salts with a melting point below 100 °C. Recently, they have engendered enormous research owing to their advantageous solvent properties, such as ultralow volatility, nontoxicity, thermal and electrochemical stability, and intrinsic high ionic conductivity. In addition, ILs that are composed of only cationic and anionic species (i.e., no solvent) can be tailored to meet specific requirements [108]. Among the many ILs that have been discovered, only a restricted number can be applied as electrolytes for battery applications [109]. Typically, ILs employed as electrolytes in batteries are made of bulky ions such as imidazolium, pyrrolidinium or phosphonium cations and fluorinated anions such as  $TFSI^-$ ,  $PF_6^-$ , and  $FSI^-$  [110]. ILs have been used as solvents, cosolvents or additives in Na-ion and Li-based batteries; however, fewer than a handful of studies have been reported for Na-S/Se batteries. An IL based on NaTFSI dissolved in N,N-diethyl-N-methyl-N-(2methoxyethyl) ammonium bis(trifluoromethanesulfonyl) amide (DEME TFSA) was employed in RT Na-S batteries, which achieved an initial capacity of 700 mAh  $g^{-1}$  with Coulombic efficiency > 95% [111]. Although the as-prepared IL demonstrated low polysulfide dissolution, capacity fading was observed during cycling, which has been attributed to the degradation of the cathode material resulting from the repetitive volume change arising from elemental sulfur conversion into Na<sub>2</sub>S.

More recently, Wei et al. introduced a 1-methyl-3-propylimidazolium-chlorate IL tethered to SiO<sub>2</sub> NPs into a 1 M NaClO<sub>4</sub> in an EC/PC liquid carbonate electrolyte [112]. The combination of the IL and  $SiO_2$  NPs stabilized the Na electrodeposition on the Na anode, thus allowing smooth Na plating/stripping processes. Furthermore, the solubility of polysulfides was extremely reduced. The introduction of an IL/SiO2 additive was clearly beneficial in terms of the specific capacity and Coulombic efficiency. Interestingly, when the percentage of IL/SiO<sub>2</sub> additive was increased from 5 wt% to 10 wt%, the battery delivered a smaller specific capacity due to the increase in viscosity. Very recently, Wang et al. introduced an IL into a  $\beta''$ -alumina solid electrolyte [113]. The IL provided high ionic conductivity, thermal and chemical stability and a wide electrochemical stability window. The dual IL-solid electrolyte enabled Na-S battery operation at an IMT of 150 °C.

#### 4.1.5 Liquid Inorganic Electrolytes

Inorganic electrolytes generally consist of liquid inorganic solvents (e.g., ammonia, SO<sub>2</sub>) in which Na salts are dissolved at high concentrations. This family of electrolytes is close to ILs. An ammonia solvate benefits high Na salt concentration (<7 M), high ionic conductivity (~ $100 \times 10^{-3}$ S cm<sup>-1</sup>) and safety (low flammability) [114]. Ruiz-Martínez et al. reported three ammoniate-based electrolytes for Na metal batteries, including NaI·3.3NH<sub>3</sub>, NaBH<sub>4</sub>·1.5NH<sub>3</sub> and  $NaBF_4 \cdot 2.5NH_3$  [115]. The ionic conductivities of these electrolytes followed the trend in the sodium concentrations (i.e.,  $NaBH_4 \cdot 1.5NH_3 > NaBF_4 \cdot 2.5NH_3 > NaI \cdot 3.3NH_3$ ). Moreover, NaBH<sub>4</sub> and NaBF<sub>4</sub> are less expensive than NaI salts. Regarding the cycling performance, the NallNa symmetric cells are promising, especially that with  $NaBF_4 \cdot 2.5NH_3$ , as the voltage hysteresis remains lower than that of the two other electrolytes. However, to further attest to the cycling performance of these electrolytes, the full cell configuration has yet to be tested. Another liquid inorganic electrolyte, more precisely a catholyte, based on SO<sub>2</sub>-solvated chloroaluminate (NaAlCl<sub>4</sub>·2SO<sub>2</sub>) was recently proposed and applied in RT Na–SO<sub>2</sub> batteries [116].

# 4.2 Liquid Electrolyte Engineering for Na Metal Anodes

The quest to find a suitable liquid electrolyte is essential for the proper operation of Na metal anodes since the electrolyte is in direct contact with the anodes and regulates the



reactions occurring at the anode/electrolyte interface. Extensive efforts in the research community have been devoted to optimizing the composition of liquid electrolytes. The nature of the solvents (e.g., carbonate, ether, IL), salts (e.g., organic, inorganic) and additives (e.g., contributing to SEI layer formation, dendrite suppression, fire extinguishment) employed in electrolytes is acknowledged to have significant impacts on the performance of Na metal batteries. **∢Fig. 8 a**, **b** In situ optical microscopy images of the gas evolution on Na in pure PC solvent and the Na<sup>+</sup>-PC solution using frontier molecular orbital theory analysis. c Frontier molecular orbital levels of PC (PC as a single molecule), PC+Sol (PC as a solvent), [NaPC] (Na atom-PC complex) and [NaPC]+ (Na+-PC complex). The red and green regions represent the positive and negative parts of the LUMO and highest occupied molecular orbital (HOMO) wave functions, respectively (isovalue: 0.02). The hydrogen, lithium, carbon, and oxygen atoms are marked with white, purple, gray, and red, respectively. d and e Schematics of the orbital hybridization between PC and a Na<sup>+</sup> ion or a Na atom. Reprinted with permission from Ref. [119]. Copyright © 2017, John Wiley & Sons. f Coulombic efficiencies of Na plating and stripping using 1 M NaPF<sub>6</sub> in various electrolyte solvents (the upper panel) and 1 M of various Na salts in diglyme (the lower panel). Reprinted with permission from Ref. [123]. Copyright © 2015, American Chemical Society. h Total gas amount generated in different electrolytes during cycling in NalNa cells. The gas amount was calculated from the first three plating/stripping cycles and is normalized with respect to the surface area of the copper mesh on which Na was plated. Reprinted with permission from Ref. [124]. Copyright © 2019, American Chemical Society

### 4.2.1 Solvent Selection

Based on the knowledge acquired on electrolytes applied in Li metal anodes, carbonate electrolytes have been investigated in Na metal anodes. However, several researchers have proven that Na anodes generally present poor electrochemical performance in carbonate-based electrolytes [117, 118]. This observation could result from the unstable SEI layer formed on Na metal anodes; however, notably, the stability of Na metal anodes in carbonate electrolytes can be improved by introducing electrolyte additives, which will be discussed in the next section. Recent studies demonstrated that vigorous decomposition of the electrolyte at the electrode surface could produce gases, which disturbed the formation of a uniform SEI. Chen et al. investigated the impact of ion-solvent complexes on the gas evolution in carbonate electrolytes [119]. As observed with in situ optical microscopy (Fig. 8a, b), the gas evolution is approximately tenfold higher in a 1.0 M NaClO<sub>4</sub> in a PC electrolyte (denoted as the Na<sup>+</sup>-PC solution, Fig. 8b) than that in pure PC solvent (Fig. 8a), attesting to more violent decomposition of the electrolyte, which can only be ascribed to the presence of Na<sup>+</sup>-PC complexes. This phenomenon was explained with frontier molecular orbital theory analysis by comparing the lowest unoccupied molecular orbital (LUMO) energy levels of PC as a single molecule, PC as a solvent (PC + Sol), the Na atom-PC complex ([NaPC]) and the Na<sup>+</sup>-PC complex ([NaPC]<sup>+</sup>).

As seen in Fig. 8c, when a PC molecule forms a complex with a Na atom, the LUMO energy level rises; however, when a PC molecule complexes with a Na<sup>+</sup> ion, the LUMO energy level sharply decreases. This explains the poorer stability of ion complexes against Na metal than that of the pure solvent. The difference in energy levels was further explained with schematics of the orbital hybridization within Na<sup>+</sup>-PC complexes (Fig. 8d) and Na atom-PC complexes (Fig. 8e). However, the impact of the anions was not taken into consideration. Other publications believe that the stability of the Na metal anode also depends on the anions present in the electrolyte since metal corrosion is generally caused by the anions [120].

Similar to Li metal anodes, ether-based electrolytes seem to be preferential for Na metal stabilization. Na<sup>+</sup> ion solvation was studied in common carbonate solvents (e.g., PC, DEC, DMC and EMC) and ether solvents (e.g., 1,2-dioxolane (DOL), THF and DME) by DFT, which established that Na<sup>+</sup> cations are more easily solvated by carbonate solvents than by ether solvents [121]. Although ether solvents are usually less polar and have a lower dielectric constant, when combined with Na salts, the ionic conductivity and viscosity of such electrolytes remain satisfactory for battery applications [122]. Wei Seh et al. investigated the reversibility of Na plating and stripping processes in both carbonate- and ether-based electrolytes [123]. Two carbonate-based electrolytes (1 M NaPF<sub>6</sub> in EC:DEC (with a volumetric ratio of 1:1) and 1 M NaPF<sub>6</sub> in EC:DMC (with a volumetric ratio of 1:1)) and three ether-based electrolytes (1 M NaPF<sub>6</sub> in diglyme, 1 M NaPF<sub>6</sub> in monoglyme and 1 M NaPF<sub>6</sub> in tetraglyme) were synthesized. The average Coulombic efficiency of Na metal plating and stripping after 300 cycles was approximately 99.9% in the ether-based electrolytes, whereas the cells with carbonate-based electrolytes could not surpass 20% Coulombic efficiency for less than 25 cycles. This confirms the excellent stability of ether-based electrolytes when in contact with Na metal anodes (Fig. 8f).

Furthermore, these authors also studied the influence of the nature of Na salts on the Na metal anode in ether solvents. According to their report, the Na salt stability against Na anodes can be ordered as  $NaPF_6 > NaSO_3CF_3$  $(NaOTF) > NaN(SO_2F)_2 (NaFSI) > NaClO_4 > NaN(SO_2CF_3)_2$ (NaTFSI) (Fig. 8g). The high stability of the Na anode in the NaPF<sub>6</sub> in glyme electrolytes arose from the formation of a stable SEI layer consisting of Na2O and NaF species. Later, Goktas et al. confirmed that NaPF<sub>6</sub> and NaOTf (NaSO<sub>3</sub>CF<sub>3</sub>) present better compatibility with Na metal anodes in diglyme electrolytes by studying the gas evolution during cycling in NallNa symmetric cells (Fig. 8h) [124]. As previously mentioned, the lower the gas release is, the higher the stability of the electrolyte during Na plating and stripping processes. Apart from NaPF<sub>6</sub> and NaOTf salts, NaBF<sub>4</sub> in an ether electrolyte also allowed reversible Na plating and stripping cycles owing to the formation of a dense and amorphous SEI layer on the Na metal anode [125]. Wang et al. further explored the effect of NaBF4 in different glyme electrolytes (e.g., diglyme, triglyme and tetraglyme) [126]. When diglyme was employed as the solvent, the electrolyte displayed superior properties, such as higher ionic conductivity,



**Fig.9** Anionic interfacial model describing the interface between electrolytes and sodium metal anodes. Model of a DME-based electrolyte using different metal salts, including **a** 1.0 M NaClO<sub>4</sub>, **c** 1.0 M NaCF<sub>3</sub>SO<sub>3</sub>, and **d** 1.0 M NaPF<sub>6</sub>. **b** Model of a 1.0 M NaClO<sub>4</sub> in a PC

electrolyte. **e** Model of a 1.0 M NaPF<sub>6</sub> in an EC/DEC electrolyte. **f** Model of a 1.0 M NaPF<sub>6</sub> in a PC electrolyte. Reprinted with permission from Ref. [120]. Copyright © 2020, American Chemical Society

lower interfacial resistance, and thinner SEI containing more B-O and NaF species, to the longer-chain glyme electrolytes (e.g., triglyme or tetraglyme).

To better understand the impact of Na salts and solvents on the plating and stripping processes of Na metal anodes, Zhou et al. extensively investigated the Na<sup>+</sup> solvation structure, with a focus on the nature and location of the anions [120]. Similar to previous results, they confirmed that both the solvents and Na salts of the electrolytes dictated the reversibility of Na plating and stripping. Furthermore, they found that the location of the anion in the solvation shell played a particular role due to the difference in the binding energy of different Na<sup>+</sup>-solvent interactions, as schematically illustrated in Fig. 9. Two aspects are essential to the bonding interaction: the first is the degree of freedom of the anion (i.e., low steric hindrance), and the second is the strength of the Na<sup>+</sup>-solvent and Na<sup>+</sup>-anion interactions. For instance,  $ClO_4^-$  with a high degree of freedom and  $CF_3SO_3^-$  with a strong interaction with Na<sup>+</sup> ions can easily migrate close to the Na metal anode regardless of the nature of the solvent, thereby causing side reactions (Fig. 9a-c). In contrast,  $PF_6^-$  stays away from the Na metal anode when DME is employed as the solvent owing to the strong bonding interaction between DME molecules and Na<sup>+</sup> ions (Fig. 9d). However, the bonding interactions between EC, DEC and PC solvents and Na<sup>+</sup> ions are not strong enough to prevent  $PF_6^-$  from contacting the Na metal anode, resulting in low Coulombic efficiency (Fig. 9e, f).

#### 4.2.2 Salt Concentration Optimization

As discussed above, the solvation structure is one of the main driving factors for high-efficiency Na plating and stripping. Increasing the salt concentration in the electrolyte dramatically changes the solvation shell of the cation owing to fewer free solvent molecules in the solution, which results in many specific electrochemical and physical properties. Several studies have investigated the solvation structure of NaFSI dissolved in several glymes (e.g., monoglyme, diglyme and triglyme) at different concentrations [127, 128]. As shown in Fig. 10a, in the dilute electrolyte, NaFSI is totally dissociated into the solvent, forming solvent-separated ion pairs (SSIPs). As the concentration of NaFSI increases, contact ion pairs (CIPs) and aggregates (AGGs) are formed due to the lack of free solvent molecules in the solution. Lee et al. further applied a highly concentrated ether-based electrolyte of 5 M NaFSI in DME in Na metal batteries [129]. Similar to previous reports, the increase in the NaFSI concentration in DME gave rise to strong interactions between Na<sup>+</sup> ions and DME solvent, as confirmed by the appearance of a new peak at 1 085 cm<sup>-1</sup> in the Fourier transform infrared (FTIR) spectrum (Fig. 10b). The morphologies of Na deposited on Cu substrates in the electrolytes of 1 M NaPF<sub>6</sub> in EC/PC, 1 M



**Fig. 10 a** Schematic representation of the three main solvated species in NaFSI-triglyme binary mixtures: solvent-separated ion pairs (SSIPs) (left), contact ion pairs (CIP I and CIP II) (middle), and aggregate structures (AGG I and AGG II) (right). Reprinted with permission from Ref. [127]. Copyright © 2018, American Chemical Society. **b** FTIR spectra of *x*M NaFSI-DME (x=0, 1, 2, 3, 4, or 5)

and pure NaFSI salt. **c** Photographs of a pristine Na metal electrode and Cu substrates after the initial Na plating at a rate of C/10. The utilized capacity of the Na metal electrode was 5.0 mAh. Reprinted with permission from Ref. [129]. Copyright © 2017, American Chemical Society.

