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### **REVIEW**

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# Inorganic solid electrolytes for all-solid-state lithium/sodium-ion batteries: recent developments and applications

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The rapid evolution in electrolyte engineering has significantly propelled the development of synthesis and the precise tailoring of the properties of inorganic solid electrolytes (ISEs). These advancements are crucial to meeting the stringent performance requirements of high-performance all-solid-state batteries (ASSBs). This review comprehensively summarizes recent progress in the synthesis techniques and electrochemical characteristics of various ISEs, including oxides, sulfides, hydroborates, antiperovskites, and halides, highlighting their applications in ASSBs. Additionally, we review key challenges in ASSB development, such as the limited compatibility between ISEs and electrodes, and the detrimental interfacial reactions. Strategies to overcome these challenges, including the use of composite cathodes and solid interface layers, are discussed. Finally, we present current ASSB models and propose emerging approaches driving the future development of ASSBs for the next generation of energy storage solutions.

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#### Introduction 1.

Battery storage technology plays an essential role in cuttingedge technologies in information and communication technology, transportation, healthcare, defense, security, and other related industries.1-4 Batteries with high energy density, the ability to operate safely at extreme temperatures, and long-term stability are essential for these applications.5-7 We can achieve



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the aforementioned objectives by concentrating on enhancing battery components with exceptional performance. Recent research interest in battery storage technology has centered on the production of solid-state electrolytes with high conductivity. Liquid electrolytes in LIBs and SIBs are less reliable at low temperatures, the viscosity of the solution increases and the ionic mobility of the liquid electrolyte decreases. Overcharging also raises the risk of explosions and fires because of the flammable electrolytes and solvents present, as well as the release of harmful gases and pollution into the environment. Hence, switching liquid electrolytes to solid-state electrolytes and broadening the utilization of all-solid-state batteries are capable of solving these problems.

All-solid-state batteries that incorporate ISEs are considered to be at the forefront of battery storage technology. These batteries provide both high energy density and dependable security systems.<sup>25,26</sup> Inorganic solid electrolytes demonstrate numerous advantages compared to liquid electrolytes. These include a broad electrochemical window, excellent air stability, high ionic conductivity at room temperature, superb electrochemical stability, good deformability, compatibility with

electrode materials, and being environmentally friendly. 1,27-29 To construct ASSBs, it is necessary to use electrolytes based on solid-state ion conductors, which must fulfill the previously mentioned requirements. Nevertheless, there is currently no solid-state ion conductor that exhibits the capability to fulfill all of these specified criteria. 30-32 The solid-state ion conductors are constructed via a combination of metal and non-metal ions, generating a polyhedral network. Polyhedral structures exhibit patterns based on the angular orientation of their positions. The crystal structure and the variety of defects formed within it determine the conductivity of the crystal.33 Some examples of solid-state ion conductors are Garnet, 34-38 NASICON, 39-41 halide,2,30,42-46 sulfide,47-49 hydroborate,50-52 and perovskite.24,31,53 High conductivity in solid-state crystals is achieved when the number of vacancies for mobile ions exceeds the number of mobile ions, the activation energy between adjacent vacancy sites is sufficiently low to allow ion diffusion, and the channels for ion diffusion are linked. 17,51,54 The challenges associated with the applications of ISEs in ASSBs include insufficient wettability and inadequate interfacial compatibility between the electrode and solid electrolytes, issues with



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in garnet-type solid electrolytes.

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electronic and ionic conductivities, fluctuations in the electrochemical window of solid electrolytes that do not meet the cathode potential, and issues related to fabrication time and cost.55

Today's goal in solid-state electrolyte research for ASSBs applications is to create solid-state electrolytes with superior performance compared to conventional liquid electrolytes.<sup>2,56–58</sup> Improved electrochemical performance, such as the electrochemical stability window and ionic conductivity of inorganic solid electrolytes, may be obtained by modifying the crystal structure framework, doping with cations and anions, and improving synthesis techniques and conditions. 16,37,42,59-64 Liang et al.56 presented the compositions and crystal structures of a ternary halide solid electrolyte,  $\text{Li}_{3-3x}\text{M}_{1+x}\text{Cl}_6$  (-0.14 <  $x \le$ 0.5, M = Tb, Dy, Ho, Y, Er, Tm). For example, the orthorhombic structure of Li<sub>2.73</sub>Ho<sub>1.09</sub>Cl<sub>6</sub> achieves a conductivity higher than that of the trigonal structure of Li<sub>3</sub>HoCl<sub>6</sub>. The change from a trigonal to an orthorhombic structure makes it much easier for Li-ions to move around and lowers the activation energy in different L-M-Cl systems. The electrochemical window for Li<sub>2,73</sub>Ho<sub>1,09</sub>Cl<sub>6</sub> and other inorganic solid electrolytes is seen in Fig. 1.

The selected method of synthesis has a significant impact on several characteristics of the ISEs, including particle size, material density, lattice parameter, and crystal defect. Furthermore, it also affects the mechanical and electrochemical properties of the solid electrolyte. 19,79,80 Ma et al.81 utilized freezedrying technology and heat treatment to produce Li<sub>3</sub>InCl<sub>6</sub> electrolytes. This method resulted in 80% of the particles having a size smaller than 200 nm, a uniform distribution of particle sizes, and the elimination of solvents during the solidification of the electrolyte solution. Additionally, this approach successfully solved the issue of particle agglomeration caused by the growth of particles and high temperatures during the thermal evaporation process of the solutions. It is believed that particle size has a significant effect on the charge transport capabilities of composite cathodes and interface contacts in ASSBs. 82,83 The growth of lithium dendrites within the pores and

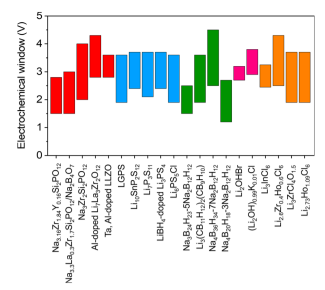


Fig. 1 Summary of electrochemical windows for the selected ISEs in ASSBs. 6,39,41,56,57,59,65-78

cracks of electrodes is a significant challenge for ASSBs and can even impair the integrity of solid electrolytes.84,85 To solve this issue, increased density and homogenization of the solid electrolyte are essential to suppressing lithium dendrite growth. 23,40,86-88 Li et al. 23 proposed a two-stage sintering technique to enhance the homogeneity and density of a sulfide-type solid electrolyte (Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub>). The initial powder is subjected to annealing and subsequently densely packed into pellets. The Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub> undergoes re-annealing by controlled modifications of the annealing temperature and time. The treatment had significant impacts on the conductivity of Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub>, leading it to increase to 8.04 mS cm<sup>-1</sup> at room temperature. Furthermore, optimal heat treatment reduces the resistance at the grain boundaries, enhances the durability from cracking, and increases the ionic conductivity of Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub>. One-step synthesis and two-step synthesis that have been applied to date include ultrafast high-temperature sintering,39 calcination + cold



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sintering process + annealing, s3 high-pressure low-temperature, s9 spark plasma sintering + annealing, s0 sol-gel + calcination, s1 liquid-phase method + wet milling + sintering, liquid-phase method + annealing, s1 ion exchange method, s9 wet chemical + low-temperature heating, s16, s2 radio frequency magnetron sputtering, s3 ball-milling + annealing/sintering, s22,24,52 two-step milling, s1 two-step annealing, s3 meltquenching + high-energy ball-milling, s4 solid-gas reaction, s5 etc.

Understanding the behavior of the electrode-ISE interface is crucial for achieving high-performance ASSBs. This includes investigating the diffusion of Li-ions at the interface, chemical reactions that occur at the interface, changes in the contact area between the electrode and electrolyte, changes in the volume of the electrode material, and concerns about poor compatibility leading to the formation of a high-resistance passive layer during the charge-discharge cycle.76,96,97 Completely eliminating the electrode/ISE interface reaction is not possible. However, the interface reaction can be controlled by employing coating layers and modifying the composition of composite electrodes.98-100 For instance, sulfide solid electrolytes exhibit excellent conductivity within the ISE class. 25,79,101 However, when utilized in ASSBs, they meet challenges such as decomposition at high voltages, interfacial reactions, the formation of a space-charge layer between the ISEs and the electrode, and a reduction in capacity and voltage due to structural deterioration on the surface and grain boundaries of cathode particles. 68,102-104 The purpose of applying a coating to the electrode surface is to prevent the spontaneous reaction between ISEs and electrodes. Additionally, the coating helps to minimize the oxidative decomposition of the ISE, prevent significant changes in the contact area and reduce the formation of cracks, maintain strong adhesion between the electrode and the ISEs, and suppress the growth of lithium dendrites. 42,73,91,97,105

The interface layers should possess excellent lithium conductivity, electron-blocking capabilities to hinder redox reactions in the ISEs, high surface energy to inhibit the formation of lithium dendrites, and compatibility with both the electrode and ISEs. 1,66 The electrode/ISE interface layers often employed include LiNbO<sub>3</sub>, 42,103 Li<sub>3</sub>PO<sub>4</sub>, 56,106 SnO<sub>2</sub>, 107 ZrO<sub>2</sub>, 108 Li<sub>3</sub>N-LiF,<sup>97</sup> ZnO-LiF,<sup>1</sup> carbon,<sup>91</sup> etc. ISEs that have been applied as an interface layer, such as Li<sub>7.5</sub>La<sub>3</sub>Zr<sub>1.5</sub>Co<sub>0.5</sub>O<sub>12</sub>,<sup>76</sup> Li<sub>6</sub>PS<sub>5</sub>-Cl, 22,42,67,75 and Li<sub>3</sub>YCl<sub>6</sub>.57 The coating method is essential to achieving a uniform coating distribution. The coating material must possess plastic properties if it covers the entire surface of the cathode and ISEs. When fabricating the electrodes using mechanical compression tools, this is necessary to prevent the coating from becoming brittle. Additional research is required to investigate the characteristics of coatings and the techniques used to apply them to electrodes, as well as their substantial influence on ASSBs.105

In ASSBs, the composite cathode is usually made up of conductive carbon, CAM, and catholyte (ISEs).<sup>50,54</sup> This composition will have a highly intricate interface system, specifically CAM/carbon, carbon/catholyte, and CAM/catholyte. The utilization of single cathode active materials like LCO, LTO, LFP, NCM622, NCM811, NCM83, NCM85, NCM88, and NCM90 is deemed disadvantageous due to the occurrence of volume

changes and structural degradation during the charge-discharge process. These factors result in a reduction in the capacity of the cathode materials. 103,109 The purpose of incorporating conductive carbon is to enhance the electrical conductivity of the electrode, improve the contact between the CAM and ISEs, and preserve the structural integrity of the electrode during the charge-discharge process. 39,41,49,78,110,111 On the other hand, the presence of ISEs in the composite cathode acts to decrease the resistance at the interface of the composite cathode, enhance Li-ion transport, and enhance the performance of the cathode. 75,112 To decrease the resistance at the anode/ISEs interface in the case of lithium or sodium anodes, a protective layer can be applied, or the metal can be modified into an alloy such as Li-In, 25,65,66,71,80 Li-Si, 75 and Na-Sn. 30

In recent years, numerous review articles have focused on inorganic solid electrolytes for ASSBs. 113-116 However, a scarcity of articles exists that provide a comprehensive review of ISEs, present designs of ASSBs, and prospective strategies therein. Through reviewing the most recent developments in the study of inorganic solid electrolytes. Herein, recent findings in the field of inorganic solid electrolytes such as oxide, sulfide, hydroborate, antiperovskite, and halide are described. This section discusses current synthesis and modification techniques that affect the crystal structure of inorganic solid electrolytes, as well as the impact of doping on electrochemical stability, ionic conductivity, and other characteristics. The stability of each ISEs toward the anode and cathode is then explained, along with several of the current techniques to reduce the shortcomings produced. In discussing the application of ISEs in all-solid-state batteries, especially for Li-ion batteries and sodium-ion batteries. Furthermore, this review summarizes the potential of ISE classes implemented as catholytes and solid interfacial layers to enhance the electrochemical stability of ASSBs and minimize electrode instability. Finally, we provide an overview of the various configuration models for ASSBs and discuss the future progress of ASSBs.

# 2. Overview of inorganic solid electrolytes (ISEs)

ISEs are regularly updated to fulfill the specifications of nextgeneration ASSBs due to their tremendous potential. This chapter discusses 5 groups of ISEs, including oxides (Garnet and NASICON), sulfides (lithium thiophosphate, thio-LISICON, and argyrodite), halides, hydroborates, and antiperovskites. The following topics will be included: an overview of ISEs, the basic structures, the impact of metal and additive doping, synthesis techniques, the chemical and electrochemical stability of ISEs, and their compatibility with anodes and cathodes.

#### 2.1. Oxide-based ISEs

Inorganic solid-state electrolytes based on oxide materials are commonly classified into several structural types, including LISICON-like, NASICON-like, perovskite, garnet, and  $\beta$ -Al<sub>2</sub>O<sub>3</sub>. $^{62,117,118}$  These oxide-based ISEs are renowned for their wide electrochemical stability, good chemical stability, and

high ionic conductivity which make them attractive candidates for advanced energy storage applications. 35,36,98,119-125 However, despite these advantageous properties, they suffer from significant limitations that hinder their overall performance. Notably, the inherent brittleness of materials, while the challenge of achieving intimate solid-solid contact at the electrolyte-electrode interface further exacerbates their performance deficiencies. 83,126 These issues are critical obstacles that must be

addressed to fully realize the potential of oxide-based ISEs in practical applications.

2.1.1. Garnet. In 2003, Thangadurai et al. 127 discovered a Listuffed garnet-type solid electrolyte,  $Li_5La_3M_2O_{12}$  (M = Nb<sup>5+</sup> or Ta<sup>5+</sup>), which exhibited a room-temperature Li<sup>+</sup> conductivity of approximately  $10^{-6}$  S cm<sup>-1</sup>. This finding marked a significant advancement in the development of solid electrolytes for Limetal batteries. Subsequently, in 2007, the material was

# Garnet solid electrolytes

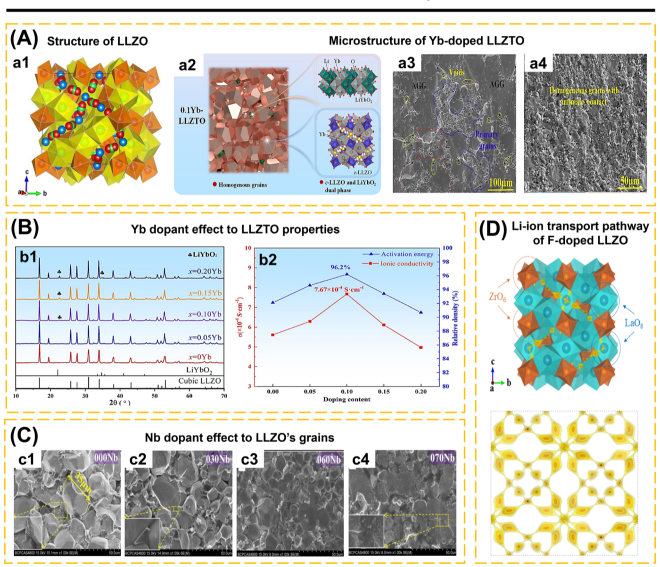


Fig. 2 (A) (a1) Crystal structure of cubic LLZO. The yellow is dodecahedrally coordinated La<sup>3+</sup> (at the Wyckoff position 24c) and the orange is octahedrally coordinated Zr<sup>4+</sup> (16a). The blue spheres correspond to tetrahedrally coordinated (24d) Li<sup>+</sup>, the green spheres to octahedrally coordinated (48g) Li<sup>+</sup>, and the red ones to distorted 4-fold coordinated (96h) Li<sup>+</sup>. Reproduced with permission from ref. 135. Copyright 2014, American Chemical Society. (a2) Microstructural schematic analysis of further Yb-doped LLZTO. SEM images of the cross-sectional grain boundary microstructure of xYb-LLZTO electrolytes: 0Yb-LLZTO (a3) and 0.10Yb-LLZTO (a4). Reproduced with permission from ref. 86. Copyright 2024, Elsevier. (B) (b1) XRD patterns and peak shift of xYb-LLZTO (x = 0, 0.05, 0.10, 0.15, 0.20) in the range of 25–32°. (b2) Ion conductivity ( $\sigma$ ) and relative density of xYb-LLZTO (x = 0, 0.05, 0.10, 0.15, 0.20) at 25 °C. Reproduced with permission from ref. 86. Copyright 2024, Elsevier. (C) SEM images of the cross-sections of the Nb-LLZO pellets. Reproduced with permission from ref. 136. Copyright 2018, Elsevier. (D) BVSE analysis of Li-ion transport pathway in the garnet structure: LLZO-F0.2. Reproduced with permission from ref. 137. Copyright 2024, American Chemical Society.

further modified by replacing the pentavalent cations (Nb<sup>5+</sup>, Ta<sup>5+</sup>) with tetravalent cation (Zr<sup>4+</sup>), leading to the formation of Li<sub>7</sub>La<sub>3</sub>Zr<sub>2</sub>O<sub>12</sub> (LLZO), a compound that has since become a cornerstone in the field of oxide solid-state electrolytes. 128 LLZO has garnered considerable attention as one of the most promising candidates for Li-metal batteries, primarily due to its combination of high ionic conductivity and chemical stability.129 Specifically, LLZO distinguishes itself with its resistance to reduction when in contact with a Li metal anode and compatibility with conventional cathode materials, a property that many other solid electrolytes lack. 128,130,131 The ionic conductivity of cubic LLZO, which can reach up to 10<sup>-3</sup> S cm<sup>-1</sup> at room temperature, is particularly noteworthy. This level of conductivity is comparable to that of commercial liquid electrolytes, underscoring LLZO's potential for practical applications in solid-state battery technology. 128,132,133 Furthermore, LLZO's thermal properties (1.45-1.55 W m<sup>-1</sup> K<sup>-1</sup>) add an additional dimension to its utility. This characteristic not only makes it suitable as a solid electrolyte but also enhances its potential use as a thermal insulation material, effectively reducing heat transfer within the battery system. Such properties underscore the versatility and promise of LLZO in advancing the performance and safety of next-generation solidstate batteries.134

The crystal structure of cubic LLZO (Fig. 2a1), exemplifies the garnet-type framework, characterized by its intricate arrangement of dodecahedral LaO<sub>8</sub> and octahedral ZrO<sub>6</sub> units. These units serve as the backbone of the structure, establishing a robust and stable matrix. Within the cubic  $Ia\bar{3}d$  space group, Li-ions are distributed across two distinct crystallographic sites: the tetrahedral (24d sites, referred to as Li1 sites) and octahedral site and off-centered octahedral site (48g/96h sites, referred to as Li2 sites). Meanwhile, La, Zr, and O occupy sites 24c, 16a, and 96h respectively. The Li1 sites demonstrate a remarkably high occupancy rate of 0.94, the highest recorded among garnet-type oxides, indicating a strong preference for Li-ions to reside within these tetrahedral sites. In contrast, the Li2 sites exhibit partial occupancy, with an occupancy value of 0.35, and are associated with significant positional disorder. This pronounced disorder at the Li2 sites is posited to play a critical role in the high ionic conductivity characteristic of cubic LLZO. Specifically, the partial occupancy and the dynamic nature of the Li2 sites likely facilitate the fast Li-ion transport within the structure, thereby enhancing the material's performance as a solid electrolyte.36,37,138

Enhancing the stability of the cubic structure, density, ionic conductivity, and other characteristics of LLZO may be accomplished by cation or anion doping. Several studies have been conducted on doping and its impact on their properties:

- (1) Cation substitution on Li sites such as Al(III), Ga(III), Fe(III),  $^{139}$  Ge(IV).  $^{140}$
- (2) Cation substitution on La sites such as Yb( $\rm III$ )-doped LLZTO.  $^{86}$
- (3) Cation substitution on Zr sites such as Ta(v), <sup>141</sup> Nb(v), <sup>136</sup> W(vI), <sup>121</sup> Sc(III), Mg(II), <sup>142</sup> Sb(v), Mo(VI), <sup>139</sup> W(VI). <sup>143</sup>
- (4) Double cation substitution on Zr sites such as Y(III), Nb(v). <sup>119</sup>

- (5) Double cation substitution on La and Zr sites such as Sr(II), Sb(V) co-doped LLZO, <sup>36</sup> Ca(II), Ta(V) co-doped LLZO, <sup>35</sup> Sr(II), Mo(VI) co-doped LLZO, <sup>144</sup> Ce(III), Bi(III) co-doped LLZO. <sup>145</sup>
- (6) Double cation substitution on Li and Zr sites such as Ga(III), Mg(II) co-doped LLZO.<sup>146</sup>
  - (7) Anion substitution on O site such as F.147

Li<sub>7-x</sub>La<sub>3</sub>Zr<sub>2-x</sub>Ta<sub>x</sub>O<sub>12</sub> (LLZTO) is a Ta-substituted LLZO derivative on the Zr site with high conductivity making it a frequently investigated SE. When x = 0 and x = 0.6, the LLZO pattern shows a single tetragonal and cubic phase, respectively. Unlike the LLZO pattern observed when  $(0.2 \le x \le 0.5)$ , which exhibits a combination of tetragonal and cubic phases. Li<sub>6.5</sub>- $La_3Zr_{1.5}Ta_{0.5}O_{12}$  (with x = 0.5) shows the highest ionic conductivity. The presence of Ta content in the range of x = 0.3to 0.5 leads to a displacement of the Li2 atom at the 96h site, resulting in a reduction of the Li-Li distance from 1.73 Å to 1.50 Å. The precise location of the Li2 atom plays a crucial role in the significant ionic conductivity observed in the LLZO structure. 64 The conductivity of Li<sub>6.5</sub>La<sub>3</sub>Zr<sub>1.5</sub>Ta<sub>0.5</sub>O<sub>12</sub> will vary according to the production process used, although having the same composition.64,86,148 The improvisation of LLZTO was carried out due to the low ionic conductivity, low lithium dendrite inhibition, and AGG during preparation which was still difficult to avoid. Lu et al.86 incorporated a Yb2O3-based dopant into the LLZTO structure to manipulate the microstructure of LLZTO (Fig. 2a2). This dopant plays a crucial role in suppressing AGG and enhancing lattice parameters by forming YbO<sub>8</sub> in the LaO<sub>8</sub> site. The Yb-O bond distance in this area is greater than that of La-O, facilitating the migration of Li-ions. Additionally, the formation of a second phase (LiYbO2) in the grain boundary region enables efficient Li-ion transport, resulting in a significant increase in the conductivity of the LLZO electrolyte. The relative density reached 96.2% and the conductivity was measured at  $7.67 \times 10^{-4} \text{ S cm}^{-1}$  when the Yb doping concentration was increased to 0.1. The microstructure noticed a significant drop in both density and conductivity when the Yb concentration reached x = 0.2. SEM images of 0Yb-LLZTO reveal that AGG is widespread and comprises disconnected grains, together with gaps between the grains, resulting in decreased density and conductivity (Fig. 2a3). Elevating Yb concentration to 1.0 results in the suppression of voids and aggregation, leading to a uniform morphological condition (Fig. 2a4). XRD elucidates the appearance of diffraction peaks corresponding to LiYbO<sub>2</sub> for Yb concentrations of 0.1 and 0.15 at 25°, and for Yb = 0.2 at 25° and 32° (Fig. 2b1). The rise in Yb concentration correlates with the elevation of LiYbO<sub>2</sub> phase concentration at the grain boundary, which reduces intergranular contact, leading to a reduction in density and ionic conductivity (Fig. 2b2).

The single and double doping of cations at  $Zr^{4+}$  sites in LLZO significantly affects its conductivity. Zhao  $et~al.^{136}$  explained that the substitution of  $Zr^{4+}$  (0.79 Å) with Nb<sup>5+</sup> (0.70 Å) resulted in a reduction of the crystal lattice from 12.9673 Å (0.1 Nb-doped LLZO) to 12.9136 Å (0.7 Nb-doped LLZO) when Nb concentration increased. The SEM photograph indicates an escalation in transgranular fractures from 000Nb to 070Nb, characterized by variation particle sizes that are interconnected without

# NASICON solid electrolytes

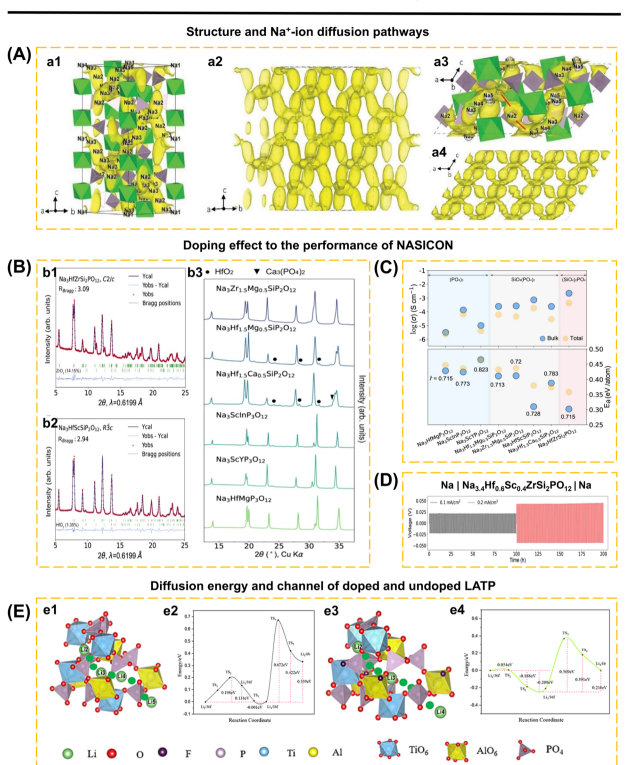


Fig. 3 (A) Na-ion diffusion pathways in monoclinic and rhombohedral Na<sub>3</sub>Zr<sub>2</sub>Si<sub>2</sub>PO<sub>12</sub> from AIMD and MEM. (a1 and a2) The Na-ion probability trajectory, and no Na2-Na2 diffusion is observed. The polyhedra are not shown in (a2). (a3 and a4) Na-ion probability density isosurfaces (yellow) of monoclinic Na<sub>3</sub>Zr<sub>2</sub>Si<sub>2</sub>PO<sub>12</sub> at 1400 K from AIMD simulation. The red lines in (a3) indicate the [101] direction, which is the rate-limiting step in the Na<sup>+</sup>-ion migration process. The polyhedra are not shown in (a4) (green:  $ZrO_6$  octahedra, light purple:  $Si(P)O_4$  tetrahedra, yellow: Na atoms). Reproduced with permission from ref. 149. Copyright 2019, Wiley-VCH. (B) XRD patterns of the as-synthesized NASICONs. Rietveld refinement results of synchrotron XRD data of Na<sub>3</sub>HfZrSi<sub>2</sub>PO<sub>12</sub> with monoclinic C2/c symmetry (b1) and Na<sub>3</sub>HfScSiP<sub>2</sub>O<sub>12</sub> with rhombohedral R3̄c

discernible voids (Fig. 2c). Elevating Nb concentration reduces the average particle size of LLZO. The magnification in Fig. 2c1c3 clarifies the transition from amorphous intergranular fracture to a smooth transgranular fracture. It is inferred that elevated Nb concentration affects particle borders, enhancing Li-ion transport inside the grain boundary area. For 0.25 Nbdoped LLZO (Li<sub>6.75</sub>La<sub>3</sub>Zr<sub>1.75</sub>Nb<sub>0.25</sub>O<sub>12</sub>) shows an ionic conductivity of  $8.07 \times 10^{-5} \text{ S cm}^{-1}$  at 30 °C, while 0.6 Nb-doped LLZO shows an ionic conductivity of  $5.22 \times 10^{-4} \, \mathrm{S \, cm^{-1}}$  at 30 °C. Gai et al.119 substituted Zr4+ sites by double doping with Nb5+ and Y<sup>3+</sup> in Li<sub>7</sub>La<sub>3</sub>Zr<sub>2-2x</sub>Nb<sub>x</sub>Y<sub>x</sub>O<sub>12</sub>. This finding indicates that Nb<sup>5+</sup> doping facilitates the formation of Li vacancies, while Y3+ enhances the concentration of Li-ions. The SEM figures indicate that the grains of undoped LLZO exceed 20 µm. Upon the introduction of Nb, the particle size of 0.25 Nb-doped LLZO  $(Li_{6.75}La_3Zr_{1.75}Nb_{0.25}O_{12})$  ranges from 5 to 20  $\mu m$ . The ionic conductivity of Li<sub>6.75</sub>La<sub>3</sub>Zr<sub>1.75</sub>Nb<sub>0.25</sub>O<sub>12</sub> is measured at 4.82  $\times$ 10<sup>-4</sup> S cm<sup>-1</sup> at 30 °C, surpassing the findings of Zhao et al., <sup>136</sup> indicating that the preparation method influences the properties of Nb-doped LLZO. This topic is further discussed in Section 3.1. In the case of 0.5 Nb/0.5 Y co-doped LLZO (Li<sub>7</sub>La<sub>3</sub>- $Zr_{1.5}Nb_{0.25}Y_{0.25}O_{12}$ ), all the grains measure less than 10 µm. It is determined that Nb and Y co-doping influences the reduction of grain size, while simultaneously impacting an increase of grain boundaries. Elevating the doping concentration of 0.5 Nb/0.5 Y co-doped LLZO (Li<sub>7</sub>La<sub>3</sub>ZrNb<sub>0.5</sub>Y<sub>0.5</sub>O<sub>12</sub>) indicates that, despite the grain size being less than 10 µm, the grains exhibit interconnectivity. Li<sub>7</sub>La<sub>3</sub>ZrNb<sub>0.5</sub>Y<sub>0.5</sub>O<sub>12</sub> has a conductivity of 8.29  $\times$  $10^{-4} \text{ S cm}^{-1} \text{ at } 30 \text{ }^{\circ}\text{C}.$ 

Anion replacement at the oxygen position in the garnet structure has been examined. Lu et al. 147 replaced the oxygen anion with fluorine in  $Li_{6.25}Ga_{0.25}La_3Zr_2O_{12}$  (LGLZO). Fluorine doping results in a stiffer LGLZO structure with decreased atomic displacement. Fluorine substitution in the anionic sublattice enhances the strength of La-(O,F) and Zr-(O,F) bonds. The Li-ion conductivity of F-LGLZO is superior to that of LGLZO, measuring 1.28 mS cm<sup>-1</sup> compared to 0.543 mS cm<sup>-1</sup>, respectively. The Li|SPE-F-LGLZO-SPE|Li symmetric cell can operate for up to 650 hours with consistent lithium plating and stripping. NPD analysis elucidates the lack of additional phases in F-doped and undoped LGLZO. Rietveld analysis verifies that La is located at 24c, Zr at 16a, and O1 at 96h. In F-doped LGLZO, F atoms are randomly allocated at the 96h site, in conjunction with O1 atoms. Li1 atoms occupy the 24d position, whereas Li2 atoms are located in the 96h position. Ga is randomly distributed with Li1 atoms. There are just 0.2 F per formula among the 12 anionic sites, which does not significantly influence the bond distance. Each atom has a small displacement, so the atoms are more tightly bound in the F-LGLZO structure. This affects the lithium transport in the F-LGLZO structure due to the smaller

displacement of the anionic sublattice. Liu et al. 137 employed BSVE to investigate the influence of Fluorine on the transport pathway of LLZO (Fig. 2d). Generally, lithium atoms occupy Li(1)O<sub>4</sub> and Li(2)O<sub>6</sub> sites. Fluorine doping at oxygen sites leads to the rearrangement of lithium atoms between Li1 and Li2, with Li(1)O<sub>4</sub> tetrahedrons linked to distorted Li(2)O<sub>6</sub> octahedrons. This process leads to a decrease in the Li-ion transport barrier energy and an enhancement in ionic conductivity. The high sintering temperature of 1200 °C for 12 hours required for the production of Li<sub>6.3</sub>La<sub>3</sub>Zr<sub>1.65</sub>W<sub>0.35</sub>O<sub>12</sub> (LLZWO) poses a challenge for the commercialization of garnet. The incorporation of CuO additive effectively reduced the sintering temperature while enhancing the properties compared to pristine LLZWO. Zhang et al.143 incorporated varying amounts of CuO into LLZWO, specifically LLZWO-x wt% CuO where x = 0, 1, 2, 5. The diffraction pattern of LLZWO pellets sintered at 1180 °C for 6 hours exhibited signals corresponding to La<sub>2</sub>Zr<sub>2</sub>O<sub>7</sub>. No impurity phase was observed in the 1 and 2 wt% CuO-LLZWO pellets sintered at 1120 °C for 6 hours. Once increasing CuO to 5 wt%, signals for Li<sub>3</sub>Cu<sub>2</sub>O<sub>4</sub> and CuO were observed. The maximum relative density attained for 2 wt%-LLZWO was 97.6%.