NaFSI in DME, and ultraconcentrated 5 M NaFSI in DME were investigated by SEM (Fig. 10c). When the highly concentrated electrolyte (5 M NaFSI in DME) was employed, the Na metal deposition layer was much smoother than that obtained in the dilute electrolytes, attesting to excellent plating/stripping processes. The superior performance of the Na anode in the highly concentrated NaFSI/DME electrolyte was explained by the stable passivation layer generated on the surface of the Na anode, which suppressed electrolyte decomposition [130].

Although increasing the salt concentration is beneficial for stabilizing Na metal anodes, it also results in high viscosity and low ionic conductivity, which causes large polarization and limited performance in high current rate cycling. Interestingly, the ionic conductivity has been reported to remain high in highly concentrated NaFSI/glyme electrolytes owing to two different charge transport mechanisms (Fig. 11a) [131]. The first mechanism is mobility through diffusion of free ions, which is found in electrolytes with long-chain glymes (e.g., triglyme and tetraglyme). The second mechanism is based on FSI<sup>-</sup> hopping through making and breaking of ion pairs and aggregates, which is observed in short-chain glymes (e.g., monoglyme and diglyme). Molecular dynamics simulations further confirmed the proposed diffusion and hopping mechanisms. In general, the ionic conductivity and viscosity are intimately linked. Rheology measurements were performed, which revealed that the viscosity of all four NaFSI/glyme electrolytes linearly increased with the salt concentration (Fig. 11b). Thus, the unusual ionic conductivity behavior of highly concentrated electrolytes cannot be only caused by the viscosity properties, which further confirms two different mechanisms related to the chain length of glyme solvents.

Highly concentrated electrolytes provide wide electrochemical windows, high thermal resistance and low volatility. However, they usually suffer from high viscosity, low ionic conductivity and high cost. Recently, a new type of electrolyte, namely, localized highly concentrated electrolytes, has been developed for Li metal anodes to resolve the abovementioned challenges. These electrolytes consist of highly concentrated electrolytes and diluents. The diluent is miscible with the solvent but interacts very weakly with the metal salt (Fig. 11c); therefore, the localized highly concentrated electrolyte preserves the same Na<sup>+</sup> ion solvation structure as the corresponding highly concentrated electrolyte. Meanwhile, the addition of the diluent improves the ionic conductivity and reduces the viscosity and cost of the electrolyte. Zheng et al. introduced bis(2,2,2-trifluoroethyl) ether (BTFE) into a highly concentrated electrolyte of NaFSI in DME [132]. They demonstrated that the as-prepared electrolyte displayed higher ionic conductivity and lower viscosity than its highly concentrated counterpart. Furthermore, similar Coulombic efficiencies of the Na plating and stripping processes on Cu substrates were obtained in localized highly concentrated electrolytes and highly concentrated electrolytes (Fig. 11d). As seen from the X-ray photoelectron spectroscopy (XPS) depth profiling in Fig. 11e, f, the atomic ratios of the SEIs formed on Na metal anodes in highly concentrated electrolytes and localized highly concentrated electrolytes are similar, which proves that the diluent has a negligible impact on the solvation shell of Na<sup>+</sup>. In contrast, the SEI formed in the dilute electrolyte contains fewer fluoride species and more oxygen species, confirming a more fragile SEI. Later, another group reported hydrofluoroether (HFE) solvent as a diluent for the electrolytes of sodium bis(trifluoromethanesulfonyl)amide (NaTFSA) in glymes (tetraglyme and pentaglyme) [133]. They discovered that the HFE solvent did not participate in the Na<sup>+</sup> ion solvation shell, and therefore, the physical properties of the localized electrolyte could be improved. The key challenge for the development of localized highly concentrated electrolytes is to find an appropriate diluent to alleviate the drawbacks faced by highly concentrated electrolytes. A better understanding of the impact of the diluent on SEI formation is also required.

#### 4.2.3 Developing IL Electrolytes for Na Metal Anodes

The ultradiverse available combination of ILs is particularly interesting for the development of liquid electrolytes for rechargeable batteries. Many ILs are safe (e.g., nonflammable, low volatility, low toxicity) and electrochemically and thermally stable and have recently been applied in Na metal batteries to reduce safety risks. However, ILs suffer from high cost, high viscosity and low ionic conductivity due to the strong interaction between the Na<sup>+</sup> ions and the anions, causing the formation of anion aggregates Na[anion], -y. Tuanan et al. theoretically investigated the Na<sup>+</sup> ion transport mechanism of 25 different ILs by molecular dynamics simulations (Fig. 12a) [134]. They demonstrated that the ionic concentration is closely related to the Na<sup>+</sup> ion transport behavior. An increase in the salt concentration resulted in an increase in the size of anion aggregates and facilitated the Na<sup>+</sup> ion hopping diffusion mechanism, thereby improving the Na<sup>+</sup> ion apparent transport number in the system. Furthermore, increasing the concentration of Na salt in ILs allowed more reversible Na metal plating and stripping. This observation was further explored by Forsyth's group, who intensively investigated the influence of the NaFSI salt concentration in an N-methyl-N-propylpyrrolidinium bis(fluorosulfonyl)imide (C<sub>3</sub>mpyrFSI) IL electrolyte [135]. They employed radial distribution function analysis to simulate the molecular structure evolution when the NaFSI concentration increases. If the concentration of Na salt becomes higher than that of the IL, then large and interconnected Na<sup>+</sup>-anion aggregates are formed. Meanwhile, an increased coordination number of the Na<sup>+</sup> ion solvation sheath with the FSI anion makes the denticity equal to one. As a result, the Na<sup>+</sup> ion diffusion dynamics increase, while the mobility of the anions and cations of the IL decreases (Fig. 12b). When applied with a Na metal anode, the superconcentrated NaFSI/C<sub>3</sub>mpyrFSI electrolyte (i.e., 50 mol% (mol% means the molar percentage) NaFSI) demonstrated a highly aggregated  $Na_r(FSI)_v$  structure near the anode, which excluded the  $[C_3mpyr]^+$  cation from the interface [136]. Therefore, such electrolytes could benefit the formation of a stable SEI and enhance the Na metal plating and stripping processes. By applying a high-voltage pretreatment, an increased concentration of aggregates was observed near the anode, thereby achieving better cycling performance. The effect of the Na salt concentration was also reported with other systems, which confirmed the benefit of increasing the salt concentration in ILs [137]. However, increasing the NaFSI salt concentration results in a sharp increase in the cost of the electrolyte. The high cost of ILs can be alleviated by



**Fig. 11 a** Representation of hopping and shuttling molecular mechanisms. **b** Conductivity (the upper panel) and viscosity (the lower panel) as a function of the molality of sodium triflate in different glymes: monoglyme (black), diglyme (red), triglyme (green), and tetraglyme (blue). Reprinted with permission from Ref. [131]. Copyright © 2018, American Chemical Society. **c** Schematic illustration of dilution from a highly concentrated electrolyte (HCE) to a localized highly concentrated electrolyte (LHCE). **d** Coulombic efficiency

of Na metal plating/stripping on Cu current collectors at 1 mA cm<sup>-2</sup> after two formation cycles at 0.2 mA cm<sup>-2</sup> with an areal capacity of 1 mAh cm<sup>-2</sup>. e Atomic ratios of C/N/O/F/Na/S elements detected at the surface (0 nm, the upper panel) and 10 nm depth (the lower panel) of the SEI layer after the 10th stripping of NallCu cells using a dilute electrolyte (1.7 M NaFSI/DME), an HCE (5.2 M NaFSI/DME), and an LHCE (2.1 M NaFSI/DME-BTFE (1:2)). Reprinted with permission from Ref. [132]. Copyright © 2018, American Chemical Society



◄Fig. 12 a Atomic structures of ILs obtained by DFT calculations. The C, O, H, S, F, and N atoms are indicated by silver, red, white, yellow, gold, and blue spheres, respectively. Reprinted with permission from Ref. [134]. Copyright © 2021, American Chemical Society. b Mean square displacements of N from both the IL cation and anion (the left panel) and the Na<sup>+</sup> ion (the right panel) at 393 K for several salt concentrations. Reprinted with permission from Ref. [135]. Copyright © 2018, American Chemical Society. c <sup>23</sup>Na MAS-NMR spectra of the SEI products on the surface of Na metal (the left panel), and high-resolution F 1s and S 2p XPS curves of the plated Na surfaces comparing the SEI composition on plated Na electrodes for dry and wet electrolytes after 100 cycles at a current density of 1.0 mA cm<sup>-2</sup> and 1 h polarization (the right panel). Reprinted with permission from Ref. [140]. Copyright © 2021, American Chemical Society

employing the dicyanamide (DCA) family of ILs, such as the methylpropyl pyrrolidinium ([ $C_3$ mpyr]) DCA IL [138]. Several fluorinated Na salts (NaFSI, NaTFSI and NaFTFSI) or nonfluorinated salts (NaDCA) were introduced at high concentrations into DCA ILs. The battery performance followed the trend of NaTFSI < NaFTFSI < NaDCA < NaFSI owing to the improved SEI formation ability of FSI anions. In addition, several reports demonstrated that less expensive IL electrolytes could be formulated by introducing additives.

In most nonaqueous electrolytes, the presence of water degrades the electrochemical performance of alkali metal batteries. Furthermore, Na metal reacts extremely violently with water. However, several publications affirm the beneficial effect of an extremely small amount of water added into ILs [139–141]. Ferdousi et al. investigated the Na metal stability in a 50 mol% NaFSI-[C<sub>3</sub>mpyrFSI] IL electrolyte with different water concentrations (up to 5 000  $\mu$ mol mol<sup>-1</sup>)] [139]. The elemental characterization of the Na metal anode after cycling in these electrolytes demonstrated the formation of an SEI containing NaF, Na<sub>2</sub>O and FSI-related species. The benefit of scarce water molecules in the electrolyte for SEI formation may be related to additional bond formation between water molecules, FSI anions, and Na<sup>+</sup> ions, along with the formation of other inorganic compounds, as demonstrated by magicangle spinning nuclear magnetic resonance (MAS-NMR) and XPS analysis (Fig. 12c).

Sun et al. developed for the first time a nonflammable chloroaluminate IL-based electrolyte for Na metal batteries (Fig. 13a, b) [142]. They employed an aluminum chloride/1-methyl-3-ethylimidazolium chloride/sodium chloride IL, into which two additives were introduced, namely, ethylaluminum dichloride and 1-ethyl-3-methylimidazolium bis(fluorosulfonyl)imide. Such an electrolyte composition contributes to high ionic conductivity (~ $9.2 \times 10^{-3}$  S cm<sup>-1</sup> at 25 °C) and robust SEI owing to the passivation effect from chloroaluminate molecules and the presence of a small amount of FSI anions. The synergistic effect of chloroaluminate and FSI species enables NallNVP@rGO and NallNVPF@rGO to achieve long-term cycling stability even at high current rates (Fig. 13c).

Other approaches consisted of introducing cosolvents into IL-based electrolytes. A carbonate-based cosolvent (i.e., a mixture of EC:PC) was introduced into an  $C_3$ mpyrTFSI IL containing 1 M NaFSI Na salt [143]. Later, Hu et al. introduced an ether cosolvent, namely, diglyme, into a 1-butyl-3-methylimidazolium tetrafluoroborate IL (vol. ratio 4:1) with 1 M NaPF<sub>6</sub> salt [144]. As seen in Fig. 13d, this hybrid electrolyte can stabilize the Na metal anode at a low temperature (i.e.,  $-40 \,^{\circ}$ C) in lean electrolyte conditions (i.e.,  $1.0 \,\mu$ L mAh<sup>-1</sup>), which contributes to achieving high energy density. Furthermore, the ether IL possesses a high ionic conductivity of  $42 \times 10^{-3}$ S cm<sup>-1</sup>, good desolvation and an SEI formation capability, which allowed stable cycling of NallNVP batteries at RT and at  $-20 \,^{\circ}$ C.

### 4.2.4 Employing Additives in an Electrolyte for Na Metal Anodes

Additives are chemical compounds that are introduced in a small amount (generally < 10%) into the electrolyte with the purpose of stabilizing the electrode/electrolyte interface through the formation of a robust SEI layer. During the first charge process, additives react and protect the Na anode, thereby avoiding further electrolyte decomposition. In particular, fluorinated additives such as FEC assist in the creation of a strong SEI layer on Li metal anodes. Therefore, many publications have focused on understanding the impact of such additives on Na metal anodes. The morphologies and gas evolution during sodium electrodeposition in NallNa batteries were investigated with several carbonate-based electrolyte systems (e.g., PC, EC, DEC) [145]. As previously discussed in Sect. 4.2.1, carbonate-based electrolytes generate a large amount of gas with poor Coulombic efficiency of Na plating and stripping. Upon FEC addition, the gassing is reduced. Meanwhile, the cycling performance of the NallNa symmetric cell is improved, even though the morphology of the SEI remains porous. To maximize the benefits, the concentration of FEC in the electrolyte should be carefully controlled, as a small amount of FEC in PC-based electrolytes improves the Na battery performance. In contrast, an increase in the concentration of the fluorinated additive has an adverse impact [146].