**2.1.2. NASICON.** NASICON or sodium ionic conductor is a type of inorganic solid electrolyte with the general structure of  $A_xM_2(BO_4)_3$ , where A represents Li or Na, M represents metal cations, and  $BO_4$  represents polyanions such as  $SiO_4^{\ 4^-}$ ,  $PO_4^{\ 3^-}$ , and  $SO_4^{\ 2^-}$ . The NASICON structure is formed by a sequence of metal octahedra and polyanion tetrahedra that are coordinated with one another through the corners, resulting in a rhombohedral or a monoclinic structure. The presence of doping and temperature variations promotes a transformation in the structure. However, the octahedra and tetrahedra arrangements remain intact, ensuring that vacancy channels are still present.  $^{29,40,41,118,149-152}$ 

Zhang et al.149 clarified the diffusion mechanism of Na-ions in NZSP-type NASICON, incorporating insights from prior research on four Na sites. 152 They introduced a novel Na5 site that significantly increases the diffusion mechanism, utilizing a combination of four methodologies: BVEL, DFT, MEM, and NPD. In the rhombohedral structure, sodium is predominantly located at the Na1 (6b) and Na2 (18e) sites, with a minor presence at the Na3 (36f) sites. The Na3 site is adjacent to Na1, preventing simultaneous occupation of both sites (Fig. 3a1). In the monoclinic structure, the Na2 (18e) site is subdivided into Na2 (4e) and Na3 (8f) sites, while the Na3 (36f) site is further divided into Na4 (8f) and Na5 (8f) sites. In the monoclinic 3D network of NZSP, Na<sup>+</sup> diffusion is observed at all five sites; however, probability density analysis indicates a limited diffusion pathway along the [101] direction that connects adjacent be planes (red lines in Fig. 3a3). The additional site Na5 functions as a crossover site, facilitating rapid Na<sup>+</sup> mobility. In the

symmetry (b2). (b3) Lab XRD patterns of other as-synthesized NASICONs. (C) Measured ionic conductivity of as-synthesized NASICONs at room temperature ( $\sim$ 25 °C). The bulk properties are indicated by blue circles, whereas the total properties are indicated by orange ones. (D) Na stripping and plating tests in a temperature chamber at 25 °C at various current rates of a Na|Na<sub>3.4</sub>Hf<sub>0.6</sub>Sc<sub>0.4</sub>ZrSi<sub>2</sub>PO<sub>12</sub>|Na symmetric cell. Reproduced with permission from ref. 40. Copyright 2023, Springer Nature. (E) (e1) Li-ion diffusion channel model in LATP and corresponding (e2) diffusion energy barrier and (e3) Li ion diffusion channel model in LATP-0.2F and corresponding (e4) diffusion energy barrier. Reproduced with permission from ref. 150. Copyright 2024, Elsevier.

NZSP rhombohedral structure, two correlated threedimensional networks have been identified: Path 1 (Na2-Na3-Na1-Na3-Na2) and Path 2 (Na2-Na3-Na3-Na2). AIMD and CI-NED calculations indicate that the correlated migration mechanism predominates over the single ion jumping mechanism. Consequently, an increased Na-ion concentration leads to a higher percentage of correlated jumping, attributed to the coulombic repulsion among Na-ions. Enhancing ionic conductivity is more effectively achieved by increasing the Naion concentration than by expanding the NASICON framework while keeping a constant Na<sup>+</sup> concentration.

The current versions of NASICON, such as LiSn<sub>2</sub>(PO<sub>4</sub>)<sub>3</sub>,<sup>153</sup> LiTi<sub>2</sub>(PO<sub>4</sub>)<sub>3</sub>,83,123 LiGe<sub>2</sub>(PO<sub>4</sub>)<sub>3</sub>,154 NaTi<sub>2</sub>(PO<sub>4</sub>)<sub>3</sub>,78 Na<sub>3</sub>V<sub>2</sub>(PO<sub>4</sub>)<sub>3</sub>,5 etc. Doping occurs when a metal cation with a radius comparable to that of the NASICON metal cation and the ability to form octahedral coordination is introduced, regardless of whether the metal cation is divalent  $(Mg^{2+}, Ca^{2+}, Cu^{2+})$ ,  $^{125,155,156}$  trivalent (Yb<sup>3+</sup>, Gd<sup>3+</sup>, Sc<sup>3+</sup>, Al<sup>3+</sup>, Fe<sup>3+</sup>), 157-162 tetravalent (Ge<sup>4+</sup>, Zr<sup>4+</sup>, Ce<sup>4+</sup>), 157, 158, 163-165 or pentavalent (Nb<sup>5+</sup>). 124 For example, doping LiGe<sub>2</sub>(PO<sub>4</sub>)<sub>3</sub> with divalent or pentavalent cations will impact the concentration of Li species and its ionic conductivity. By depositing divalent cations of varying ionic radius sizes into LiGe<sub>2</sub>(PO<sub>4</sub>)<sub>3</sub>, the ionic conductivity will differ due to the hindrance of Li diffusion caused by the larger size of the doping metal cation.

The selection of doping has a significant role in expanding the diffusion channel of Na<sup>+</sup> or Li<sup>+</sup>, enhancing the density of the solid electrolytes, inhibiting the impurities formation at high temperatures, lowering the sintering temperature, and increasing the ionic conductivity. However, not all doping agents and methods comply with NASICON compliance standards due to their inability to increase conductivity, contaminant growth, and lack of economic feasibility. 60,125,155 Wang et al.40 examined the electrochemical characteristics of eight recently discovered NASICON compounds. The ionic conductivity of these compounds can increase up to a certain point with the average size of the metal. Sodium concentration, activation barrier, and substitution of the polyanion PO<sub>4</sub><sup>3-</sup> with SiO<sub>4</sub><sup>4-</sup>, these factors collectively contribute to the enhancement of ionic conductivity. Out of the eight recently discovered NASICON compounds, seven compounds exhibit a rhombohedral structure  $(R\bar{3}c)$ , while the remaining one has a monoclinic structure (C2/c). Fig. 3b1-b3 displays the Rietveld refinement results for Na<sub>3</sub>HfZrSi<sub>2</sub>PO<sub>12</sub> (monoclinic) and Na<sub>3</sub>HfScSiP<sub>2</sub>O<sub>12</sub> (rhombohedral), together with XRD patterns of additional NASICON compounds. The value of the lattice parameter a is influenced by the species of metal present, whereas the value of the lattice parameter c is determined by the type of polyanion present. The compounds Na<sub>3</sub>ScInP<sub>3</sub>O<sub>12</sub> and Na<sub>3</sub>Hf<sub>1.5</sub>Ca<sub>0.5</sub>SiP<sub>2</sub>- $O_{12}$  exhibit a relative density of  $\sim$ 87%, and the remaining compounds are in the range of  $\sim$ 93% to  $\sim$ 95%. Electrochemical impedance spectroscopy analysis performed at room temperature revealed that the ionic conductivity of NASICON increased as the concentration of silicate increased, irrespective of the cation type (Fig. 3c). NASICON exhibits conductivity values ranging from  $10^{-6}$  to  $10^{-5}$  S cm<sup>-1</sup> for the pure phosphate group, from  $10^{-5}$  to  $10^{-4}$  S cm<sup>-1</sup> for the  $(SiO_4)(PO_4)_2$  group, and

a maximum conductivity of  $4.4 \times 10^{-4} \, \mathrm{S \, cm^{-1}}$  for the  $(\mathrm{SiO}_2)_2 \mathrm{PO}_4$ group. The ionic conductivity of Na<sub>3</sub>HfZrSi<sub>2</sub>PO<sub>12</sub> was enhanced by doping it with Sc3+, which increased the average cation radius and modification of the composition. Na<sub>3.4</sub>Hf<sub>0.6</sub>Sc<sub>0.4</sub>-ZrSi<sub>2</sub>PO<sub>12</sub> exhibits an increase in the average cation size from 0.715 Å to 0.722 Å. It has a completely rhombohedral structure and achieves an ionic conductivity of 1.2 mS cm<sup>-1</sup>, which is higher than that of the pristine Na<sub>3</sub>HfZrSi<sub>2</sub>PO<sub>12</sub> compound. Na<sub>3.4</sub>Hf<sub>0.6</sub>Sc<sub>0.4</sub>ZrSi<sub>2</sub>PO<sub>12</sub> demonstrated stability in Na stripping and plating for 200 h (Fig. 3d).

Yin et al. 150 assessed the impact of partially substituting fluorine for oxygen at the 36f1 and 36f2 sites within the LATPtype NASICON structure. The fluorine atom exhibits a propensity to substitute the oxygen atom in the Al-O-P configuration, shifting closer to the phosphorus atom (P-F, 1.60 Å) and away from the aluminum atom (Al-F, 2.49 Å). Doping leads to the elongation of the Al-O and P-O bonds, accompanied by the distortion of the octahedron (from AlO<sub>6</sub> to AlO<sub>5</sub>F) and tetrahedron. The doping effect enlarges the crystal lattice while preserving the  $R\bar{3}c$  crystal structure. The analysis of Li vacancies elucidates the impact of F doping. The incorporation of a single fluorine atom (LATP-0.1F) reduces the vacancy energy of lithium ions in Li4, while simultaneously increasing the vacancy energy in Li1. In LATP-0.3F, the vacancy energy in Li3 and Li4 is reduced. LATP-0.2F demonstrates the ability to maintain vacancy energy in Li1, Li2, and Li3 at significantly lower energy levels compared to LATP-0.1F and LATP-0.3F. The vacanciesassisted diffusion mechanism of LATP and LATP-0.2F is represented by the following pathways: Li2  $\rightarrow$  Li3  $\rightarrow$  Li4  $\rightarrow$  Li5 (36f  $\rightarrow$  36f  $\rightarrow$  36f  $\rightarrow$  6b) for LATP (Fig. 3e(1)) and Li2  $\rightarrow$  Li3  $\rightarrow$  Li4  $(36f \rightarrow 36f \rightarrow 6b)$  for LATP-0.2F (Fig. 3e(3)). The graph indicates that the initial energy barriers for both are minimal. LATP requires crossing multiple barriers, specifically from the octahedral [TiO<sub>6</sub>] to the tetrahedral [PO<sub>4</sub>], then to the octahedral [AlO<sub>6</sub>], and finally back to the tetrahedral [PO<sub>4</sub>]. This outcome is attributable to LATP containing four lithium ions. Lithium migrates from 36f to 36f with minimal energy consumption (0.001 eV). The diffusion of 36f  $\rightarrow$  36f in LATP-0.2F exhibits a low energy level of 0.054 eV. The transition from the [AlO<sub>6</sub>] octahedron to [AlO<sub>5</sub>F] results in a modification of the Al-O bond length (1.93 Å) to the Al-F bond length (2.49 Å), thereby facilitating the passage of the initial diffusion barrier with minimal required energy (Fig. 3e2). The 36f  $\rightarrow$  6b diffusion barrier for LATP-0.2F (0.578 eV) is lower than that for LATP (0.672 eV) due to the shorter Al-O bond length of 1.85 Å in the [AlO<sub>5</sub>F] octahedron compared to 1.96 Å in the [AlO<sub>6</sub>] octahedron (Fig. 3e4). F doping is concluded to facilitate the opening of diffusion channels and to reduce the diffusion energy barrier. The ionic conductivities of LATP and LATP-0.2F are measured at 0.142 mS cm<sup>-1</sup> and 0.358 mS cm<sup>-1</sup>, respectively. Li *et al.*<sup>122</sup> presented another study related to Cl doping in LATP. Chlorine atoms are distributed randomly between two oxygen positions (36f), with occupancy rates of 12.3% for position 36f<sub>1</sub> and 7.2% for position 36f2. This suggests that chlorine predominates in occupying the O site (36f1). This influences the asymmetry of the metal-oxygen bond length (M = Li, Al, or Ti). The bond length of Li (36f)–O/Cl (36f<sub>1</sub>) increases from 2.145 Å to 2.304 Å, whereas

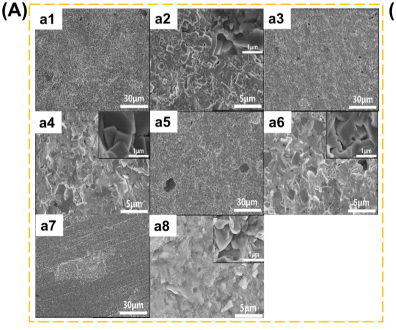
the Li (36f)–O/Cl (36f<sub>2</sub>) decreases from 2.069 Å to 1.874 Å. The variation in Li–O bond distance influences the diffusion rate of Li-ions. The conductivity of LATP-0.4Cl is approximately 0.423 mS cm<sup>-1</sup>, whereas pristine LATP exhibits a conductivity of 0.139 mS cm<sup>-1</sup>.

Wu et al. 166 enhanced the stability and mobility of Na-ions in Na<sub>3</sub>V<sub>2</sub>(PO<sub>4</sub>)<sub>2</sub>F<sub>3</sub> (NVPF) by the doping of K<sup>+</sup> cations, which partially substitute Na<sup>+</sup> in the NVPF structure. The unique structure of NVPF consists of the interconnection of octahedra [VO<sub>4</sub>F<sub>2</sub>] via F atoms and the interconnection of tetrahedra [PO<sub>4</sub>] via O atoms, resulting in a three-dimensional NASICON framework with Na<sup>+</sup> migration channels oriented in the [110] direction. NVPF has three pairs of redox potentials (about 3.4, 3.7, and 4.2 V) that influence structural deformation, modify the diffusion pathway, and reduce the mobility of Na<sup>+</sup> during the discharging process compared to the charging process. Cation doping at the V site of NVPF effectively enlarges the crystal lattice parameters when the doping cation's size exceeds that of V, thereby enhancing Na<sup>+</sup> migration and improving the structural and physicochemical stability of the crystal, as the doping has no impact on the electrochemical process. K+ doping at the Na<sup>+</sup> site results in a reduction of bandgap energy from 3.31 eV to 2.70 eV, hence enhancing electrical conductivity from  $7.95 \times 10^{-6} \text{ S cm}^{-1}$  (NVPF/C) to  $3.78 \times 10^{-5} \text{ S cm}^{-1}$ (NKVPF/C). The optimization of  $K^+$  doping (x = 0.08, N<sub>0.92</sub>K<sub>0.08</sub>VPF/C) promotes a reduction in particle size and enhances crystallinity, resulting in a lattice spacing of 0.322 nm.  $K^+$  doping does not influence the  $V^{3+}$  valence state in NKVPF. The CV profiles and dQ/dV plot indicate that NKVPF/C significantly reduces polarization and voltage hysteresis throughout the charge–discharge operation.  $N_{0.92}K_{0.08}VPF/C$  has a highly reversible phase shift during sodiation and desodiation. EIS study indicates that K doping enhances  $Na^+$  transport, leading to improved cell performance.

Wang et al.41 demonstrated that the addition of 3 wt%  $Na_2B_4O_7$  (NBO<sub>3</sub>) to  $Na_{3.3}La_{0.3}Zr_{1.7}Si_2PO_{12}$  (NLZSP) can solve deficiencies in the synthesis and properties of NASICON. This addition facilitates a reduction in sintering temperature to 950 ° C, enhances NLZSP grain connection, decreases grain boundary resistance  $(R_{\rm sh})$  from 135  $\Omega$  cm<sup>2</sup> for NLZSP to 8  $\Omega$  cm<sup>2</sup> for NLZSP (NBO<sub>3</sub>), and increases ionic conductivity to 1.81 mS cm<sup>-1</sup> with an activation energy of 0.22 eV. Fig. 4a1-a8 shows the SEM of NLZSP (NBO<sub>3</sub>) sintered from 850 to 1150 °C. At 850 °C, NLZSP (NBO<sub>3</sub>) has many tiny holes and cracks, but at 950 °C, it exhibits small grains with different morphologies and a higher density. These variables are shown to impact the differences in  $R_{g}$ ,  $R_{gh}$ , and Rt values in NLZSP (NBO), Fig. 4b. In another case, Li<sub>1.5</sub>-Al<sub>0.5</sub>Ge<sub>0.5</sub>P<sub>3</sub>O<sub>12</sub> (LAGP) with 0.5 wt% LiBF<sub>4</sub> improved conductivity to  $3.21 \times 10^{-4} \, \mathrm{S \, cm^{-1}}$ , reduced grain boundary impedance from 162.2  $\Omega$  to 35.2  $\Omega$ , raised relative density from 93.6% to 96.5%, eliminated the impurity phase, and reduced the sintering temperature from 800 to 700 °C. The Li|LAGP-LiBF<sub>4</sub>|LFP cell has a discharge capacity of 164.2 mA h g<sup>-1</sup> at 0.2C and 117.5 mA h  $\rm g^{-1}$  after 100 cycles. The decrease in ionic

# NASICON solid electrolytes

### Sintering and additive effects to morphologies and conductivities



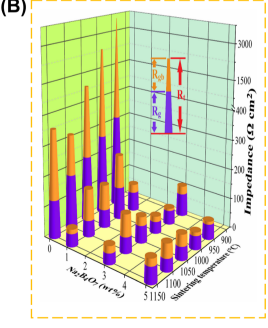


Fig. 4 (A) Cross-sectional SEM images of NLZSP (NBO<sub>3</sub>) ceramic electrolyte sheets sintered at (a1 and a2) 850 °C, (a3 and a4) 950 °C, (a5 and a6) 1050 °C and (a7 and a8) 1150 °C. (B)  $R_{\rm g}$ ,  $R_{\rm gb}$ , and  $R_{\rm t}$  of NLZSP and NLZSP (NBO) ceramic electrolyte. Reproduced with permission from ref. 41. Copyright 2023, Elsevier.

# LPS solid electrolytes

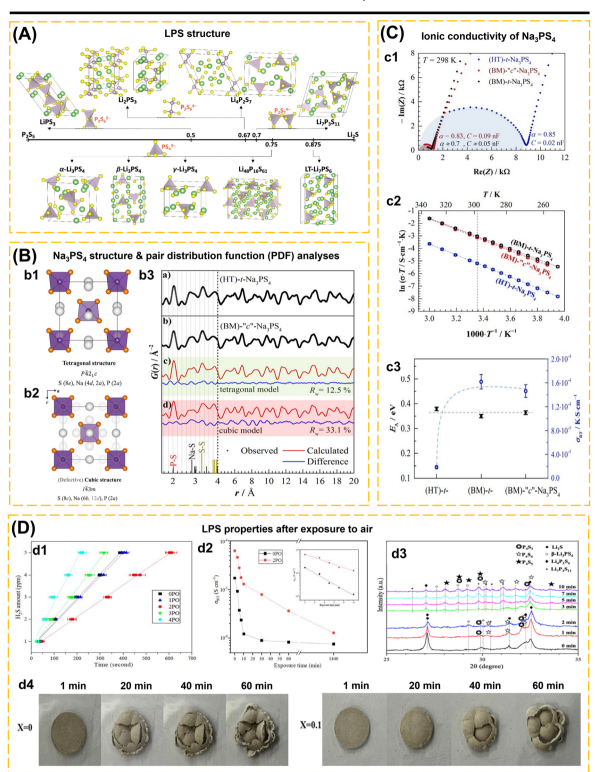


Fig. 5 (A) Crystal structures of LPS compositions on and near the  $Li_2S-P_2S_5$  composition line (Li: green; S: yellow; P: purple). Reproduced with permission from ref. 174. Copyright 2022, American Chemical Society. (B) Crystal structure of (b1) tetragonal and (b2) cubic Na<sub>3</sub>PS<sub>4</sub> projected in the (010) plane. The perfectly cubic phase (i.e., no occupancy of the 12d positions) shows  $PS_4^{3-}$  tetrahedra in a body-centered lattice. In the tetragonal modification, a minor rotation of the tetrahedra leads to a splitting of the Na positions and an elongation of the c-lattice parameter. (b3) Experimentally obtained G(r) data for (a) HT-t-Na<sub>3</sub>PS<sub>4</sub> and (b) BM-"c"-Na<sub>3</sub>PS<sub>4</sub> showing that there is no significant difference in the local structure. BM-"c"-Na<sub>3</sub>PS<sub>4</sub> was fitted using a (c) tetragonal model, shaded in green and a (d) cubic model shaded in red. Experimental data are shown as black points. The red line denotes the calculated pattern, and the difference profile is shown in blue. A fit to the low-r range (1.5–4 Å)

conductivity can be caused by the increase in temperature during the synthesis of NASICON because it has an impact on the decrease in the concentration of mobile ions (Li<sup>+</sup> and Na<sup>+</sup>). This is followed by grain cracking, which disrupts the ionic transport pathway, the formation of by-products that inhibit ionic transport, and the formation of unfavorable pores and becomes a place for the growth of Li dendrites during the cycle. Modification in sintering techniques is the solution to obtain high-density materials, without additional phases, with high conductivity, and efficiency in energy consumption and cost. Common synthesis methods of NASICON that have been applied to date include solid-state reaction, 40,41,60,78,125 solgel, 168,169 spark plasma sintering, 90 stereolithography, 170 melt-casting technique, 123 CSP followed by post-annealing, 83 and ultrafast high-temperature sintering, 39

#### 2.2. Sulfide-based ISEs

In this section, sulfides are divided into several groups: lithium thiophosphate, thio-LISICON, and argyrodite. In the ISE class, sulfides show good ductility and high ionic conductivity. 47,80,82,88,171–173 The properties of each class are described further.

2.2.1. Lithium thiophosphates. Lithium thiophosphates (LPS) or  $(\text{Li}_2\text{S})_x(\text{P}_2\text{S}_5)_{1-x}$  are a sulfide electrolyte with superionic conductivity (10<sup>-3</sup> to 10<sup>-2</sup> S cm<sup>-1</sup>), soft mechanical properties, and low grain boundary resistance. 109,174 However, the instability of LPS with Na/Li metal and oxide-based electrodes, coupled with its high moisture sensitivity, necessitates delicate handling in an inert environment and the use of techniques such as electrode composites and interface layers, as detailed in Section 3.2.175 Guo et al.174 presented the crystal structure of glass-ceramic LPS based on anionic species  $P_x S_y^{n-}$  (Fig. 5a) such as ortho-thiophosphate (PS<sub>4</sub><sup>3-</sup>), pyro-thiophosphate (P<sub>2</sub>S<sub>7</sub><sup>4-</sup>), hypo-thiodiphosphate (P<sub>2</sub>S<sub>6</sub><sup>4-</sup>), meta-thiodiphosphate (P<sub>2</sub>S<sub>6</sub><sup>2-</sup>). Different anion species affect the ionic sites and conductivity of lithium ions. Even for  $Li_3PS_4$  polymorphs,  $\alpha$ -,  $\beta$ -, and  $\gamma$ - $Li_3PS_4$ , they contain the same PS<sub>4</sub><sup>3-</sup> anion but differ in the arrangement of cation sites and orientation of PS<sub>4</sub><sup>3-</sup> anions. The study showed that the paddle-wheel effect was found in the structure of  $\beta$ -Li<sub>3</sub>PS<sub>4</sub> and the orientation of PS<sub>4</sub><sup>3-</sup> enhanced the diffusion of lithium ions in  $\beta$ -Li<sub>3</sub>PS<sub>4</sub>. <sup>174,178</sup>

Sodium-based thiophosphate, Na<sub>3</sub>PS<sub>4</sub>, exhibits two types of crystal structures: cubic and tetragonal. The cubic structure exhibits higher conductivity compared to the tetragonal structure (Fig. 5b1 and b2). In the cubic structure, PS<sub>4</sub><sup>3-</sup> polyhedra are arranged within a body-centered cubic lattice, with one Na<sup>+</sup>

occupying the octahedral sites (6b). The tetragonal structure results from the rotation of the PS<sub>4</sub><sup>3-</sup> polyhedra towards the Na<sup>+</sup> positions (4d and 2a), resulting in an increased ratio of the lattice parameters c/a. Krauskopf et al. 176 examined the influence of the crystal lattice of Na<sub>3</sub>PS<sub>4</sub>, which was prepared through ball-milling (BM-"c"-Na<sub>3</sub>PS<sub>4</sub>) and high temperature (HT-t-Na<sub>3</sub>PS<sub>4</sub>). The cubic structure of (BM)-"c"-Na<sub>3</sub>PS<sub>4</sub> determined by Bragg diffraction exhibited no significant change when compared to the local tetragonal structure derived from PDF analysis. Despite the brief annealing of BM-"c"-Na<sub>3</sub>PS<sub>4</sub>, the resulting product, (BM)-t-Na<sub>3</sub>PS<sub>4</sub>, exhibited structural parameters comparable to those of HT-t-Na<sub>3</sub>PS<sub>4</sub> (Fig. 5b3). This indicates that varying preparation conditions lead to distinct crystal structures, and the ionic conductivity of Na<sub>3</sub>PS<sub>4</sub> is independent of the crystal structure. Despite the sintering of the HT-t-Na<sub>3</sub>PS<sub>4</sub> (500 °C for 10 hours) to decrease grain boundary resistance, SEM results indicated a smooth and dense surface; however, the conductivity exhibited no significant change. The conclusion indicates that conductivity is influenced by variations in defect concentration rather than by the grain boundary (Fig. 5c1-c3). The milling conditions result in a reduced grain size, which enhances conductivity, while simultaneously increasing defect concentration.

Park *et al.*<sup>179</sup> demonstrated the hydrolysis stability of sulfides ( $\text{Li}_3\text{PS}_4$ , LGPS, and  $\text{Li}_6\text{PS}_5\text{Cl}$ ) in the presence of moisture. The reaction energies for  $\text{Li}_3\text{PS}_4$ , LGPS, and  $\text{Li}_6\text{PS}_5\text{Cl}$  were measured at 49, 50, and 92 meV per atom, respectively. This suggests that sulfides easily interact with  $\text{H}_2\text{O}$ , resulting in the formation of  $\text{H}_2\text{S}$ ,  $^{177,179}$  as demonstrated by the reaction:

$$0.9412H_2O + 0.1176Li_{10}GeP_2S_{12} \rightarrow 0.2353Li_3PO_4 \\ + 0.1176Li_4GeS_4 + 0.9412H_2S$$
 
$$0.5H_2O + 0.5Li_6PS_5Cl \rightarrow 0.125Li_3PO_4 + 0.375Li_3PS_4 + LiHS \\ + 0.5LiCl$$
 
$$0.9412H_2O + 0.2353Li_3PS_4 \rightarrow 0.9412H_2S + 0.2353Li_3PO_4$$

Research and development focus mainly on synthesis and modification using addictive materials to reduce metal and moisture instability, inhibit lithium dendrites, and boost conductivity.  $^{49,65,180,181}$  Mi and Hall<sup>177</sup> developed a glass-ceramic electrolyte containing  $(100 - x)(0.75 \text{Li}_2\text{S} \cdot 0.25 \text{P}_2\text{S}_5) \cdot x \text{P}_2\text{O}_5$  (mol%) (x = 0, 1, 2, 3 and 4). A glass-ceramic based on  $98(0.75 \text{Li}_2\text{S} \cdot 0.25 \text{P}_2\text{S}_5) \cdot 2 \text{P}_2\text{O}_5$  (2PO) has strong air stability, producing 5 ppm of  $\text{H}_2\text{S}$  gas after 10 minutes of exposure at 20–

was independently performed and leads to larger scale factors, thereby indicating an additional fraction of amorphous content or phase with a low coherence length. (C) (c1) Nyquist plots of t-Na<sub>3</sub>PS<sub>4</sub> and "c"-Na<sub>3</sub>PS<sub>4</sub> synthesized using the different synthetic approaches. Data points are only displayed up to 2 MHz. (c2) Arrhenius plots for all compounds. (c3) Activation energy and room temperature ionic conductivity for each of the Na<sub>3</sub>PS<sub>4</sub> compounds, showing that while the activation barrier for ionic motion remains unchanged the ball-milling procedure leads to much higher conductivities irrespective of the crystal structure. Reproduced with permission from ref. 176. Copyright 2018, American Chemical Society (D) LPS properties after exposure to air: (d1) amounts of H<sub>2</sub>S released from 0PO, 1PO, 2PO, 3PO, and 4PO after exposure to air (left). (d2) The ionic conductivity (RT) of 0PO and 2PO electrolytes at various exposure times (middle). (d3) XRD patterns of 2PO exposed to air for 1 min, 2 min, 3 min, 7 min, and 10 min in the  $2\theta$  of 25° to 35° range. Reproduced with permission from ref. 177. Copyright 2023, Royal Society of Chemistry. (d4) Optical images of Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub> and Li<sub>7</sub>P<sub>2.9</sub>Ge<sub>0.05</sub>S<sub>10.75</sub>O<sub>0.1</sub> electrolyte at 1, 20, 40, and 60 min exposure in the open air at RT. Reproduced with permission from ref. 65. Copyright 2022, American Chemical Society.

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25 °C and 70% humidity (Fig. 5d1). 2PO has 3.7 times better ionic conductivity than undoped P2O5 and is stable in air (Fig. 5d2). The ss-NMR data revealed that phases with high conductivity in 2PO (β-Li<sub>3</sub>PS<sub>4</sub> and Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub>) were sustained, resulting in high conductivity. 2PO EDS analyzes the stability of 2PO in terms of element composition. At 10 minutes of exposure, the conductivity of 2PO decreases more slowly than that of 0PO and serves several levels higher. The XRD investigation demonstrates that 2PO only underwent minor changes after being exposed to air for several days; nonetheless, an S<sub>8</sub> layer was produced, which preserves the material's stability against air and delays deterioration (Fig. 5d3).

Lv et al. 65 developed  $\text{Li}_7 P_{3-x} Ge_{0.5x} S_{11-2.5x} O_x$  (x = 0, 0.1, 0.2, and 0.3) glass-ceramic electrolytes. Under 30% humidity at 30  $^{\circ}$ C for 60 minutes,  $\text{Li}_7\text{P}_3\text{S}_{11}$  and  $\text{Li}_7\text{P}_{2.9}\text{Ge}_{0.05}\text{S}_{10.75}\text{O}_{0.1}$  produced 0.5472 cm<sup>3</sup> g<sup>-1</sup> and 0.3168 cm<sup>3</sup> g<sup>-1</sup> of H<sub>2</sub>S, respectively. XRD analysis before and after air exposure shows that there is no

structural change in Li<sub>7</sub>P<sub>2.9</sub>Ge<sub>0.05</sub>S<sub>10.75</sub>O<sub>0.1</sub>. Fig. 5d4 depicts the conditions of  $\text{Li}_7\text{P}_3\text{S}_{11}$  and  $\text{Li}_7\text{P}_{2.9}\text{Ge}_{0.05}\text{S}_{10.75}\text{O}_{0.1}$  solid electrolytes exposed to open air. In electrochemical stability tests, Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub> exhibited anodic fluctuations in the range of 3 to 5 V (vs.  $\text{Li/Li}^+$ ), whereas  $\text{Li}_7\text{P}_{2.9}\text{Ge}_{0.05}\text{S}_{10.75}\text{O}_{0.1}$  continues to be stable up to 5 V. The electrochemical stability of Li/Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub>/Li symmetric cell shows an overpotential of ~48 mV followed by a voltage drop after the 7th cycle caused by short circuit due to the growth of lithium dendrites. While the Li/Li<sub>7</sub>P<sub>2.9</sub>Ge<sub>0.05</sub>- $S_{10.75}O_{0.1}/Li$  cell shows a small and stable overpotential ( $\sim$ 5 mV) for 100 h @ 0.1 mA cm<sup>-2</sup>/0.1 mA h cm<sup>-2</sup> at room temperature condition. The presence of GeO<sub>2</sub> to Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub> structure improves the ionic conductivity, stability in the air, electrochemical stability, and compatibility with Li anodes.

2.2.2. Thio-LISICON. Thio-LISICON is a sulfide derivative of LISICON, where the oxygen ions (O<sup>2-</sup>) are replaced with sulfur ions (S<sup>2-</sup>). The sulfur ions have a larger size and higher

# Thio-LISICON solid electrolytes

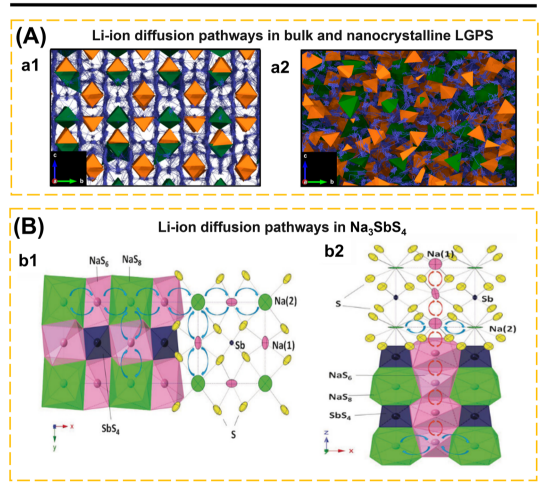


Fig. 6 (A) Li-ion diffusion pathways in bulk and nanocrystalline LGPS. Diffusion density plots of Li ions (blue) overlaid on GeS<sub>4</sub> (green) and PS<sub>4</sub> (orange) tetrahedra in (a1) bulk and (a2) nanocrystalline LGPS with a particle volume of 10 nm $^3$  at 300 K (2.4  $\times$  3.5 nm cross-section). Reproduced with permission from ref. 183. Copyright 2023, American Chemical Society. (B) Li-ion diffusion pathways in LGPS: (b1) [001] and (b2) [010] views of the framework structure of  $Na_3SbS_4$ . The 2D planar  $Na^+$  transport tunnels in the xy-plane are formed by alternatively arranged  $NaS_6$  and  $NaS_8$ polyhedra, which share common faces. Planar tunnels are connected along the z-axis through chains formed by NaS<sub>6</sub> octahedra, which share common edges. Reproduced with permission from ref. 184. Copyright 2016, Wiley-VCH.

### Argyrodite solid electrolytes

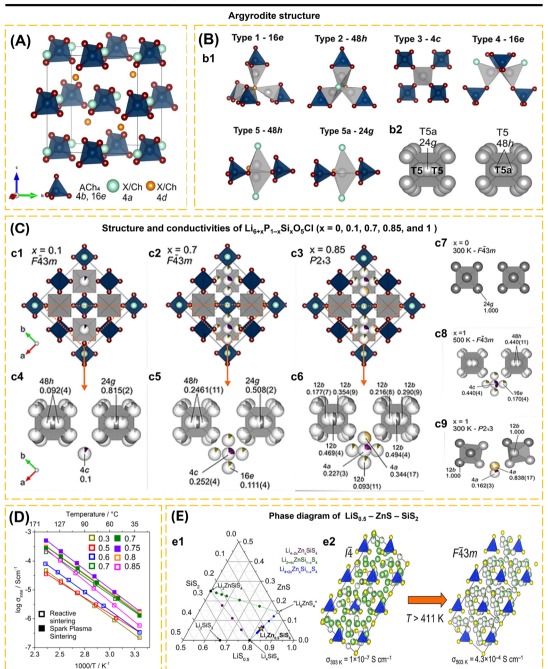


Fig. 7 (A) Unit cell of Li<sub>6</sub>ACh<sub>5</sub>X (A = P, Si, and Al; Ch = O, S, and Se; X = Cl, Br, and I); Ch/X anions are tetrahedrally close-packed on Wyckoff site (4a, 4d, and 16e) forming 136 tetrahedral voids; four are occupied by A cations on the 4b site defining Ach<sub>4</sub> tetrahedra (Ch: 16e). Lithium atoms are not shown. (B) (b1) Panels showing five types (T1, T2, T3, T4, and T5) of possible interstitial tetrahedral sites available for lithium occupancy and the trigonal bipyramidal T5a site. Lithium atoms are shown in gray. (b2) Octahedral Li<sup>+</sup>-ion cages consisting of T5 and T5a sites surrounding central anions on the 4d site; due to the proximity of T5 and T5a sites, the total occupancy over the two sites is constrained to unity. Reproduced with permission from ref. 88. Copyright 2022, American Chemical Society. (C) Structures of x = 0, 0.1, 0.7, 0.85, and 1 in  $\text{Li}_{6+x}\text{P}_{1-x}\text{Si}_x\text{O}_5\text{Cl}$ highlighting the different Li sites occupied and associated symmetry settings as a function of composition and temperature in Li<sub>7</sub>SiO<sub>5</sub>Cl; atom and polyhedra colors: SiO<sub>4</sub> tetrahedra (dark blue), O (dark red), Cl (light blue), Li-T5a site (dark gray), Li-T5 site (light gray), Li-T3 site (purple), Li-T4 site [green (16e/12b), and yellow (4a)]. Unit cell for (c1) x = 0.1, (c2) x = 0.7, and (c3) x = 0.85 (T5 and T5a Li atoms are omitted for clarity with octahedral cages shown in gray); orange triangle highlights the structural subsection shown in panels (c4-c6); (c7-c9) octahedral Li ion cages (T5 and T5a) surrounding central oxide anions and interstitial T3 and T4 sites with their site occupancy factor for (c4) x = 0.1, (c5) x = 0.7, (c6) x = 0.7, (c6) x = 0.7, (c7) x = 0.7, (c8) x = 0.7, (c8) x = 0.7, (c9) x = 0.70.85, (c7) x = 0, (c8) x = 1, (500 K), and (c9) x = 1 (300 K). (D) Arrhenius plots for  $\text{Li}_{6+x}P_{1-x}\text{Si}_xO_5\text{Cl}$  for samples prepared *via* reactive sintering (RS) (x = 0) (RS) (x = 0) (C8) x = 0) (RS) (x = 0) (C8) x = 0) (C9) x = 0) (RS) (x = 0) (C9) x = 0) (C9) (x = 0) (x= 0.3, 0.5, 0.6, 0.7, 0.8, 0.85) and SPS (x = 0.7, 0.75); reproduced with permission from ref. 88. Copyright 2022, American Chemical Society. (E) (e1) Pseudoternary  $LiS_{0.5}$ –ZnS– $SiS_2$  phase field was explored by synthesizing the compositions shown. Known phases are shown as black circles and are labeled. Purple, green, and blue circles show compositions explored on the  $\text{Li}_{4-2x}\text{Zn}_x\text{SiS}_4$ ,  $\text{Li}_{2+4x}\text{ZnSi}_{1-x}\text{S}_4$ , and  $\text{Li}_{4+2x}\text{Zn}_x\text{Si}_{1-x}\text{S}_4$  tie lines,

polarization, which leads to an increase in the lithium transport channel and thus enhances the ionic conductivity. The LGPStype lithium conductor, Li<sub>10</sub>GeP<sub>2</sub>S<sub>12</sub>, has a tetragonal crystal structure consisting of PS<sub>4</sub><sup>3-</sup> and GeS<sub>4</sub><sup>4-</sup> tetrahedra that are surrounded by Li ions in both tetrahedral and octahedral coordination. 182,183 Experimental and theoretical investigations demonstrate that the most straightforward route for Li-ion diffusion is along the c direction, as opposed to the a and b directions (Fig. 6a1). The examination of MSD indicates that the diffusion in the c direction is fivefold greater than the diffusion in the a and b plane directions. The diffusion pathway for nanocrystalline LGPS exhibits an equitable distribution of ion diffusion along the c-axis and within the ab plane, as the reduction in particle size and enhancement of surface area provide shorter ion diffusion pathways. This enhancement facilitates ion diffusion in all directions, resulting in a broad diffusion area on the surface of LGPS particles, which correlates with an increase in ionic conductivity (Fig. 6a2). Wang et al. 184 explained two sodium sites in Na<sub>3</sub>SbS<sub>4</sub>, specifically Na(1) located within the distorted NaS<sub>6</sub> octahedron, and Na(2) located in the NaS<sub>8</sub> dodecahedron. In the [001] plane, the NaS<sub>6</sub> and NaS<sub>8</sub> sites are linked to the same surface, generating a planar Na(1)-Na(2) diffusion pathway. In the [010] plane, the NaS<sub>6</sub> is interconnected to form a Na(1)-Na(1) diffusion pathway. The Na(1)-Na(1) interplane diffusion pathway is key in linking neighboring Na(1)-Na(2) planar diffusion pathways, thereby forming a three-dimensional diffusion network within the Na<sub>3</sub>SbS<sub>4</sub> framework (Fig. 6b).

Lu et al.3 fabricated a glass-ceramic LGPS using Li<sub>2</sub>S, GeS<sub>2</sub>, and P2S5. This was achieved by subjecting the mixture to highenergy ball milling for a duration of 40 to 520 minutes. The formation of glassy-ceramic LGPS may be elucidated by considering three distinct stages: the vitrification zone refers to the stage when the mixture of Li<sub>2</sub>S, GeS<sub>2</sub>, and P<sub>2</sub>S<sub>5</sub> becomes uniform and transitions into an amorphous state. In the intermediary zone, intermediate substances such as Li<sub>4</sub>GeS<sub>4</sub> and Li<sub>3</sub>PS<sub>4</sub> are formed. Finally, in the product stage, glassyceramic LGPS is generated. Nevertheless, the EIS investigation revealed that LGPS-HBM 520 min had the greatest ionic conductivity, 1.07 mS cm<sup>-1</sup>. The percentage of LGPS is 40%. The ionic conductivity of LGPS can be increased by post-heating at 575, 600, and 625 °C for 8 h. The highest ionic conductivity at room temperature was attained by LGPS 600 °C, 3.27 mS cm<sup>-1</sup>, with an LGPS concentration of 91.1%. Luo et al.66 replaced Ge with Sn to reduce costs and synthesized a solid electrolyte based on Li<sub>10</sub>SnP<sub>2</sub>S<sub>12</sub>. Li<sub>10</sub>SnP<sub>2</sub>S<sub>12</sub> has an ionic conductivity of 4.79 mS cm<sup>-1</sup> and electronic conductivity of  $2.6 \times 10^{-8}$  S cm<sup>-1</sup> at room temperature. LiNi<sub>0.6</sub>Mn<sub>0.2</sub>Co<sub>0.2</sub>O<sub>2</sub>||Li-In cells exhibit limited discharge capacity and poor SE compatibility with the electrode. The use of Li<sub>3</sub>InCl<sub>6</sub> as a catholyte and a cathode protective layer results in an initial discharge capacity of 186.9 mA h g<sup>-1</sup> at 60 °C. The combination of Li<sub>3</sub>InCl<sub>6</sub> may

separate the direct contact between ISE and cathode, reducing space charge effects and side reactions.

The incorporation of additive Al<sub>2</sub>O<sub>3</sub> into the LGPS structure has been examined by Yabuzaki et al. 80 The XRD study indicated no significant alterations in lattice parameters and Ge/P ratio for LGPS and LGPS-Al<sub>2</sub>O<sub>3</sub>. Increase the concentration of Al<sub>2</sub>O<sub>3</sub> (3, 10, and 30 wt%) did not considerably increase the Al<sub>2</sub>O<sub>3</sub> particles inside LGPS grains; nevertheless, when Al<sub>2</sub>O<sub>3</sub> reached the maximum level in LGPS, the particles occupied the voids in LGPS. The ionic conductivity remained almost unchanged with varying Al<sub>2</sub>O<sub>3</sub> concentrations: LGPS (5.4 mS cm<sup>-1</sup>); LGPS-3 wt%  $Al_2O_3$  (5.7 mS cm<sup>-1</sup>); LGPS-10 wt%  $Al_2O_3$  (4.8 mS cm<sup>-1</sup>); and LGPS-30 wt%  $Al_2O_3$  (4.9 mS cm<sup>-1</sup>). Section 2.2.1 discusses the reactivity of sulfide to moisture. 179 According to HSAB theory, oxygen is harder base than sulfur, binds with hard acid phosphorous and releases H2S. The ionic conductivity retention of LGPS-Al<sub>2</sub>O<sub>3</sub> following air exposure surpasses that of LGPS. The substitution of oxide in LGPS enhances moisture stability. The moisture stability of LGPS-Al<sub>2</sub>O<sub>3</sub> is attributed to the Al<sub>2</sub>O<sub>3</sub> filling the voids of LGPS, which inhibits moisture penetration at the grain boundaries. Additionally, the reactivity of Al<sub>2</sub>O<sub>3</sub> with H<sub>2</sub>O forms a passivation layer of Al(OH)3, inhibiting water residue and reactions at the LGPS interface. The electrolyte composite approach has the potential to serve as an option for preserving moisture stability while maintaining good LGPS conductivity.

**2.2.3.** Argyrodite. Argyrodite with the formula  $\text{Li}_{7-x}$  $ACh_{6-x}Hal_x$  (A = P, Si, Sb, Sn, and Ge; Ch = S, Se, and O; Hal = Cl, Br, and I) exhibits excellent ionic conductivity, good mechanical properties, and is easy to prepare. 88,185,186 The closepacked tetrahedral arrangement of chalcogenides at Wyckoff sites 4a, 4d, and 16e in Argyrodite results in a cubic phase structure (Fig. 7a). The anion framework creates 136 interstitial tetrahedral gaps in each unit cell for cations to occupy. Four of them are filled by A cations (P, Si, Sb, Sn, and Ge) at site 4b, known as the ACh<sub>4</sub> tetrahedron. The other 132 tetrahedral sites function as Li<sup>+</sup> occupancy and are categorized according to the quantity of Ch ions shared with the ACh<sub>4</sub> tetrahedra (Fig. 7b1). Tetrahedral sites are split into five groups: type 1 (T1 - 16e) and type 2 (T2 - 48h) share faces and edges with ACh<sub>4</sub>, respectively; type 3 (T3 - 4c) and type 4 (T4 - 16e) share four and three corners with ACh<sub>4</sub>, respectively; type 5 (T5 - 48h) shares two corners with the ACh<sub>4</sub> tetrahedra, and type 5a (T5a - 24g) shares face with two neighboring T5 tetrahedra in a bipyramidal trigonal system. The T5 and T5a sites define an octahedral cage around the central anion (4d) that forms a 3D network throughout the structure. Full occupancy of the T5 or T5a sites will cause the arrangement of the  $Li^+$  atoms and deviation of the  $F\bar{4}3m$ structure (Fig. 7b2). In most argyrodites, Li<sup>+</sup> atoms occupy disordered T5 (48h) and/or T5a (24g) sites. Recent research suggests that partial occupancy of T2, T3, or T4 sites might improve ionic conductivity by providing shorter and more favorable pathways for Li<sup>+</sup> diffusion, as occurs when P<sup>5+</sup> is partly substituted by other cations.

 $respectively. \ The \ orange \ star \ indicates \ the \ new \ Li_7Zn_{0.5}SiS_6. \ (e2) \ High-temperature \ (\mathcal{T}=448 \ K) \ and \ room-temperature \ structures \ of \ Li_7Zn_{0.5}SiS_6. \ (e2) \ High-temperature \ (\mathcal{T}=448 \ K) \ and \ room-temperature \ structures \ of \ Li_7Zn_{0.5}SiS_6. \ (e2) \ High-temperature \ (e2) \ High-temperature \ (e3) \ High-temperature \ (e3)$ show cubic close-packed layers of SiS<sub>4</sub><sup>4-</sup> polyanions stacked along the [111] and [011] directions, respectively. Atom colors: Li, green; Zn, gray; S, yellow; SiS<sub>4</sub> tetrahedra, blue. Reproduced with permission from ref. 185. Copyright 2022, American Chemical Society.

Morscher et al.88 synthesized a Li-rich oxide argyrodite, Li7-SiO<sub>5</sub>Cl, with more than 6 Li atoms per formula unit. By substituting the P5+ by Si4+ in Li6+xP1-xSixO5Cl, to understand the distribution behavior of Li<sup>+</sup> on the T5 (48h), T5a (24g), T3 (4c), and T4 (16e) sites which resembles the behavior of Li<sub>7</sub>-SiO<sub>5</sub>Cl. The oxide was produced by ball milling a mixture of Li<sub>4</sub>SiO<sub>4</sub>, Li<sub>3</sub>PO<sub>4</sub>, Li<sub>2</sub>O, and LiCl under an argon environment, followed by heat treatment at 823 K for 3 hours. In addition,  $\text{Li}_{6+x}\text{P}_{1-x}\text{Si}_x\text{O}_5\text{Cl}$  (x = 0.7, 0.75, and 0.8) was prepared by SPS. The cubic argyrodite structure is adopted by all compositions in  $\text{Li}_{6+x}\text{P}_{1-x}\text{Si}_x\text{O}_5\text{Cl}$ . In  $\text{Li}_{6+x}\text{P}_{1-x}\text{Si}_x\text{O}_5\text{Cl}$ , the O and Cl anions are arranged in a regular pattern without mixing between the O<sup>2-</sup> (4d) and Cl<sup>-</sup> (4a) anions, which is different from the commonly reported arrangement of S2- and Cl- anions in argyrodite sulfide. It is known that  $\text{Li}_{6+x}P_{1-x}\text{Si}_xO_5\text{Cl}$  has 4 Li sites (48h, 24g, 4c, and 16e) compared to argyrodite sulfide. For x = 0.1 ( $F\bar{4}3m$ ), the Li<sup>+</sup> atom occupies three separate sites, 4c, 48h, and 24g (Fig. 7c1). Most of the Li<sup>+</sup> atoms occupy the 24g and 48h sites, the remaining Li<sup>+</sup> atoms occupy the 4c site (Fig. 7c4). While for x = 0, Li<sub>6</sub>PO<sub>5</sub>Cl, has only one site, 24g (Fig. 7c7). When x = 0.3-0.75, the occupancy of 24g and 48h sites increases, resulting in the disorder of Li<sup>+</sup> sites in the octahedral cage and an increase in ionic conductivity in argyrodite. Li<sub>6+x</sub>P<sub>1-x</sub>Si<sub>x</sub>O<sub>5</sub>Cl was the first to investigate the disorder of Li<sup>+</sup> sites in argyrodite oxide at room temperature. Additional Li<sup>+</sup> atoms introduced by cation substitution will occupy the 16e position sporadically (Fig. 7c2 and c5). This composition remains at 500 K with the crystal structure  $F\overline{4}3m$  (Fig. 7c8). At x = 0.8, the transition from  $F\overline{4}3m$  to  $P2_13$  occurs (Fig. 7c3). For x = 0.8 to x = 0.85, site 4a (T4) shows a stronger occupancy tendency than site 12b (T4) (Fig. 7c6). The  $Li^+$  site arrangement for x = 0.8 and 0.85 (Fig. 7c3 and c6) is similar to that of Li<sub>7</sub>SiO<sub>5</sub>Cl at 300 K (Fig. 7c9), where only site 4a (T4) is occupied. Thus, controlling the composition of Li<sub>6+x</sub>-P<sub>1-x</sub>Si<sub>x</sub>O<sub>5</sub>Cl may explain the behavior of Li<sub>7</sub>SiO<sub>5</sub>Cl. Based on EIS analysis (Fig. 7d and Table 1), the Li<sub>6.75</sub>P<sub>0.25</sub>Si<sub>0.75</sub>O<sub>5</sub>Cl sample prepared via SPS shows the highest conductivity of 1.82  $\times$  $10^{-6} \,\mathrm{S} \,\mathrm{cm}^{-1}$  at 303 K with an activation energy of 0.522 eV. The Li<sub>6.75</sub>P<sub>0.25</sub>Si<sub>0.75</sub>O<sub>5</sub>Cl-SPS sample demonstrates electrochemical stability against Li metal, showing its capacity to transport Liions at a rate of 20 mA cm<sup>-2</sup> (298 K) for 60 hours. Addition-

ally, the sample exhibits stability at the Li/electrolyte interface.

 ${\rm Li_6PS_5Cl}$  shows instability after exposure to air for 1 hour. In contrast to  ${\rm Li_{6.7}P_{0.3}Si_{0.7}O_5Cl}$  which does not show significant changes after contact with air for 1 hour. After exposure for 60 hours,  ${\rm Li_{6.7}P_{0.3}Si_{0.7}O_5Cl}$  diffraction data indicates the presence of  ${\rm Li_2CO_3}$ . The superior air stability of oxide argyrodite opens up opportunities for commercialization in ASSBs devices and the replacement of sulfide argyrodite electrolytes.

Leube et al. 185 first reported the argyrodite type of LiS<sub>0.5</sub>-ZnS-SiS<sub>2</sub> phase mixture via solid-state reaction. According to the pseudoternary phase diagram of LiS<sub>0.5</sub>-ZnS-SiS<sub>2</sub> (Fig. 7e1), the synthesis of Li<sub>4-2x</sub>Zn<sub>x</sub>SiS<sub>4</sub> (purple line) does not result in the formation of a new phase. Instead, only Li<sub>4</sub>SiS<sub>4</sub> and Li<sub>2</sub>ZnSiS<sub>4</sub> are formed. Likewise, the Li<sub>2+4x</sub>ZnSi<sub>1-x</sub>S<sub>4</sub> line (green line) only consists of a mixture of Li<sub>2</sub>ZnSiS<sub>4</sub>, Li<sub>4</sub>SiS<sub>4</sub>, Li<sub>2</sub>S, and ZnS. A new phase can be observed in  $Li_{4+2x}Zn_xSi_{1-x}S_4$  (blue line) at x =0.267, despite the presence of certain impurities of Li<sub>2</sub>S and  $\text{Li}_4\text{SiS}_4$ . By increasing the composition to x=0.33, a pure Li<sub>4.67</sub>Zn<sub>0.33</sub>S<sub>0.67</sub>S<sub>4</sub> phase is obtained, without the presence of Li<sub>2</sub>S and Li<sub>4</sub>SiS<sub>4</sub>. The composition may be expressed as Li<sub>7</sub>-Zn<sub>0.5</sub>SiS<sub>6</sub>, which represents a novel argyrodite phase with an I4 crystal structure (Fig. 7e2). This crystal structure is tetragonal under room temperature conditions. At a temperature of 411.1 K, the crystal structure transforms into an  $F\bar{4}3m$  structure (cubic). In structure I4, five Li sites (T5, T2, T4, T1, and T2a) are observed. In structure  $F\bar{4}3m$ , four Li sites (T5, T5a, T2, and T4) are observed. The substitution of Zn2+ in the lattice creates a jumping variation. At high temperatures, Zn<sup>2+</sup> occupies the T5-48h position in the  $F\overline{4}3m$  structure, whereas at room temperature, it only partially fills the T5 position in the I4 structure. The transition from the ordered I4 structure to the disordered  $F\bar{4}3m$  is associated with a significant decrease in the activation energy to 0.34(1) eV above 411 K, while the conductivity increases from 1.0(2)  $\times$  10<sup>-7</sup> S cm<sup>-1</sup> (RT) to 4.3(4)  $\times$  $10^{-4} \text{ S cm}^{-1} (503 \text{ K}).$ 

Cation and anion doping of argyrodite increase ionic conductivity while also improving argyrodite stability with Li anodes and moisture stability. Arnold *et al.* Feported F-doped Li<sub>5+y</sub>PS<sub>5</sub>F<sub>y</sub> argyrodite and double dopants (F<sup>-</sup>/Cl<sup>-</sup> and F<sup>-</sup>/Br<sup>-</sup>) synthesized using a solution-based method. Li<sub>5+y</sub>PS<sub>5</sub>F<sub>y</sub> argyrodites with various F levels (y = 0.1, 0.2, 0.5, 1.0, and 2.0) exhibit diffraction at  $2\theta = 25.5^{\circ}$ , 30°, and 31.2°, indicating

Table 1 Extracted total ionic conductivities and activation energies for  $\text{Li}_{6+x}\text{P}_{1-x}\text{Si}_x\text{O}_5\text{Cl}$ ; samples prepared *via* reactive sintering (RS) (x=0.1,0.3,0.5,0.6,0.7,0.75,0.8,0.85) and spark plasma sintering (SPS) (x=0.7,0.75,0.8)<sup>88</sup>

Composition	Sintering method	Relative density (%)	Ionic conductivity (S cm <sup>-1</sup> )	Activation energy (eV)
0.1	RS	73.4	${\sim}10^{-9}$	0.687(13)
0.3	RS	79.0	$2.23(1)  imes 10^{-7}$	0.496(4)
0.5	RS	74.5	$3.53(4) \times 10^{-7}$	0.433(4)
0.6	RS	77.1	$4.72(3) \times 10^{-7}$	0.502(3)
0.7	RS	79.7	$1.16(4) \times 10^{-6}$	0.478(11)
0.7	SPS	96.0	$1.63(4) \times 10^{-6}$	0.50(1)
0.75	SPS	95.1	$1.82(1) \times 10^{-6}$	0.522(5)
0.8	RS	85.0	$1.54(2) \times 10^{-6}$	0.491(7)
0.8	SPS	96.5	$1.42(5) \times 10^{-6}$	_ ``
0.85	RS	78.4	$4.93(4) \times 10^{-7}$	0.564(8)

### Argyrodite solid electrolytes

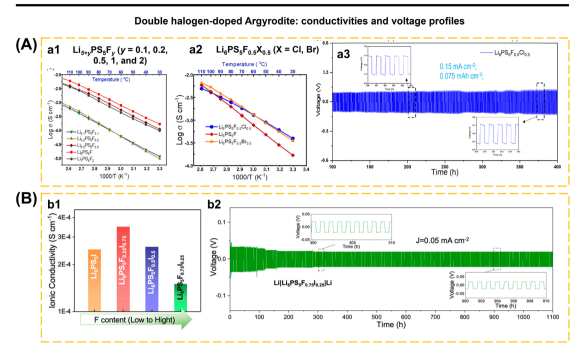


Fig. 8 Double halogen-doped argyrodite. (A) (a1) Arrhenius plots of F-incorporated lithium argyrodites  $Li_{5+v}PS_5F_v$  (y = 0.1, 0.2, 0.5, 1, and 2) that were synthesized from a solvent-based method. (a2) Arrhenius plots of  $\text{Li}_6\text{PS}_5\text{F}_{0.5}\text{X}_{0.5}$  samples (X = Cl, Br), in comparison with  $\text{Li}_6\text{PS}_5\text{F}$ . (a3) Voltage profiles of Li|Li<sub>6</sub>PS<sub>5</sub>F<sub>0.5</sub>Cl<sub>0.5</sub>|Li symmetric cell cycling under current density of 0.15 mA cm<sup>-2</sup> (0.075 mA h cm<sup>-2</sup>). Reproduced with permission from ref. 16. Copyright 2022, American Chemical Society. (B) (b1) Composition dependent ionic conductivity of Li<sub>6</sub>PS<sub>5</sub>I and hybriddoped Li<sub>6</sub>PS<sub>5</sub>F<sub>8</sub>I<sub>1-x</sub> argyrodites at RT. (b2) Voltage profiles of Li<sub>6</sub>PS<sub>5</sub>F<sub>0.75</sub>I<sub>0.25</sub>-based symmetric cell cycling up to 1100 h (current density of 0.05 mA cm<sup>-2</sup>). Reproduced with permission from ref. 92. Copyright 2023, American Chemical Society.

a cubic crystal phase ( $F\bar{4}3m$ ). For  $\gamma > 0.5$ , diffraction at  $2\theta = 38.6^{\circ}$ and 65.6° indicates the existence of LiF. The Raman spectra of  $\text{Li}_7\text{PS}_6$  and  $\text{Li}_{5+\nu}\text{PS}_5\text{F}_{\nu}$  (y = 0.1, 0.2, 0.5, 1.0, and 2.0) show a strong peak at 421–426 cm<sup>-1</sup> indicates the stretching vibration of the P-S bond in PS<sub>4</sub><sup>3-</sup> (ortho-thiophosphate), a shift slightly owing to the replacement of the F<sup>-</sup> anion at the 16e site. The introduction of 9.5 wt% LiF in Li<sub>6</sub>PS<sub>5</sub>F (y = 1) shows the conductivity of  $2.3 \times 10^{-4} \text{ S cm}^{-1}$  (Fig. 8a1). Li<sub>6</sub>PS<sub>5</sub>F<sub>0.5</sub>X<sub>0.5</sub> with double halogen (X = Cl, Br) were synthesized using the same method. XRD analysis of  $\text{Li}_6\text{PS}_5F_{0.5}X_{0.5}$  (X = Cl, Br) samples revealed 7.9 wt% LiF in Li<sub>6</sub>PS<sub>5</sub>F<sub>0.5</sub>Cl<sub>0.5</sub>. At room temperature, Li<sub>6</sub>PS<sub>5</sub>F<sub>0.5</sub>Cl<sub>0.5</sub> and Li<sub>6</sub>PS<sub>5</sub>F<sub>0.5</sub>Br<sub>0.5</sub> possessed ionic conductivities of  $3.5 \times 10^{-4} \, \mathrm{S \, cm^{-1}}$  and  $3.2 \times 10^{-4} \, \mathrm{S \, cm^{-1}}$ , respectively. It concluded that double doping of Li<sub>6</sub>PS<sub>5</sub>F<sub>0.5</sub>X<sub>0.5</sub> increased the ionic transport rate more than single doping (Fig. 8a2). Li<sub>6</sub>-PS<sub>5</sub>F<sub>0.5</sub>Cl<sub>0.5</sub> is stable to Li metal for up to 300 cycles (Fig. 8a3). Fig. 8b1 shows the ionic conductivity of Li<sub>6</sub>PS<sub>5</sub>I and hybriddoped Li<sub>6</sub>PS<sub>5</sub>F<sub>x</sub>I<sub>1-x</sub> argyrodites at RT. Li<sub>6</sub>PS<sub>5</sub>F<sub>0.25</sub>I<sub>0.75</sub> displays the highest ionic conductivity, 40% higher than that of Li<sub>6</sub>PS<sub>5</sub>I. The increase in ionic conductivity is due to the anion disorder caused by the double dopants F- and I-. The stability test of  $\text{Li}_6\text{PS}_5\text{I}$ ,  $\text{Li}_6\text{PS}_5\text{F}_{0.75}\text{I}_{0.25}$ , and  $\text{Li}_6\text{PS}_5\text{F}_{0.25}\text{I}_{0.75}$  with lithium anode at 0.05 mA cm<sup>-2</sup> showed good cycle stability. The interfacial reaction between Li<sub>6</sub>PS<sub>5</sub>I and Li metal caused an increase in polarization-voltage up to the 100th cycle. While  $Li_6PS_5F_{0.75}I_{0.25}$ and Li<sub>6</sub>PS<sub>5</sub>F<sub>0.25</sub>I<sub>0.75</sub> had stable polarization-voltage. The

concentration of F dopant impacts the stability of Li<sub>6</sub>PS<sub>5</sub>F<sub>0.75</sub>- $I_{0.25}$  in Li symmetric cells, demonstrating stability for up to 1100 hours (Fig. 8b2).92