Recently, Han et al. employed sophisticated cryo-TEM analysis to carefully observe the SEI composition and morphology on Na metal anodes in carbonate electrolytes (NaPF<sub>6</sub> in EC/DMC) with and without an FEC additive [147]. When FEC is absent, the SEI layer is mainly composed of Na<sub>2</sub>CO<sub>3</sub> and Na<sub>3</sub>PO<sub>4</sub>, which cannot efficiently protect the Na metal anode. The electrolyte continuously reacts with the anode upon cycling, as schematically illustrated in



**Fig. 13 a** Schematic illustration of the battery configuration and electrolyte composition of IL electrolytes. **b** Flammability tests using a buffered IL (the left picture) and conventional electrolytes (1.0 M NaClO<sub>4</sub> in EC:DEC with 5 wt% FEC) (the right panel). Scale bars = 1 cm. **c** Rate performance (at various current densities from 20 to 500 mA g<sup>-1</sup>) and cycling stability (current density of 150 mA g<sup>-1</sup>) of a NalNVP@rGO cell using a buffered IL electrolyte (the left panel), and cycling stability of a NalNVPF@rGO cell using a buff-

Fig. 14a. In contrast, the cryo-TEM images of the anodes retrieved from the cells containing the FEC additive displayed a compact SEI layer composed of a  $Na_3PO_4$  inner layer and an amorphous NaF-rich outer layer (Fig. 14b). The NaF species in the outer layer of the SEI prevent dissolution of  $Na_3PO_4$  and other amorphous compounds, avoiding direct contact of fresh Na metal with the electrolytes,

ered IL electrolyte at 300 mA g<sup>-1</sup> (the right panel). Reprinted with permission from Ref. [142]. Copyright © 2019, Nature. **d** Electrochemical performance of NallNa symmetric cells at various low temperatures with a cycling capacity of 1 mAh cm<sup>-2</sup> at 0.5 mA cm<sup>-2</sup> (the left panel), and voltage–time profiles of NallNa symmetric cells with a cycling capacity of 50 mAh cm<sup>-2</sup> at 2 mA cm<sup>-2</sup> (the right panel). Reprinted with permission from Ref. [144]. Copyright © 2020, John Wiley & Sons

which results in effective protection of the Na metal anode. Reducing the solubility of the SEI species in the electrolyte is more important in Na-based batteries than in Li-based batteries. Fluorinated additives, such as FEC or difluoroethylene carbonate (DFEC), were confirmed to be able to similarly limit, to some extent, the dissolution of the SEI layer [148]. Zhu et al. proposed a nonflammable electrolyte



**Fig. 14 a** Schematic of SEI formation in the FEC-free EC/DMC electrolyte in the 1st cycle and after 10 cycles, and the corresponding cryo-TEM image after 10 cycles. **b** Schematic of SEI formation in the EC/DMC/FEC electrolyte in the 1st cycle and after 10 cycles, and the corresponding cryo-TEM image after 10 cycles. Reprinted with permission from Ref. [147]. Copyright © 2021, Nature. **c** Flammability tests for the EC/PC carbonate electrolyte (the red panel) and TMP/FEC/DTD-E electrolyte (the purple panel). High-resolution F 1s, P 2p and S 2p XPS depth profiling of the Na metal anode retrieved from cells after 50 cycles in **d** TMP/FEC/DTD-E and **e** 

for Na metal batteries based on TMP solvent with FEC and 1,3,2-dioxathiolane 2,2-dioxide (DTD) additives (Fig. 14c) [149]. TMP solvent has been previously reported as a non-flammable alternative to flammable carbonate solvents owing to the phosphorous functional groups that can scavenge hydrogens [11]. As explained above, FEC facilitates the creation of a strong SEI layer containing NaF species. Meanwhile, the contribution of DTD to the SEI was identified with S-containing compounds (e.g., Na<sub>2</sub>S, Na<sub>2</sub>SO<sub>3</sub> and organic S-containing salt), as observed from the XPS depth profiling spectra (Fig. 14d, e).

TMP/FEC-E electrolytes. Reprinted with permission from Ref. [149]. Copyright © 2021, Elsevier. **f** Atomic structure and binding energy between the Na<sup>+</sup> cation and the SEI components (Na-EDC—sodium ethylene decarbonate; Na-EG—sodium ethylene glycol; Na-DMDT, Na<sub>2</sub>-DMDT<sup>+</sup>, Na<sub>3</sub>-DMDT-CH<sub>3</sub>—organic sodium sulfide salts from the decomposition of TMTD). **g** Nyquist plots of the symmetric cells in the TMTD-added electrolyte (the upper panel) and in the blank electrolyte (the lower panel) after 10, 20 and 40 cycles (red, orange and blue lines, respectively). Reprinted with permission from Ref. [151]. Copyright © 2020, American Chemical Society

Although adding FEC into the electrolyte efficiently enhances the performance of Na metal batteries, a higher polarization of the Na metal anode was observed in halfcells, which encouraged the search for other additives [150]. To reduce the interfacial resistance, an organosulfur compound, namely, tetramethylthiuram disulfide (TMTD), was introduced into 1 M NaPF<sub>6</sub> in EC:PC [151]. The SEI layer formed in this system rich in organic sulfide salts can efficiently protect the Na anode; meanwhile, the Na<sup>+</sup> cation can easily penetrate the SEI layer owing to the lower binding energy between SEI compounds (e.g., sodium



**Fig. 15 a** Simulated electric field intensity distribution and schematic of Li/Na plating mechanisms. Reprinted with permission from Ref. [153]. Copyright © 2018, Wiley–VCH. **b** Schematics of the stabilization effects on the Na metal anode of the cation additives in the NaOTf electrolyte without any additives (the upper left panel), with the TFSI<sup>-</sup> additive (the lower left panel), with the K<sup>+</sup> additive (the upper right panel), and with both the K<sup>+</sup> and TFSI<sup>-</sup> additives (the lower right panel). Reprinted with permission from Ref. [154]. Copyright © 2018, Wiley–VCH. **c** Comparison of the LUMO energy level decreases and binding energy of ion-DME complexes. **d** Calculated and experimental electrode potentials of the considered metal/cat-

dimethyldithiocarbamate (DMDT)) and Na<sup>+</sup> (Fig. 14f, g). Wang et al. introduced  $Na_2S_6$  as an additive into an etherbased electrolyte, which contributed to S-containing SEI layer formation on the Na anode [152]. Interestingly, when the  $Na_2S_6$ -NaNO<sub>3</sub> coadditive was mixed with the electrolyte, the NallNa symmetric battery performance degraded, unlike in the Li metal system.

Another approach of exploring cation additives to stabilize Na metal anodes through electrostatic shielding has been demonstrated to be effective in inhibiting electrolyte decomposition on Na metal anodes. This method has been applied with cations such as  $Li^+$  and  $K^+$ , where the presence of cation additives near the tip of the Na dendrites forces Na<sup>+</sup> ions to deposit on a smoother area of the Na anode

ion pairs. Reprinted with permission from Ref. [155]. Copyright © 2020, Elsevier. e Schematics of SEI formation on a Na metal anode cycled in a regular carbonate electrolyte (the upper panel) and in a SnCl<sub>2</sub>-added electrolyte with the formation of a Na–Sn alloy layer plus a NaCl-rich SEI (the lower panel). f Pictures of Na metal before and after reaction with the SnCl<sub>2</sub>-added carbonate electrolyte. Crosssectional SEM and energy-dispersive X-ray spectroscopy (EDX) mapping images of SnCl<sub>2</sub>-treated Na metal with Na, Sn, and Cl elemental signals. Reprinted with permission from Ref. [156]. Copyright © 2019, American Chemical Society

(Fig. 15a, b) [153, 154]. Recently, Chen et al. reviewed this strategy by proposing three principles for cation additive design [155]. First, the newly formed cation-solvent complex must be stabler than the Na<sup>+</sup>-solvent complex to suppress electrolyte decomposition. Second, the cation additive should present a lower redox potential than that of Na/Na<sup>+</sup> (i.e., avoid reaction with the Na anode). However, the reduction of certain cation additives creates an alloy protective layer on Na metal, which will be discussed in the next paragraph. Third, to ensure suppression of cation additive reduction on the Na metal anode, the cation additives should have a larger binding energy than Na<sup>+</sup>. As shown in Fig. 15c, d, cation additives with these properties are Li<sup>+</sup>, K<sup>+</sup> and

 $Ca^{2+}$ , of which the Li<sup>+</sup> cation additive is the most promising cation additive.

Na metal anodes can also be protected by introducing reduction/alloy-type additives. The formation of an alloy phase on the surface of the Na metal anode was demonstrated to reduce the Na<sup>+</sup> ion diffusion barrier, thereby lowering the interfacial impedance and achieving smoother Na metal plating [156–158]. Zheng et al. introduced SnCl<sub>2</sub> into an EC/PC carbonate electrolyte. The reduction/alloying compound was chemically reduced on the surface of the Na metal anode, forming an SEI with a Sn-containing inner layer and a NaCl-rich outer layer (Fig. 15e, f) [156].

# 4.3 Gel Polymer Electrolytes (Quasi-Solid-State Electrolytes)

Gel polymer electrolytes (GPEs), also known as quasi-solidstate electrolytes (QSEs), are a class of electrolytes with an intermediate phase between liquid and solid. With the application of liquid electrolytes as a plasticizer, a polymer matrix as a mechanical scaffold, and sodium salt and additives, GPEs can minimize the safety risks by locking the liquid solvent inside the polymer matrix to form a quasi-solid-state structure and maintain a high ionic conductivity at the same time [159]. Furthermore, GPEs offer several advantages, including a high concentration of charge carriers, superior interfacial contact between the electrolytes and electrodes, free-standing consistency, and good mechanical properties (e.g., strength, flexibility) [160, 161]. Moreover, GPEs have been reported to be beneficial when applied with Na metal anodes in terms of safety assurance owing to their ability to prevent leakage and reduce Na dendrite growth [162].

However, there are also some challenges that need to be solved for the future practical application of GPEs. For instance, to ensure high ionic conductivity, a high liquid electrolyte content is inevitably required, which will reduce the mechanical strength of the GPE and increase the safety risk due to leakage [161]. Some other challenges, including interfacial instability on the cathode side, poor dimensional stability, and poor liquid retention capacity, are also nonnegligible in GPE development [160]. Exploring new preparation methods and optimizing the composition and structures of GPEs have been demonstrated to be promising pathways to overcome the abovementioned challenges.

In 2004, Tian et al. reported a type of gel electrolytes from a copolymer of polyvinylidene fluoride and hexafluoropropylene (PVDF-HFP), which achieved a good sodium ion conductivity of  $1 \times 10^{-4}$  S cm<sup>-1</sup> at RT [163]. After that, Ryu et al. applied this type of GPE in Li–S batteries, achieving 72% sulfur utilization with a high capacity of 1 206 mAh g<sup>-1</sup> during the first discharge [164]. In 2006, Park et al., for the first time, reported the application of a GPE combined with a sulfur cathode and a Na anode in quasi-solid-state RT Na–S batteries [165]. They used PVDF-HFP comprising a glyme solvent and NaCF<sub>3</sub>SO<sub>3</sub> salt to synthesize a GPE, which offered a good sodium ion conductivity of  $5.1 \times 10^{-4}$ S cm<sup>-1</sup> and provided an initial capacity of 489 mAh g<sup>-1</sup> at RT. However, significant cycling decay could not be avoided, and only 40 mAh g<sup>-1</sup> remained after 20 cycles at 0.144 mAh cm<sup>-2</sup> [165]. The potential of using PVDF-HFPbased GPEs in RT Na–S batteries was further demonstrated by Kim et al. [166]. However, there is a suggestion that the nucleophilic polysulfides decompose the polymer via hydrodefluorination.

Later, many improvement strategies were applied to enhance the performance of GPEs: (1) manipulating the morphology of GPEs by introducing fillers, new types of liquid electrolytes, or novel polymer matrices, (2) in situ formation of GPEs to offer higher binding energy, (3) SEI formation optimization by introducing additives into the liquid part of the electrolyte, (4) ion flux regulation by creating ionic channels in the electrolyte, and (5) fabrication of a robust electrolyte with enhanced mechanical properties (i.e., high Young's modulus) in which dendrite penetration is inhibited.

Kumar et al. optimized the PVDF-HFP GPE by introducing SiO<sub>2</sub> NPs (15 wt%) into it, which induced a more porous morphology (Fig. 16a) and thus offered a better capacity to hold the liquid solvent, free-standing consistency and a highly transparent character [167]. Due to the change in morphology, the ionic conductivity of PVDF-HFP-15% SiO<sub>2</sub> was also significantly higher than that of conventional PVDF-HFP GPEs (Fig. 16b), which provided a satisfactory electrochemical potential window of ~4.0 V (from -2.0 to 2.0 V). However, the Na-S battery with the PVDF-HFP-15% SiO<sub>2</sub> GPE only delivered an initial capacity of ~165 mAh  $g^{-1}$  in the first discharge, which remained at ~21 mAh  $g^{-1}$ after 8 cycles due to the formation of irreversible short-chain Na polysulfides. Kumar et al. subsequently continuously improved the GPE by introducing an IL (1-ethyl-3-methylimidazolium trifluoromethanesulfonate (EMITf)) with EC-PC as a liquid solvent and NaCF<sub>3</sub>SO<sub>3</sub> as a Na salt [160]. The NaCF<sub>3</sub>SO<sub>3</sub>/EC:PC:EMITf/PVDF-HFP gel electrolyte delivered an excellent ionic conductivity of ~ $8.4 \times 10^{-3}$  S cm<sup>-1</sup> at RT and a wide electrochemical stability window (ESW) of ~4.6 V (from – 2.3 to 2.3 V). However, this design can only provide a capacity of ~ 267 mAh  $g^{-1}$  in the first discharge, and rapid cycling decay is still observed after the first charge/discharge cycle.

In addition to using PVDF-HFP as the polymer matrix, other polymer materials, such as polyacrylonitrile (PAN) [168, 169], poly(methyl methacrylate) (PMMA) [170], poly(ethylene glycol) divinyl ether with flame-retardant polysulfonamide (PDE/PSA) [171], poly(vinylene carbonate) and polymeric sodium tartaric acid borate with polyethylene terephthalate (PVCA/PSTB/PET) [172], poly(ethylene



glycol) diglycidyl ether with diamino-poly(propylene oxide) (PEGDE/DPPO) [173], poly(vinyl chloride) (PVC) [174], poly(dimethyldiallylammonium) (polyDADMA) [175], and pentaerythritol tetraacrylate-tris[2(acryloyloxy)ethyl] isocyanurate (PETEA-THEICTA) [176], can also be applied to fabricate GPEs for RT Na–S batteries. Lim et al. immersed a PAN electrospun nanofiber membrane (ES-NFM) into poly(ethylene glycol) dimethyl ether (PEGDME) with sodium triflate (NaCF<sub>3</sub>SO<sub>3</sub>) salt, forming a GPE for use in the RT Na–S battery system [168]. Due to the high porosity of the nanofiber membrane (Fig. 16c), the PAN membrane GPE provided a good ionic conductivity of ~ $6.0 \times 10^{-4}$  S **∢Fig. 16 a** SEM images of the PVDF-HFP GPE compared with PVDF-HFP with 15% SiO<sub>2</sub>. Reprinted with permission from Ref. [167]. Copyright © 2011, Elsevier B.V. b Ionic conductivity of the PVDF-HFP GPE with 0%, 15%, and 25% SiO<sub>2</sub>. Reprinted with permission from Ref. [167]. Copyright © 2011, Elsevier B.V. c SEM image of the porous PAN membrane. Reprinted with permission from Ref. [168]. Copyright © 2018, Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim. d Cycling performance of the Na-S battery with the PAN membrane GPE compared to the Na-S battery with a commercial Whatman<sup>TM</sup> separator. Reprinted with permission from Ref. [168]. Copyright © 2018, Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim. e Schematic of the in situ formation of the cross-linked (PETEA-THEICTA)-based GPE combined with the poly(S-PETEA)@C cathode for the quasi-solid-state RT Na-S battery. Reprinted with permission from Ref. [176]. Copyright © 2018, Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim. f High binding energy between the GPE components (THEICTA monomer and ester group-rich PETEA) and Na<sub>2</sub>S<sub>6</sub>. Reprinted with permission from Ref. [176]. Copyright © 2018, Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim

 $cm^{-1}$  at 30 °C, and the Na–S battery with the GPE electrolyte delivered a much higher capacity and a lower overvoltage polarization than the Na–S battery using the commercial separator (Fig. 16d).