#### 2.3. Hydroborate-based ISEs

Hydroborate-type solid electrolytes provide attractive properties such as outstanding stability when combined with sodium and lithium metals, high thermal and chemical stability, soft mechanical properties, low crystallographic density, ease of processing, and low toxicity. In addition, hydroborates have an electrochemical stability window of around 4.0 V.6,51,52,59,69,70,84 Jin et al. 70 presented a novel conjuncto-hydroborate compound, Na<sub>3</sub>B<sub>24</sub>H<sub>23</sub>, which exhibits a wide electrochemical stability window. The  $\left[B_{24}H_{23}\right]^{3-}$  anion is produced by oxidizing  $[B_{12}H_{12}]^{2-}$  using the ion-exchange technique.  $Na_3B_{24}H_{23}$ xNa<sub>2</sub>B<sub>12</sub>H<sub>12</sub> was synthesized using the ball-milling method under the N<sub>2</sub> atmosphere. The Na<sub>3</sub>B<sub>24</sub>H<sub>23</sub>-5Na<sub>2</sub>B<sub>12</sub>H<sub>12</sub> composite, consisting of a double anion hydroborate, has an excess conductivity of 1.42 mS cm<sup>-1</sup> at room temperature. It also demonstrates a Na-ion transference amount of 0.97 and a wide electrochemical window of 5.8 V vs. Na/Na<sup>+</sup>. These properties make it a promising candidate for use in high-voltage all-solid-state batteries. Braun et al.59 conducted an investigation that mixed LiCB<sub>11</sub>H<sub>12</sub> and LiCB<sub>9</sub>H<sub>10</sub> and examined how different ratios of these compounds affected the oxidative

## Hydroborate solid electrolytes

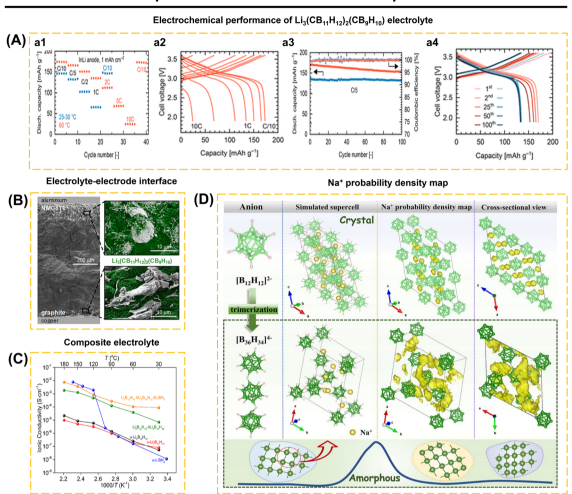


Fig. 9 (A) Solid-state cell performance at room temperature (blue) and at 60 °C (red) with Li<sub>3</sub>(CB<sub>11</sub>H<sub>12</sub>)<sub>2</sub>(CB<sub>9</sub>H<sub>10</sub>) solid electrolyte, NCM811 composite cathode, and In-Li anode, in terms of (a1 and a2) rate capability, (a3 and a4) cycling stability at C/5. (B) SEM images of the cell crosssection with schematically added current collectors: aluminum current collector, cathode composite with NCM811 and solid electrolyte, solid electrolyte separator, anode composite with graphite and solid electrolyte, and copper current collector (top to bottom). Enlarged views show SEM images of NCM811 particles and vapor-grown carbon fibers (top) and graphite flakes (bottom) embedded in Li<sub>7</sub>(CB<sub>11</sub>H<sub>12</sub>)<sub>2</sub>(CB<sub>0</sub>H<sub>10</sub>) solid electrolyte (false-colored green). Current collectors are not drawn to scale. Reproduced with permission from ref. 59. Copyright 2024, American Chemical Society. (C) Lithium ionic conductivities  $\sigma(\text{Li}^+)$  as a function of the temperature of  $\text{Li}_2\text{B}_{12}\text{H}_{12}$  –  $5\text{Li}_2\text{B}_{10}\text{H}_{10}$  –  $6\text{LiBH}_4$ , compared to the  $\text{Li}^+$ conductivity of pristine  $\alpha$ -Li<sub>2</sub>B<sub>12</sub>H<sub>12</sub>,  $\alpha$ -Li<sub>2</sub>B<sub>10</sub>H<sub>10</sub>, o-LiBH<sub>4</sub>, and Li<sub>2</sub>B<sub>12</sub>H<sub>12</sub>-5Li<sub>2</sub>B<sub>10</sub>H<sub>10</sub>. Reproduced with permission from ref. 52. Copyright 2023, American Chemical Society. (D) Proposed structural models of  $Na_4B_{36}H_{34}$  and  $Na_2B_{12}H_{12}$  ( $[B_{36}H_{34}]^{4-}$ , green;  $[B_{12}H_{12}]^{2-}$ , light green;  $Na^+$ , yellow) and the corresponding Na-ion probability density maps (yellow isosurface) from AIMD simulations at 500 K. All H atoms are omitted for clarity except for the B-H-B bridges. Reproduced with permission from ref. 69. Copyright 2024, American Chemical Society.

stability of the electrolyte.  $Li_3(CB_{11}H_{12})_2(CB_9H_{10})$  with a ratio of 2:1 shows a stability window of  $\sim$ 3.9 V vs. Li/Li<sup>+</sup> is lower than  $LiCB_9H_{10}$ -rich 1:2 ratio, with stability windows of  $\sim$ 3.6 V vs. Li/ Li<sup>+</sup>. An appropriate electrolyte can be fully integrated with a high-voltage NCM811 cathode. Meanwhile, Li-ion conductivity reaches 1.5 mS cm<sup>-1</sup> at RT and 29.4 mS cm<sup>-1</sup> at 60 °C. Electrochemical analyses of the NCM811||LiIn cell show an initial discharge capacity of 145 mA h  $g^{-1}$  (RT) and 175 mA h  $g^{-1}$  (60  $^{\circ}$ C) at C/10. This cell showed capacity retention of 98% (RT) and 90% (60 °C) after 100 cycles at C/5. Excellent cycle stability was achieved using Li-In and graphite-based anodes. In addition, the formation of anion hydroborate dimers and TiO<sub>2</sub> coating on

the surface of NCM811 also contribute to the stability of the electrode-electrolyte interface (Fig. 9A and B).

Li<sub>2</sub>B<sub>12</sub>H<sub>12</sub> and Li<sub>2</sub>B<sub>10</sub>H<sub>10</sub> in the closo-borates class show relatively low conductivity of 10<sup>-8</sup> S cm<sup>-1</sup> at RT. However, after compositing, Li<sub>2</sub>B<sub>12</sub>H<sub>12</sub>-5Li<sub>2</sub>B<sub>10</sub>H<sub>10</sub>, its conductivity is still relatively low (10<sup>-5</sup> S cm<sup>-1</sup>). In addition, fast decay and low coulombic efficiency are the disadvantages of the closo-borates class. Zhou et al.52 solved the ionic conductivity and instability problem of the  $Li_2B_{12}H_{12}$ – $5Li_2B_{10}H_{10}$  type *closo*-borate composite by introducing LiBH<sub>4</sub>. This modification increases the Li-ion conductivity to  $1.0 \times 10^{-4} \ \text{S cm}^{-1}$  at RT (Fig. 9C) and the electrochemical window reaches  $\sim$ 3.0 V. This is supported

## Antiperovskite solid electrolytes

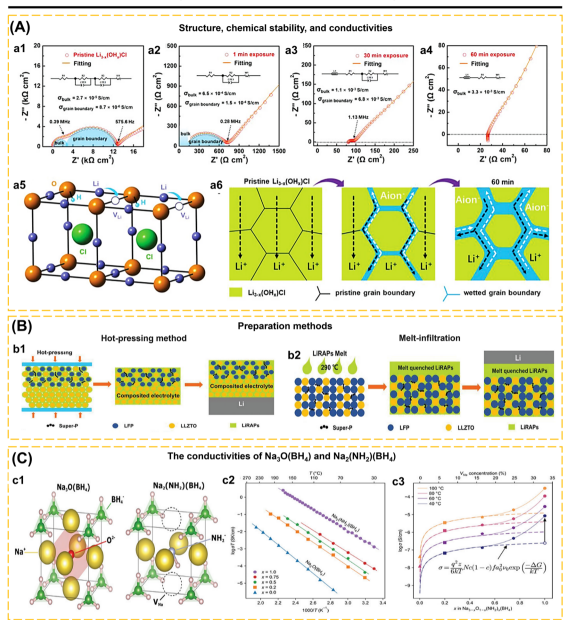


Fig. 10 (A) The effects of hygroscopicity on ionic conductivity. (a1-a4) EIS profiles of Li<sub>3-x</sub>(OH<sub>x</sub>)Cl with different expose durations: (a1) pristine, (a2) 1 min, (a3) 30 min, (a4) 60 min. (a5) Crystal structure of  $\text{Li}_{3-x}(\text{OH}_x)\text{Cl.}$  (a6) Temporal evolution of Li-ion conducting mechanisms in  $\text{Li}_{3-x}(\text{OH}_x)$ Cl. (B) Schematics of the battery assembling through hot-pressing (b1) and melt-infiltration (b2). Reproduced with permission from ref. 24. Copyright 2023, Wiley-VCH. (C) Sodium ion transport in  $Na_{3-x}O_{1-x}(NH_2)_x(BH_4)$  (x = 0-1) upon systematic substitution of  $O^{2-}$  by  $NH_2^-$ , with corresponding increase in Na<sup>+</sup> vacancy concentration. (c1) Powder X-ray diffraction (PXRD) shows that the whole series of compositions  $Na_{3-x}O_{1-x}(NH_2)_x(BH_4)$  form cubic antiperovskites with space group  $Pm\overline{S}m$  at room temperature. Between the endmember compositions  $Na_3O(BH_4)$  and  $Na_2(NH_2)(BH_4)$ , the  $Na^+$  vacancy concentration increases from zero to one-third at the full substitution of  $O^{2-}$  by  $NH_2^-$ . In the cubic unit cell Na<sub>3</sub>O(BH<sub>4</sub>) (c1, left), Na-ions at the face-centered sites comprise the vertices of the Na<sub>6</sub>O octahedra, and (BH<sub>4</sub><sup>-</sup>) cluster anions occupy the corner sites. In the cubic unit cell  $Na_2(NH_2)(BH_4)$  (c1, right), two of the six Na-ions are missing, and the  $O^{2-}$  ion residing at the center of the pseudo-octahedron is replaced with an  $NH_2^-$  anion. (c2) Temperature-dependent ionic conductivity of  $Na_{3-x}O_{1-x}(NH_2)_x(BH_4)$  (where x=0, 0.2, 0.5, 0.75, and 1.0) exhibits Arrhenius behavior above the room temperature region and ionic conductivity increases with the substitution of  $O^{2-}$  by NH<sub>2</sub><sup>-</sup>. Activation energies are 0.358, 0.299, 0.301, 0.278, and 0.268 eV, respectively. The endmember composition Na<sub>2</sub>(NH<sub>2</sub>)(BH<sub>4</sub>) with the highest Na $^+$  vacancy and NH $_2^-$  cluster anion concentration shows 10 $^4$  higher ionic conductivity than that of the endmember Na $_3$ O(BH $_4$ ) without the substitution of NH2-. (c3) The dependence of ionic conductivity on the concentration of Na+ vacancy and NH2- cluster ion concentration at several temperatures between 40 °C and 100 °C is fitted as the dashed line using the vacancy-dependent ionic conductivity model (inset equation). At low  $x \leq 0.75$ ), the ionic conductivity can be fit well by the classical vacancy-concentration model. At high  $x \leq 0.75$ , the ionic conductivity of  $Na_{3-x}O_{1-x}(NH_2)_x(BH_4)$  exhibits a large positive deviation from the expected classical vacancy-dependent conductivity. It is correlated with not only high vacancy concentration but also the  $NH_2^-$  cluster anion concentration, suggesting a paddle-wheel effect. Reproduced with permission from ref. 26. Copyright 2022, Wiley-VCH.

by <sup>7</sup>Li MAS NMR spectra data showing that the full-width at half maximum of  $\text{Li}_2\text{B}_{12}\text{H}_{12}$ – $5\text{Li}_2\text{B}_{10}\text{H}_{10}$ – $6\text{LiBH}_4$  (2.4 ppm) is smaller than Li<sub>2</sub>B<sub>12</sub>H<sub>12</sub>-5Li<sub>2</sub>B<sub>10</sub>H<sub>10</sub> (5.2 ppm), which demonstrates the rapid diffusion of Li-ion in solid-state electrolytes based on  $\text{Li}_2\text{B}_{12}\text{H}_{12}$ -5 $\text{Li}_2\text{B}_{10}\text{H}_{10}$ -6 $\text{LiBH}_4$ . In a recent study by Jin et al.<sup>69</sup> a nonflammable compound called Na<sub>4</sub>B<sub>36</sub>H<sub>34</sub> was produced. This compound was synthesized by oxidizing  $[B_{12}H_{12}]$ . At high temperatures, the compound forms the trimeric anion [B<sub>36</sub>H<sub>34</sub>]<sup>4-</sup> exhibiting an electrochemical stability range of up to 6.9 V. Density functional theory (DFT) investigation verified that the polyanion  $[B_{36}H_{34}]^{4-}$  may lead to the expansion of interstitial sites, creating a favorable channel for the diffusion of sodium ions. This was further corroborated by the paddle-wheel effect of the polyanion  $[B_{36}H_{34}]^{4-}$  (Fig. 9D). The Na<sub>4</sub>B<sub>36</sub>H<sub>34</sub>-7Na<sub>2</sub>B<sub>12</sub>H<sub>12</sub> solid electrolyte exhibits numerous benefits within the hydroborate-SE class. These include low electronic conductivity  $(9.3 \times 10^{-12} \text{ S cm}^{-1})$ , Na<sup>+</sup> transference number of 0.97, ionic conductivity of  $1.02 \times 10^{-3}$  S cm<sup>-1</sup>, and an electrochemical stability window of 5.5 V. In a symmetric cell configuration (Na|Na<sub>4</sub>B<sub>36</sub>H<sub>34</sub>-7Na<sub>2</sub>B<sub>12</sub>H<sub>12</sub>|Na), it can maintain stability for over 200 hours at 0.1 mA cm<sup>-2</sup>. Additionally, it possesses favorable properties such as good compressibility, non-flammability, and decent mechanical properties. The application of Na<sub>4</sub>B<sub>36</sub>H<sub>34</sub> as solid-state electrolytes is advantageous for high-voltage all-solid-state batteries.

#### 2.4. Antiperovskite-based ISEs

Antiperovskite, reverse-charged versions of perovskite, with the general structure (X3AB, where X is an alkali cation, A and B are anions), is a highly potential solid electrolyte because it can be processed at low temperatures, has a wide electrochemical stability window, high ionic conductivity, good reduction melting point.189-192 Lithium-rich and low antiperovskite-type solid electrolytes have garnered significant interest due to their large ion vacancy area and low activation energy barrier.32 Li-rich antiperovskite (LiRAP) materials demonstrate excellent ionic conductivity and high electrochemical stability when used with Li metal anodes.24,74,94 Antiperovskite has features that make it suitable for use as a protective layer for Li metal anodes, solid electrolyte films in thin film batteries, and solid electrolytes in all-solid-state batteries. 189 Li<sub>3-x</sub>(OH<sub>x</sub>)Cl is an example of LiRAPs that exhibit instability in the presence of air and undergo decomposition to form LiOH and LiCl.24 Li3-x(OHx)Cl has a tetragonal structure at room temperature but transitions to a cubic form at 40 °C, this structure arises from the presence of lithium vacancies in the absence of cation or anion doping (Fig. 10a5). The  $Li_{3-x}(OH_x)Cl$ compound has a conductivity of around 7.1  $\times$  10<sup>-6</sup> S cm<sup>-1</sup> when not exposed to air (Fig. 10a1). After being subjected to ambient air conditions (25 °C, 25  $\pm$  5% RH) for 1 minute, the conductivity rises by  $1.2 \times 10^{-4} \text{ S cm}^{-1}$  ( $E_a = 0.45 \text{ eV}$ ) and reaches  $3.3 \times 10^{-3} \text{ S cm}^{-1}$  after 60 minutes (Fig. 10a2-a4). However, non-Arrhenius behavior occurs as the temperature goes over 60 °C, causing a reduction in the ionic conductivity of  $\text{Li}_{3-x}(\text{OH}_x)\text{Cl.}$  Fig. 10a6 indicates that prior to being exposed to open air, Li<sub>3-x</sub>(OH<sub>x</sub>)Cl exhibits significant resistance in both its

bulk and grain boundaries. By extending the duration of exposure, moisture may be absorbed via the grain borders, resulting in a reduction in grain boundary resistance. After 60 minutes, all grain boundaries are fully wetted, and the Nyquist plot only shows the Warburg resistance, while the bulk resistance is attributed to ionic migration in the aqueous solution.  $\text{Li}_{3-x}(-\text{OH}_x)\text{Cl}$  was determined to be unstable in the presence of air. The hygroscopic features can be decreased by introducing 0.1 molar F on the Cl site. This weakens the hydrogen bonds between LiRAPs and  $\text{H}_2\text{O}$ , resulting in the formation of  $\text{Li}_{3-x}(-\text{OH}_x)\text{Cl}_{0.9}\text{F}_{0.1}$ . This compound exhibits resistance to moisture, with a water content of 0.27 wt% for pellets and 0.93 wt% for powder. Unlike  $\text{Li}_{3-x}(\text{OH}_x)\text{Cl}_{0.9}\text{F}_{0.1}$  exhibits an enhancement in conductivity until a duration of 60 minutes  $(2.9 \times 10^{-5} \text{ S cm}^{-1})$ .

The mechanical softness and low melting point (274 °C) of  $\text{Li}_{3-x}(\text{OH}_x)\text{Cl}_{0.9}\text{F}_{0.1}$  antiperovskite make it applicable as a composite material for ASSBs electrodes in two ways. The first technique used is the hot-pressing method, where a layer of antiperovskite material is applied onto the surfaces of LFP and LLZTO (Li<sub>6.75</sub>La<sub>3</sub>Zr<sub>1.75</sub>Ta<sub>0.25</sub>O<sub>12</sub>). Antiperovskite may function as an adhesive, enhancing the contact between LFP and LLZTO, and reducing strain on the cathode. The cathode and electrolyte are combined using the hot-pressing technique at a temperature of 290 °C. This is followed by the addition of Li metal on the opposite side (Fig. 10b1). The second technique involves the melt-infiltration method, in which the antiperovskite is heated to 290 °C and then poured in excess onto the cathode composite mixture to fill the cathode cavity. This results in the formation of a solid antiperovskite membrane over the cathode composite. The membrane is subsequently collected along with Li metal (Fig. 10b2). The initial discharge capacity of the two manufacturing processes is competitive, with 1st-type ASSBs exhibiting a capacity of about 83.6 mA h  $g^{-1}$  and 2nd-type ASSBs exhibiting a capacity of approximately 70.8 mA h  $g^{-1}$  at a rate of 0.05C.

The ionic conductivity of  $Li_2OHCl_{1-x}Br_x$  (x = 0, 0.1, 0.3, 0.5,0.7, 0.9, and 1) antiperovskite electrolyte was investigated by Lee et al.31 The crystal structure of Li<sub>2</sub>OHCl<sub>1-x</sub>Br<sub>x</sub> is significantly influenced by the Br to Cl ratio, as proven by X-ray diffraction studies. This ratio directly affects the conductivity of Li ions inside the grains. It can be deduced that both the Li<sub>2</sub>OHCl and Li<sub>2</sub>OHCl<sub>0.9</sub>Br<sub>0.1</sub> structures are defect-rich and have a distorted structure, as shown by the reduction in the Li-O distance around the Schottky defect. In the Li<sub>2</sub>OHCl structure, the Li-O distance decreases from 1.926 Å to 1.814 Å (a decrease of 0.112 Å), while in the Li<sub>2</sub>OHCl<sub>0.9</sub>Br<sub>0.1</sub> structure, the Li-O distance decreases from 1.929 Å to 1.839 Å (a decrease of 0.09 Å). The Li<sub>2</sub>OHCl<sub>0.9</sub>Br<sub>0.1</sub> compound exhibited a higher ionic conductivity  $(2.52 \times 10^{-3} \text{ mS cm}^{-1})$  compared to Li<sub>2</sub>OHCl  $(1.93 \times 10^{-3}$ mS cm<sup>-1</sup>). While the replacement of Br may maintain the cubic crystal structure under room temperature conditions, a high ratio of Br to Cl might hinder the migration of Li-ions owing to the larger size of the Br ion. After 7 days, EIS data indicates that the interfacial resistance between the solid electrolyte and Li metal in the Li|Li<sub>2</sub>OHCl<sub>0.9</sub>Br<sub>0.1</sub>|Li cell has increased to 705  $\Omega$ cm<sup>2</sup> at 80 °C. After 200 hours at 80 °C, the voltage profile of the

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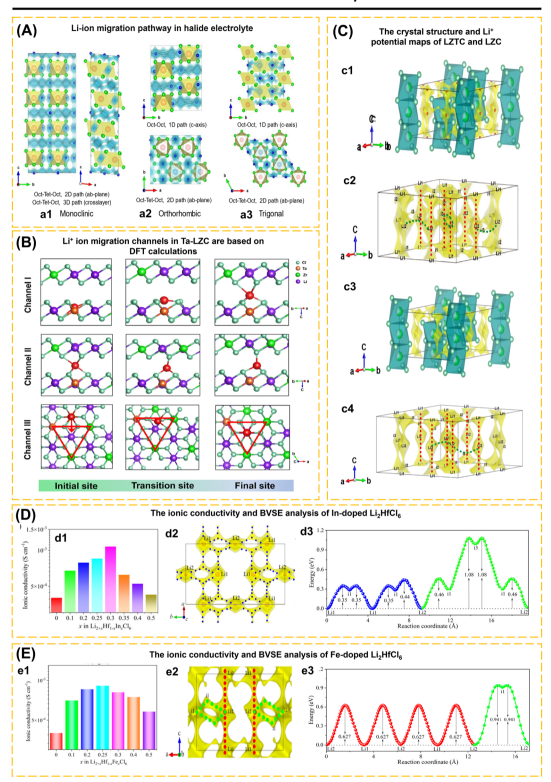


Fig. 11 (A) Li-ion migration pathway in (a1) monoclinic, (a2) orthorhombic, and (a3) trigonal Li<sub>3</sub>MCl<sub>6</sub> obtained using the BVSE method. Blue atom and yellow polyhedral correspond to Li and MCl<sub>6</sub>, respectively, while the isosurface with light blue corresponds to the ionic migration path. The monoclinic phase (CCP anion sublattice) exhibits two-dimensional intra-layer and three-dimensional cross-layer paths between octahedral sites via tetrahedral interstitial sites (Oct-Tet-Oct), while trigonal and orthorhombic phase (HCP anion sublattice) exhibit anisotropic paths including one dimensional path along c-axis between octahedral sites (Oct-Oct) and two-dimensional migration path along ab plane between octahedral sites via interstitial tetrahedral sites (Oct-Tet-Oct). Reproduced with permission from ref. 179. Copyright 2020, American Chemical Society. (B) The key local environment fragments of three Li<sup>+</sup> ion migration channels in Ta-LZC are based on DFT calculations. The red ball is highlighted to

Li-Li symmetric cell remains constant and does not show any increase in polarization.

Tsai et al.,26 Ahiavi et al.,193 and Sun et al.194 elucidate the "paddle-wheel" effects, which enhance the movement of cations in solid electrolytes. Structures that include cluster anions, also known as polyanions, complex anions, or rotor groups, such as sulfate (SO<sub>4</sub><sup>2-</sup>), nitrite (NO<sub>2</sub><sup>-</sup>), borohydride (BH<sub>4</sub><sup>-</sup>), phosphate (PO<sub>4</sub><sup>3-</sup>), aluminum hydride (AlH<sub>4</sub><sup>-</sup>), thiophosphates (PS<sub>4</sub><sup>3-</sup>), closo-borates and carba-closo-borates, etc. 195 These structures have a larger ion radius and create a "paddle-wheel" effect, where the rotation of the anion group facilitates the movement of cations with a lower energy barrier for migration, resulting in enhanced cation mobility. Cluster anions can enhance the conductivity of cations compared to configurations containing single atoms that are similar in nature. Systems containing numerous types of anion clusters are believed to exhibit various ways of motion using multiple paddle wheels and cations. Antiperovskite is a suitable framework for investigating the "paddle-wheel" phenomenon due to its ability to include various ionic substitutions, including cluster anions. The research published by Tsai et al.26 showcased the utilization of sodium oxide borohydride (Na<sub>3</sub>O(BH<sub>4</sub>)) with a unique antiperovskite structure consisting of a single umbrella wheel. In this structure, the oxide anion fraction (O<sup>2-</sup>) was substituted with an amide cluster anion (NH<sub>2</sub><sup>-</sup>) (Fig. 10c1). The compound  $Na_{3-x}O_{1-x}(NH_2)_x(BH_4)$  (where x = 0, 0.2, 0.5, 0.75, 1.0) was synthesized using a solid-state reaction method. The PXRD analysis reveals that the primary phases of Na<sub>3</sub>O(BH<sub>4</sub>) and  $Na_{3-x}O_{1-x}(NH_2)_x(BH_4)$  exhibit a cubic crystal structure ( $Pm\bar{3}m$ ). Furthermore, when the value of x rises, the lattice parameters likewise increase. Borohydride cluster anions are located at the corners of a cube, whereas amide/oxide anions are located in the center of the cube. Sodium cations are positioned at the center of each face of the cube, forming an octahedron. At x = 1, there are two vacant octahedral sites, and all body center sites are occupied by amide cluster anions. An increase in the concentration of the amino group anion (NH2 ) leads to a corresponding rise in both the ionic conductivity and the concentration of cation vacancies. The NH<sub>2</sub><sup>-</sup> and BH<sub>4</sub><sup>-</sup> species have a Na-ion conductance that is 100 times greater when x = 1(Fig. 10c2). Despite the conductivity values plotted with the sodium vacancy concentration exhibiting no correlation between the two (Fig. 10c3). Extensive investigation and comprehension of the interactions between cluster anions and cations are crucial for the advancement of solid-state electrolytes in the future.

#### 2.5. Halide-based ISEs

Halides exhibit high conductivity (>1 mS cm<sup>-1</sup> (a) RT), oxidation stability (~4 V), better performance under high voltage, and temperature processability. 43,196 Several articles have described the division of halide electrolytes. Based on the type of metal element, halides are divided into three categories, i.e., Li<sub>a</sub>MX<sub>b</sub> halide with group 3 elements (Sc, Y, La-Lu), Li<sub>a</sub>MX<sub>b</sub> halide with group 13 elements (M = Al, Ga, In), and  $Li_aMX_b$  halide with divalent metal elements (M = Ti, V, Cr, Mn, Fe, Co, Ni, Cu, Zn, Cd, Mg, Pb).45 Huang et al.43 describes the division of halide division, which is  $\text{Li}_3\text{M}^{3+}\text{X}_6$  (M<sup>3+</sup> = In, Sc, Ti, Y, Ho, Er, Yb and X = Cl, Br, I),  $\text{Li}_2 M^{4+} X_6$  ( $M^{4+} = \text{Zr}$ , Hf and X = Cl, Br, I), and  $LiM^{5+}Cl_4O$  ( $M^{5+}=Nb$ , Ta). Based on the sublattice structure of halogen anions for ternary halides: CCP (cubic  $Fd\bar{3}m$  symmetry and monoclinic C2/m symmetry) and HCP (trigonal  $P\bar{3}m1$ symmetry and orthorhombic Pnma symmetry). 197 Park and coworker<sup>179</sup> investigated the Li-ion transport pathways in Li<sub>3</sub>MCl<sub>6</sub> (where M = Bi, Dy, Er, Ho, In, Lu, Sc, Sm, Tb, Tl, Tm, and Y) utilizing BSVE and AIMD methods. The analysis revealed that the monoclinic structure exhibited the lowest migration energy barrier, approximately 0.5 eV, compared to the orthorhombic and trigonal structures. AIMD simulations indicated that the orthorhombic and trigonal structures exhibited slow ion migration within the 2D plane, leading to low ionic diffusion and increased activation energy. The monoclinic structure shows an Oct-Tet-Oct transport mechanism along both the 2D (ab-plane) and 3D (cross-layer) pathways. In the trigonal and orthorhombic structures, an Oct-Oct transport mechanism is observed along the 1D path, while an Oct-Tet-Oct mechanism is present along the 2D path (Fig. 11a1-a3).

The ionic conductivity enhancement of halide-based ISEs is affected by the types of substituted aliovalent cations and their respective substitution ratios. 43,45,56,198-200 A novel halide structure,  $\text{Li}_{2-x}\text{Zr}_{1-x}\text{Ta}_x\text{Cl}_6$  (0  $\leq x \leq$  0.7), was synthesized through high-energy ball-milling.46 The LZC structure consists of two layers: layer A (Li<sub>A</sub>), which is occupied by ZrCl<sub>6</sub><sup>2-</sup> and LiCl<sub>6</sub><sup>5-</sup> octahedra, and layer B (Li<sub>B</sub>), positioned between layer A. The migration route of Li-ions can be explained through three primary channels: channel I represents the conduction path of Li-ions in the c direction; channel II denotes the conduction path along the a and b directions within layer B; and channel III indicates the conduction path in the a and b directions of layer A. The integration of the three channels establishes the threedimensional migration pathway in LZTC (Fig. 11b). The [Li1-Li2-Li1] transport pathway in the c direction is the most favorable for LZC and LZTC. Nonetheless, the migration

illustrate the migrations of Li<sup>+</sup> ions in three different channels. Concisely, channel I for Li<sup>+</sup> ion conduction pathway in c direction. Channel II for Li<sup>+</sup> ion conduction pathway in a and b directions of layer A. (C) (c1) The crystal structure of LZTC superimposed with the Li<sup>+</sup> potential map. (c2) Li<sup>+</sup> migration pathways of LZTC. (c3) The crystal structure of LZC is superimposed with the Li<sup>+</sup> potential map. (c4) Li<sup>+</sup> migration pathways of LZC. Reproduced with permission from ref. 46. Copyright 2024, Elsevier. (D) (d1) The Li-ion conductivities of Li<sub>2+x</sub>Hf<sub>1-x</sub>In<sub>x</sub>Cl<sub>6</sub> (0  $\leq$  x  $\leq$  0.5). (d2) Li-ions migration pathways within the LHIC crystal structure along with the ab plane. (d3) Energy barrier profile of various migration pathways within the LHIC crystal structure, and i1 and i3 represent the different interstitial tetrahedra (Tet) sites. Reproduced with permission from ref. 198. Copyright 2023, Elsevier. (E) (e1) the Li-ion conductivities Li<sub>2+x</sub>Hf<sub>1-x</sub>Fe<sub>x</sub>Cl<sub>6</sub> (0  $\leq$  x  $\leq$  0.5). (e2) Diffusion pathways of the Li-ion within LHFC crystalline structure through the face-sharing lithium-centered octahedron along the c axis. (e3) Energy barrier profile of different Li-ion diffusion pathways within the LHFC crystalline structure. Reproduced with permission from ref. 199. Copyright 2024, American Chemical Society.

barriers will vary as a result of Ta doping. Ta<sup>5+</sup> doping in LZC decreases the migration barriers for Li-ions along the [Li1-Li2-Li1] pathway in the c direction and the [Li2–Li3–Li2] pathway in the ab plane. The migration barriers of LZTC are lower than those of LZC by 0.849 eV and 0.988 eV, respectively. The substitution of Ta<sup>5+</sup> is concluded to decrease the migration barriers of Li<sup>+</sup> and enhance the transport of Li<sup>+</sup> within the crystal lattice (Fig. 11c1-c4).

Li<sub>2</sub>HfCl<sub>6</sub> type halides were synthesized through mechanical milling, with partial substitution of Hf4+ atoms by M3+ ions (Fe<sup>3+</sup>, V<sup>3+</sup>, Cr<sup>3+</sup>, and In<sup>3+</sup>). 198,199 The variation in atomic radius significantly influences the structure of the bare LHC, specifically Hf<sup>4+</sup> (71 pm), Fe<sup>3+</sup> (65 pm), Cr<sup>3+</sup> (61.5 pm), V<sup>3+</sup> (64 pm), and In<sup>3+</sup> (94 pm). The optimal percentage and ionic conductivity of  $M^{3+}$  substituted Li<sub>2+x</sub>Hf<sub>1-x</sub>M<sub>x</sub>Cl<sub>6</sub> are as follows: Cr<sup>3+</sup> (x = 0.2, 0.81 mS cm<sup>-1</sup>),  $V^{3+}$  (x = 0.2, 0.64 mS cm<sup>-1</sup>),  $In^{3+}$  (x = 0.3, 1.05 mS cm<sup>-1</sup>) (Fig. 11d1) and Fe<sup>3+</sup> ( $x = 0.25, 0.91 \text{ mS cm}^{-1}$ ) (Fig. 11e1). The substitution of  $Hf^{4+}$  with  $M^{3+}$  (where  $rHf^{4+}$  <  $rM^{3+}$ ) will result in an expansion of the Li-ion diffusion barrier. Conversely, if  $rHf^{4+} > rM^{3+}$ , the pure LHC crystal lattice will contract, leading to a disruption in the distribution of Li-ions. Optimization of the concentration of substituted M<sup>3+</sup> is essential; excessive amounts can reduce ionic conductivity due to a deficiency of vacancies in the LHC structure. The diffraction pattern of LHC is identical to that of Li<sub>3</sub>YCl<sub>6</sub> (P3m1). Substituting Fe  $(0 \le x \le 0.5)$  into LHC results in trigonal  $\text{Li}_3\text{YCl}_6$ . In the case of  $\text{In}^{3+}$  substitution in LHC, if  $\text{In}^{3+}$  (x > 0.3) leads to a monoclinic Li<sub>3</sub>InCl<sub>6</sub> structure. The In<sup>3+</sup>-substituted LHC exhibits a diffusion path of [Li1-Tet.1-Li1] in the ab plane with an energy barrier of 0.35 eV. Furthermore, an additional alternative path [Li1-Tet.1-Li2] is linked with the [Li2-Tet.1-Tet.3-Tet.1-Li2] chain to establish a three-dimensional network (1.08 eV) (Fig. 11d2 and d3). In the case of LHC, an energy barrier of 1.31 eV exists between two adjacent ab planes. The Fe<sup>3+</sup>-substituted LHC exhibits a diffusion path of [Li2-Li1-Li2] along the c direction, characterized by an energy barrier of 0.627 eV. This path is interconnected with the [Li2-Tet.1-Li2] path, resulting in a three-dimensional diffusion network with an energy barrier of 0.941 eV (Fig. 11e2 and e3). The energy remains lower than that of pure LHC and In<sup>3+</sup>-doped LHC. Aliovalent substitution induces a rearrangement of Li-ions and an anisotropic distortion of the local structure, resulting in a reduction of migration barrier energy. This process facilitates the formation of efficient Li-ion diffusion pathways and enhances ionic conductivity.

 $\text{Li}_{3-x}Y_{1-x}\text{Hf}_x\text{Cl}_6$  (0  $\leq x < 1$ ) were synthesized through mechanical milling.200 Pristine LYC and Hf-doped LYC exhibited a uniform and interconnected particle size of less than 10 μm, characterized by a porous structure. The diffraction pattern of LYC closely resembled the space group P3m1. The soft nature and low crystallinity of LYHC facilitated its processing into dense pellets through cold pressing. LYHC exhibited an ionic conductivity of 1.49 mS cm<sup>-1</sup>, surpassing that of pristine LYC, which measured 0.139 mS cm<sup>-1</sup>. In LYHC, the radius of the doping ion  $Hf^{4+}$  (71 pm) is smaller than that of  $Y^{3+}$  (90 pm). The substitution of  $Hf^{4+}$  (x > 0.6) in LYC results in a narrowing of the crystal lattice and ionic migration channels, leading to

a reduced concentration of Li-ions. Analysis of the Bond Valence Energy Landscape indicates Li<sup>+</sup> migration LYHC. The transition from Li1 (6h) to Li2 (6g) may occur along the c-axis. The Li1 site exhibits six-fold coordination perpendicular to the z-axis and is situated between the Y1-Cl<sub>6</sub> octahedra. Li2 atoms are linked to other Li2 atoms in the a or b direction. The percolation energy calculated along the three axes is uniform at 2.74 eV. Hf-doped LYC exhibits a percolation energy of approximately 2.8 eV, facilitating the three-dimensional diffusion of Li-ions.

The combination of good deformability, high Ionic conductivity, and electrochemical stability allows for the synthesis of a novel group of halides with heterogeneous structures by combining high- and low-coordinated halide frameworks. Fu et al. 30 synthesized a compound by combining the UCl3-type halide framework, which is based on a highcoordination framework with a coordination number of more than 6, such as  $Na_{3x}M_{2-x}Cl_6$  (M = La, Sm), and a lowcoordination framework represented by scattered [MCl<sub>6</sub>]<sup>x-</sup> octahedra, such as NaTaCl6 (LCF-Ta). The arrangement of Na<sub>0.75</sub>Sm<sub>1.75</sub>Cl<sub>6</sub> (HCF-Sm) and Na<sub>0.75</sub>La<sub>1.75</sub>Cl<sub>6</sub> (HCF-La) is shown as a tricapped trigonal prism, with the Na atom occupying the octahedral site and the La or Sm atoms occupying the other sites (Fig. 12a1). The LCF-Ta crystal consists of an octahedral framework [TaCl<sub>6</sub>] positioned at the corner of the unit cell  $(P2_1/c)$ . This framework shares an edge with two  $[NaCl_7]^{6-}$ closed trigonal prisms. The mixing of HCF and LCF leads to the formation of new heterogeneous structures, the study revealed that the optimal ionic conductivity was 2.7 mS cm<sup>-1</sup> for the combination of 0.62[HCF-Sm]·0.38[LCF-Ta], and 1.8 mS cm<sup>-1</sup> for the combination of 0.57[HCF-La]·0.43[LCF-Ta], both at a temperature of 25 °C.

A novel class of zeolite-like halide framework called SmCl<sub>3</sub> was introduced by Fu et al.2 The ionic radius of metal atoms in Li-M-Cl halides, such as Li<sub>3</sub>InCl<sub>6</sub>, Li<sub>3</sub>YCl<sub>6</sub>, and Li<sub>2</sub>ZrCl<sub>6</sub>, are in close proximity. Specifically, Zr<sup>4+</sup> (72 pm), Li<sup>+</sup> (76 pm), In<sup>3+</sup> (80 pm), and  $Y^{3+}$  (90 pm) form an octahedral framework  $[MCl_6]^{x-}$ . When the ionic radius of the atom M is raised, specifically for M = Sm $^{3+}$  (113.2 pm) and La $^{3+}$  (121.6 pm), the halide structure undergoes a transition from 6-fold coordination to either 8-fold or 9-fold coordination. This transition occurs in a P6<sub>3</sub>/m crystal lattice and is accompanied by the formation of 1D vacancy channels along the *c*-axis. The obtained pore size is comparable to the pore size of zeolite (Fig. 12a2). Zeolite-like SmCl<sub>3</sub> framework has a one-dimensional channel that is surrounded by [SmCl<sub>9</sub>]<sup>6-</sup> structures, allowing for a small distance of 2.08 Å between two octahedral structures, enabling the transit of Liions. AIMD simulations indicate that the Li<sup>+</sup> diffusion is facilitated by the presence of several tunnels (between distorted octahedral) promoting efficient diffusion pathways for mobile ions (Fig. 12b1). BM-SmCl<sub>3</sub>, produced through the ball-milling method, can be combined with halide species like LiCl, LiF, LiBr, LiI, Li<sub>2</sub>ZrCl<sub>6</sub>, LiFeCl<sub>4</sub>, LiAlCl<sub>4</sub>, and Li<sub>2</sub>HfCl<sub>6</sub> to improve the ionic conductivity (Fig. 12b2). The solid electrolyte BM-SmCl<sub>3</sub>-·0.5Li<sub>2</sub>ZrCl<sub>6</sub>, which exhibits the highest ionic conductivity, is suitable for use as a solid electrolyte in ASSBs and this solid

### Halide solid electrolytes

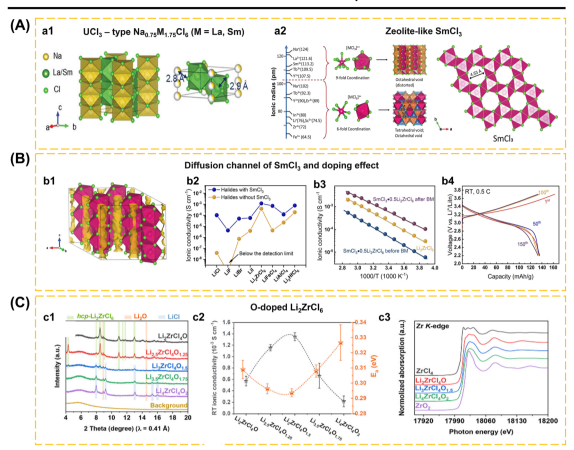


Fig. 12 (A) (a1) The crystal structure of UCl<sub>3</sub>-type Na<sub>0.75</sub>M<sub>1.75</sub>Cl<sub>6</sub> (M = La, Sm). Reproduced with permission from ref. 30. Copyright 2023, Wiley-VCH. (a2) Porous framework structure based on halides. Relationship between the ionic radius of M and the coordination of [MCl<sub>x</sub>] polyhedra. The black arrows on the crystal structures indicate the  $Li^+$  diffusion pathway. Top view of the SmCl<sub>3</sub> lattice along the c axis to show the existing plentiful channels with an inner diameter of 4.53 Å. Reproduced with permission from ref. 2. Copyright 2023, American Chemical Society. (B) (b1) Crystal structure of SmCl<sub>3</sub> superimposed with Li<sup>+</sup> probability density (yellow iso-surface) from AIMD simulations of Li0.17SmCl<sub>3</sub> at 300 K. Reproduced with permission from ref. 2. Copyright 2023, American Chemical Society. (b2) Ionic conductivities of the SmCl<sub>3</sub> framework coupled with different halide adsorbents at 30 °C. (b3) Arrhenius-plots of Li<sub>2</sub>ZrCl<sub>6</sub> and the SmCl<sub>3</sub>·0.5Li<sub>2</sub>ZrCl<sub>6</sub> (before and after ball-milling). (b4) Charge and discharge voltage profiles of the ASSLIB using LiNi<sub>0.83</sub>Mn<sub>0.06</sub>Co<sub>0.11</sub>O<sub>2</sub> (NCM83) as the cathode and SmCl<sub>3</sub>·0.5Li<sub>2</sub>ZrCl<sub>6</sub> as the ISE layer at different cycles (1st, 50th, 100th, and 150th). Reproduced with permission from ref. 2. Copyright 2023, American Chemical Society. (C) (c1) SXRD patterns of the  $Li_{2+2x}ZrCl_4O_{1+x}$  (x=0, 0.25, 0.5, 0.75, and 1). (c2) Comparison of ionic conductivities and activation energies of the  $Li_{2+2x}$ - $ZrCl_4O_{1+x}$ . (c3) XANES spectra of the  $Li_2ZrCl_4O$ ,  $Li_3ZrCl_4O_{1.5}$ , and  $Li_4ZrCl_4O_2$  at the Zr K-edge, respectively.  $ZrCl_4$  and  $ZrO_2$  are reference samples. Reproduced with permission from ref. 57. Copyright 2024, American Chemical Society.

electrolyte demonstrates stability for up to 150 cycles (Fig. 12b3 and b4).

Zhang et al. 57 investigated the impact of the amorphous structure on the mechanical deformability and Li<sup>+</sup> conduction of the quasi-crystalline Li<sub>2</sub>ZrCl<sub>6</sub> structure, which was partly replaced by O at the Cl site. The  $\text{Li}_{2+2x}\text{ZrCl}_4\text{O}_{1+x}$  ISE (where x=0, 0.25, 0.5, 0.75, and 1) is generated by adjusting the proportion of ZrCl<sub>4</sub> and Li<sub>2</sub>O in the synthesis process. Raising the quantity of Li<sub>2</sub>O decreases the formation of crystalline hcp-Li<sub>2</sub>ZrCl<sub>6</sub>. When the value of x is 0.5, the level of amorphization is 89.5%, and there are minor impurities, including hcp-Li<sub>2</sub>ZrCl<sub>6</sub>, LiCl, and Li<sub>2</sub>O. The substitution of O reaches 42.9%. The Li<sub>3</sub>ZrCl<sub>4</sub>O<sub>1.5</sub> solid electrolyte has the highest ionic conductivity, measured at  $(1.35 \pm 0.07) \times 10^{-3}$  S cm<sup>-1</sup>, and the lowest activation energy of  $0.294 \pm 0.003$  eV (Fig. 12c2). Li $_3 Zr Cl_4 O_{1.5}$  ISE has a much lower electronic conductivity of  $7.10 \times 10^{-10} \text{ S cm}^{-1}$  compared to  $\text{Li}_2\text{ZrCl}_6$ , which has an electronic conductivity of 5.61  $\times$ 10<sup>-9</sup> S cm<sup>-1</sup>. Zr K-edge FT-EXAFS spectroscopy is utilized to quantitatively determine the coordination system around the Zr atom in  $\text{Li}_{2+2x}\text{ZrCl}_4\text{O}_{1+x}$  ISEs (x = 0, 0.5, and 1). Increasing the value of x leads to an increase in Zr-O intensity, indicating a higher degree of anion replacement of Cl by the O anion (Fig. 12c3). Consequently, the Cl anion combines with the Li cation to form a LiCl compound, which is in agreement with the SXRD results (Fig. 12c1).

Li et al.54 reported a crystal structure called ternary halide, which is formed by combining LaCl<sub>3</sub> and CeCl<sub>3</sub>. This crystal structure follows the UCl<sub>3</sub> type structure with space group P6<sub>3</sub>/

m. It has one-dimensional hexagonal channels that may hold monovalent cations such as Li<sup>+</sup>, Na<sup>+</sup>, K<sup>+</sup>, Cu<sup>+</sup>, and Ag<sup>+</sup>. The UCl<sub>3</sub>type superionic chloride SE is composed of LiCl, LaCl<sub>3</sub>, CeCl<sub>3</sub>, AlCl<sub>3</sub>, TaCl<sub>5</sub>, ZrCl<sub>4</sub>, and HfCl<sub>4</sub>. The ionic conductivity values for different combinations of halides are as follows: Li-LaCeHfAlTa-Cl (1.3 mS cm<sup>-1</sup>), Li-LaCeZrHfTa-Cl (1.8 mS cm<sup>-1</sup>), Li-LaCeZrAlTa-Cl (1.0 mS cm<sup>-1</sup>), Li-LaCeZrHfAlTa-Cl (1.1 mS cm<sup>-1</sup>). The activation energy of certain UCl<sub>3</sub>-type chloride solid-state electrolytes is quite low. Examples include Li-LaCeZrHfTa-Cl (0.318 eV), Li-LaCeZrAlTa-Cl (0.336 eV), Li-LaCeHfAlTa-Cl (0.364 eV), Li-LaCeZrHfAlTa-Cl (0.383 eV), Li-LaZr-Cl (0.404 eV), and Li-La-Cl (0.511 eV). The modification is achieved by substituting LiCl with Li2O and introducing monovalent cations, namely Na<sup>+</sup>, K<sup>+</sup>, Cu<sup>+</sup>, and Ag<sup>+</sup>. These cations are denoted as Li-LaCeZrHfTa-OCl, Na-LaCeZrHfTa-Cl, K-LaCeZrHfTa-Cl, Cu-LaCeZrHfTa-Cl, and Ag-LaCeZrHfTa-Cl. The Li-LaCeZrHfTa-OCl sample exhibits an ionic conductivity of approximately 1.02 mS cm<sup>-1</sup> at 25 °C. In comparison, the Na-LaCeZrAlTa-Cl sample has a conductivity

of 0.88 mS cm<sup>-1</sup> at the same temperature and an activation energy of 0.326 eV. The K-LaCeZrAlTa-Cl sample, on the other hand, has a conductivity of  $1.32 \times 10^{-3}$  mS cm<sup>-1</sup> at a temperature of 55 °C and an activation energy of 0.552 eV. The Cu-LaCeZrAlTa-Cl sample demonstrates a conductivity of 7.13 mS cm<sup>-1</sup> at 25 °C with an activation energy of 0.283 eV. Lastly, the Ag-LaCeZrAlTa-Cl sample displays a conductivity of 7.75 mS cm<sup>-1</sup> at 25 °C and an activation energy of 0.264 eV.

Wang et al.87 investigated the air stability characteristics of Li<sub>3</sub>InCl<sub>6</sub>-based halides. Pelletized halides exhibit a greater relative density compared to their powdered forms. Applying pressure ranging from 100 to 400 MPa during pelletizing enhances relative density and stability in air. The water absorption rate of halides in air correlates with their relative density. Li<sub>3</sub>InCl<sub>6</sub> absorbs water and will form hydrate crystals and partially decompose into InCl3 and LiCl. Subsequent hydrolysis of InCl<sub>3</sub> results in the formation of corrosive compounds. The pH value of LiCl indicates no significant change; however, the hydrolysis of In<sup>3+</sup> results in the formation

Table 2 Recently reported Na-based ISEs with synthetic method and electrochemical properties

Solid electrolyte	Synthesis method	Ionic conductivity	$E_{\mathrm{a}}$	Ref.
Na <sub>3</sub> HfZrSi <sub>2</sub> PO <sub>12</sub>	Solid-state reaction	$4.4 \times 10^{-4} \ \mathrm{S \ cm^{-1}}$ @ RT	0.358 eV	40
$Na_{3.4}Hf_{0.6}Sc_{0.4}ZrSi_2PO_{12}$	Solid-state reaction	$1.2 \times 10^{-3} \text{ S cm}^{-1}$ @ RT	_	40
NZSP	UHS (20 A, 60 s)	$2.62 \times 10^{-4} \text{ S cm}^{-1}$ @ RT	0.28 eV	39
NZSP	Sol-gel method + calcinating (950 °C, 10 h)	$4.43 \times 10^{-3} \text{ S cm}^{-1} \ \text{@ } 50 \ ^{\circ}\text{C}$	_	201
Al-doped NZSP	Sol-gel method + calcinating (950 °C, 10 h)	$1.06 \times 10^{-3} \text{ S cm}^{-1}$ (a) 50 °C	_	201
$Na_{3,2}Zr_{1,9}Mg_{0,1}Si_{2}PO_{12}$	Solid-state reaction	$1.16 \times 10^{-3} \text{ S cm}^{-1}$ @ RT	0.25 eV	202
Ca-doped NZSP	Sol–gel method + heat treatment (600 °C, 6h and 950 °C, 12 h)	$1.67 \times 10^{-3} \text{ S cm}^{-1}$ (a) RT	0.29 eV	156
NZSP-ATO	Calcinating (1100 °C, 9 h, air) + sintering (1100 °C, 6 h)	$1.43 \times 10^{-3} \text{ S cm}^{-1}$	_	60
NLZSP-NBO <sub>3</sub>	Solid-state reaction (950 °C, 6 h)	$1.81 \times 10^{-3} \text{ S cm}^{-1}$ (a) RT	0.22 eV	41
Mg-NZSP-0.128	Solid-state reaction (1250 °C, 5 h, air)	$3.2 \times 10^{-3} \text{ S cm}^{-1}$ (a) RT	0.247 eV	155
Cu-doped NZSP	Solid-state reaction (1150 °C, 12 h)	$5.57 \times 10^{-4} \text{ S cm}^{-1}$ (a) RT	_	125
r	, ,	$1.42 \times 10^{-3} \text{ S cm}^{-1}$ (a) 60 °C		
$Na_3Zr_2Si_2PO_{12}$	SPS (1050 °C, 20 min) + annealing (1100 °C, 10 h)	$8.2 \times 10^{-4} \text{ S cm}^{-1}$ @ 75 °C	_	90
Na <sub>5</sub> GaS <sub>4</sub>	Gas passing	$4.0 \times 10^{-7} \text{ S cm}^{-1}$ @ RT	0.28 eV	203
$Na_{2.9}Sb_{0.9}W_{0.1}S_4$	Solid-state reaction (823 K, 20 h)	$41 \times 10^{-3} \text{ S cm}^{-1}$	0.19 eV	79
Na <sub>3</sub> SbS <sub>4</sub>	Solid-state reaction (823 K, 20 h)	$1.3 \times 10^{-5}  \mathrm{S \ cm^{-1}}$	0.25 eV	79
Ca-doped c-Na <sub>3</sub> PS <sub>4</sub>	Mechanochemical milling (500 rpm, 3–5 h) + heat treatment (700 °C, 12 h)	$0.94 \times 10^{-3} \text{ S cm}^{-1}$ (a) 25 °C	0.49 eV	101
$NaB_3H_8 \cdot xNH_3$ @ $NaB_3H_8$	Mechanical milling	$0.84 \times 10^{-3} \text{ S cm}^{-1}$ @ RT $20.64 \times 10^{-3} \text{ S cm}^{-1}$ @ 318 K	1.46 eV @ RT 0.18 eV @ 318 K	50
$Na_2B_{20}H_{18}$	Wet chemical method	$2.5 \times 10^{-6} \text{ S cm}^{-1}$ @ 25 °C	_	51
Na <sub>2</sub> B <sub>10</sub> H <sub>10</sub>	Thermolysis (185 °C) + wet chemical method	$1.7 \times 10^{-7} \text{ S cm}^{-1}$ @ 25 °C	_	51
$Na_2B_{20}H_{18}-4Na_2B_{12}H_{12}$	Ball-milling (400 rpm, 3 h, N <sub>2</sub> atmosphere)	2.8 × 10 <sup>-4</sup> S cm <sup>-1</sup> (a) 25 °C >10 <sup>-3</sup> S cm <sup>-1</sup> (a) 50 °C	_	51
$Na_4B_{20}H_{18}$ – $3Na_2B_{12}H_{12}$	Ball-milling (400 rpm, 3 h, N <sub>2</sub> atmosphere)	$2.2 \times 10^{-4} \text{ S cm}^{-1}$	0.449 eV	6
Na <sub>4</sub> B <sub>20</sub> H <sub>18</sub>	Wet chemical method	$3.8 \times 10^{-6}  \mathrm{S \ cm^{-1}}$	_	6
$Na_3B_{24}H_{23}-5Na_2B_{12}H_{12}$	Ball-milling (400 rpm, 3 h, N <sub>2</sub> atmosphere)	$1.42 \times 10^{-3} \text{ S cm}^{-1}$ (a) 25 °C	0.385 eV	70
Na <sub>3</sub> B <sub>24</sub> H <sub>23</sub>	Ion-exchange method	$2.3 \times 10^{-5} \text{ S cm}^{-1}$ @ 25 °C	0.59 eV	70
NaBH <sub>4</sub> @Na <sub>2</sub> B <sub>12</sub> H <sub>12</sub>	Solid-gas reaction (10 MPa of H <sub>2</sub> , 150 °C, 4 h)	$4 \times 10^{-6} \text{ S cm}^{-1}$ (a) RT	0.57 eV	95
$Na_4B_{36}H_{34}-7Na_2B_{12}H_{12}$	Ball-milling (400 rpm, 3 h, Ar atmosphere)	$1.02 \times 10^{-3} \text{ S cm}^{-1}$	_	69
Na <sub>4</sub> B <sub>36</sub> H <sub>34</sub>	Ion exchange method	<u> </u>	0.55 eV	69
NaNbCl <sub>6</sub>	Ball-milling (500 rpm) + heat treatment (200 °C, 24 h)	$3.1 \times 10^{-6} \text{ S cm}^{-1}$ (a) 30 °C	0.48 eV	43
$Na_{1.5}Nb_{0.5}Zr_{0.5}Cl_6$	Ball-milling (500 rpm) + heat treatment (200 °C, 24 h)	$2.3 \times 10^{-5} \text{ S cm}^{-1}$ (a) 30 °C	$\sim$ 0.40 eV	43
NaTaCl <sub>6</sub>	Ball-milling (500 rpm) + heat treatment (200 °C, 24 h)	$5.2 \times 10^{-5} \text{ S cm}^{-1}$ (a) 30 °C	0.39 eV	43
Na <sub>1.5</sub> Ta <sub>0.5</sub> Zr <sub>0.5</sub> Cl <sub>6</sub>	Ball-milling (500 rpm) + heat treatment (200 °C, 24 h)	$\sim$ 7 × 10 <sup>-5</sup> S cm <sup>-1</sup> (a) 30 °C	$\sim$ 0.40 eV	43
Na <sub>3</sub> YCl <sub>6</sub>	Ball-milling (500 rpm) + heat treatment (200 °C, 24 h)	$5.7 \times 10^{-8} \text{ S cm}^{-1}$ (a) 30 °C	_	43

 Table 3
 Recently reported Li-based ISEs with synthetic method and electrochemical properties