The previous GPEs for RT Na-S batteries were mostly prepared by using the conventional solution casting method [160, 167]. Recently, Zhou et al. prepared a new type of (PETEA-THEICTA)-based GPE by using an in situ radiation-initiated synthesis route. This remarkable GPE used the PETEA and THEICTA monomers combined with the 2-hvdroxy-2-methylpropiophenone (HMPP) photoinitiator dissolved in PC-FEC with 1 M NaTFSI as a precursor solution. The precursor solution was irradiated by ultraviolet (UV) light to trigger C = C bond radical polymerization of the monomers in a glass fiber separator. Subsequently, the (PETEA-THEICTA)-based GPE with a cross-linked structure was formed in situ (Fig. 16e). The formed GPE delivered an excellent ionic conductivity of  $\sim 3.85 \times 10^{-3}$  S cm<sup>-1</sup> at 25 °C, and it could also strongly suppress dissolution and diffusion of Na polysulfides due to the higher binding energy between the inner components (THEICTA monomer and ester group-rich PETEA) and Na<sub>2</sub>S<sub>6</sub> compared to FEC-Na<sub>2</sub>S<sub>6</sub> (-1.22 eV) and PC-Na<sub>2</sub>S<sub>6</sub> (-1.57 eV) (Fig. 16f). By combining this GPE with a novel poly(S-PETEA)@C cathode, the RT Na-S battery system avoided significant capacity fading and offered good cycling stability with a high capacity of 877 mAh  $g^{-1}$  in the first discharge, which remained at 736 mAh  $g^{-1}$  after 100 cycles at 0.1 C [176].

Similar to liquid electrolytes, SEI formation in Na metal battery systems with GPEs can prevent side reactions and dendrite growth. A series of additives have been introduced into the liquid part of GPEs to generate a protective SEI on the Na metal anode [177, 178]. Xu et al. synthesized a multifunctional GPE by an in situ polymerization process for rechargeable dual-ion sodium metal batteries [177]. The electrolyte precursor solution contained 1,3-propanesultone (PS) as an additive in a 0.5 M NaPF<sub>6</sub> in PC/EMC/FEC solvent (1:1:1 by volume) with 1.5 wt% ethoxylated pentaerythritol tetraacrylate (EPTA) monomer and 0.1 wt% AIBN initiator. Subsequently, the precursor solution was polymerized in situ at 60 °C. FEC and PS molecules exhibit a lower LUMO energy level than PC and EMC molecules. Thus, FEC and PS will be preferentially reduced on the Na metal anode, forming a protective SEI (Fig. 17a, b). Moreover, the gel structure (i.e., matrix arrangement) contributes to regulating the ion flux, thereby achieving smooth plating and stripping of Na<sup>+</sup> ions and Na.

Depending on the monomer chemical structure, the gel matrix can also facilitate ionic dissociation and provide anionic anchoring points through chemical interactions, thus improving the ionic conductivity of the gel electrolyte [179, 180]. Lei et al. confirmed that the creation of ionic channels in a GPE enabled smooth Na deposition on the anode [181]. They synthesized a PVDF-HFPcontaining cross-linked beta alumina nanowire membrane, which allowed ion transport along and through the nanowires (Fig. 17c). To prepare the GPE, the synthesized porous membrane was soaked in a liquid electrolyte (1 M NaClO<sub>4</sub> in EC:DEC with 5% FEC). The as-prepared GPE displayed smooth Na deposition on the Cu current collector (Fig. 17d), as well as high cycling performance in a NallNVP full cell (Fig. 17e). Notably, the commonly employed glass fiber separator was replaced by the crosslinked beta alumina nanowire gel polymer electrolyte (AN-GPE), which played the roles of both the electrolyte and separator, proving the structural robustness of the QSE.

Fabricating a mechanically robust (i.e., high Young's modulus) GPE is another strategy to reduce dendrite generation by mechanically preventing dendrite growth [182]. Glass fiber filters are largely used as separators owing to their high electrolyte intake capability and inexpensive feature. However, their low Young's modulus and large pores benefit dendrite propagation. Glass fiber separators can be strengthened by copolymerization inside the glass fiber network, thereby allowing better electrode/electrolyte contact, inhibiting dendrite growth and easing smooth Na plating and stripping processes [173]. Luo et al. significantly increased the Young's modulus of a PVDF-HFP membrane by simply tailoring the amount of graphene oxide (GO) in the PVDF-HFP-GO membrane [182]. Subsequently, the membrane was immersed in a  $1.0 \text{ M} \text{ NaClO}_4$ in EC/PC (with a volumetric ratio of 1:1) electrolyte to form the GPE. The high structural strength of the GPEs enabled smooth Na plating and stripping (Fig. 17f).

In general, the properties of GPEs can be easily tailored by introducing additives, manipulating the preparation methods, or adapting the monomer/polymer structure. In particular, GPEs constitute great candidates for flexible



◄Fig. 17 a Schematic illustration of the full cell using a 0.5 M NaPF<sub>6</sub> in PC/EMC liquid electrolyte (left) or a GPE (right). b HOMO and LUMO energy comparison of PC, EMC, FEC, and PS molecules. Reprinted with permission from Ref. [177]. Copyright © 2020, Elsevier. c Schematic of NallNVP batteries using the AN-GPE (the upper panel) and glass fiber (GF)-GPE (the lower panel) with the structure and Na-ion transport mechanism in the electrolytes. d SEM images of Na metal deposition on a Cu current collector using the GF-liquid electrolyte (LE), GPE, GF-GPE and AN-GPE at current densities of 0.5 mA cm<sup>-2</sup> and 60 °C. Areal capacity of 3 mAh cm<sup>-2</sup>. eCycling performance of NallNVP batteries using the GF-LE, GPE, GF-GPE, and AN-GPE at 1 C under 25 °C (the left panel) and 60 °C (the right panel). Reprinted with permission from Ref. [181]. Copyright © 2019, Springer Nature. f SEM images of 2-GO+PVDF-HFP (upper left) and PVDF-HFP (lower left), surface roughness of 2-GO+PVDF-HFP (upper middle) and PVDF-HFP (lower middle) and Young's modulus of 2-GO+PVDF-HFP (upper right) and PVDF-HFP (lower right) gel electrolytes. Reprinted with permission from Ref. [182]. Copyright © 2021, Wiley-VCH

batteries owing to their superior safety and high flexibility [183].

### 4.4 Solid-State Electrolytes

The solid-state electrolytes (SSEs) applied in Na metal batteries are identified as two classes: solid inorganic electrolytes (SIEs) and solid polymer electrolytes (SPEs). SSEs require the following main properties to be competitive with other electrolyte systems in Na metal battery applications: (1) high ionic conductivity (>  $10^{-3}$  S cm<sup>-1</sup>) with minor anionic and electronic contributions, (2) high mechanical strength to suppress Na dendrite growth, (3) superior interfacial contact with electrodes, (4) easy and inexpensive manufacturing, and (5) excellent electrochemical and thermal stability [184, 185].

### 4.4.1 SIEs

The SIEs applied in Na metal batteries include Na superionic conductor (NASICON) electrolytes, the  $\beta$ -alumina ( $\beta$ "-Al<sub>2</sub>O<sub>3</sub>) solid electrolyte (BASE), and sulfide-based electrolytes.

NASICON belongs to the rhombohedral system. The octahedral  $ZrO_6$  and tetrahedral  $(Si,P)O_4$  inside NASI-CON are connected by their apex angles to build up the 3D network [186]. As the bottlenecks formed by  $ZrO_6$  octahedra and  $(Si,P)O_4$  tetrahedra are larger than 4.8 Å (1 Å = 1 × 10<sup>-10</sup> m), the Na<sup>+</sup> ions at the interstitial sites of the 3D network can pass through the bottlenecks and diffuse in this material (Fig. 18a) [187].  $\beta''$ -Al<sub>2</sub>O<sub>3</sub> belongs to the hexagonal system. As one of the sodium aluminates,  $\beta''$ -Al<sub>2</sub>O<sub>3</sub> has a layered structure consisting of tightly packed Al-O spinel blocks and loosely packed sodium conduction planes that allow diffusion of Na<sup>+</sup> ions (Fig. 18b) [188]. Sulfide-based electrolytes such as Na<sub>3</sub>SbS<sub>4</sub> belong to the tetragonal system. The tetrahedral SbS<sub>4</sub><sup>3-</sup> inside Na<sub>3</sub>SbS<sub>4</sub> is connected with the adjacent tetrahedra by Na<sup>+</sup> ions. According to the bond valence sum mapping (BVSM) (Fig. 18c), the 3D framework built-up by tetrahedral SbS<sub>4</sub><sup>3-</sup> offers Na<sup>+</sup> ion conduction routes in all three dimensions (along the *a*-, *b*- and *c*-axes), and this type of structure connotes good ionic conductivity for the solid electrolyte [189]. The structural analysis of a sulfide-based electrolyte (Na<sub>3</sub>SbS<sub>4</sub>) is shown in Fig. 18d.

4.4.1.1 NASICON-Type SIEs The first NASICON structure  $(Na_{1+x}Zr_2P_{3-x}Si_xO_{12}, 0 < x < 3)$  was discovered by Hong and Goodenough, and it displayed a decent sodium ion conductivity of  $\sim 1.0 \times 10^{-4}$  S cm<sup>-1</sup> [190]. However, its conductivity is still more than one order of magnitude lower than that of conventional liquid electrolytes. Furthermore, the high interfacial resistance between NASICON-type electrolytes and metallic Na anodes is an issue limiting the performance of NASICON-type electrolytes in Na metal batteries. Heteroatom doping can improve the ionic conductivity of NASI-CON-type electrolytes. For example, Guin et al. reported a type of NASICON structure material by replacing Zr with Sc  $(Na_{1+r}Sc_2P_{3-r}Si_rO_{12})$ , and it gave an enhanced sodium ion conductivity of  $6.9 \times 10^{-4}$  S cm<sup>-1</sup> at 25 °C for x = 0.4 [191]. Song et al. synthesized a novel NASICON-type electrolyte by doping Mg into NASICON (Na<sub>3.1</sub>Zr<sub>1.95</sub>Mg<sub>0.05</sub>Si<sub>2</sub>PO<sub>12</sub>), and this electrolyte offered an excellent conductivity of  $3.5 \times 10^{-3}$  S cm<sup>-1</sup> with a wide ESW of 4.5 V at RT [192].

Introducing an interlayer between the NASICON electrolytes and electrodes can be a pathway toward alleviating the large interfacial resistance. Yu et al. applied a conventional NASICON-type SIE (Na<sub>3</sub>Zr<sub>2</sub>Si<sub>2</sub>PO<sub>12</sub>) coated by a layer of a polymer with intrinsic nanoporosity (PIN) that significantly enhanced the interfacial conduction between the NASICONtype SIE and the sodium metal anode (Fig. 18e) [193]. Moreover, this NASICON-type SIE with the PIN retained the advantages of SSEs, such as suppressing the shuttle effect and inhibiting Na dendrite growth. Thus, it effectively improved the cycling stability of RT Na-S batteries compared to the RT Na-S cell with a Celgard separator and offered a good ionic conductivity of  $1 \times 10^{-3}$  S cm<sup>-1</sup> at RT. Combining the Na metal anode with the carbon nanofiber (CNF)/sulfur cathode, the NalPIN-Na<sub>3</sub>Zr<sub>2</sub>Si<sub>2</sub>PO<sub>12</sub>||CNF/S RT Na-S cell delivered an initial discharge capacity of 700 mAh g<sup>-1</sup>, which remained at 550 mAh g<sup>-1</sup> after 100 cycles at 0.2 C.

**4.4.1.2 BASE-Type SIEs** Another type of SIE that has been applied in Na metal batteries is BASE. The BASE already widely applied in HT Na–S batteries can offer a comparably high ionic conductivity of  $\sim 2 \times 10^{-3}$  S cm<sup>-1</sup> at RT [194]. However, the limited wettability of BASE with Na is still a problem under RT conditions, which makes BASE mainly



**∢Fig. 18 a** Schematic of the Na<sup>+</sup> ion transport path inside the NASI-CON structure, where the rhombohedral symmetry represents the ZrO<sub>6</sub> and (Si,P)O<sub>4</sub> polyhedra, circles with dark shading and circles with light shading represent the Na(1) and Na(2) positions, respectively, and the curved line with an arrow is the conduction route of sodium ions. Reprinted with permission from Ref. [187]. Copyright © 1999, Elsevier Science B.V. b Schematic of the crystal structure of BASE. Reprinted with permission from Ref. [188]. Copyright  $\odot$  2016, Elsevier. c 3D bond valence map isosurfaces of Na<sub>3</sub>SbS<sub>4</sub> (isovalue:  $\pm 0.3$  v.u.). **d** Crystal structure with the unit cell outlined of Na<sub>3</sub>SbS<sub>4</sub>. Reprinted with permission from Ref. [189]. Copyright © 2016, Wiley-VCH Verlag GmbH & Co. e Schematic of the solidstate RT Na-S batteries applying the NASICON electrolyte coated with the PIN. Reprinted with permission from Ref. [193]. Copyright © 2019. Elsevier Inc. f Comparison between the electrolyte with and without BASE in RT Na-S batteries. Reprinted with permission from Ref. [194]. Copyright © 2013, Elsevier B.V. g Schematic of applying an in situ formation method to prepare Na<sub>3</sub>PS<sub>4</sub> SIEs and the Na<sub>2</sub>S-Na<sub>3</sub>PS<sub>4</sub>-CMK-3 cathode. Reprinted with permission from Ref. [78]. Copyright © 2018, American Chemical Society. h Combination of Na<sub>3</sub>SbS<sub>4</sub> with an organic electrolyte to form a thin and transferable hybrid electrolyte layer between the solid electrolyte and electrode to stabilize the interface. Reprinted with permission from Ref. [199]. Copyright © 2020, American Chemical Society

used in combination with liquid electrolytes. Wenzel et al. reported an RT Na-S battery with a liquid/solid electrolyte by applying BASE with 1 M sodium triflate (NaCF<sub>3</sub>SO<sub>3</sub>) in DOL/DME (with a volumetric ratio of 1:1) and compared it with the liquid electrolyte (LE) without BASE, as shown in Fig. 18f [194]. Compared with the electrolyte without BASE (starting from 450 mAh  $g^{-1}$  and remaining at ~80 mAh  $g^{-1}$  after 40 cycles at 0.148 mA cm<sup>-2</sup>), the electrolyte with BASE gives better capacity and cycling performance by suppressing shuttling of Na polysulfides and thus preventing the formation of an irreversible layer with increased resistivity on the Na anode surface (starting from 475 mAh  $g^{-1}$  and remaining at ~200 mAh  $g^{-1}$  after 40 cycles at  $0.148 \text{ mA cm}^{-2}$ ). Kim et al. applied BASE with TEGDME, which also provided better cycling stability and higher capacity (855 mAh  $g^{-1}$  to 521 mAh  $g^{-1}$ ) than the cell using the porous separator with the LE (starting from 350 mAh  $g^{-1}$  and remaining at 100 mAh  $g^{-1}$  after 2 cycles at 1/64 C) [195]. BASE provides total physical shielding to inhibit shuttling of Na polysulfides compared to the bare LEs and conventional porous separators.