Solid electrolyte	Synthesis method	Ionic conductivity	$E_{ m a}$	Ref.
		6 4 - 1		;
Yb-doped $L_{16.5}La_3Zr_{1.5}Ta_{0.5}O_{12}$	Solid-state reaction and ceramic pressure-less sintering	$7.67 \times 10^{-2} \text{ s cm}^{-1}$ (a) 25 °C	1	98
$\mathrm{Li_7La_3Zr_2O_{12}}$	Solid-state, cold pressing, 1100 °C, 24 h	$9.14 \times 10^{-7} \mathrm{~S~cm}^{-1}$	0.431 eV	144
$\mathrm{Li}_{\mathrm{E},\mathrm{g}}\mathrm{La}_{3}\mathrm{Zr}_{1,8}\mathrm{Mo}_{0,2}\mathrm{O}_{1,2}$	Solid-state, cold pressing, 1100 °C, 24 h	$3.17 \times 10^{-4}  \mathrm{S  cm^{-1}}$	0.26 eV	144
Lie er La or Srong Zr o Mon O1		$6.43 \times 10^{-4}  \mathrm{S  cm^{-1}}$	0.25 eV	144
Lie ar Ale ar La Zr.O.	Solid-state reaction (950 °C. 12 h + 1200 °C. 12 h)	$3.72 \times 10^{-4}  \mathrm{S  cm^{-1}}$	0.31 eV	139
Lie or Gao or Las Zr.O.	Solid-state reaction (950 °C. 12 h + 1200 °C. 12 h)	$13.0 \times 10^{-4}  \mathrm{S  cm^{-1}}$	0.26 eV	139
LiFPLa-Zr-O	Solid-state reaction (950 °C 12 h + 1200 °C 12 h)	$11.2 \times 10^{-4} \mathrm{s} \mathrm{cm}^{-1}$	V 9 CC 0	139
Li I 3 7r T3 O	Solid-state reaction (950 °C, 12 II   1200 °C, 12 II)	$_{2.28}^{+1.2} \times _{10}^{+1}$ 5 28 $_{2.00}^{+1.4}$ 5 cm $^{-1}$	0.22.CV	130
1: 1 7: 31 0.5 012	30.11:3 at the manufact (950.00 to 12.11 + 1200 to 12.11)	2.30 × 10 3 CIII	0.37 50	139
$L_{16.5}L_{43}Zr_{1.5}Nb_{0.5}Q_{12}$	Solid-state reaction (950 °C, 12 h + 1200 °C, 12 h)	$1.91 \times 10^{-3}$ S cm $^{-1}$	0.44 eV	139
${ m Li_{6.5}La_3Zr_{1.5}Sb_{0.5}O_{12}}$	Solid-state reaction (950 °C, 12 h + 1200 °C, 12 h)	$3.41 \times 10^{-4}  \mathrm{S \ cm^{-1}}$	0.41 eV	139
${ m Li_{6.5}La_3Zr_{1.75}W_{0.25}O_{12}}$	Solid-state reaction (950 $^{\circ}$ C, 12 h + 1200 $^{\circ}$ C, 12 h)	$5.43  imes 10^{-4}  ext{ S cm}^{-1}$	0.37 eV	139
${ m Li_{6.5}La_3Zr_{1.75}Mo_{0.25}O_{12}}$	Solid-state reaction (950 $^{\circ}$ C, 12 h + 1200 $^{\circ}$ C, 12 h)	$3.03  imes 10^{-4}  ext{ S cm}^{-1}$	0.36 eV	139
$\text{Li}_7 \text{La}_{2.5} \text{Ce}_{0.5} \text{Zr}_{1.625} \text{Bi}_{0.3} \text{O}_{12}$	Sol-gel method and heat treatment (1150 $^{\circ}$ C, 6 h)	$5.12  imes 10^{-4}  ext{ S cm}^{-1}  ext{ (a) RT}$	0.1083 eV	145
${ m Li_7La_3Zr_{1.625}Bi_{0.3}O_{12}}$	Sol-gel method and heat treatment (1150 °C, 6 h)	$8.04  imes 10^{-5}  \mathrm{S \ cm}^{-1} \ \mathrm{(B) \ RT}$	0.1750 eV	145
Li <sub>2</sub> La <sub>3</sub> Zr <sub>1</sub> z <sub>5</sub> Ce <sub>0</sub> z <sub>5</sub> O <sub>1</sub>	Solid-state reaction	$2.2  imes 10^{-4}  \mathrm{S \ cm}^{-1} \ (\mathrm{a})  \mathrm{RT}$	0.30 eV	34
Lie 55Ga0 15La3Zr2O12	Solid-state reaction	$0.13  imes 10^{-3}  ext{ S cm}^{-1}$ (a) RT	I	146
Lie es Gan 15 Las Zr. 105 Mgn 105 O12	Solid-state reaction	$1.13 \times 10^{-3}  \mathrm{S  cm^{-1}} \stackrel{\odot}{=} \mathrm{RT}$	I	146
CuO-doped	Solid-state reaction (1120 °C. 6 h)	$1.88 \times 10^{-4}~{ m S~cm}^{-1}$ (a) 25 °C	0.34 eV	143
$\mathrm{Li}_{6.3}\mathrm{La_3Zr_{1.65}W_{0.35}O_{12}}$		)		
$\text{Li. }_{2}\text{Alo.}_{2}\text{Ti. }_{7}\text{(PO_{4})},$	Calcinating (800 °C, 2 h) + CSP (250 MPa. 250 °C, 1 h) + annealing (900 °C, 1 h)	$4.29 \times 10^{-4} \mathrm{Scm}^{-1}$ (a) RT	I	83
LATP	HPI.T (200 °C. 2 min. 1 GPa)	$6.15 \times 10^{-5}  \mathrm{S  cm^{-1}}$	I	68
Te-doned I AGD	$Sol_{-\infty}$ interestintered (500 of A h) + coloinated (850 of 5 h)	$6.33 \times 10^{-4}  \mathrm{G  cm}^{-1}$	VA 00 0	6 6
I ACD	So $[Set]$ presument (500 °C, 4 m) - cardinated (650 °C, 5 m)	1.04 × 10-4 C 2m-1	0.25 CV	7 5
LAGE	501-get pre-sintered (500 °C, 4 II) + carcinated (650 °C, 5 II)	1.94 × 10 S CIII	0.33 eV	91
L <sub>17</sub> P <sub>3</sub> S <sub>11</sub> glass-ceramic	Mechanical milling (270 °C, 3 h)	× 10 ×	0.22 eV	48
GeO <sub>2</sub> -doped L <sub>7</sub> P <sub>3</sub> S <sub>11</sub>	Ball-milling (510 rpm, 40 h @ RT) + annealing (240–290 °C, 4 h)	$4.77  imes 10^{-3}  ext{ S cm}^{-1}  ext{ (@) RT}$	$18.66 \text{ kJ mol}^{-1}$	65
$(\mathrm{Li_7P_{2.9}Ge_{0.05}S_{10.75}O_{0.1}})$		ı	•	
$P_2O_5$ -doped $75Li_2S \cdot 25P_2S_5$		$6.3  imes 10^{-3}   ext{S cm}^{-1}   ext{(@ RT)}$	$20 \text{ kJ mol}^{-1}$	177
CaS-doped $\mathrm{Li}_7\mathrm{P}_3\mathrm{S}_{11}$	Liquid-phase synthesis (50 °C, 72 h + 80 °C, 6 h + 270 °C, 1 h)	$0.95  imes 10^{-3}~{ m S~cm}^{-1}~{ m (@~30~^{\circ}C)}$	$34 \text{ kJ mol}^{-1}$	171
$({ m Li}_{6.72}{ m Ca}_{0.14}{ m P}_3{ m S}_{11})$				
$0.98 \text{Li}_7 \text{P}_3 \text{S}_{11} - 0.02 \text{CaI}_2$		$0.78  imes 10^{-3}~{ m S~cm}^{-1}~{ m (@~29~^{\circ}C}$	$36 \text{ kJ mol}^{-1}$	171
$95\text{Li}_7\text{P}_3\text{S}_{11}-5\text{CaI}_2$	Liquid-phase synthesis (50 °C, 24 h + 270 °C, 2 h)	$1  imes 10^{-3}  ext{ S cm}^{-1}  ext{ (a) RT}$	$30 \text{ kJ mol}^{-1}$	181
		$10.1  imes 10^{-3}~\mathrm{S~cm}^{-1}$ ® 70 °C		
$90 \mathrm{Li_7P_3S_{11}}$ - $5 \mathrm{CaI_2}$ - $5 \mathrm{LiI}$	Liquid-phase synthesis (50 °C, 72 h + 270 °C, 1 h)	$3.1 \times 10^{-3}  \mathrm{S \ cm}^{-1}  (\mathrm{\hat{a}})  \mathrm{RT}$ $13 \times 10^{-3}  \mathrm{S \ cm}^{-1}  (\mathrm{\hat{a}})  70   \mathrm{°C}$	$25 \text{ kJ mol}^{-1}$	181
(1-x)Li <sub>3</sub> PS <sub>4</sub> -2xLiBH <sub>4</sub> $(x=0.54)$	Two-step milling (213 rpm, 45 h and 510 rpm, 15 h)	$11  imes 10^{-3}  ext{ S cm}^{-1}  imes  ext{RT}$	$34.4 \text{ kJ mol}^{-1}$	71
$\mathrm{Li_4PS_4I}$	Solid-state reaction (500 °C, 24 h)	$2.2 \times 10^{-5}  \mathrm{S \ cm}^{-1} \ (\mathrm{a}) \ \mathrm{RT}$	0.45 eV	204
$\mathrm{Li}_{4/3}\mathrm{P}_{0/7}\mathrm{Si}_{0/3}\mathrm{S}_{4}$	Solid-state reaction (700 °C, 5 h)	$1.46 \times 10^{-3}  \mathrm{S \ cm}^{-1}  (\mathrm{a})  \mathrm{RT}$	0.32 eV	205
Li.PS.I	Solid-state reaction (700 °C. 5 h)	$4.6 \times 10^{-5}  \mathrm{S  cm^{-1}}   \mathrm{@  RT}$	0.44 eV	205
1-20, [1]	Mechanochemical process (510 rnm 45 h) and heat treatment (550 °C 2 h Ar)	$3.8 \times 10^{-7}  \mathrm{S  cm}^{-1}$ 9 RT	$43 \text{ kI mol}^{-1}$	47
LingePoS.	Solid-state reaction (680 °C. 8 h)	$6.2 \times 10^{-3} \text{ S cm}^{-1}$	ioni fi	49
ShCldoned LGPS	Solid-state reaction (680 °C 8 h)	$6.8 \times 10^{-3} \text{ S cm}^{-1} \otimes 25 \text{ oC}$	0.196 eV	49
Li-GaS.	Gas nascing	1.8 × 10 <sup>-7</sup> S cm <sup>-1</sup> ® RT	0.54 PV	203
Liscus4	Gus pussing Two-stan hall milling (100 mm for 3 h ± 550 mm for 19 h) and annealing	$1.35 \times 10^{-4}  \mathrm{c}  \mathrm{cm}^{-1}  \mathrm{@ DT}$	0.275.07	180
$\mathbf{L}_{14,025}\mathbf{D}_{10.975}\mathbf{D}_{10.025}\mathbf{D}_{4}$	1800-802 Dan mining (180 tpm 101 Z n + 350 tpm 101 18 n) and anneaning (300 °C; 1.5 h Ns)	1.55 × 10 S CIII @ KI	0.2/3 eV	180
$\text{Li}_{10}$ $^{\circ}\text{P}_{2}$ $^{\circ}\text{S}\text{Sb}_{0}$ $^{\circ}\text{S}_{1}$ $^{\circ}\text{I}$	High-energy ball milling (220 °C, 2 h, Ar)	$5.9 \times 10^{-3}  \mathrm{S \ cm^{-1}} \ (3) \ 25  ^{\circ}\mathrm{C}$	0.28–0.29 eV	111
	3	)		

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Solid electrolyte	Synthesis method	Ionic conductivity	$E_{ m a}$	Ref.
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$\mathrm{Li_{10}P_{3}S_{12}I}$	High-energy ball milling (220 $^{\circ}$ C, 2 h, Ar)	$5.06 \times 10^{-3} \ \mathrm{S \ cm^{-1}} \ \mathrm{@} \ 25 \ \mathrm{^{\circ}C}$	0.28-0.29 eV	111
$\text{Li}_{3.68}\text{Ge}_{0.69}\text{Sb}_{0.31}\text{S}_4$	Solid-state reaction (773 K, 8 h)	$1.23 \times 10^{-4}  \mathrm{S  cm^{-1}} \ \mathrm{(@\ 303\ K)}$	$\sim\!0.28~\mathrm{eV}$	25
Li <sub>1</sub> GeS <sub>2</sub>	Solid-state reaction (973 K)	$3.98 \times 10^{-6}  \mathrm{S  cm}^{-1}  (3)  303  \mathrm{K}$		25
Li.snP.S.,	High-energy hall-milling (550 °C. 4 h)	$4.79 \times 10^{-3} \text{ S cm}^{-1}$ (a) RT	0.15 eV	99
$\frac{1}{10000000000000000000000000000000000$	Solid-state reaction (823 K. 8 h. Ar)	$5.7 \times 10^{-3} \text{ S cm}^{-1}$ (a) RT		80
1 GPS	Tiquid-phase method (mix solvents stirring time 30 min) + heat treatment	1.6 × 10 <sup>-3</sup> S cm <sup>-1</sup> @ P.T.	I	206
	f550 °C 6 h)			0
	(550 C) 0.11	20 0 1 10 -3 c1 0 pm	,	ć
LGPS	High-energy ball milling 520 min + neat treatment	$3.27 \times 10^{-5} \text{ cm}^{-3} \text{ (a) KI}$	0.28 eV	S
${ m Li}_{10}{ m P}_{2.8}{ m S}_{10}{ m O}_2$	Liquid-phase method (2-propanol solvent) + annealing (300 °C, 2 h)	$1.6 \times 10^{-4}~{ m S~cm}^{-1}$ (a) $25~{ m ^{\circ}C}$	$20 \text{ kJ mol}^{-1}$	21
LSiPSCl	Liquid-phase method (ACN solvent) + wet-milling (300 rpm, 20 h) + sintering	$6.6  imes 10^{-3}  ext{ S cm}^{-1}  imes  ext{RT}$	I	7
	(748 K, 8 h, Ar)			
LSiPSCI	Ball-milling (380 rpm, 40 h) + sintering (748 K, 8h, Ar)	$8.8  imes 10^{-3}~{ m S~cm}^{-1}~{ m (@~RT)}$	I	7
Lie 75P0 25Si0 75O5Cl	Spark plasma sintering	$1.82 \times 10^{-6}  \mathrm{S  cm}^{-1}  (3)  303  \mathrm{K}$	0.522 eV	88
LizznosSiSe	Solid-state reaction (973 K. 24 h)	$1 \times 10^{-7} \text{ S cm}^{-1}$ (a) RT	0.66 eV (I4)	185
		$4.3 \times 10^{-4} \text{ S cm}^{-1}$ (a) 503 K	$0.34 \text{ eV} (F\overline{4}3m)$	
Li-GeS-Br	Ball-milling (1 h 250 mm + 10 h 450 mm)	$0.63 \times 10^{-3} \text{ S cm}^{-1} \gg 298 \text{ K}$	0.43 eV	207
I : DC Cl	VII milling and next annualing (500 of 4 b)	$3.03 \times 10^{-3} \text{ cm}^{-1}$	0.45 5 7	204
1: p.c. ol	Wet IIIIIIIII and postaniicaniiig (500 °C, 4 II)	2.5 × 10 S CIII	0.243 eV	90
LiePS5CI	Solid-state reaction (550 °C, 6 n)	$3.19 \times 10^{-5}$ s cm		102
$ m Li_6SbS_5I$	Mechanochemical process (510 rpm, 45 h) and heat treatment (400-450 °C,	$2.1 imes10^{-9}~\mathrm{Scm}^{-1}$ (a) $25~\mathrm{^oC}$	$35 \text{ kJ mol}^{-1}$	47
	2 h, Ar atmosphere)			
${ m Li_{6.5[P_{0.25}Si_{0.25}Ge_{0.25}Sb_{0.25}]S_5I}}$	High-energy milling and post-annealing at 500 $^{\circ}\mathrm{C}$	$13.2 \times 10^{-3}  \mathrm{S \ cm^{-1}} \ \mathrm{(@ \ RT)}$	0.19 eV	208
Li <sub>5.6</sub> PS <sub>4.6</sub> I <sub>1.4</sub> glass–ceramic	Ultimate-energy mechanical alloying (1500 rpm, 1 h + 550 °C, 2 h)	$2.04  imes 10^{-3}~{ m S~cm}^{-1}$	0.31 eV	209
$\mathrm{Li_6PS_5F_{0.5}Cl_{0.5}}$	Wet chemical (ethanol solvent, Ar atmosphere, and heating 200 °C, 1 h)	$3.5  imes 10^{-4}  \mathrm{S \ cm^{-1}}$	0.32 eV	16
LiePS-F	Wet chemical (ethanol solvent, Ar atmosphere, and heating 200 °C, 1 h)	$2.4  imes 10^{-4}  \mathrm{S  cm^{-1}}$	0.45 eV	16
Li-DS-F	Wet chemical (ethanol solvent Ar atmosphere and heating 200 °C 1 h)	$3.5 \times 10^{-4}  \mathrm{S  cm^{-1}}$	0 34 eV	60
T; DC E I	Wet chemical (ethanol colvent, Ar atmosphere, and heating 200 C 1 h)	$\frac{1}{15} \times \frac{10^{-4}}{10} \times \frac{10^{-4}}$	0.277 oV	20
L16F35F0.75L0.25	wet chemical (ethanol solvent, Ar aumosphere, and heating 200 °C, 1 II)	1.3 × 10 S CIII	0.377 eV	92
$L_{16}PS_{5}I$	Wet chemical (ethanol solvent, Ar atmosphere, and heating 200 °C, 1 h)	$2.5 \times 10^{-2} \text{ s cm}^{-2}$	0.387 eV	92
${ m Li_2B_{12}H_{12}}{ m -5Li_2B_{10}H_{10}} - { m 6LiBH_4}$	Ball-milling (350 rpm, 30 min) + annealing (280 $^{\circ}$ C, 1 h, 20 MPa of H <sub>2</sub> )	$\sim\!\!1.0 imes10^{-4}~\mathrm{S~cm}^{-1}$ @ RT	0.16 eV	52
${ m Li_2B_{12}H_{12}}$ – ${ m 5Li_2B_{10}H_{10}}$	Ball-milling (350 rpm, 30 min) + annealing (380 $^{\circ}$ C, 1 h)	$6.92  imes 10^{-6}  ext{ S cm}^{-1}  imes 30  ext{ °C}$	0.51 eV	52
${ m Li}_3({ m CB}_{11}{ m H}_{12})_2({ m CB}_9{ m H}_{10})$	Ball-milling (ratio of 2:1, Ar atmosphere)	$1.5  imes 10^{-3}~{ m S~cm}^{-1}~{ m (@~RT)}$	I	59
		$29.4  imes 10^{-3}~{ m S~cm}^{-1}$ (a) $60~{ m ^{\circ}C}$		
Li <sub>3</sub> OCl	Melt-quenching and high-energy ball-milling (350 °C, 6 h)	$3.21  imes 10^{-5}  ext{ S cm}^{-1}$ @ RT	0.52 eV	94
$\mathrm{Li}_{2}\mathrm{OHCl}_{0.9}\mathrm{Br}_{0.1}$	Solid-state reaction (350 °C, 30 min)	$2.52 \times 10^{-6}  \mathrm{S \ cm}^{-1} \ \mathrm{(a) \ RT}$	0.52 eV	31
Li <sub>2</sub> OHCl	Solid-state reaction (350 °C, 30 min)	$1.93 \times 10^{-6}  \mathrm{S \ cm}^{-1} \ \mathrm{(a) \ RT}$	0.83 eV	31
$ ilde{ ext{Li}_3 ext{SI}}$	High-energy ball-milling (400 rpm, 24 h) + heat treatment (150–550 °C, 3 h)	$2.2  imes 10^{-5}  ext{ S cm}^{-1}  ext{ (a) RT}$	$27.6 \text{ kJ mol}^{-1}$	32
$\mathrm{Li}_{\mathrm{i}_{2}-\mathrm{v}}(\mathrm{OH}_{\mathrm{v}})\mathrm{Cl}_{\mathrm{o}}{}_{\mathrm{o}}\mathrm{F}_{\mathrm{o}}{}_{\mathrm{1}}$	Ball-milling (400 rpm, 24 h) + sintering (350 °C, 5 h)	$9.0  imes 10^{-6}  ext{ S cm}^{-1} \stackrel{\odot}{ ext{(a)}} 25 \ ^{\circ} ext{C}$	,	24
Lj.OHBr	Mechanochemical method (700 mm. 48 h)	$1.6 \times 10^{-6} \text{ S cm}^{-1} \ \text{@ 25 °C}$	I	74
11-00	Podio framework magnetron contraring $(D-1-2) \times 10^{-3}$ mhar BF namer —	$1.7 \times 10^{-8} \text{ cm}^{-1} \odot 5.0$	0 93 AV	, L
Lison	reducing inequency inagine on spancing $(r-1.2\times 10^{-1100})$ , at power = 22 W. Ar atmosphere)	1.27 × 10 3 cm (d/23 - C	0.93 6 6	CC
Li,OHCl	Wet mechanical milling (hexane solvent, 400 rpm, 10 h)	$4.27 \times 10^{-6}  \mathrm{S  cm^{-1}}$ (a) 25 °C	1.03 eV	210
LioHCl	Solid-state reaction (360 °C. 2 h)	$2.37 \times 10^{-7} \text{ S cm}^{-1}$ (a) 25 °C	0.52 eV	210
(LioH), a.K. a.Cl	Solid-state reaction (400 °C: 1 h. Ar atmosphere)	$4.5 \times 10^{-6} \text{ S cm}^{-1} \otimes 25 \text{ °C}$	0.56 eV	72.
1 10	Woter remaind route (100,000)	16 × 10 <sup>-3</sup> C cm <sup>-1</sup> © 25 C		1 5
1: H2 C	Water removal route (100-200 °C)	1.0 × 10 3 cm $^{-1}$ @ 298 N 1.2 × 10 $^{-3}$ S cm $^{-1}$ @ p.T.	I	7 7 2
L12.73H01.09Cl6	Solid-state reaction (550 $^{\circ}$ C, 4 II + anneaning 10f 24 II)	1.3 $\times$ 10 S cm (a) K1	;	30
SmCl <sub>3</sub> ·0.5LiCl	_	$1.2  imes 10^{-4}  ext{ S cm}^{-1}  ext{ (a) } 30  ext{ °C}$	<0.4 eV	0 0
SmCl <sub>3</sub> ·0.5LiCl	Co-melting (650 $^{\circ}$ C, 10 h + 50 $^{\circ}$ C, 1 h)	$3.1 \times 10^{-5} \text{ cm}^{-1}$ (a) 30 °C	I	7

Table 3 (Contd.)

Solid electrolyte	Synthesis method	Ionic conductivity	$E_{ m a}$	Ref.
HCF-Sm	Ball-milling	$1.1 \times 10^{-4} \mathrm{S}\mathrm{cm}^{-1}$	ı	30
HCF-La	Ball-milling	$5.5  imes 10^{-5}  \mathrm{S  cm^{-1}}$	I	30
LCF-Ta	Ball-milling	$5.0  imes 10^{-5}  \mathrm{S  cm^{-1}}$	0.387 eV	30
LCF-Zr	Ball-milling	$\sim \! \! 1.0  imes 10^{-5}  \mathrm{S \ cm^{-1}}$	I	30
LCF-Hf	Ball-milling	$\sim \! \! 1.0  imes 10^{-5}  \mathrm{S \ cm^{-1}}$	I	30
$0.62[\mathrm{HCF\text{-}Sm}] \cdot 0.38[\mathrm{LCF\text{-}Ta}]$	Ball-milling	$2.7  imes 10^{-3}  \mathrm{S \ cm^{-1}} \ \mathrm{(B)} \ 298 \ \mathrm{K}$	0.305 eV	30
$0.62[\mathrm{HCF\text{-}Sm}] \cdot 0.38[\mathrm{LCF\text{-}Hf}]$	Ball-milling	${\sim}2.0\times10^{-4}~\mathrm{S~cm^{-1}}$	I	30
$0.62[\mathrm{HCF\text{-}Sm}] \cdot 0.38[\mathrm{LCF\text{-}Zr}]$	Ball-milling	$\sim \! \! 2.0  imes 10^{-4}  \mathrm{S \ cm^{-1}}$	I	30
$0.57[\mathrm{HCF}\text{-La}] \cdot 0.43[\mathrm{LCF}\text{-Ta}]$	Ball-milling	$1.8  imes 10^{-3}  \mathrm{S \ cm^{-1}} \ \mathrm{(B) \ 298 \ K}$	0.309 eV	30
0.57[HCF-La]·0.43[LCF-Hf]	Ball-milling	$\sim \! 1.0  imes 10^{-4}  \mathrm{S \ cm^{-1}}$	1	30
0.57[HCF-La]·0.43[LCF-Zr]	Ball-milling	$\sim \! 1.0  imes 10^{-4}  \mathrm{S \ cm^{-1}}$	1	30
Li <sub>3</sub> ScCl <sub>6</sub>	Ball-milling (480 rpm, 12 h) + heat treatment (450 °C, 12 h)	$0.6  imes 10^{-3}  \mathrm{S \ cm^{-1}}$	0.367 eV	211
$\mathrm{Li}_{2.6}\mathrm{Sc}_{0.6}\mathrm{Zr}_{0.4}\mathrm{Cl}_{6}$	Ball-milling (480 rpm, 12 h) + heat treatment (450 $^{\circ}$ C, 12 h)	$1.61  imes 10^{-3}  ext{ S cm}^{-1}$	0.326 eV	211
$\mathrm{Li_{2.6}Sc_{0.6}Hf_{0.4}Cl_{6}}$	Ball-milling (480 rpm, 12 h) + heat treatment (450 °C, 12 h)	$1.33 \times 10^{-3}  \mathrm{S \ cm^{-1}}$	0.323 eV	211
$\mathrm{Li}_{2.7}\mathrm{In}_{0.7}\mathrm{Zr}_{0.3}\mathrm{Cl}_{6}$	Two-step annealing (450 °C, 24 h + 450 °C, 8 h)	$2.02  imes 10^{-3}  ext{ S cm}^{-1}  ext{ (a) RT}$	$\sim \!\! 0.28 \; \mathrm{eV}$	93
$\mathrm{Li}_{2.4}\mathrm{Zr}_{0.6}\mathrm{Ho}_{0.4}\mathrm{Cl}_{6}$	Solid-state reaction (450 °C, 48 h)	$\sim \! 1.8  imes 10^{-3}  \mathrm{S \ cm^{-1}}$	$\sim\!\!0.34~{ m eV}$	75
$\mathrm{Li}_{2.4}\mathrm{Zr}_{0.6}\mathrm{Lu}_{0.4}\mathrm{Cl}_{6}$	Solid-state reaction (450 °C, 48 h)	$\sim \! 1.8  imes 10^{-3} \mathrm{~S~cm}^{-1}$	$\sim\!\!0.34~\mathrm{eV}$	75
$Li_3HoCl_6$	Solid-state reaction (650 °C, 48 h)	${\sim}0.1\times10^{-3}~\mathrm{S~cm^{-1}}$	I	75
${ m Li_3LuCl_6}$	Solid-state reaction (650 °C, 48 h)	$\sim \!\! 0.1  imes 10^{-3}  \mathrm{S \ cm^{-1}}$	I	75
$\mathrm{Li}_3\mathrm{ZrCl}_4\mathrm{O}_{1.5}$	Ball-milling $(100 \text{ rpm, } 2 \text{ h} + 500 \text{ rpm, } 10 \text{ h})$	$1.35  imes 10^{-3}~{ m S~cm}^{-1}~{ m (@~25~oC)}$	0.294 eV	57
$\text{Li}_{2.375}\text{Sc}_{0.375}\text{Zr}$ $_{0.625}\text{Cl}_{6}$	Ball-milling (400 h, 10 h) + annealing (300 °C, 5 h)	$2.2  imes 10^{-3}  ext{ S cm}^{-1}  ilde{ ext{ (a)}} 25  ext{ °C}$	0.31 eV	22
${ m Li}_{2.375}{ m Sc}_{0.375}{ m Hf}_{0.625}{ m Cl}_{6}$	Ball-milling (400 h, 10 h) + annealing (300 °C, 5 h)	$1.1  imes 10^{-3}  ext{ S cm}^{-1}  ilde{ ext{ (a)}} 25  ext{ °C}$	0.33 eV	22
Li <sub>3</sub> ScCl <sub>6</sub>	Ball-milling (400 h, 10 h) + annealing (300 °C, 5 h)	$\sim 4  imes 10^{-4}  ext{ S cm}^{-1}  ilde{ ext{ (a)}}  ext{ 25 °C}$	0.44 eV	22
${ m Li_{2.1}Zr_{0.83}Ta_{0.27}Cl_6}$	Ball-milling (550 rpm, 48 h)	$1.68  imes 10^{-3}~{ m S~cm}^{-1}$ (a) 25 $^{ m oC}$	0.28 eV	46
$\mathrm{Li}_{2.3}\mathrm{Hf}_{0.7}\mathrm{In}_{0.3}\mathrm{Cl}_{6}$	Ball-milling	$1.05  imes 10^{-3}~{ m S~cm}^{-1}$ (a) 30 °C	0.337 eV	198
$\mathrm{Li}_{2.4}\mathrm{Y}_{0.4}\mathrm{Hf}_{0.6}\mathrm{Cl}_{6}$	Ball-milling	$1.49  imes 10^{-3}~{ m S~cm}^{-1}$ @ RT	0.331 eV	200
Li <sub>3</sub> YCl <sub>6</sub>	Ball-milling	$1.39  imes 10^{-4}  ext{ S cm}^{-1}$ (a) RT	1	200
${ m Li}_2{ m ZrCl}_6$	Ball-milling (500 rpm, 45 h)	$8.08 \times 10^{-4}  \mathrm{S \ cm^{-1}} \ \mathrm{at} \ 25  \mathrm{^{o}C}$	0.35 eV	212
${ m Li}_{2.25}{ m Hf}_{0.75}{ m Fe}_{0.25}{ m Cl}_{6}$	Ball-milling	$0.91  imes 10^{-3}  ext{ S cm}^{-1}  ilde{ ext{ (a)}} 30  ext{ °C}$	0.34 eV	199
$\mathrm{Li}_{2.2}\mathrm{Hf}_{0.8}\mathrm{V}_{0.2}\mathrm{Cl}_{6}$	Ball-milling		0.351 eV	199
$\mathrm{Li}_{2.2}\mathrm{Hf}_{0.8}\mathrm{Cr}_{0.2}\mathrm{Cl}_{6}$	Ball-milling	$0.81  imes 10^{-3}~{ m S~cm}^{-1}$ @ $30~{ m ^{\circ}C}$	0.342 eV	199
$ ext{Li}_2 ext{HfCl}_6$	Ball-milling	$0.39  imes 10^{-3}~{ m S~cm}^{-1}$ (a) $30~{ m ^{\circ}C}$	1	199

# ISEs preparation methods

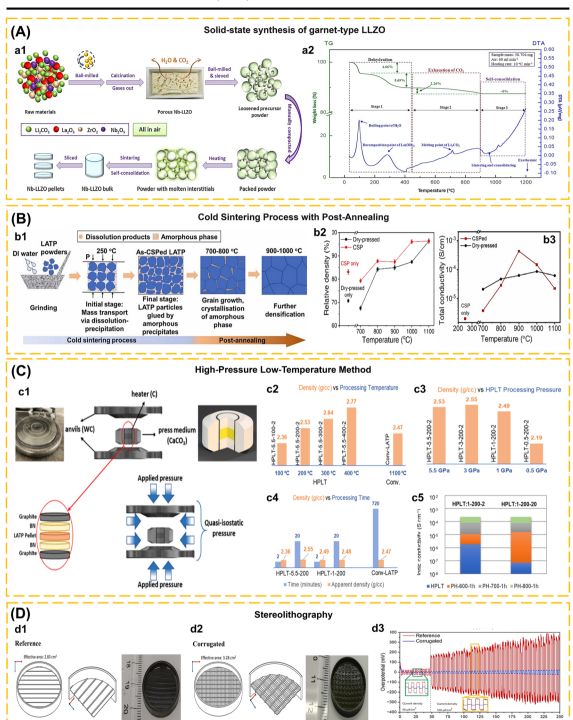


Fig. 13 (A) (a1) Schematic illustration for the preparation of Nb-LLZO pellets by self-consolidation method. (a2) TG/DTA curves of the precursor 060Nb-LLZO powder before sintering. Reproduced with permission from ref. 136. Copyright 2018, Elsevier. (B) (b1) Schematic illustration of the microstructure evolution in LATP ceramics during CSP and subsequent post-annealing. (b2) Effect of annealing temperatures on the relative density of LATP samples produced by CSP and dry-pressing. All samples were annealed for 1 h. (b3) Total ionic conductivity (measured at room temperature) of the CSPed and dry-pressed LATP samples as a function of annealing temperature. Reproduced with permission from ref. 83. Copyright 2023, Elsevier. (C) (c1) The schematic representation of components for the HPLT technique. Density of LATP pellets densified by HPLT processing and conventional sintering. From top to down: (c2) LATP density vs. HPLT processing temperature for pellets processed at 5.5 GPa and 2 minutes compared to conventional sintering, (c3) LATP density vs. HPLT processing pressure for pellets processed at 200 °C and for 2 min, (c4) LATP density vs. HPLT processing time for pellets processed at 200 °C and under high (5.5 GPa) and low (1 GPa) processing pressures compared to conventional sintering conditions. (c5) Evolution of Li-ion conductivity with increase in PH temperature: HPLT-1-200-2 (b) pellet:

of  $H^+$ , as illustrated by the reaction:  $In^{3+} + 3H_2O \leftrightarrow In(OH)_3 + 3H^+$ . The stability of halides in air can be enhanced by applying a coating of aluminum oxide  $(Al_2O_3)$  to their surface. The water absorption rate of  $Li_3InCl_6@Al_2O_3$  is 25% that of  $Li_3InCl_6$ . The stability duration of  $Li_3InCl_6@Al_2O_3$  is sevenfold superior to that of  $Li_3InCl_6$ . Overall, Tables 2 and 3 summarize several examples of Li/Na-based ISEs and their characteristics.

# 3. ISEs preparations and interface of ISEs/electrode

#### 3.1. ISEs preparations

The method of preparation impacts the mechanical and electrochemical characteristics of solid electrolytes, which consequently impacts the commercialization of these electrolytes in the assembly of all-solid-state batteries. <sup>7,64,82,90,102,126,210,212-215</sup> The following overview outlines various preparation methods that have been presented to date, including solid-state synthesis, liquid-phase synthesis, and other advanced techniques. Solid-state reactions can be categorized into three distinct methods: mechanochemical milling, mechanical milling combined with heat treatment, and direct solid-state sintering. <sup>216</sup>

In oxide electrolytes, attention must be given to preparation conditions to achieve high density, ensure appropriate stoichiometry, and prevent the formation of impurity phases.39,40,64,89,126,133 In general, these electrolytes necessitate high sintering temperatures and extended holding time required to achieve high-density oxides. Conversely, high-temperature conditions can lead to the evaporation of Li2O, and prolonged exposure to severe conditions may result in the decomposition of the oxide. 119 Lithium deficiency resulting from Li<sub>2</sub>O evaporation inhibits oxide electrolyte formation and promotes the development of secondary oxide phases. High-temperature sintering can create microstructural defects that block ion transport.83,89 Therefore, understanding the preparation conditions is invaluable for ISE development. Utilizing considerably temperatures and durations may manufacturing yields while minimizing energy consumption, thus promoting a more environmentally sustainable approach.

Garnet-type Nb-doped LLZO is generally synthesized via solid-state reaction.  $^{119,136,217,218}$  Li $_{7-x}$ La $_3$ Zr $_{2-x}$ Nb $_x$ O $_{12}$  (Nb-LLZO, x=0.25) was synthesized by Ohta and coworkers using a conventional solid-state reaction method that included ball-milling, calcination (950 °C, 12 hours), pressing, and sintering (1200 °C, 36 hours).  $^{218}$  All LLZO and Nb-LLZO-type garnets were synthesized in a cubic structure, achieving ionic conductivities of  $\sim$ 0.8 mS cm $^{-1}$  for 0.25Nb-LLZO and  $\sim$ 0.2 mS cm $^{-1}$  for LLZO at 25 °C, with relative densities between 89% and 92%. Zhao  $et~al.^{136}$  synthesized 0.25Nb-LLZO via a self-consolidation method (Fig. 13a1), including ball-milling (400 rpm, 6 h),

calcination (1150 °C, 10 h), and ball-milling (500 rpm, 6 h). The Nb-LLZO powder was manually compacted using vibration and sintered at 1150 °C for 15 hours. The material properties attained include relative density, crystal phase, and ionic conductivity for LLZO (93%, tetragonal phase, 0.0564 mS cm<sup>-1</sup> at 30 °C) and 0.25Nb-LLZO (91%, cubic phase, 0.0807 mS cm<sup>-1</sup> at 30 °C). The maximum relative density and ionic conductivity were attained by 0.6Nb-LLZO, 94% and 0.522 mS cm<sup>-1</sup> at 30 °C. Gai et al.119 synthesized 0.25Nb-LLZO by solid-state reaction, which included ball-milling (8 h), pressing, calcination (900 °C, 8 h), further ball-milling (8 h), and sintering (1230 °C, 15 h). The material properties obtained from this approach include relative density, crystal phase, and ionic conductivity for LLZO (84.1%, tetragonal phase, 0.00734 mS cm<sup>-1</sup> at 30 °C) and 0.25Nb-LLZO (90%, cubic phase, 0.482 mS cm $^{-1}$  at 30 °C). The solid-state reaction exhibits a consistent sequence with modifying temperature, sintering duration, and pellet densification by the pressing process.

The TGA/DTA analysis for the Nb-LLZO sample (Fig. 13a2)136 highlights the thermal gravimetry mechanism into three distinct stages. The first stage involves the dehydration of H<sub>2</sub>O, shown by an endothermic peak at 97 °C, and the decomposition of La(OH)3, seen at an endothermic peak of 280 °C. The second stage is the decomposition process of Li<sub>2</sub>CO<sub>3</sub>, characterized by an endothermic peak at 714 °C. The third stage is the sintering process. There is no weight loss within the temperature range of 880 to 1020 °C. This region has an exothermic peak. While weight loss is not evident in the third stage, heat treatment at temperatures up to 1200 °C results in volume shrinkage, yielding dense sintered bulk. Zhao et al. 136 explain that significant shrinkage occurs during the sintering process, thereby necessitating careful adjustment of the heating rate. The heating rate from room temperature to 800 °C was set at 10 °C min<sup>-1</sup> to eliminate remaining gas components, while the rate from 800 °C to 1150 °C was set at 1 °C min<sup>-1</sup> to mitigate significant volume shrinkage. Subsequently, hold the sintering temperature to get a dense sintered bulk. LLZO sintered at 1150 °C for 15 hours achieved a relative density of 93%, 136 while at 1230 °C for 15 hours, the relative density was 84.5%.119 The 0.25Nb-LLZO exhibits competing relative densities of 91% at 1150 °C (ref. 136) and 90% at 1230 °C.119 The elevated sintering temperature in the LLZO sintering process is likely the cause. The LLZO synthesized by Zhao et al.136 and Gai et al.119 has a tetragonal structure. Elevating the sintering temperature reduces the stability of the tetragonal structure, hence leading to material decomposition and/or a reduction in the volume of LLZO. Doping Nb into LLZO leads to a stable cubic structure. Ohta et al.218 achieved a relative density, 89-92% using a sintering temperature of 1200 °C for 36 hours. Despite using a longer sintering holding time that may lead to lithium

HPLT-1-200-20. Reproduced with permission from ref. 89. Copyright 2024, Elsevier. (D) 3D CAD drawings, details of the cross-sections, and photos of the as-printed membranes: reference (d1) and corrugated (d2). (d3) Galvanostatic cycling of the printed electrolytes in a Li|Ge-LAGP-Ge|Li configuration. Reproduced with permission from ref. 170. Copyright 2023, Royal Society of Chemistry.

### ISEs preparation methods

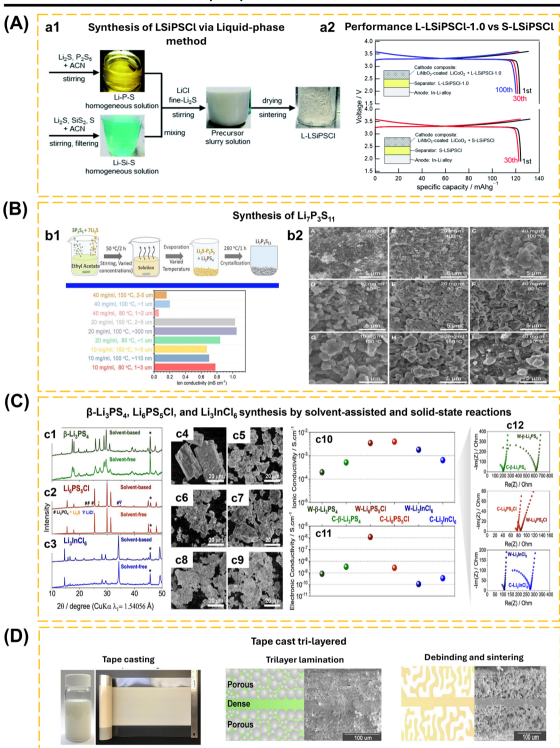


Fig. 14 (A) (a1) Schematic representation of the LSiPSCl synthesized via a liquid-phase process. (a2) Charge-discharge curves of the all-solidstate cells prepared using L-LSiPSCI-1.0 or S-LSiPSCI as the solid electrolyte. The current density applied to the cell was 0.096 mA cm<sup>-2</sup>, which corresponds to a 0.2C rate. Reproduced with permission from ref. 7. Copyright 2022, Royal Society of Chemistry. (B) (b1) A schematic synthesis mechanism of  $Li_7P_3S_{11}$  via ethyl acetate solvent. Ionic conductivity of samples prepared under varying conditions of precursor concentration (10 to 40 mg mL $^{-1}$ ) and evaporation temperature (80 to 150 °C). (b2) The SEM images of Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub> samples synthesized at different concentrations and evaporation temperatures followed by heating at 260 °C: (A) 10-100, (B) 20-100, (C) 40-100, (D) 10-80, (E) 20-80, (F) 40-80, (G) 10-150, (H) 20-50, (I) 40-150. Reproduced with permission from ref. 82. Copyright 2022, Elsevier. (C) Powder X-ray diffraction patterns of (c1) W- $\beta$ -Li<sub>3</sub>PS<sub>4</sub> (dark green),  $C-\beta-Li_3PS_4$  (green), (c2)  $W-Li_6PS_5Cl$  (wine),  $C-Li_6PS_5Cl$  (red), (c3)  $W-Li_3InCl_6$  (navy), and  $C-Li_3InCl_6$  (blue). Asterisks indicate Be

evaporation or product degradation, the density of LLZO remains notably high.

Nonetheless, it is difficult to ascertain whether density correlates to ionic conductivity. In the case of Zhao et al. 136 LLZO and 0.25Nb-LLZO exhibit comparable densities (93% and 91%) and ionic conductivities (0.05 and 0.08 mS cm<sup>-1</sup>). In the case of Gai et al. 119 LLZO and 0.25Nb-LLZO exhibit significantly higher densities (84.5% and 90%) and ionic conductivities (0.00734 and 0.482 mS cm<sup>-1</sup>). SEM analysis from Zhao et al. 136 indicate that the average particle size for LLZO and 0.25Nb-LLZO remains unchanged at around 12 µm, with a maximum particle size of 50 µm. In the case of Gai et al., 119 the reduction in particle size is significant for LLZO (>20 µm) and 0.25Nb-LLZO (5-20 μm). The reduction in grain size due to preparation conditions will impact the increase of grain boundaries and ionic conductivity of LLZO. Although the grain sizes vary but are interconnected, this has an impact on the stability of Li-ion transport in Nb-LLZO. Huge voids are seen in LLZO, affecting the ionic conductivity. Gai et al. 119 clarified that the doping of Y and Nb in LLZO effectively enhanced the ionic conductivity.  $\text{Li}_7\text{La}_3\text{Zr}_{2-2x}\text{Nb}_x\text{Y}_x\text{O}_{12}$  (x=0.5) sintered at 1230 °C for 15 hours exhibited a particle size reduction to below 10 µm, demonstrating excellent inter-grain connectivity compared to 0.25Nb-LLZO. The sintering of 0.5Nb/0.5Y co-doped LLZO at 1230 °C resulted in the formation of a secondary phase, Y2O3, which infiltrated the grain boundaries, enhanced grain connectivity, reduced grain boundaries resistance, and enhanced Li-ion transport pathways.

Cai et al.83 reported that the combination of the cold sintering process (CSP) technique and annealing is highly successful in removing amorphous phase residues in NASICON. Additionally, this approach allows for the control of LATP growth and the decrease of impurity phases (Fig. 13b1). The proposed CSP presents initial densification facilitated by dissolution precipitation. This aimed to regulate grain growth. The CSP was optimized at 250 °C under 250 MPa for 1 hour. The density of the as-CSPed sample reached 83%, which is slightly lower than that achieved through conventional sintering, but superior to the 70% observed in dry-pressed samples. The as-CSPed sample exhibited a higher density compared to the drypressed sample, with a particle size range of 30 to 180 nm. The as-CSPed LATP samples were post-annealed at 700-1100 °C for 1 h. The as-CSPed sample achieved optimal density at 1000  $^{\circ}$ C (96%), with a particle size ranging from 0.4 to 1.7  $\mu$ m, comparable to the dry-pressed samples, 0.3 to 1.8 µm (Fig. 13b2). Raising the temperature resulted in variable grain size formation, characterized by an inhomogeneous microstructure and the presence of micro defects. The best ionic conductivity of the as-CSPed sample after annealing at 900 °C for 1 h  $(4.29 \times 10^{-4} \, \text{S cm}^{-1})$  which is 213 times higher than that

of the CSP-only samples,  $2.01 \times 10^{-6}$  S cm<sup>-1</sup>. While the drypressed sample achieved the best conductivity at annealing at 1000 °C for 1 h ( $8.51 \times 10^{-5}$  S cm<sup>-1</sup>). While the conductivity obtained remains slightly below that of traditional methods, it measures at  $1.09 \times 10^{-3}$  S cm<sup>-1</sup> after 40 hours of mechanical milling followed by sintering at 900 °C for 6 hours (Fig. 13b3).<sup>214</sup> The applied CSP method offers benefits related to reduced processing temperature and duration, regulation of microstructure, and enhanced energy efficiency.

Valiyaveettil-SobhanRaj and co-workers89 enhanced the density and ionic conductivity of NASICON-type LATP by applying the HPLT processing technique followed by post-heat treatment. The HPLT method involves positioning the pellet at the center of a toroid, which is a container made of CaCO<sub>3</sub>. This setup is sealed within a graphite tube that has been coated on the interior with hexagonal boron nitride (h-BN), effectively preventing any contact between the graphite and the sample. The graphite tube provides effective thermal conductivity. The toroid is situated between two anvils and functions to regulate the pressure (Fig. 13c1). Varying the temperature, pressure, and processing time will affect the microstructure and ionic transport processes. Sample nomenclature was used in the format: HPLT-(pressure)-(temperature)-(time). For example, in the case of the LATP pellet densified by HPLT at 5.5 GPa, 200 °C for 2 min, the nomenclature is HPLT-5.5-200-2. The density of conventional LATP (2.47 g cm<sup>-3</sup>) achieved at 1100 °C for 720 min can be compared with that of 2.77 g cm<sup>-3</sup> (for HPLT-5.5-400-2) and 2.55 g cm<sup>-3</sup> (for HPLT-3-200-2 and HPLT-5.5-200-20) (Fig. 13c2-c4). For HPLT-5.5 GPa-2 min under temperature 100-400 °C shows low conductivity ( $\sim 10^{-8}$  S cm<sup>-1</sup>). Lowering pressure for HPLT-200 °C-2 min increases conductivity 2 orders of magnitude higher ( $\sim 10^{-6} \, \mathrm{S \, cm^{-1}}$ ). The effect of holding time (20 min) for HPLT-5.5 GPa-200 °C and HPLT-1 GPa-200 °C samples show conductivity at  $\sim 10^{-7}$  S cm<sup>-1</sup>. The conductivity of HPLT pellets is improved through post-heat treatment (PH) in the range of 600 to 800 °C and a duration of between 1 minute to 1 hour. Samples HPLT-1 GPa-200 °C-2 min and HPLT-1 GPa-200 °C-20 min exhibited conductivity greater than 10<sup>-4</sup> S cm<sup>-1</sup> after PH treatment at 800 °C, a temperature lower than that used in conventional sintering at 1000 °C (Fig. 13c5). The duration of PH treatment significantly affects conductivity. Samples sintered at 800 °C with HPLT-3 GPa-200 °C for holding times of 2, 10, and 60 minutes exhibited conductivities of  $1.9 \times 10^{-5}$  S cm<sup>-1</sup>,  $5.1 \times 10^{-5}$  S cm<sup>-1</sup>, and  $1.5 \times 10^{-4} \, \mathrm{S \ cm^{-1}}$ , respectively. The study of HPLT-based solid electrolyte synthesis is still relatively new, and further improvisation is needed to increase ionic conductivity, but a significant increase in density is a promising achievement for lowtemperature sintering.

reflections from the sample holder. SEM images of (c4) W- $\beta$ -Li<sub>3</sub>PS<sub>4</sub>, (c5) C- $\beta$ -Li<sub>3</sub>PS<sub>4</sub>, (c6) W-Li<sub>6</sub>PS<sub>5</sub>Cl, (c7) C-Li<sub>6</sub>PS<sub>5</sub>Cl, (c8) W-Li<sub>3</sub>InCl<sub>6</sub>, and (c9) C-Li<sub>3</sub>InCl<sub>6</sub> as synthesized. Comparison of (c10) ionic and (c11) electronic conductivities of studied ISEs and (c12) Nyquist plots from up to down:  $\beta$ -Li<sub>3</sub>PS<sub>4</sub>, Li<sub>6</sub>PS<sub>5</sub>Cl, and Li<sub>3</sub>InCl<sub>6</sub> prepared through solvent-engineered and solvent-free methods. Reproduced with permission from ref. 219. Copyright 2021, American Chemical Society. (D) (d1) Photographs of a tape-casting slurry in a glass vial and after casting/drying on a mylar sheet. (d2) Diagram and SEM cross-sectional image of the trilayer in the green unfired state after lamination of the porous and dense tapes. (d3) Diagram and cross-sectional SEM of the trilayer after debinding and sintering. Reproduced with permission from ref. 220. Copyright 2018, Elsevier.

Sabato et al.170 have researched enhancing the surface reactivity of glass-derived LAGP NASICON-type electrolytes by modifying their surface morphology. They employed a 3D printing process called stereolithography, which was helped by CAD. The LAGP ceramic powder was evenly dispersed in an acrylate-based resin that contained a monofax additive. The ratio of the components was 65 wt% LAGP ceramic powder, 34.35 wt% acrylate-based resin, and 0.65 wt% monofax additive. A temperature of at least 600 °C guarantees a structure that closely resembles the crystal structure of LiGe<sub>2</sub>(PO<sub>4</sub>)<sub>2</sub> (PDF#1-080-1924), with no observable impurities such as AlPO4 or GeO<sub>2</sub>. By utilizing CAD technology, the solid electrolyte is constructed in two forms: reference-planar (Fig. 13d1) and corrugated (Fig. 13d2). The wrinkling effect enhances the active surface area of the membrane by 15%. The EIS results showed that the corrugated structures in symmetric cells, Au|LAGP|Au configuration, can decrease the total area-specific resistance by 15%. Fig. 13d3 shows the cycling test results of the Li|Ge-LAGP-Ge|Li associated corrugated structure, indicating a consistently low overpotential for up to 250 hours.

Solid-state methods for synthesizing sulfide-type solid electrolytes have drawbacks, including significant energy consumption and limited large-scale production capacity. Liquid-phase synthesis offers a more economical method for large-scale production. The solution state divides liquid-phase synthesis into two categories: solution synthesis and suspension synthesis. 91,145,147,171,181,206 Hikima et al. 206 described a simple suspension synthesis method to produce LGPS using liquid phase shaking. LGPS with ionic conductivity of 1.6  $\times$  $10^{-3} \text{ S cm}^{-1}$ , electrical conductivity of 9.8  $\times$   $10^{-9} \text{ S cm}^{-1}$ , and low impurities such as GeS2, Li3PO4, β-Li3PS4, and Li2S were obtained by improving the synthesis conditions utilizing a combination of solvents such as ACN, THF, and ethanol, a shaking temperature of 45 °C, a stirring time of 30 minutes, excess sulfur, and heat-treatment at 550 °C for 6 hours. Ito et al.7 reported the preparation of LGPS-type LSiPSCl using the liquidphase method (Fig. 14a1). Solution 1 contains Li, P, and S and solution 2 contains Li, Si, and S, each prepared separately with ACN solvent. ACN as a solvent is an aprotic polar, which is not easily decomposed by precursors. The two solutions are mixed to produce a Li-Si-P-S solution containing Li<sub>2</sub>S: P<sub>2</sub>S<sub>5</sub>: SiS<sub>2</sub> with a molar ratio of 2.1:1.1:2.0. This ratio is lower than the target molar ratio for LSiPSCl,  $\text{Li}_2\text{S}: \text{P}_2\text{S}_5: \text{SiS}_2 = 6.6: 1.1: 2.0.$  To meet the precursor target, fine-Li<sub>2</sub>S was added to the precursor slurry solution. During the drying process of the slurry at 453 K under vacuum conditions, a certain amount of sulfur evaporates, and to compensate for the lost sulfur, excess sulfur is added and adjusted to achieve optimal conditions following the ratio Li<sub>2</sub>S:  $P_2S_5: SiS_2: LiCl: S = 6.6: 1.1: 2.0: 0.4: X (X = 0, 1.0, and 2.9).$ The sample was sintered at 748 K for 8 hours under an Argon atmosphere. LSiPSCl is prepared by a liquid-phase synthetic route and a solid-state reaction is coded L-LSiPSCl and S-LSiPSCl, respectively. L-LSiPSCl-1.0 shows minimal impurity phases, while its ionic conductivity of 6.6 mS cm<sup>-1</sup> at 298 K is higher than L-LSiPSCl-X (X = 0 and 2.9) and slightly lower than S-LSiPSCl of 8.8 mS cm<sup>-1</sup>. The electrochemical performance of ASSB was tested in LCO||Li-In cells using L-LSiPSCl-1.0 and S-

LSiPSCl electrolytes. The electrochemical stability of both electrolytes competes with each other (Fig. 14a2). The particle size factor of solid electrolytes has a significant impact on particle defect reduction, ionic conductivity, and long-term stability. Adjusting particle size can be accomplished by selecting a synthesis method; at this point, liquid-phase synthesis is considered to be capable of controlling particle size. Zhou et al. 82 reported the synthesis of Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub> using ethyl acetate solvent (Fig. 14b1). Adjusting the precursor concentration (10 to 40 mg mL $^{-1}$ ) and solvent evaporation temperature (80 to 150  $^{\circ}$ C) results in a particle size of  $\sim$ 100 nm and a maximum ion conductivity of 1.05 mS cm<sup>-1</sup>. Fig. 14b2 shows the SEM photograph of the Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub> sample produced under optimum conditions. Compared to solvents like ACN, THF, and DME, ethyl acetate has superior Li<sub>2</sub>S and P<sub>2</sub>S<sub>5</sub> dissolving systems. The selection of solvent is an important aspect in liquid-phase synthesis to prevent Li<sub>2</sub>S-P<sub>2</sub>S<sub>5</sub> insolubility, side reactions, crystalline-amorphous phase mixing, and non-uniform particle

Koc et al.<sup>219</sup> compared the properties of β-Li<sub>3</sub>PS<sub>4</sub>, Li<sub>6</sub>PS<sub>5</sub>Cl, and Li3InCl6 were prepared using solvent-assisted and solidstate reactions, indicated by the prefixes W- and C-, respectively. Both methods indicated that the XRD patterns of all samples exhibited pure phases. Solvent-assisted Li<sub>6</sub>PS<sub>5</sub>Cl exhibited impurities, including Li<sub>3</sub>PO<sub>4</sub>, Li<sub>2</sub>S, and LiCl (Fig. 14c1-c3). The ball-milling method facilitated the formation of nano-sized ISE particles and resulted in the formation of microstructural defects. For Li<sub>3</sub>InCl<sub>6</sub>, no morphological differences were observed between the two synthesis routes (Fig. 14c4-c9). The ionic and electronic conductivities at room temperature indicated distinct characteristics of the ISEs (Fig. 14c10-c12), with C-β-Li<sub>3</sub>PS<sub>4</sub> exhibiting higher ionic conductivity compared to W-β-Li<sub>3</sub>PS<sub>4</sub>. Similar to prior research by Krauskopf et al.,176 the defect concentration influences the ionic conductivity of Na<sub>3</sub>PS<sub>4</sub>. The ionic conductivity of Li<sub>6</sub>PS<sub>5</sub>Cl remains consistent across both synthesis routes. The solventassisted synthesis of Li<sub>3</sub>InCl<sub>6</sub> demonstrates superior ionic conductivity.

Balijapelly et al.203 successfully synthesized ternary alkali ion thiogallates, A<sub>5</sub>GaS<sub>4</sub> (A = Li and Na), utilizing a gas-passing synthesis route. The synthesis of Li<sub>5</sub>GaS<sub>4</sub> using a combination of Li<sub>2</sub>S, Ga, and S is hindered by the formation of a secondary phase, LiGaS<sub>2</sub>. Li<sub>5</sub>GaS<sub>5</sub> was produced through the combination of Li<sub>2</sub>CO<sub>3</sub> and Ga<sub>2</sub>O<sub>3</sub> in a precisely controlled tube furnace. A carrier gas was used to flow CS2 gas into a furnace filled with oxides for sulfidation at high temperatures.

The electrolytes  $Na_3B_{24}H_{23}$  and  $Na_4B_{36}H_{34}$  were synthesized through the ion-exchange method. 69,70,225 The anion [B<sub>24</sub>H<sub>23</sub>]<sup>3-</sup> was derived from the oxidation of  $[B_{12}H_{12}]^{2-}$ . The precursor Na2B12H12 underwent treatment with a strong acid ion exchange column and was subsequently neutralized using Me<sub>4</sub>NCl, resulting in the formation of [Me<sub>4</sub>N]<sub>3</sub>B<sub>24</sub>H<sub>23</sub>. The product underwent processing via an H<sup>+</sup> ion exchange column and was subsequently neutralized using NaOH to yield  $Na_3B_{24}H_{23}$ . The hybrid  $Na_3B_{24}H_{23}$ . $xNa_2B_{12}H_{12}$  was synthesized by the mechanical ball-milling of Na<sub>3</sub>B<sub>24</sub>H<sub>23</sub> and Na<sub>2</sub>B<sub>12</sub>H<sub>12</sub>.

Luo et al.95 developed a novel solid electrolyte with a coreshell structure using a gas-solid reaction method by controlling the concentration of B2H6 to form a Na2B12H12 layer on the surface of the NaBH<sub>4</sub> particles. The NaBH<sub>4</sub>/B<sub>2</sub>H<sub>6</sub> ratio of 2:5 results in NaBH<sub>4</sub>@Na<sub>2</sub>B<sub>12</sub>H<sub>12</sub> with an ionic conductivity of 10<sup>-4</sup> S cm<sup>-1</sup> at 115 °C, which makes it highly suitable for ASSBs. Solid-state <sup>11</sup>B NMR shows peaks of -42 ppm attributed to  $[BH_4]^-$  in NaBH<sub>4</sub>, -21 and -15 ppm attributed to  $[B_9H_9]^{2-}$  and  $[B_{12}H_{12}]^{2-}$ . Formation of  $[B_9H_9]^{2-}$  consequence of partial decomposition of NaBH<sub>4</sub>. FTIR spectroscopic investigations revealed that the presence of OH vibrations in the 3600, 3200, and 1614 cm<sup>-1</sup> regions was associated with the hygroscopic nature of Na<sub>2</sub>B<sub>12</sub>H<sub>12</sub> and NaBH<sub>4</sub>. The XPS results of NaBH<sub>4</sub>@-Na<sub>2</sub>B<sub>12</sub>H<sub>12</sub> show two peaks in the B 1s spectrum region that shift to a higher energy level, indicated as an increase in boron in NaBH<sub>4</sub>@Na<sub>2</sub>B<sub>12</sub>H<sub>12</sub>.

Advanced fabrication techniques for LLZO membranes have also gained significant attention. For instance, the sintering of tape-cast trilayered LLZO frameworks represents a promising strategy to produce membranes with desirable properties.<sup>220</sup> Another innovative method involves sintering a freeze-cast, vertically aligned LLZO scaffold, which offers potential improvements in the microstructural control of the electrolyte (Fig. 14d).<sup>226,227</sup> These sintering techniques, while promising, face considerable challenges in scaling up, particularly in achieving high manufacturing yields of thin, high-performance LLZO-based electrolytes. Consequently, the widespread commercial adoption of these materials may be delayed for years, if not decades, until these production hurdles are

overcome. Looking forward, research on LLZO must prioritize several key areas to accelerate the practical application of LLZO-based solid electrolytes. First, engineering the cathode–LLZO interface is critical to improving overall battery performance and longevity. Second, efforts should be directed toward scaling up the production of LLZO-based sheet electrolytes, ensuring that manufacturing processes are both cost-effective and capable of producing consistent, high-quality SEs. Finally, addressing the persistent issue of Li dendrite growth is essential to enhance the safety and reliability of solid-state batteries.

#### 3.2. Interface of ISEs/electrode

The compatibility of interfaces between ISEs and electrodes is a critical factor in achieving high performance in ASSBs, alongside ionic conductivity and electrochemical stability. <sup>120</sup> This section summarizes the behavior of ISEs at the anode and cathode interfaces, as well as the approach to controlling these interfaces through the implementation of artificial SEI and CEI layers, referred to as anode and cathode interface layers in this review. This may decrease interfacial resistance, mitigate ISEs decomposition, and enhance battery performance.