**4.4.1.3 Sulfide-Based SIEs** The last type of SIE is sulfidebased electrolytes, which can also provide an acceptable ionic conductivity at RT (e.g.,  $Na_3SbS_4$ ,  $10^{-4}$ – $10^{-3}$  S cm<sup>-1</sup>). However, the interfacial resistance, parasitic reaction when in contact with the metallic sodium anode, and potential brittle cracking due to volume change during charge/discharge are still challenges that need to be solved for the application of sulfide-based SIEs. Several methods, such as using an in situ formation method [196–198] or a novel cathode material [198] or combining sulfide-based SIEs with other electrolytes [199], were adopted to solve these issues, which improved the electrochemical performance of sulfide-based SIEs in Na metal batteries.

The conventional cold-pressing process to fabricate sulfide-based SIEs induces poor interfacial contact between the electrolyte and electrodes. The in situ fabrication of a sulfide solid electrolyte with cathode active materials can ensure high ionic and electronic conductivity and sufficient interfacial contact. Yue et al. used the Na<sub>3</sub>PS<sub>4</sub>-Na<sub>2</sub>S-C nanocomposite as the cathode and contained Na<sub>3</sub>PS<sub>4</sub> SIEs inside the nanocomposite to solve the contact problem between the electrodes and SSE [196]. The RT Na-S batteries with Na<sub>3</sub>PS<sub>4</sub> as sulfide-based SIEs offered good cycling stability and a high capacity at 60 °C (starting from 869.2 mAh  $g^{-1}$ , which remained at 438.4 mAh  $g^{-1}$  after 50 cycles at 50 mA g<sup>-1</sup>). Tanibata et al. employed an amorphous S-P<sub>2</sub>S<sub>5</sub> electrode on the cathode side, which was in situ reacted with the Na metal anode to form a nanosized Na<sub>3</sub>PS<sub>4</sub> and Na<sub>2</sub>S crystal mixture [197]. The crystal mixture precipitated on the surface of the cathode and endowed the Na-S battery with a high initial discharge capacity of approximately 1 250 mAh  $g^{-1}$  at 0.013 mA cm<sup>-2</sup>. Recently, a type of Na<sub>3</sub>SbS<sub>4</sub>-based SIEs reported by Wan et al. could provide an excellent conductivity of  $1.14 \times 10^{-3}$  S cm<sup>-1</sup> at RT along with a S-Na<sub>3</sub>SbS<sub>4</sub>-C cathode and a Na metal anode [198]. They used a mechanochemical process (liquid-phase reaction + mechanical milling) to trigger in situ formation of S-Na<sub>3</sub>SbS<sub>4</sub>, which can provide both microscale primary and nanoscale secondary pathways for electronic/ionic transformation. In addition to offering stabler interfacial contact between the SSE and cathode, the in situ formation method also reduces the cathode material stress/strain and leads to superior rate and cycling performance of the Na-S battery (showing a high initial capacity of 1 504.3 mAh  $g^{-1}$ at 50 mA  $g^{-1}$ , which remains at 468.1 mAh  $g^{-1}$  after 100 cycles at 1 000 mA  $g^{-1}$ ).

More recently, Ren et al. applied Na<sub>3</sub>SbS<sub>4</sub> combined with an organic electrolyte (pentaerythritol tetraacrylate (PETEA)) by using an in situ cross-linking method and generated a thin and transferable hybrid electrolyte that can offer a stable interface with sodium during cycling (Fig. 18h) [199]. This inorganic/organic hybrid sulfide-based solid electrolyte provides a stabler electrochemical performance by preventing parasitic reactions between the SIEs and electrode, and the discharge capacity gradually increases from 0.32 to 0.52 mAh cm<sup>-2</sup> over 90 cycles due to potential pore formation and delamination. However, the ionic conductivity of Na<sub>3</sub>SbS<sub>4</sub>/PETEA is only  $0.47 \times 10^{-4}$  S cm<sup>-1</sup> at RT, which is ~ 22% of that of Na<sub>3</sub>SbS<sub>4</sub> and requires further improvement in future research.



# 4.4.2 SPEs

SPEs are commonly assembled by dissolving metal salts in a polymer matrix. SPEs can be easily achieved by using hot molding, solvent casting, or extrusion techniques without the participation of liquid solvents as plasticizers [200]. For the polymer matrix in SPEs, poly(ethylene oxide) (PEO) is the most widely researched material, and it is a type of semic-rystalline polymers with both crystalline regions and amorphous regions (Fig. 19a) [201]. The stable interfacial contact

◄Fig. 19 a Schematic of the ion transport mechanism in the amorphous parts of PEO-based SPEs. Reprinted with permission from Ref. [201]. Copyright © 2016, The Royal Society of Chemistry. b Schematic illustration of the introduction of non-ion-conductive filler SiO<sub>2</sub> into the PEO matrix. c TEM image of PEO combined with SiO<sub>2</sub> fillers. Reprinted with permission from Ref. [204]. Copyright © 2015, American Chemical Society. d Schematic drawing of the Na<sup>+</sup> ion transport mechanism inside PEO with the addition of NASICON. Reprinted with permission from Ref. [207]. Copyright © 2017, Elsevier B.V. e SEM images of the Na metal surface after 50 cycles in Na–SnS<sub>2</sub> cells with the Na<sub>3</sub>PS<sub>4</sub> and Na<sub>3</sub>PS<sub>4</sub>-PEO-NaClO<sub>4</sub> electrolytes. f Comparison of the cycling performance of the Na–SnS<sub>2</sub> batteries with the Na<sub>3</sub>PS<sub>4</sub> electrolyte and with the Na<sub>3</sub>PS<sub>4</sub>-PEO-NaClO<sub>4</sub> electrolyte after 40 cycles at 20 mA g<sup>-1</sup>. Reprinted with permission from Ref. [208]. Copyright © 2019, Elsevier B.V. and Science Press

with electrodes and the excellent solubility for sodium salts make PEO quite attractive in sodium metal battery research. However, the operation temperature range for PEO is quite limited since the crystalline regions of PEO offer very limited ionic conductivity (the amorphous regions of the PEO matrix are the main ion transport pathways) [202]. On one hand, when the temperature is below 65 °C, the PEO chains will crystallize and deliver a very poor ionic conductivity of  $10^{-8}$ – $10^{-7}$  S cm<sup>-1</sup>. On the other hand, when the temperature is higher than 80 °C, the mechanical properties of PEO may be degraded, limiting the ESW of the solid electrolyte, which should also be noted [161, 200].

Modification strategies, including using optimized concentrated sodium salts, introducing inorganic fillers, and employing novel cathode materials in PEO, need to be applied to ensure satisfactory mechanical properties, decent ionic conductivities (> $1.0 \times 10^{-4}$  S cm<sup>-1</sup>), and wide ESWs. Park et al. embedded NaCF<sub>3</sub>SO<sub>3</sub> salt into the PEO matrix and first tested it in an all-solid-state RT Na–S battery system [203]. The PEO/NaCF<sub>3</sub>SO<sub>3</sub> electrolyte delivered a high ionic conductivity of  $3.38 \times 10^{-4}$  S cm<sup>-1</sup> at 90 °C, and the Na–S battery with this SPE provided a high initial capacity of ~ 505 mAh g<sup>-1</sup>, which remained at 166 mAh g<sup>-1</sup> after 10 cycles at 0.144 mA cm<sup>-2</sup>.

There are two classes of fillers that can be introduced into the PEO matrix. The first type is non-ion-conductive fillers (e.g., SiO<sub>2</sub>, TiO<sub>2</sub>), which aim to suppress the crystallization of PEO and increase cation transport (Fig. 19b) [204–206]. The surface groups on the SiO<sub>2</sub> particles can incur a local structural change in the PEO matrix (Fig. 19c), which effectively prevents the formation of crystalline regions and accelerates cation transport by increasing the free cations. Zhu et al. introduced 1% TiO<sub>2</sub> filler into the PEO electrolyte with NaFSI salt, and the PEO-NaFSI-1% TiO<sub>2</sub> solid electrolyte achieved a high conductivity of  $4.89 \times 10^{-4}$  S cm<sup>-2</sup> at 60 °C because of the extension of the amorphous regions [205].

The other type of filler is ion-conductive fillers (e.g., NASICON,  $Na_3PS_4$ , MIL-53(Al)), which can offer

additional ion diffusion pathways to improve the ionic conductivity of PEO-based SPEs (Fig. 19d) [207-209]. Zhang et al. applied NASICON as an ion-conductive filler for the PEO matrix [207]. PEO can increase the Na vacancies on the NASICON surface and accelerate ion transport along the surface region. Meanwhile, the NASICON ceramic bulk can provide a competing ionic conductivity. Therefore, the composite SSE resulted in a high ionic conductivity of approximately  $2.8 \times 10^{-3}$  S cm<sup>-1</sup> at 80 °C. Xu et al. coated PEO with NaClO<sub>4</sub> salt onto Na<sub>3</sub>PS<sub>4</sub> particles to form a solid electrolyte, which provided a high ionic conductivity of  $9.4 \times 10^{-5}$ S cm<sup>-1</sup> at RT and  $2.5 \times 10^{-4}$  S cm<sup>-2</sup> at 60 °C [208]. This coating strategy effectively stabilized the surface contact between the solid electrolyte and electrodes (Fig. 19e). Thereby, the all-solid-state RT Na-SnS2 battery offered an enhanced cycling performance of approximately 230 mAh  $g^{-1}$  after 40 cycles at 20 mA  $g^{-1}$  (Fig. 19f). The addition of ion-conductive fillers is similar to combining SIEs and SPEs to form composite electrolytes, which can synergize the advantages of both SIEs (e.g., sufficient mechanical strength, good ionic conductivity) and SPEs (e.g., high flexibility, good interfacial contact between electrodes and the electrolyte) to improve the electrochemical performance of the all-solid-state RT Na-S battery system. However, these types of composite SSEs still have not attracted enough interest, and this field can be a promising pathway for future SSE development for RT Na-S batteries.

In general, the ionic conductivity, mechanical strength, interfacial conduction, and electrochemical and thermal stability for both SIEs and SPEs can be manipulated by introducing other electrolytes or fillers, adapting the synthesis method and cathode structure, or coating the surface of SSEs. However, the Na metal electrode, which is much more unstable than Li metal electrodes, will incur more serious contact problems when using SSEs. SSEs were first thought to be able to suppress dendrite formation owing to the high Young's modulus. However, battery failure due to short circuit was observed, attesting to dendrite penetration through the SSE. The mechanism of dendrite formation in SSEs is still under investigation, with two predominant mechanisms being mostly accepted. (1) Preferential Na plating where the local current density is higher (similar to dendrite formation in LEs). The dendrites grow through the SSE and propagate via defects or voids, which present higher energy. Usually, such dendrites are bulky with a large diameter, which can generate cracks in the SSE and lead to battery failure. (2) Dendrite growth through the grain boundaries due to electronic leakage. In this situation, the dendrites are very thin and follow the grain boundaries. The very small diameter of these dendrites results in a stronger electric field at the tip of the dendrites, which in turn further promotes their growth. Therefore, dendrite growth is even severer than that in LEs or GPEs [210]. Due to the above issues, the contact stability



**∢Fig. 20 a** Time-of-flight secondary-ion mass spectrometry 3D illustration of NaO<sup>−</sup>, BO<sub>3</sub><sup>−</sup> and PO<sub>3</sub><sup>−</sup> along the depth of the sample, and **b** corresponding depth profiles of SiO<sub>4</sub><sup>−</sup>, PO<sub>4</sub><sup>−</sup>, NaO<sup>−</sup> and BO<sub>3</sub><sup>−</sup> signals. **c** SEM image of the cross-section and **d** EDS mapping of NASICON with a Na<sub>2</sub>B<sub>4</sub>O<sub>7</sub> additive/Na interface disassembled from a symmetric cell cycled at 0.3 mA cm<sup>−2</sup> for approximately 2 500 h. Reprinted with permission from Ref. [217]. Copyright © 2021, Elsevier. **e** Schematic illustration of the Na<sup>+</sup> ion mobility in PEO and optimized PEO-PPC-NASICON electrolytes. **f** Cycling performance of the NalPEO-PPC-NASICONINa symmetric cell at a current density of 0.1 mA cm<sup>−2</sup>. **g** SEM images of the surface morphologies of Na anodes after cycling in PEO-PPC-NASICON (the upper panel) and PEO (the lower panel) electrolytes. Reprinted with permission from Ref. [218]. Copyright © 2021, Royal Society of Chemistry

between the Na metal and SSEs needs to be given more focus along with enhancement strategies in the all-solid-state Na metal battery system, which will be investigated in the next section.

# 4.4.3 Enhancement of the Interfacial Contact of Na Metal/ SSEs

The Na metal/SSE interface can be categorized into three types: (1) a thermodynamically stable interface, (2) a mixed ionic and electronic conducting interphase, and (3) an SEI with ionic conductivity and electronic insulation [211, 212]. Only the former and the latter type of interfaces can withstand long cycling.

SIEs usually suffer from poor electrode/electrolyte contact, which further deteriorates during the plating/stripping processes due to the creation of voids and accumulation of dead Na at the interface. The void formation is even more pronounced at high current densities, where larger voids are created during stripping and harder to fill during plating [213]. Applying an external pressure to the cell enhances the electrode/electrolyte contact and efficiently reduces the creation of voids during cycling. However, the external force applied must be carefully chosen to prevent mechanical failure [214]. Generally, Na metal is directly pressed on an SSE to achieve high solid—solid contact. However, undesirable functional groups with high resistance can form on the surface of solid electrolytes, thereby leading to poor contact with the Na anode.

To improve the Na metal electrode/SSE contact, several advanced approaches have been investigated, such as surface polishing, heat treatment and the introduction of additives. Heat treatments have been employed to clean the solid electrolyte surface. For instance, by thermal treatment of the  $Na_3Zr_2Si_2PO_{12}$  surface, the Na wettability was improved, and the interfacial resistance decreased [215]. Recently, an annealing process of a  $Na_3Zr_2Si_2PO_{12}$  NASICON solid electrolyte surface enabled the removal of the side products on the surface of the solid electrolyte as well as partial reduction, thereby reducing the interfacial impedance [216].

Zhao et al. proposed another method: the introduction of the sintering additive  $Na_2B_4O_7$  into a NASICON solid, where  $NaO^-$  and  $BO_3^-$  would reside on the surface of the sample (Fig. 20a, b) [217]. The as-prepared electrolyte allowed a high ionic conductivity of  $1.72 \times 10^{-3}$  S cm<sup>-1</sup> at RT owing to the facile Na<sup>+</sup> ion migration across the grain boundaries. Furthermore, the additive enabled the formation of a Na<sup>+</sup>-rich interface on the Na metal anode, which blocked electronic conductivity but was permeable to Na<sup>+</sup> ions, thereby promoting high cycling performance (Fig. 20c, d).

SPE systems can also be tailored to further enhance the interfacial solid-solid contact stability. The polypropylene carbonate (PPC) polymer undergoes nucleophilic attacks in which Na<sup>+</sup> randomly breaks C–O bonds in the PPC backbone; therefore, PPC is not a suitable SPE for Na metal batteries. However, the reacted PPC could be employed as a highly conductive interfacial wetting layer on the Na metal anode, which would be formed in situ on the anode (Fig. 20e). The interfacial resistance observed between Na metal and an SPE mixture of PEO and PPC was smaller than that of Na/PEO owing to the decomposition of PPC on the Na metal, which enhanced the electrode/electrolyte contact [218]. The NalNa symmetric cell was cycled for more than 1 000 h, and the surface morphology of Na after cycling remained smooth in comparison with the cell cycled in the PEO electrolyte (Fig. 20f, g). PPC was also introduced into amorphous phases in a polymer blend structure, thereby contributing to higher ionic conductivity. The ionic conductivity of this electrolyte was further improved by introducing a small amount of NASICON as an inorganic filler. As a result, the optimized PEO-PPC-NASICON (48-37-15 wt%) SPE exhibited high chemical stability, high ionic conductivity and resistance to dendrite growth owing to the superior tensile strength and Young's modulus of the polymer electrolyte.

### 5 Nanostructured Na Metal Anodes

To enable the practical application of Na–S and Na–Se batteries, we need to achieve high-performance Na metal anodes with long-term cyclability and good safety. As mentioned in the previous section, Na metal is more reactive than Li metal. The severe reactions between Na metal and the organic electrolyte generate a thick SEI between them, which causes a significant increase in the charge transfer resistance, inducing large voltage hysteresis and low energy efficiency. In addition, the infinite volume change of the Na metal anode during repeated plating and stripping is another major challenge that needs to be resolved to achieve a stable Na metal anode. Therefore, nanostructure engineering, including protective layer engineering and skeleton design, has been a focus of this research area, and tremendous outcomes have been achieved thus far.

### 5.1 Artificial Protective Layers for Na Metal Anodes

Constructing protective layers on Na metal anodes is beneficial for stabilizing the SEI during the repeated plating/ stripping process [219]. The protective layers can isolate the highly reactive Na metals from the LEs. In addition, a protective layer with high ionic conductivity can regulate the Na<sup>+</sup> ion flux on the anode surface, leading to uniform metallic Na plating. The in situ formation of protective layers or an SEI via electrolyte engineering has been summarized in the previous section. Therefore, in this section, we will focus on the recently developed ex situ artificial protective layers. The ex situ artificial layers should possess superior ionic conductivity and abundant Na<sup>+</sup> ion transport channels to ensure that electrochemical deposition of Na metal occurs underneath the artificial layers. In addition, the protective layers should possess good flexibility to adapt to the large volume change and sufficient mechanical strength to withstand dendrite penetration.

#### 5.1.1 Constructing Protective Layers via Solution Methods

Traditional solution methods have been used to form artificial protective layers with controlled composition on Na metal anodes. By exposing Na metal anodes to solutions containing certain chemicals, an artificial protective layer can be formed on Na anodes via chemical reactions (Fig. 21a, b). Tian et al. successfully constructed a NaI layer via a Wurtz reaction. The Na metal anodes were first exposed to a solution containing 2-iodopropane or 1-iodopropane. Then, a NaI layer was formed on the Na metal anode after one electron from the Na metal was transferred to the iodine, which endowed the Na metal anodes in Na-I<sub>2</sub> batteries with significantly enhanced electrochemical performance [220]. By using the same reaction mechanism, a thin layer of NaBr was constructed on a Na metal anode through the reaction of metallic Na with 1-bromopropane solution [221]. A joint DFT study of the SEI demonstrated that the energy barrier for adatom diffusion on NaBr is as low as that of magnesium, which is known to form spherical nuclei upon charging. Compared to pristine Na metal, the reduced energy barrier for ion transport (0.02 eV per atom) of NaBr promoted uniform Na plating and prevented unwanted parasitic reactions between the electrode and electrolyte. Electrochemical testing in symmetric cells showed that NaBr-coated Na metal anodes achieved an almost 10 times longer cycle life than unprotected Na metal anodes.

Slurries containing organics and/or inorganics have been coated on the surface of Na metal anodes through a variety of methods, such as tape casting and spin coating. After evaporation of the solvents, uniform protective layers can be conformally coated on the surfaces of Na metal anodes. Hou et al. prepared PVDF@Cu, PVDF-HFP@Cu, and PEO@Cu current collectors by casting solutions with the corresponding polymers on Cu current collectors [222]. Among all the as-prepared current collectors, the PVDF@Cu current collector exhibited the best electrochemical performance, with an ultralong cycling life (1 200 h) and high average Coulombic efficiency (approximately 99.91%) at 1 mA cm<sup>-2</sup>. By using a similar preparation method, an inorganic-organic hybrid protective layer (NaF-PVDF) was coated on commercial Cu current collectors [223]. The organic component, a flexible PVDF matrix, can maintain the integrity of the protective interface against volume change. Meanwhile, the inorganic component, NaF particles, can enhance the mechanical strength and increase the Na<sup>+</sup> ion conductivity, which is beneficial for suppressing dendrite growth. As a result, the hybrid interface-coated Cu current collector showed a tenfold higher lifespan than the bare Cu current collector. Qian et al. also fabricated several MOF-coated Cu electrodes using the tape casting method. Slurries containing MOF particles and PVDF binders in N-methyl-2-pyrrolidone (NMP) solvent were cast on Cu foil by a doctor blade. After drying in an oven, MOF-coated Cu electrodes were obtained [224]. The porous and robust MOF protective layer acted as a resilient shield and a homogenized metal ion concentration layer, which could physically suppress Li/Na dendrite growth.

# 5.1.2 Interface Engineering via Advanced Nanotechnologies

Highly uniform and compact artificial protective layers constructed by nanomaterials, especially ultrathin 2D materials, can be realized via the chemical vapor deposition (CVD) method. The thickness of the protective layer based on 2D materials can be controlled at the atomic scale while maintaining high mechanical strength. However, many factors influence the physical and chemical properties of the 2D material-based protective layer, such as the defects, bond length and metal proximity of the 2D materials [225]. Increasing the defects and reducing the thickness of the 2D materials benefit the ionic conductivity but show negative effects on the mechanical strength of the protective layer. Therefore, finding a balance between the electrochemical performance and the mechanical properties is essential for the design of an effective artificial protective layer with 2D materials. For instance, free-standing graphene films with different thicknesses were synthesized via the CVD method and used as the artificial protective layer for Na metal anodes (Fig. 21e) [226]. A few-layer graphene (FLG) film with a thickness of approximately 2.3 nm favors



**Fig. 21 a** Schematic illustration of the formation of NaI-protected Na metal anodes. **b** SEM image of NaI-coated Na metal anodes. Reprinted with permission from Ref. [220]. Copyright © 2021, Elsevier. **c** Schematic illustration of the preparation of an MOF (MOF-199)-coated Cu electrode. **d** Photographs of pristine Cu foil and MOF-199-coated Cu foil. Reprinted with permission from Ref. [224]. Copyright © 2021, Elsevier. **e** Synthesis process of graphene film-protected Na metal anodes. **f** Cycling performance of Na metal

anodes with and without a graphene film at different current densities. Reprinted with permission from Ref. [226]. Copyright © 2017, American Chemical Society. **g** Schematic illustration of the ALD of Al<sub>2</sub>O<sub>3</sub> on Na metal anodes. Reprinted with permission from Ref. [13]. Copyright © 2020, Wiley–VCH. **h** Schematic illustration of the Na plating/stripping behavior with and without an ALD-Al<sub>2</sub>O<sub>3</sub> protective layer. Reprinted with permission from Ref. [227]. Copyright © 2017, Wiley–VCH good electrochemical performance at a lower current density (Fig. 21f). In contrast, a multilayer graphene (MLG) film with a thickness of approximately 5 nm shows the best performance at a higher current density (Fig. 21f). Other 2D materials, such as hexagonal boron nitride (h-BN), have already been used in Li metal anodes, have achieved significantly improved performance, and could be potential candidates for Na metal anode protection.

Other advanced thin film fabrication nanotechnologies, including atomic layer deposition (ALD) and molecular layer deposition (MLD) technologies, were introduced to form atomic-scale artificial protective layers on Na metal anodes (Fig. 21g). Through ALD and MLD technologies, the thickness and composition of the protective film can be precisely controlled. For instance, an inorganic layer, such as an Al<sub>2</sub>O<sub>3</sub> layer, was directly coated on Na metal anodes via plasma-enhanced ALD (PE-ALD), which demonstrated a significant positive effect on the cycling stability of Na metal anodes in carbonate electrolytes [13, 227, 228]. To further enhance the mechanical properties of the artificial protective layer (i.e., alucone), an inorganic and organic protective layer was prepared by using the MLD method [229, 230]. The inorganic and organic layer (MLD-alucone)-coated Na metal anodes showed much-improved cycling stability compared to the inorganic layer (ALD-Al<sub>2</sub>O<sub>3</sub>)-coated electrode and the pristine electrode. The enhanced flexibility has been assigned as the major contributor to maintaining the integrity of the protective film against repeated volume changes during the Na plating/stripping process.

In brief, employing nanotechnology has been demonstrated to be an effective strategy to fabricate uniform artificial protective layers at the atomic scale. The good chemical stability, high ionic conductivity, and enhanced mechanical properties facilitate uniform Na plating. The major challenges of using these sophisticated methods are how to reduce the processing cost and simplify the fabrication process for large-scale applications.

### 5.2 Skeleton Design for Sodium Metal Anodes

As "hostless" anodes, Na metal anodes suffer from severe volume change during repeated plating/stripping processes. Designing a suitable 3D "host" has been demonstrated to be a promising strategy to avoid very large volume change and guide homogeneous Na metal deposition [231, 232]. To meet the requirements of practical applications, some selection criteria, including the mechanical strength, electronic conductivity, surface area, chemical stability and cost, need to be carefully considered to achieve a suitable "host" for metallic Na to achieve high electrochemical performance. To date, a variety of skeletons have been developed to form composite anodes with Na metal, such as porous metal skeletons, alloy-based skeletons, carbon-based nanostructure

skeletons, and polymer skeletons. Therefore, in this section, we will discuss the recent progress in skeleton design for sodium metal anodes.

### 5.2.1 Metallic Skeletons

Metallic materials, such as Cu or Al foils, have been used as standard substrates to investigate Na metal plating/stripping behavior. Severe dendrite formation has been observed on planar metallic substrates due to their low surface area and poor sodiophilic property. The first approach to improve the Na metal plating/stripping behavior is to increase the surface areas of the dense metallic foils to reduce the local current density [233]. The lower local current density can facilitate homogenous Na metal plating owing to the minimal Na-ion concentration gradient near the surface of Na metal anodes. For instance, constructing a porous Al skeleton to increase the surface area of the substrate has been demonstrated to be an effective strategy to suppress Na dendrite formation and enhance the cycling stability of Na metal anodes [234]. The interconnected porous structure of porous Al foils increases the available sites for Na nucleation and regulates the Na<sup>+</sup> ion flux distribution, which facilitates uniform Na deposition without obvious dendrite formation (Fig. 22a). For the LilAl half-cell with porous Al foil as the working electrode, a low and stable voltage hysteresis and an average plating/stripping Coulombic efficiency of above 99.9% were achieved for over 1 000 cycles, which is a significant improvement compared to planar Al foil. In addition, a Na-O<sub>2</sub> full cell with a low amount of metallic Na predeposited on porous Al foil as the anode showed superb cycling stability for 200 cycles.

Constructing 3D conductive frameworks on Cu foils to increase the surface area of the current collectors has also been reported to develop high-performance Na metal anodes. Lu et al. in situ grew Cu nanowires on commercial Cu foil through a hydrothermal process in absolute alcohol solvent [233]. The obtained Cu nanowires showed much thinner diameters of 40 nm compared with previously reported Cu nanowires (> 300 nm) for Li metal anodes. The significantly increased surface area of the 3D conductive skeleton with Cu nanowires effectively regulated the electric field distribution and thus facilitated reversible Na plating and stripping with high Coulombic efficiencies. The full cells with the Na predeposited 3D Cu nanowires as the anode and  $Na_3V_2(PO_4)_3@C$  as the cathode showed extremely stable cycling performance and enhanced high-rate capabilities, with an energy density approaching  $314.5 \text{ Wh kg}^{-1}$ . In addition, Xu et al. electrodeposited porous Ni on Cu foil via a facile hydrogen bubble dynamic template method [235]. The 3D Ni@Cu substrate showed a unique honeycomb-like porous structure, which could accommodate both metallic Li and Na deposition in the porous networks and prevent large volume expansion. In the half-cell Coulombic efficiency



**Fig.22 a** Schematic illustration of Na deposition on planar and porous Al foils, and corresponding SEM observation at different magnifications. Reprinted with permission from Ref. [234]. Copyright © 2017, American Chemical Society. **b** Schematic illustration of the fabrication process of in situ growth of Cu nanowires on Cu foam. **c** Long-term time vs. voltage curves of Na plating and stripping on pristine Cu foam and CuNW-Cu foam. Reprinted with permission

from Ref. [236]. Copyright © 2021, Elsevier. **d** Schematic illustration of the Na nucleation and deposition behavior on pristine Cu foam and ZnO nanorod-decorated Cu foam. **e** COMSOL simulation results of the local current density distribution and Na<sup>+</sup> flux at the interfaces of ZnO nanorod-decorated Cu foam (CF@ZnO) and bare Cu foam. Reprinted with permission from Ref. [237]. Copyright © 2021, Elsevier

testing, the 3D Ni@Cu substrate achieved excellent stability of Na reversible deposition without dendrite formation under high current density and capacity limit.

Modifying commercial Cu foams as conductive substrates for Na metal anodes has also attracted significant attention recently. As shown in Fig. 22b, through in situ growth of Cu nanowires on 3D Cu form, a hierarchical porous Cu substrate (CuNW-Cu) showed an excellent confinement effect to reduce the volume change during Na plating/stripping [236]. The enlarged surface area and nanoporous architecture facilitated homogenous Na<sup>+</sup> ion flux, which promoted uniform Na metal deposition without dendrite formation. The controlled half-cell cycling test results demonstrated repeated and uniform Na metal deposition with a capacity limit of up to 12 mAh cm<sup>-2</sup> for over 1 000 h (Fig. 22c). In addition, Yang et al. reported in situ growth of ZnO nanorods on Cu foam via a simple chemical precipitation method and used this material as a 3D skeleton for a dendrite-free Na metal anode (Fig. 22d) [237]. The excellent sodiophilic property of ZnO nanorods provided abundant nucleation sites during the initial Na deposition process, which was reflected by the reduced nucleation overpotential compared with the pristine Cu foam skeleton. The COMSOL Multiphysics simulation results in Fig. 22e indicated that the rationally designed core-shell cylindrical structure of the nanorod arrays could reduce the local current density and facilitate uniform Na deposition. Furthermore, Wang et al. treated a commercial Cu foam through a facile chemical engineering process (e.g., oxidation or sulfurization) to form a cylindrical core-shell skeleton with a good sodiophilic property [238]. The chemical interaction between Na and chemically processed Cu foam, especially oxygen-treated Cu foam, facilitated uniform molten Na impregnation during the infusion process and confined metallic Na within the matrix pores during the repeated plating/stripping process. The galvanostatic cycling tests in symmetric cells with two Na-infused O-treated Cu foam (Na@O-CF) electrodes showed exciting cycling stability in widely used carbonate electrolytes, which promotes their future practical application.

For the future development of metallic skeletons as the current collectors for Na metal batteries, there are still several challenges that need to be resolved. The optimization of the multiscale porous structure design needs to effectively accommodate the Na metal while maintaining the robust mechanical strength. Meanwhile, how to reduce the cost for feasible mass production needs to be considered.

### 5.2.2 Alloy Skeletons

Alloying Na metal with other metals (e.g., Li, Na, K, Sn and Sb) to form Na–M alloy anodes is another strategy to improve the electrochemical performance of Na metal anodes. The alloying process will change the physical and chemical properties of Na metal-based anodes, such as the melting points, mechanical properties, chemical reactivities, and plating behavior.

Li-Na alloy anodes have been used in oxygen batteries and have achieved improved cycling stability [239]. A robust and flexible SEI was formed on Li-Na anodes via an in situ reaction with a 1,3-dioxolane additive, which could buffer the volume change of the alloy anodes and thus prevent cracking and further electrolyte consumption. In addition, the continuous release of Li<sup>+</sup> ions into the electrolyte will help form a strong electrostatic shield that can effectively suppress dendrite formation. The Li-Na alloy-oxygen cells with optimized Li-Na alloy anodes achieved good cycling stability up to 137 cycles at a current density of 200 mA  $g^{-1}$ with a capacity limit of 1 000 mAh  $g^{-1}$ . Similarly, Yu et al. infused Li-Na alloy into a carbon cloth (CC) host and used it as the anode in Li-Na alloy-iodine batteries [240]. The Li-Na alloy-CC anodes showed not only excellent cycling stability in symmetric cells but also significantly improved cycling stability in Li-Na alloy-iodine batteries.

Owing to their liquid state at RT, Na–K alloys have also been employed in rechargeable batteries. Replacing the solid–liquid interface with a liquid–liquid interface completely avoided dendrite formation. Na–K alloys can be used as the anodes for both Na-based and K-based batteries, depending on the reaction mechanism of the counter cathode electrode. If Na<sup>+</sup> ions are involved in the redox reaction of the cathode materials, then the alloys can be used as the anodes for Na-ion batteries. Otherwise, they serve as the anodes for K-ion batteries.

### 5.2.3 Carbonaceous Skeletons

Carbonaceous materials are considered alternative candidates for synthesizing nanostructure frameworks due to their high electronic conductivity, low cost, low density, and diverse nanostructures. Many nanostructure frameworks have been developed to improve the electrochemical performance and safety of alkali metal anodes, including 1D nanotubes and nanofibers, 2D graphene, and 3D porous carbon materials.

**5.2.3.1 Carbon Nanotube (CNT) Skeletons** As one of the most used carbon materials, CNTs have been massively produced in industry and comprehensively used to obtain 3D conductive skeletons due to their excellent conductivity, unique 1D structure, and good mechanical robustness. The interconnected conductive networks of CNT skeletons have large specific surface areas, which can reduce the local current density and facilitate homogenous Na plating. Meanwhile, the abundant voids formed between the CNTs effectively alleviate the volume change during the metallic Na plating/stripping process. Through a facile mechanical

mixing process, a CNT-Na composite was fabricated and showed significantly reduced nucleation overpotential during Na plating (Fig. 23a). The CNT content in the CNT-Na composite significantly influenced the nucleation overpotential; therefore, the optimized 11.8CNT-Na electrode exhibited the best cycling performance, which was approximately 5 times longer than that of the pristine Na metal anodes in symmetric cell testing [241].

Due to the "sodiophobic" property of pristine CNTs, a variety of surface modification strategies have been reported to enhance their "sodiophilic" property. Nitrogen-doped carbon nanotubes (NCNTs) rationally grown on carbon paper (CP) have been used as a 3D porous host for Na metal anodes (Fig. 23b). The vertical NCNTs serve as "sodiophilic" sites, which effectively decrease the contact angle between CP and liquid metal Na. The interconnected conductive network of CNTs on CP increases the surface area and reduces the local current density, facilitating smooth Na nucleation and deposition. After the Na infusion process, the as-prepared Na@CP-NCNT composite anodes demonstrate excellent cycling stability at different current densities with a capacity limit of up to 3 mAh cm<sup>-2</sup> [242]. Introducing oxygenfunctional groups onto CNTs via oxygen microwave plasma is another method to enhance their "sodiophilic" property (Fig. 23c, d) [243]. DFT calculations showed that the binding energy between Na and oxygen-functionalized CNTs is stronger than that with pristine CNTs and Cu current collectors. Therefore, the oxygen-functionalized CNT matrix exhibited homogeneous Na nucleation and dendrite-free growth behavior with a remarkable electrochemical performance (e.g., a stable Coulombic efficiency of over 99.5% for more than 680 cycles at 10 mA h  $cm^{-2}$  with a current density of 5 mA cm<sup>-2</sup>). In addition, Sun et al. introduced N and S-codoped CNT (NSCNT) paper as a highly "sodiophilic" matrix for dendrite-free Na metal anodes (Fig. 23e) [244]. The NSCNT paper showed very good mechanical strength with excellent flexibility and could be bent and punched into different shapes (Fig. 23f). DFT calculations proved that the codoped CNTs showed a better "sodiophilic" property than the singly doped and pristine CNTs. As a result, the NSCNT paper can guide uniform Na<sup>+</sup> ion flux and facilitate dendritefree Na plating at high current densities and capacity limits. Furthermore, the Na/NSCNT composite anodes can be used to improve the cycling performance of RT Na-O<sub>2</sub> batteries, showing great potential for use in RT Na-S and Na-Se batteries.

Although great improvements have been achieved through the advantages of CNT skeletons, further in-depth studies must still be conducted with the help of advanced characterization and testing methods, such as cryogenic electron microscopy, in situ optical microscopy, and titration gas chromatography, to study the mechanism of the enhanced electrochemical performance of Na metal anodes. 5.2.3.2 Graphene Skeletons As one of the most popular 2D materials for rechargeable batteries, graphene attracts great attention in alkali metal anode research owing to its high electronic conductivity, good chemical stability, and excellent mechanical properties at the atomic scale. Graphene nanosheets have been used as building blocks to construct a variety of porous structures, which can be used as promising hosts for metallic Na. A reduced graphene oxide aerogel (rGa) was synthesized via hydrothermal reduction followed by an oriented freeze-drying treatment (Fig. 24a) [245]. After infusing molten sodium into the porous rGa, a Na/rGa composite anode was obtained and showed excellent flexibility (Fig. 24a). The large specific surface area and uniform pore structure of the rGa can effectively reduce the local current density and inhibit Na dendrite formation (Fig. 24b). Due to the ultralight weight of the porous graphene host, the Na/rGa anodes obtained a high specific capacity of 1 064 mAh  $g^{-1}$ . In symmetric cell testing, the Na@rGa anode exhibited a low overpotential of approximately 50 mV and a stable cycling performance for 1 000 cycles at a high current density of 5 mA  $cm^{-2}$  in a carbonate electrolyte (1.0 M NaClO<sub>4</sub> in EC/DEC). For full cell application, Na@rGa presented enhanced cycling stability when paired with the Na<sub>0.67</sub>Ni<sub>0.25</sub>Mn<sub>0.75</sub>O<sub>2</sub> cathode. Free-standing molten Na-infused reduced graphene oxide (Na@rGO) was also demonstrated to be a promising anode for highenergy Na-based batteries [246]. A porous GO film was first synthesized via a freeze-drying method (Fig. 24c). Then, the gray GO film was immediately reduced to dark black rGO after parallel connection with molten Na, resulting in a more porous structure. The final Na@rGO was obtained after fast Na infusion by simply placing the edge of the rGO film into the molten Na. The Na@rGO anode showed excellent mechanical strength and could be easily processed to assemble flexible batteries. No obvious Na dendrites were observed after cycling for 300 h at a high current density  $(5 \text{ mA cm}^{-2})$  and a high capacity  $(5 \text{ mAh cm}^{-2})$  in a carbonate electrolyte.

The doping strategy can also be used to enhance the "sodiophilic" property of graphene toward Na plating. For instance, boron-doped graphene (BG) was synthesized via the CVD method [247]. DFT calculation results showed that the binding energy of BG is higher than that of N-doped graphene, bare Cu and pristine graphene. The enhanced "sodiophilic" property through heteroatom doping greatly reduced the Na nucleation barrier and facilitated uniform Na plating. The obtained BG/Na anode showed a very good cycling stability up to 300 h at a current density of 5 mA cm<sup>-2</sup> with a capacity limit of 2 mAh cm<sup>-2</sup>. Later, by employing lightweight N-doped graphene microcubes in Na metal anodes, super-stable cycling stability and ultrahigh-rate capability were achieved. With only 3 wt% of the N-doped graphene microcube additive, excellent cycling stability was realized



Fig. 23 a Schematic illustration of the fabrication processes of CNT-Na electrodes, and mechanisms of Na nucleation and stripping on CNT-Na and bare Na electrodes. Reprinted with permission from Ref. [241]. Copyright © 2021, Elsevier. b Schematic diagram of the fabrication procedure of Na@CP-NCNTs and the Na plating/stripping behavior on the surface of pristine Na foil and Na@CP-NCNT skeletons. Reprinted with permission from Ref. [242]. Copyright © 2018, Wiley-VCH. c Schematic illustration of uniform Na deposition on an oxygen-functionalized CNT skeleton. d Na growth behavior on oxygen-functionalized CNT and conventional Cu substrates obtained via DFT calculations. Reprinted with permission from Ref. [243]. Copyright © 2019, Wiley-VCH. e Schematic illustration of Na deposition on the pristine Na metal anode and Na/NSCNT anode. f Digital photos of bent NSCNT paper. g SEM observation of the Na anode and Na/NSCNT anode before and after 20 cycles. h Reprinted with permission from Ref. [244]. Copyright © 2018, Wiley-VCH

at a high current density of 20 mA cm<sup>-2</sup> with 10 mAh cm<sup>-2</sup> in additive-free carbonate electrolytes. When paired with the Na<sub>3</sub>V<sub>2</sub>(PO<sub>4</sub>)<sub>3</sub> cathode, the full cell could maintain a high capacity of approximately 95.2 mAh g<sup>-2</sup> after 1 000 cycles at 5 C (1 C = 110 mA g<sup>-2</sup>).

Introducing metal oxides into graphene skeletons to form composites is another effective way to increase the "sodiophilic" property of porous hosts. Through theoretical modeling, Jin et al. identified the driving force of the "sodiophilic" property-guided Na plating and the influence of interfacial interactions on the cyclability of the sodiated NaF/SnO<sub>2</sub>@rGO system [248]. Furthermore, through experimental investigation, they discovered that the intermediate phases of Na2O and/or Na15Sn4 reduced the nucleation overpotentials and guided uniform Na plating owing to the strong binding energy to metallic Na. In addition, the strong interaction between the SnO<sub>2</sub> nanodots and graphene skeletons reduced the loss of the sodiophilic sites, benefiting the improved cycling stability of Na plating/stripping. This interfacial design strategy provides an effective way to enhance Na metal anodes in Na-based batteries.

# 6 Separator Modification and Interlayer Design

Commercialized polyolefin separators produced by Celgard® and glass fiber (GF) separators are commonly used for RT Na–S and Na–Se batteries. The long-chain sodium polysulfide and polyselenide intermediates formed during discharging are prone to dissolve in conventional LEs and diffuse across both types of separators due to their micron-sized porous structures (Fig. 25a). Polysulfide and polyselenide shuttling leads to severe self-discharge, deterioration of the metal anode and loss of active materials at the cathode, manifesting as a low Coulombic efficiency and rapid capacity decay [251]. Kaskel et al. first achieved an enhanced cycling performance in RT Na-S batteries by adopting a sodiated Nafion (Na-Nafion) coating on a polypropylene (PP) backbone as a cationselective separator [252]. Compared to the porous PP membrane, the nonporous Na-Nafion separator with sufficient Na<sup>+</sup> conductivity has much smaller channels that can efficiently block polysulfide permeation (Fig. 25b) [105]. As shown in Fig. 25c, a Nafion membrane is composed of a hydrophobic poly(tetrafluoroethylene) (PTFE) backbone and a hydrophilic region (4-5 nm) with fixed SO<sup>3-</sup> clusters, which are connected to each other through a hydrophilic channel (1–2 nm in diameter) [253]. These sulfonic groups provide a negatively charged environment, enabling the counter Na<sup>+</sup> to transfer through the hydrophilic pores and blocking the negatively charged polysulfides due to the "charge repulsion" of the Donnan potential (Fig. 25d). Therefore, the RT Na-S batteries using the Na-Nafion separator showed improved cycling stability.

Additionally, most of the inorganic host materials introduced above can be employed to modify separators to improve the electrochemical performance of Na-S and Na-Se batteries. In general, functional materials are directly coated on one side of PP or GF separators via a drop-coating or filtration-assisted method (the obtained coating layer is denoted as an interlayer). The additional interlayers cover the large pores of commercial separators and can directly block the dissolved sodium polysulfides or polyselenides from transferring across the separators. Furthermore, some interlayers containing polar materials can chemically trap intermediates and enable them to be reutilized as active materials [254]. For example, Xu et al. prepared hollow carbon spheres decorated with MoS<sub>2</sub> (HCS/MoS<sub>2</sub>) as both a sulfur host and a separator modifier [255]. The dual cathodeelectrolyte defense system efficiently prevents polysulfide shuttling (Fig. 25e). A similar tactic has also been applied to improve the performance of Na–Se batteries [256, 257].

To maintain the electronic insulating property of separators, the strategies mentioned above are all focused on modification of the cathode side of batteries. The exploitation of the anode side of the separator to further enhance the electrochemical performance has been overlooked. Recently, our group designed and prepared a Janus separator with a singleion-conducting polymer-grafted side and a functional lowdimensional material coated side by a grafting-filtering route for Na–S batteries [103]. A poly(1-[3-(methacryloyloxy) propylsulfonyl]-1-(trifluoromethanesulfonyl)imide sodium) (PMTFSINa) layer was first grafted on one side of the PP separator by a UV-induced surface polymerization method. Subsequently, a defective  $Ti_{3-r}CNT_{v}$  MXene (DN-MXene) was adhered to the other side of the separator through filtration (Fig. 25f). PMTFSINa remarkably enhances the electrolyte wettability, prevents polysulfide diffusion via electrostatic repulsion, and suppresses Na dendrite growth.



◄Fig. 24 a, b Schematic illustration of the fabrication and plating/ stripping behavior of Na@rGa anodes. Reprinted with permission from Ref. [245]. Copyright © 2019, Elsevier. c SEM images of GO film, rGO film and rGO-Na anodes. Reprinted with permission from Ref. [246]. Copyright © 2017, National Academy of Sciences, USA. d Schematic illustrations of Na plating on different matrices, including Cu foil, rGO and partially reduced GO (PrGO). e Tension illustrations of Na plating on planar rGO and PrGO films obtained through mechanical simulation. Reprinted with permission from Ref. [249]. Copyright © 2020, American Chemical Society. f Schematic illustration of the Na nucleation, growth, and further plating process on BG. Reprinted with permission from Ref. [247]. Copyright © 2019, Wiley-VCH. g Schematic of CVD growth of N-doped graphene microcubes on cubic NaCl crystals, and schematic of the Na stripping/plating process with PN-G microcubes as solid additives. Reprinted with permission from Ref. [250]. Copyright © 2020, Elsevier. h Schematic illustrations of Na deposition on NaF/SnO<sub>2</sub>@rGO. Reprinted with permission from Ref. [248]. Copyright © 2020, Elsevier

Meanwhile, the DN-MXene next to the cathode electrocatalytically boosts the polysulfide conversion kinetics. In brief, an ideal separator for Na–S and Na–Se batteries should possess excellent Na<sup>+</sup> selectivity and conductivity, superior isolation of polysulfides and polyselenides and good sodiophilicity (which is discussed in detail in Sect. 6).

# 7 Conclusions and Perspectives

### 7.1 Conclusions

We have comprehensively reviewed the recent progress regarding RT Na–S and Na–Se battery systems. These include the reaction mechanisms, the design of the electrode materials, i.e., sulfur cathodes and Na metal anodes, the optimization of the LEs, and the exploration of novel (quasi-) SSEs. Although significant achievements have been reported, many issues still need to be overcome to realize future commercialization.

The slow reaction kinetics and the shuttle effect of soluble sodium polysulfides/polyselenides are the major challenges in obtaining high-performance cathodes. The rational design of cathode hosts with multifunctionalities can overcome these challenges, including physical and chemical confinement to capture dissolved high-order polysulfides/polyselenides, superior electrocatalytic activity to accelerate the conversion from long-chain to short-chain Na polysulfide/ selenide species, and high conductivity to remedy the low conductivity drawback of the sulfur/selenium element. Several cathode engineering strategies have been demonstrated to be effective for realizing this purpose: (1) constructing a hierarchical porous architecture to tackle the volume change and provide excellent physical confinement, (2) loading precious metal NPs or SAs onto a highly conductive carbonbased host to offer an electrocatalytic function, (3) utilizing polar functional or polymeric sulfur composites to provide strong chemical confinement, and (4) using other sulfurbased composites, which can bypass the use of elemental S to increase the structural stability of the electrode. Notably, a combination of the strategies mentioned above that can synergize individual advantages is highly desirable for achieving the enhanced performance of RT Na–S and Na–Se batteries.

Electrolyte modulation is also key to realizing highperformance Na-S/Se batteries, and here, we generalize the challenges in electrolyte exploration and offer guidelines for future research directions as follows (1) The understanding of the SEI formation on Na metal in LEs should be deepened, and more compatible solvents, electrolyte additives, or Na salts that can assist the formation of a resistant SEI should be identified. In this way, Na dendrites would be largely suppressed, leading to long cycling stability and high Coulombic efficiency. (2) The cathode species dissolution into the LE should be reduced. Several strategies can be further explored, such as developing catholytes and electrolytes in which polysulfides/polyselenides present a low solubility and form a cathode-electrolyte interphase (CEI) layer. (3) Some LEs, i.e., highly concentrated ILs, exhibit high cycling performance and Coulombic efficiency; however, these LEs are usually too expensive for the industrial scale. Reducing the cost of the electrolyte is key for commercialization. (4) An LE can be gelled into a quasi-solid electrolyte to lower cathode species dissolution and dendrite growth propagation. However, polymer synthesis is sometimes complex and costly. Alternative synthesis routes should be explored to enable polymer electrolyte synthesis at a large scale and at an adequate cost. (5) Ultimately, SSEs can resolve the above issues but face other challenges, such as low ionic conductivity at RT, which prevents a high cycling rate, narrow electrochemical stability, which impedes the utilization of high-voltage cathodes, and/or high cost. (6) Additional effort should also be devoted to the safety and eco-friendliness of electrolyte synthesis by prioritizing nonflammable, nontoxic and nonenvironmentally hazardous chemicals. (7) With specific and advanced experiments, such as XPS depth profiling and cryo-TEM, as well as theoretical calculations, further progress is expected to support an in-depth understanding of the electrode interfacial chemistry and electrolyte solvation structure, thereby facilitating electrolyte design.

Reducing parasitic reactions and suppressing dendrite formation are still great challenges in achieving stable and high-performance Na metal anodes. More fundamental and experimental studies are critical to reveal the reaction mechanisms on Na metal anodes, including, practically, the Na plating/stripping behavior on different substrates and the reactions with the dissolved byproducts from the cathodes. Constructing an artificial protective layer on Na metal



**Fig. 25 a, b** SEM images of a porous Celgard separator and a Nafion membrane. Reprinted with permission from Ref. [105]. Copyright © 2015, John Wiley & Sons. **c, d** Architecture and schematic of the ion selectivity of the Nafion membrane. Reprinted with permission from Ref. [253] Copyright © 2016, American Chemical Society. **e** Schematic of the S/C composite (top) and S@HCS/MoS<sub>2</sub> electrode with

modified GFs (bottom) during discharging. Reprinted with permission from Ref. [255]. Copyright © 2019, WILEY-VCH Verlag GmbH & Co. KGaA, Weinheim. **f** Illustration of the fabrication of PMTF-SINa-graftedIDN-MXene-coated Janus separators for RT Na–S batteries. Reprinted with permission from Ref. [103]. Copyright © 2020, John Wiley & Sons

anodes has attracted significant attention and will become even more popular via coating other innovative materials with high ionic conductivity and good mechanical strength. Advanced surface coating technologies such as ALD and MLD will provide more options to improve the coating quality of the artificial protective layers. In addition, designing suitable hosts to accommodate metallic Na is also important to reduce the volume change during continuous Na plating/ stripping. Rational structure design combined with surface functionalization will facilitate Na plating inside these hosts to minimize the volume change.

The low-cost and high-energy properties of Na metalbased rechargeable batteries hold great promise for largescale renewable energy storage. The currently developed sulfur cathodes, modified separators, optimized electrolytes and rationally designed Na anodes show significant improvement toward high-performance and long-cycle-life Na RT Na–S and Na–Se batteries. Although there is still a long way to go to before the practical application of RT Na–S and Na–Se batteries, the fast-developing advanced technologies would definitively accelerate this process.

### 7.2 Perspectives

According to the above discussion, significant progress has been made in the development of high-performance RT Na–S(Se) batteries for grid-scale SEES systems. However, there are still many challenges that need to be overcome before their practical application. Here, we propose a few potential research directions to promote the development of these systems.

(A) Increasing the sulfur loading and sulfur utilization.

Currently, the sulfur loading and cycling performance of RT Na-S batteries are contradictory. Normally, the sulfurbased cathode used in RT Na-S batteries can only have an approximately 35%-50% sulfur loading to achieve high electrochemical performance. With a higher sulfur loading, the cycling performance will be worse for the RT Na-S batteries because the state-of-the-art cathode hosts still cannot confine a higher amount of sulfur to prevent the shuttling of Na polysulfides. Many cathode hosts for sulfur-based cathodes used in RT Na–S batteries were taken from the Li–S battery system. These hosts may be good enough to confine the Li polysulfides. However, they cannot effectively limit the longchain soluble Na polysulfides that have much more active chemical characteristics. The cathode hosts, whether based on 3D porous carbon or a 2D material, for RT Na-S batteries need to be further developed in terms of the sulfur confinement function to realize a high sulfur loading, which is very important for reducing the cost and increasing the volumetric energy density of the batteries. Another aspect that needs to be mentioned is sulfur utilization. Some cathode hosts with good physical or chemical confinement of sulfur will combine with sulfur or Na polysulfides and no longer release them. This factor directly leads to loss of the active material (sulfur) and low initial Coulombic efficiency of the RT Na-S batteries. Hence, the development of a cathode host for RT Na-S battery cathodes in future will need to have the ability to confine sulfur and Na polysulfide as well as the ability to release them during the charging process.

(B) Boosting the conversion reaction kinetics of sodium polysulfides.

Another research path is increasing the reaction kinetics of short-chain sodium polysulfides. Unlike active longchain Na polysulfides (e.g.,  $Na_2S_4$ ,  $Na_2S_6$ ), short-chain solidstate Na polysulfides (e.g.,  $Na_2S_2$ ,  $Na_2S$ ) are very passive and hard to transform back to long-chain Na polysulfides, which will also incur the loss of the specific capacity. If the redox kinetics of short-chain sodium polysulfides can be increased, especially  $Na_2S_2 \leftrightarrow Na_2S$ , then the reaction overpotential of the RT Na–S batteries can be greatly reduced and the capacity and cycling performance thus significantly improved. This improvement could be realized with further development of cathode hosts with effective electrocatalytic functions. Moreover, catalytic functions to accelerate the reaction kinetics of short-chain Na polysulfides can also be developed by adding additives or optimizing the composition of the electrolyte, which will also be a promising direction in future.

(C) Developing solid-state RT Na-S batteries.

The development of solid-state RT Na–S batteries will be a promising strategy to fulfill the safety expectations of stationary energy storage. The safety concerns arising from LEs, such as leakage, fire, or even explosion, which may contaminate the environment or threaten communities, require configuration optimization of the SEES system. RT Na–S batteries with SSEs can prevent all these risks and effectively enhance the practical application of RT Na–S batteries in different extreme environments, such as at extremely high temperatures or under abuse usage. However, many challenges, such as the sluggish ion transport inside the SSE and the uneven solid-to-solid interfacial contact between the solid electrolytes and electrodes, still need to be overcome to ensure the stable operation of solid-state RT Na–S batteries before real application.

#### (D) Optimization of the separator, interlayer and binder.

Compared with Li-based batteries, research on other components, including separators, interlayers, and binders, in RT Na-S(Se) batteries is still in its infancy. Currently, separators based on GFs are mainly used in the RT Na-S battery system. To prevent the shuttle effect and enhance the confinement of sulfur, novel types of separators or interlayers based on 2D materials or 3D frameworks with electrocatalytic functions can be potential alternatives to GFs. Moreover, the solvent NMP used with the PVDF binder is toxic and nonenvironmentally friendly. The development of biocompatible and aqueous binders derived from, for example, wood or fertilizer that are based on a water-based slurry can effectively reduce the cost, confine the Na polysulfides on the cathode side, act as a flame retardant to reduce the safety risks, and be environmentally benign, which is worth further investigation in future. These research paths of separators, interlayers, and binders all have a bright future to further enhance the performance of the RT Na–S(Se) battery system.

Although there are still many challenges that need to be overcome for RT Na–S(Se) batteries, considering that research of RT Na–S(Se) batteries has already become one of the hottest topics and the extremely inexpensive and abundant raw materials, as well as the high operating safety, RT Na–S(Se) batteries are predicted to have significant breakthroughs in the near future and obtain a considerable part of the market share in the grid-scale energy storage field.

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Zefu Huang graduated from the University of Sydney and obtained a Master of Professional Engineering (Chemical and Biomolecular) degree in 2019. Currently, he is a Ph.D. candidate in the Centre for Clean Energy Technology (CCET) at the University of Technology Sydney (UTS), Australia. His research interests include sodium-ion batteries, sodiumsulfur batteries, and 2D materials



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Dong Zhou is an assistant professor and Ph.D. supervisor at Tsinghua Shenzhen International Graduate School. He received his Ph.D. degree from Tsinghua University in 2017, and then worked as a postdoctoral research associate in the University of Technology Sydney, Australia and the University of Tokyo, Japan. His research topic mainly focuses on the design and synthesis of specialized solid polymer electrolytes and nanoelectrode materials for various advanced energy storage devices

and investigation of the interfacial mechanisms. He is the recipient of the First Prize of Guangdong Natural Science in 2020, the Discovery Early Career Researcher Award (DECRA) and the Japan Society for the Promotion of Science (JSPS) Fellowship.



Pauline Jaumaux is currently a Ph.D. candidate at the Center for Clean Energy Technology at the University of Technology Sydney. Her research focuses on electrolyte engineering including nonflammable liquid, quasi-solid electrolytes and solid-state electrolytes for alkali and multivalent metal-based batteries.





Bing Sun received his Ph.D. degree in 2012 at the University of Technology Sydney (UTS), Australia. Currently, he is a research associate in the Centre for Clean Energy Technology at UTS. His research interests focus on the development of next-generation battery materials and technology for lithium-based batteries and sodium-based batteries. He is the recipient of Australian Research Council (ARC) Discovery Early Career Researcher Award (DECRA).

Xin Guo is currently a Research Fellow at the Centre for Clean Energy Technology, University of Technology Sydney (UTS). He received his Bachelor's degree in materials physics from Lanzhou University in 2011 and completed his Ph.D. degree from the Faculty of Science at UTS in 2019. His research interest mainly focuses on nanostructured functional materials for alkali-ion batteries, Li-O2 batteries, and Na-S/Se batteries.



composite buffer layers for anode/cathode protection, SEI building additives and sacrificial salts for post-lithium batteries.



Michel Armand received his Ph.D. degree in Physics in 1978 from the University of Joseph Fourier. He has been the Directeur de Recherche at Centre National de la Recherche Scientifique (CNRS) since 1989 and Professor at the University of Montreal (1995-2004). He pioneered several theoretical concepts and practical applications in the field of energy-related electrochemistry. He ushered the concept of intercalation compounds (inorganic 1972, organic 1996) and the proposition of

polymer electrolytes for battery application (1978), followed by the introduction of new families of highly conductive salts (perfluoroimides like TFSI and FSI) for liquid and polymer electrolytes (1986). Since 2013 he is a Group Leader at CIC Energigune.

Devaraj Shanmukaraj obtained his Ph.D. degree (Physics) degree with a specialization on Solid State Ionics in 2008 from Pondicherry University, India, after completing his Master degree in Materials Science and Technology. From 2013 till date, he is working as a Senior Researcher Scientist at CIC Energigune, Spain, to carry out research on solid-state Li/Na metal polymer batteries and Li/S batteries. His expertise includes novel polymer, hybrid and liquid

electrolytes for Li/Na batteries,



Teofilo Rojo received his Ph.D. degree in chemistry from the University of the Basque Country in 1981. He became Full Professor of Inorganic Chemistry at the UPV/EHU in 1992. His research has been focused on Solid State Chemistry and Materials Science. Since 2010 he is the Scientific Director of CIC Energigune developing materials research for advanced batteries (lithium, sodium, etc.). In 2015 he was appointed as an Academic Member of the Royal Spanish Academy of Exacts,

Physical and Natural Sciences, and in 2016 he was named member of the Working Party on Chemistry and Energy of EuCheMS (European Chemical Science).



Guoxiu Wang is the Director of the Centre for Clean Energy Technology and a Distinguished Professor at the University of Technology Sydney (UTS), Australia. His research interests include lithium-ion batteries, lithium-air batteries, sodium-ion batteries, lithium-sulfur batteries, supercapacitors, hydrogen storage materials, fuel-cells, 2D materials, and electrocatalysis for hydrogen production. Professor Wang has published more than 650 refereed journal papers with an H-index of 134. His pub-

lications have attracted over 59 800 citations. He has been listed as a highly cited researcher in both Materials Science and Chemistry by Web of Science/Clarivate Analytics in 2018 and a highly cited researcher in Materials Science in 2019, 2020 and 2021.