**3.2.1. Interface of ISEs/anode.** At the ISE-Anode interface, three distinct behaviors may manifest when both components are in contact: (1) a thermodynamically stable ISEs/anode interface. (2) A spontaneous reaction between ISEs and anode forms an interphase compound that is a good ionic conductor and a poor electronic conductor, called a solid electrolyte interphase (SEI). (3) A spontaneous reaction between ISEs and

Table 4 Phase equilibria for lithiation and delithiation (sodiation and desodiation) reactions of Li/Na-based ISEs with metal anode based on Li/Na grand potential phase diagrams

ISEs	Anode	Phase equilibria at low voltage	Phase equilibria at high voltage	Ref.
Li <sub>3</sub> PS <sub>4</sub>	Li	P, Li <sub>2</sub> S @ 1.72 V	$P_2S_5$ , S @ $\sim$ 2.4 V	230
		$\text{Li}_3\text{P}$ , $\text{Li}_2\text{S}$ @ 0.5 V		
Li <sub>4</sub> GeS <sub>4</sub>	Li	Ge, Li <sub>2</sub> S @ 1.62 V	$\text{Li}_2\text{GeS}_3$ , S @ $\sim$ 2.4 V	230
		$\text{Li}_{15}\text{Ge}_4$ , $\text{Li}_2\text{S}$ @ 0 V	$GeS_2$ , S @ > 2.5 V	
Li <sub>6</sub> PS <sub>5</sub> Cl	Li	P, Li <sub>2</sub> S, Li <sub>3</sub> P, LiCl @ 1.71	S, P <sub>2</sub> S <sub>5</sub> , LiCl @ 2.01	172 and 231
$\text{Li}_{7}\text{P}_{3}\text{S}_{11}$	Li	Li <sub>3</sub> PS <sub>4</sub> , P <sub>4</sub> S <sub>9</sub> @ 2.28 V	S, P <sub>2</sub> S <sub>5</sub> @ 2.31	231
$\text{Li}_7\text{P}_2\text{S}_8\text{I}$	Li	P, Li <sub>2</sub> S, Li <sub>3</sub> P, LiI @ 1.71	S, P <sub>2</sub> S <sub>5</sub> , LiI @ 2.31	231
LGPS	Li	Li <sub>4</sub> GeS <sub>4</sub> , P, Li <sub>2</sub> S @ 1.71 V	Li <sub>3</sub> PS <sub>4</sub> , S, GeS <sub>2</sub> @ 2.14 V	120
		Li <sub>2</sub> S, Li <sub>15</sub> Ge <sub>4</sub> , Li <sub>3</sub> P @ 0 V	$P_2S_5$ , S, $GeS_2$ @ 2.31 V	
LLZO	Li	Li <sub>2</sub> O, Zr <sub>3</sub> O, La <sub>2</sub> O <sub>3</sub> @ 0.05 V	Li <sub>2</sub> O <sub>2</sub> , Li <sub>6</sub> Zr <sub>2</sub> O <sub>7</sub> , La <sub>2</sub> O <sub>3</sub> @ 2.91 V	120
		Li <sub>2</sub> O, Zr, La <sub>2</sub> O <sub>3</sub> @ 0.004 V	O <sub>2</sub> , Li <sub>6</sub> Zr <sub>2</sub> O <sub>7</sub> , La <sub>2</sub> O <sub>3</sub> @ 3.3 V	
Ta-doped LLZO	Li	Li <sub>2</sub> O, Zr <sub>3</sub> O, La <sub>2</sub> O <sub>3</sub> , Ta @ 0.05 V	Li <sub>2</sub> O <sub>2</sub> , Li <sub>6</sub> Zr <sub>2</sub> O <sub>7</sub> , Li <sub>5</sub> TaO <sub>5</sub> , La <sub>2</sub> O <sub>3</sub> @ 2.91 V	120
		Li <sub>2</sub> O, La <sub>2</sub> O <sub>3</sub> , Zr, Ta @ 0.004 V	$O_2$ , $La_3TaO_7$ , $La_2Zr_2O_7$ , $La_2O_3$ @ 3.3 V	
Al-doped LLZO	Li	Li <sub>2</sub> O, Zr <sub>3</sub> O, La <sub>2</sub> O <sub>3</sub> , Zr <sub>3</sub> Al @ 0.05 V	Li <sub>2</sub> O <sub>2</sub> , LiAlO <sub>2</sub> , Li <sub>6</sub> Zr <sub>2</sub> O <sub>7</sub> , La <sub>2</sub> O <sub>3</sub> @ 3.04 V	120
		Li <sub>2</sub> O, La <sub>2</sub> O <sub>3</sub> , Zr, Zr <sub>3</sub> Al@0.004 V	$O_2$ , $La_4Al_2O_9$ , $La_2Zr_2O_7$ , $La_2O_3$ @ 3.3 V	
LLTO	Li	$\text{Li}_4\text{Ti}_5\text{O}_{12}$ , $\text{Li}_{7/6}\text{Ti}_{11/6}\text{O}_4$ , $\text{La}_2\text{Ti}_2\text{O}_7$ @ 1.75 V	$O_2$ , $TiO_2$ , $La_2Ti_2O_7$ @ 3.71 V	231
LATP	Li	P, LiTiPO <sub>5</sub> , AlPO <sub>4</sub> , Li <sub>3</sub> PO <sub>4</sub> @ 2.17 V	O <sub>2</sub> , LiTi <sub>2</sub> (PO <sub>4</sub> ) <sub>3</sub> , Li <sub>4</sub> P <sub>2</sub> O <sub>7</sub> , AlPO <sub>4</sub> @ 4.21 V	231
LAGP	Li	Ge, GeO <sub>2</sub> , Li <sub>4</sub> P <sub>2</sub> O <sub>7</sub> , AlPO <sub>4</sub> @ 2.70 V	$O_2$ , $Ge_5O(PO_4)_6$ , $Li_4P_2O_7$ , $AlPO_4$ @ 4.27 V	231
Li <sub>3</sub> ScCl <sub>6</sub>	Li	Sc <sub>5</sub> Cl <sub>8</sub> , LiCl @ <0.91 V	ScCl <sub>3</sub> , Cl <sub>2</sub> @ >4.26 V	232
		Sc, LiCl @ 0.V		
Na <sub>3</sub> SbS <sub>4</sub>	Na	Sb, Na <sub>2</sub> S @ 0.66 V	Sb <sub>2</sub> S <sub>3</sub> , Na <sub>2</sub> S @ 2.19 V	104 and 233
		Na <sub>3</sub> Sb, Na <sub>2</sub> S @ 0 V	Sb <sub>2</sub> S <sub>3</sub> , S (a) >3.4 V	
Na <sub>3</sub> PS <sub>4</sub>	Na	Na <sub>2</sub> PS <sub>3</sub> , Na <sub>2</sub> S @ <1.55 V	Na <sub>2</sub> PS <sub>3</sub> , Na <sub>2</sub> S @ >2.25 V	234
		Na <sub>3</sub> P, Na <sub>2</sub> S @ 0 V	P <sub>2</sub> S <sub>7</sub> , S @ 3.0 V	
Na <sub>3</sub> PSe <sub>4</sub>	Na	Na <sub>2</sub> PSe <sub>3</sub> , Na <sub>2</sub> Se @ <1.80 V	Na <sub>2</sub> PSe <sub>3</sub> , Se @ >2.15 V	234
		Na <sub>3</sub> P, Na <sub>2</sub> Se @ 0 V	PSe, Se @ 3.0 V	

### ISEs/anode interface

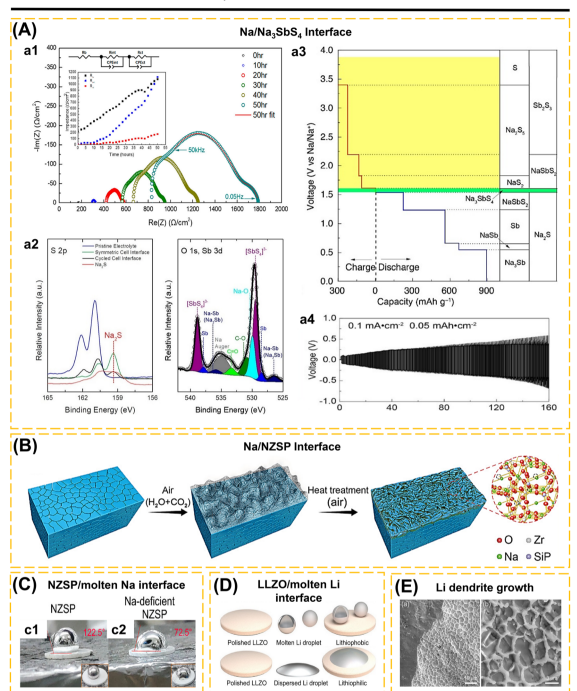


Fig. 15 (A) (a1) Nyquist plot of the Na|Na<sub>3</sub>SbS<sub>4</sub>|Na symmetric cell from 0 to 50 h, showing the increasing impedance with time. Inset: equivalent circuit used to fit the data with a corresponding fit plotted on the 50 h curve. Inset: the fitted impedance components with time. (a2) S 2p region scan of the pristine NAS, anode interface of the cell cycled to completion, the symmetric cell, and the Na<sub>2</sub>S precursor, overlaid on top of each other. O 1s/Sb 3d region scan of the Na|NAS SSEI from the cell cycled to completion. Reproduced with permission from ref. 104. Copyright 2018, American Chemical Society. (a3) Calculated equilibrium voltage profile and phase equilibria for sodiation and desodiation reactions of Na<sub>3</sub>SbS<sub>4</sub>. (a4) Galvanostatic voltage profile of the cell cycled at a current density of 0.1 mA cm<sup>-2</sup> for 0.5 h Na deposition and 0.5 h Na stripping during each cycle at 60 °C after 1 h rest at the beginning of cycling. Reproduced with permission from ref. 233. Copyright 2019, American Chemical Society. (B) Moisture leading to the byproduct layer formed on the surface of NZSP and its thermal stability verified by the HT. Schematic illustration of the NZSP surface structure change in moisture (H<sub>2</sub>O + CO<sub>2</sub>) and further HT. First, a byproduct layer was formed on the NZSP surface; secondly, decomposition of the byproduct layer during HT; at last, a Na-deficient surface formed after decomposition of Na compounds contained in the byproduct layer. (C) (c1 and c2) Contact angle measurement of the molten metallic Na on NZSP SSE and Na-deficient NZSP. Reproduced with permission from ref. 229. Copyright 2020, American Chemical Society. (D) Effects of coating methods on Li wetting behavior with LLZO: stacking-coating (upper panel) and rub-coating (lower panel).

anode forms a mixed ion-electron conducting (MIEC) interphase.<sup>228,229</sup> This part will clarify the behavior of the ISE interface with the anode and propose solutions for enhancing the interface by means of surface impurity removal (polishing), the application of molten Na/Li, alloys, and porous surface engineering.

Sulfide exhibits reactivity with Na and Li metals, leading to the spontaneous formation of an SEI layer. The stable formation of the solid electrolyte interphase at the anode/ionic solid electrolyte interface enhances solid-solid contact, minimizes interfacial resistance, and facilitates the migration of lithium ions.120 The phase diagram of lithium grand potential aids in identifying phase equilibria across various potentials and determining the thermodynamically dominant products influenced by these potentials (Table 4). The SEI layers, including LiF, Li<sub>2</sub>O, Li<sub>2</sub>S, Li<sub>3</sub>N, and Li<sub>3</sub>P, exhibit stability in relation to Li metal.235 Although NASICONs are superior for high voltage operation and are stable in air, they have poor chemical and mechanical compatibility with Li anodes, resulting in inhomogeneous Li<sup>+</sup> transfer and reduction of the active material, such as the reduction of Ge4+ to Ge2+ in LAGP, which occurs frequently when in contact with Li metal. 236,237 For garnet-type oxides, LLZO at 2.91 V undergoes reduction into Li<sub>2</sub>O<sub>2</sub>, Li<sub>6</sub>Zr<sub>2</sub>O<sub>7</sub>, and La<sub>2</sub>O<sub>3</sub>. The increase in voltage discovered for O<sub>2</sub> and La<sub>2</sub>Zr<sub>2</sub>O<sub>7</sub> resulted from the decomposition of Li<sub>2</sub>O<sub>2</sub> and Li<sub>6</sub>Zr<sub>2</sub>O<sub>7</sub>, respectively. At a low voltage of 0.05 V, LLZO undergoes reduction to form Li<sub>2</sub>O, Zr<sub>3</sub>O, La<sub>2</sub>O<sub>3</sub>, and Zr<sub>3</sub>O. Zr<sub>3</sub>O undergoes reduction to Zr at potentials below 0.004 V. The electrochemical stability remains relatively unaltered for Aldoped LLZO (Li<sub>6.28</sub>La<sub>3</sub>Zr<sub>2</sub>Al<sub>0.24</sub>O<sub>12</sub>) and Ta-doped LLZO (Li<sub>6.75</sub>-La<sub>3</sub>Zr<sub>1.75</sub>Ta<sub>0.25</sub>O<sub>12</sub>). As voltages near 0 V, Ta-doped LLZO undergoes reduction to Ta metal, while Al-doped LLZO is transformed into Al-Zr alloy. 120 The expectation is that Ta/Aldoped materials can be reduced at slightly high potentials; however, the low concentration of dopants results in insufficient contribution to enhancing stability. Additional investigation is required regarding the Lithium grand potential phase diagram of Ta/Al-doped LLZO as the concentration of dopants increases. The stable SEI does not consistently prevail at the interface, as the interface layer also includes decomposition products that serve as ionic and electronic conductors. During charge-discharge cycles, interface products inevitably form, leading to the accumulation of unstable SEI. This accumulation deteriorates interface contact, elevates interfacial resistance, and promotes dendrite formation due to the presence of electronic conductors at the interface. The resulting dendrites compromise the ISE structure and ultimately reduce battery performance.238

The Na<sub>3</sub>SbS<sub>4</sub>/Na interface has been evaluated by Hu *et al.*<sup>233</sup> and Wu *et al.*<sup>104</sup> The symmetric Na/Na<sub>3</sub>SbS<sub>4</sub>/Na cell's Nyquist plot indicates an increase in both bulk resistance and charge transfer as Na ions are progressively consumed by Na<sub>3</sub>SbS<sub>4</sub> to

form the solid electrolyte interphase (Fig. 15a1).104 The grand potential phase stability plot demonstrates that in the anodic region, Na<sub>3</sub>SbS<sub>4</sub> transforms into Na<sub>3</sub>Sb and Na<sub>2</sub>S, whereas in the cathodic region, Sb<sub>2</sub>S<sub>3</sub> and S are present (Fig. 15a3). 104,233 The XPS S 2p spectra show that all conditions exhibit a peak at 159.4 eV, which is confirmed to be Na<sub>2</sub>S. The XPS Sb spectra exhibit a separation of 9.39 eV in the 3d orbital, specifically observed in the Sb 3d<sub>5/2</sub> (529.5 and 528.5 eV) and Sb 3d<sub>3/2</sub> (538.9 and 537.9 eV) regions (Fig. 15a2). The galvanostatic voltage profiles for 160 cycles at a current density of 0.1 mA cm<sup>-2</sup>, with 0.5 h allocated for Na plating and 0.5 h for Na stripping, indicate a significant increase in the overpotential of the cell (Fig. 15a4).<sup>233</sup> In a related study, Wu et al.<sup>104</sup> discussed how the presence of Cl dopant in Na<sub>3</sub>PS<sub>4</sub> influences the stability of the Na/Na<sub>3</sub>PS<sub>4</sub> interface. The ASR values of Na<sub>3</sub>PS<sub>4</sub>, Cl-doped Na<sub>3</sub>PS<sub>4</sub> (6.25%), and Cl-doped Na<sub>3</sub>PS<sub>4</sub> (12.5%) exhibited increases from 1986 to 3740  $\Omega$  cm<sup>2</sup> (88.3%), 992 to 2139  $\Omega$  cm<sup>2</sup> (115.6%), and 6372 to 7163  $\Omega$  cm<sup>2</sup> (12.5%), respectively. The findings indicate that adjusting the dopant concentration from 6.25% to 12.5% results in enhanced cycling stability and ionic conductivity. Na<sub>2</sub>S and Na<sub>3</sub>P are both solid electrolyte interphases generated from Na<sub>3</sub>PS<sub>4</sub> and Cl-doped Na<sub>3</sub>PS<sub>4</sub>. Furthermore, the NaCl peak is observed as a decomposition product of Cl-doped Na<sub>3</sub>PS<sub>4</sub>. The NaCl passivation layer serves as an electronic insulator, effectively inhibiting electronic percolation across ISEs and interfaces.

Byproduct layers were also found to form periodically as a result of air exposure on the NZSP surface. Gao  $et~al.^{229}$  studied the heat treatment (HT) technique that effectively eliminates surface contaminants, thereby enhancing the wettability of the NZSP surface (Fig. 15B). The contact angle of the as-prepared NZSP was measured at 122.5°, in contrast to the 72.5° observed for HT-NZSP. The reduction in contact angle following the elimination of the impurity layer led to a surface deficient in Na ions, thereby enhancing the contact power with the molten Na (Fig. 15C). HT did not produce any side effects by altering the structure of NZSP and facilitating the formation of the stable SEI layers (Na<sub>2</sub>O) to inhibit the further decomposition of NZSP. The total resistance measured of Na/NZSP/Na and Na/HT-NZSP/Na is about 13 725 and 680  $\Omega$  cm².

A major challenge with LLZO is its propensity to react with atmospheric moisture and CO<sub>2</sub>, resulting in the formation of surface contaminants such as LiOH and Li<sub>2</sub>CO<sub>3</sub>. These contaminants significantly hinder the material's ability to wet molten Li, which is critical for its application as a solid electrolyte in Li-metal batteries. Several mechanisms have been proposed to explain the interactions between LLZO and ambient air.<sup>240</sup> The reactions leading to the formation of carbonate and hydroxide impurities in the presence of CO<sub>2</sub> and H<sub>2</sub>O, respectively, can be described by the following equations:

 $2\text{Li}_7\text{La}_3\text{M}_2\text{O}_{12} + 7\text{H}_2\text{O} \rightleftharpoons 14\text{LiOH} + 3\text{La}_2\text{O}_3 + 4\text{MO}_2$ 

Reproduced with permission from ref. 239. Copyright 2022, Wiley-VCH. (E) Li dendrite growth phenomenon and potential mechanisms. SEM images of intergranular Li dendrite growth along grain boundaries of polycrystalline LLZO. Reproduced with permission from ref. 38. Copyright 2016, Elsevier.

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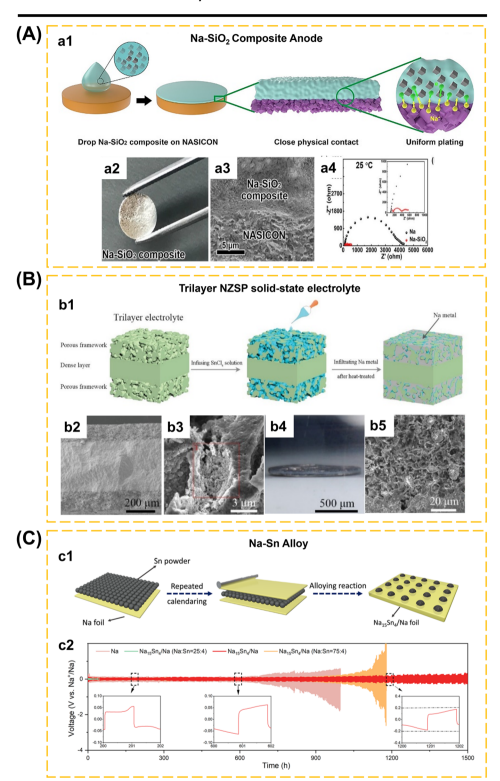


Fig. 16 (A) (a1) the intimate contact between Na-SiO<sub>2</sub> composite and NASICON SE facilitates a close physical contact and a stabilized solidsolid interface. (a2) Digital photo of Na-SiO<sub>2</sub>|NASICON interface. (a3) SEM images of the cross-section of Na-SiO<sub>2</sub>|NASICON interface, which indicates drastically enhanced physical contact at the interface. (a4) EIS spectra of symmetric cells using Na metal and Na-SiO<sub>2</sub> as electrodes. Reproduced with permission from ref. 260. Copyright 2020, American Chemical Society. (B) (b1) Schematic of the trilayer NZSP solid-state electrolyte. (b2) SEM images of the trilayer membrane at low magnification. (b3) SnO<sub>2</sub> modified porous electrolyte surface. (b4) Photos of the SnO<sub>2</sub> modified-trilayer membranes with melting sodium. (b5) SEM images of the Na-impregnated porous layer. Reproduced with permission from ref. 156. Copyright 2019, Wiley-VCH. (C) (c1) Schematic diagram of the preparation process of the  $Na_{15}Sn_4/Na$  composite foil. (c2) Voltage

$$2\text{Li}_{7}\text{La}_{3}\text{M}_{2}\text{O}_{12} + 7\text{CO}_{2} \rightleftharpoons 7\text{Li}_{2}\text{CO}_{3} + 3\text{La}_{2}\text{O}_{3} + 4\text{MO}_{2}$$

where M=Zr, Sn, and  $Hf.^{241}$  These surface impurities are the primary obstacle preventing LLZO from effectively wetting molten Li. To mitigate this issue, various strategies have been proposed, such as the introduction of an interface layer or the modification of the Li anode composition to enhance wettability.242-245 An alternative approach was reported by Zheng et al., 239 who found that a rubbing process can disrupt the impurity layers on the surfaces of both LLZO and molten lithium, thereby significantly enhancing the lithiophilicity of LLZO (Fig. 15D). The effect of sandpaper size (60, 600, and 2000 grits) during garnet polishing greatly affects the surface contact properties with Li metal. Ma et al.246 demonstrated the SEM of LALZOBr<sub>0.15</sub>-600 which showed great contact with lithium metal with interfacial resistance of 20.9  $\Omega$  cm<sup>-2</sup> compared to LALZOBr<sub>0.15</sub>-60 (53.9  $\Omega$  cm<sup>-2</sup>) and LALZOBr<sub>0.15</sub>-2000 (41.2  $\Omega$  $cm^{-2}$ ).

Interfacial instability at the Li/LLZO interface is a critical challenge, primarily due to lithiophobicity and the propensity for Li dendrite formation. 38,120,239,247 The poor wettability between Li and LLZO results in inadequate interfacial contact, leading to an uneven current distribution, which can facilitate the nucleation and growth of Li dendrites (Fig. 15E). 248,249 While substantial experimental and theoretical research suggests that grain boundaries play a pivotal role in Li penetration behavior, it is important to recognize that this is not the sole mechanism by which Li penetration occurs. Evidence of Li penetration has been documented in both amorphous and single-crystalline solid electrolytes, which inherently lack grain boundaries. 247,250,251 Recent studies have also proposed that residual stresses within the solid electrolyte may significantly contribute to Li penetration, particularly through mechanisms analogous to stress-corrosion cracking, where these stresses are amplified by chemical interactions with Li.252 These residual stresses may originate from grain boundaries, impurities, or processinginduced defects, and could potentially be correlated with the observed heterogeneous stresses associated with polymorphism in LLZO.253 Moreover, the role of electronic conductivity in Li penetration within ISEs has garnered considerable attention in recent literature. Several reports suggest that even trace levels of electronic conductivity, whether located in the grain interior or at grain boundaries, may facilitate the nucleation of metallic Li within the solid electrolyte matrix. 254-256 This underscores the complexity of Li penetration phenomena, indicating that multiple, interrelated factors contribute to the interfacial instability at the Li/LLZO interface. Furthermore, previous research has focused on enhancing the compatibility between garnet and lithium by using coating layers such as SnO2,257 LIPON,258 and graphite.259

Fu et al.<sup>260</sup> proposed a solution to the instability of the Na metal and NASICON interface by using Na–SiO<sub>2</sub> as the anode,

effectively decreasing the interfacial resistance from 1658 to 101  $\Omega$  cm<sup>2</sup>. At the Na/NASICON interface, the ability of molten Na to wet the NASICON electrolyte surface is limited. The molten Na-SiO<sub>2</sub> completely covers the NASICON surface (Fig. 16a1-a4). The high surface tension and poor wettability of planar NZSP present challenges for the application of molten Na on the NZSP surface. The adoption of monolithic electrolyte architecture is attributed to its unique characteristics that integrate solid and porous layers (Fig. 16b1). 156,201 NZSP electrolyte is composed of a solid layer (300 μm) sandwiched by a porous layer (100 μm) (Fig. 16b2). The SnO<sub>2</sub> additive is applied to the NZSP surface to improve the sodiophilic properties and facilitate the wetting of the NZSP surface (Fig. 16b3). The integration of monolithic architecture with high sodiophilicity results in an extensive contact area, enhancing the solid-solid interaction of NZSP-Na (Fig. 16b4 and b5). This approach effectively reduces the surface resistance significantly, decreasing from 10 000  $\Omega$  cm<sup>2</sup> to 275  $\Omega$ cm<sup>2</sup>, outperforming planar NZSP. 156

The use of metal alloy reduces the reactivity of ISEs with metal anodes. The Na/Na<sub>12</sub>Sn<sub>4</sub> electrode was fabricated using a mechanical folding and calendaring technique, as detailed by Wang et al.261 An alloy with a thickness of 300 μm was produced by spreading Sn granules on Na foil and subsequently rolling it repeatedly (Fig. 16c1). The electrochemical behavior of Na alloy (Na/Sn = 25:4, 45:4, and 75:4) and pure Na were tested in symmetrical cells Na<sub>15</sub>Sn<sub>4</sub>/Na|NZSPF3-PVDF-HFP|Na<sub>15</sub>Sn<sub>4</sub>/Na and Na|NZSPF3-PVDF-HFP|Na at 50  $\mu$ A cm<sup>-2</sup> (Fig. 16c2). Na/Sn alloy (Na/Sn = 45:4) showed low polarization voltage and cycle stability reaching 1500 hours. This indicates that Na alloy is more stable facilitating Na deposition and suppressing dendrite growth. Na/Sn alloy (Na:Sn = 25:4) experienced faster cell failure due to the lack of Na as a binder. The cell was subsequently damaged by the irregular sodium deposition of the Na/ Sn alloy (Na: Sn = 75:4). The voltage fluctuations observed in the Na anode indicated that the discharging mechanism was unstable. The ratio of Na/Sn = 45:4 as a Na/Sn alloy with low reactivity, good sodiophilic properties, and stable precipitation. After the plating/stripping process, the Na metal anode exhibited morphological irregularities and cracks, as determined by SEM analysis. While the Na/Sn alloy demonstrated stability under the same measurement conditions. The Na/Sn alloy is electrochemically stable and devoid of dendrites due to the fact that the surface remains stable despite the presence of precipitation in certain regions.30,101

**3.2.2. Interface of ISEs/cathode.** The performance of cathodes in constructing ASSBs is evaluated based on their excellent reversible capacity, high-voltage operation, and ease of preparation. Nonetheless, with ongoing charging and discharging, the performance of the cathode material deteriorates owing to structural degradation and interactions at the electrode–electrolyte interface. Park *et al.*<sup>179</sup> investigated the stability of halide and sulfide-based ISEs using different cathodes. Halide–ISE

profiles of symmetric coin cells with  $Na_{15}Sn_4$  alloys/Na with various atomic ratios of Na/Sn = 25 : 4, 45 : 4, and 75 : 4 and bare Na = 15 and the inset shows the enlarged voltage profiles of the symmetric cells with  $Na_{15}Sn_4/Na$  electrode (Na/Sn = 45 : 4) during 200-202, 600-602, and 1200-1202 h, respectively. Reproduced with permission from ref. 261. Copyright 2024, American Chemical Society.

# ISEs/cathode interface

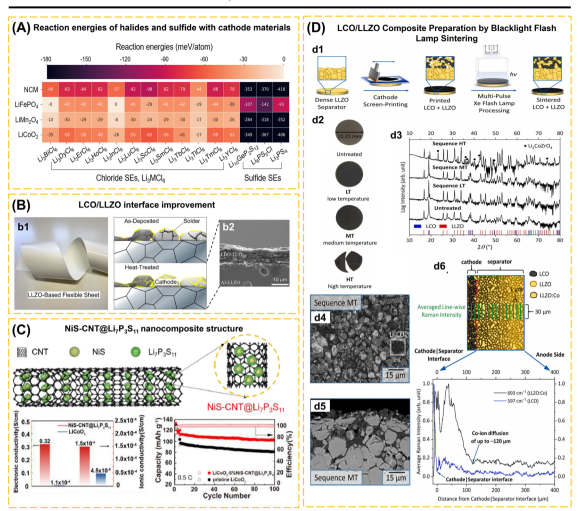


Fig. 17 (A) Heat map of the maximum reaction energy between 12 lithium chloride and three lithium sulfide SEs and four cathode materials (LCO, LMO, LFP, and NMC). Reproduced with permission from ref. 179. Copyright 2020, American Chemical Society. (B) (b1) Strategies to overcome the contact issue between the electrode and LLZO solid electrolyte: LLZO-based flexible sheet electrolyte for improved contact between LLZO and cathode layer (left). Reproduced with permission from ref. 61. Copyright 2020, American Chemical Society. (b2) A solid electrolyte solder-coated cathode layer is deposited onto an LLZO substrate by aerosol deposition; the cracks and voids are filled by the solid electrolyte solder after heating. Reproduced with permission from ref. 73. Copyright 2022, American Chemical Society. (C) NiS-CNT@Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub> nanocomposite structure. Electronic and ionic conductivities of the LCO and NiS-CNT@Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub> at room temperature. Cycling stability of pristine LCO and LCO-5%NiS-CNT@Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub> at current densities of 0.5C. Reproduced with permission from ref. 110 Copyright 2021, Elsevier. (D) (d1 and d2) Schematic representation of LLZO-LCO preparation and images of the cathode surface in the untreated condition and after processing of 3 different pulse sequences. (d3) XRD patterns of the processed and untreated composite cathodes with the reference pattern of LLZO (red, ICSD 422259) and LCO (blue, ICDD 010702685). (d4) Top-view SEM images (BSE) of the flash lamp processed composite cathodes (sequence MT). (d5) Polished cross-sections of the flash lamp processed half-cells (seguence MT). (d6) Scheme of the Raman measurement of the sintered half-cell crosssection processed with sequence MT to detect Co-ion diffusion into the LLZO separator as a function of distance from the cathode|separator interface. The measurement was performed in a range of 30 μm over the whole half-cell in 1 μm steps (x and y) and the resulting signals were averaged line-wise and normalized; (b) Raman spectra with the LLZO: Co photoluminescence signal at 693 cm<sup>-1</sup> (black) and the LCO phase signal at 597 cm $^{-1}$  (blue) are shown as a function of distance from the cathode|separator interface (0  $\mu$ m). Reproduced with permission from ref. 77. Copyright 2023, Elsevier.

exhibits stability to LMO. Halide has greater stability at the cathode compared to sulfide, indicating that decomposition products are more likely to accumulate at the sulfide-cathode interface; nonetheless, these decomposition products provide a passivation layer that inhibits further decomposition of sulfide.179 The decomposition products and the energy levels of the ISE chemical reaction with the cathode are shown in Fig. 17a and Table 5.

The interface between cathode active materials and crystalline LLZO presents significant challenges during the fabrication and operation of solid-state batteries. A primary concern lies in the chemical instability at the cathode/LLZO interface, which

Table 5 Decomposition products of Li/Na-based ISEs with cathode materials

ISEs	Cathode	Decomposition product	Ref.
Li <sub>3</sub> PS <sub>4</sub>	NMC	Co(NiS <sub>2</sub> ) <sub>2</sub> , Li <sub>3</sub> PO <sub>4</sub> , Li(MnS <sub>2</sub> ) <sub>2</sub> , Co <sub>2</sub> NiS <sub>4</sub> , MnS <sub>2</sub> , Li <sub>2</sub> S	179
	LFP	$\text{Li}_4\text{P}_2\text{O}_7$ , FePS, FeS <sub>2</sub>	
	LMO	$Li_3PO_4$ , $Li(MnS_2)_2$	
	LCO	$Co_9S_8$ , $Li_3PO_4$ , $Li_2SO_4$ , $Li_2S$	
Li <sub>6</sub> PS <sub>5</sub> Cl	NMC	Li <sub>3</sub> PO <sub>4</sub> , Co <sub>2</sub> NiS <sub>4</sub> , Li(MnS <sub>2</sub> ) <sub>2</sub> , Li <sub>2</sub> S, Co(NiS <sub>2</sub> ) <sub>2</sub> , MnS <sub>2</sub> , LiCl	179
	LFP	P <sub>4</sub> S <sub>7</sub> , Li <sub>3</sub> PO <sub>4</sub> , FePS, FeS <sub>2</sub> , LiCl	
	LMO	Li <sub>3</sub> PO <sub>4</sub> , Li(MnS <sub>2</sub> ) <sub>2</sub> , Li <sub>2</sub> S, LiCl	
	LCO	Li <sub>3</sub> PO <sub>4</sub> , Li <sub>2</sub> SO <sub>4</sub> , Li <sub>2</sub> S, Co <sub>9</sub> S <sub>8</sub> , LiCl	
$Li_{10}GeP_2S_{12}$	NMC	Co(NiS <sub>2</sub> ) <sub>2</sub> , MnO, Li <sub>2</sub> MnGeO <sub>4</sub> , Li <sub>3</sub> PO <sub>4</sub> , Li <sub>2</sub> SO <sub>4</sub> , Co <sub>2</sub> NiS <sub>4</sub> , Li <sub>2</sub> S	179
	LFP	Li <sub>4</sub> P <sub>2</sub> O <sub>7</sub> , Li <sub>3</sub> PO <sub>4</sub> , FePS, GeS <sub>2</sub> , FeS <sub>2</sub>	
	LMO	Li(MnS <sub>2</sub> ) <sub>2</sub> , Li <sub>3</sub> PO <sub>4</sub> , Li <sub>2</sub> MnGeO <sub>4</sub> , MnS <sub>2</sub> , Li <sub>2</sub> S	
	LCO	Li <sub>4</sub> GeO <sub>4</sub> , Co <sub>9</sub> S <sub>8</sub> , Li <sub>3</sub> PO <sub>4</sub> , Li <sub>2</sub> SO <sub>4</sub> , Li <sub>2</sub> S	
Li <sub>3</sub> YCl <sub>6</sub>	NMC	Li <sub>2</sub> MnCo <sub>3</sub> O <sub>8</sub> , YClO, Li <sub>2</sub> Mn <sub>3</sub> NiO <sub>8</sub> , NiCl <sub>2</sub> , Co <sub>3</sub> O <sub>4</sub> , LiCl	179
	LFP	Fe <sub>2</sub> PClO <sub>4</sub> , YPO <sub>4</sub> , LiCl	
	LMO	$MnO_2$ , $Mn_8Cl_3O_{10}$ , $YMn_2O_5$ , LiCl	
	LCO	YClO, LiClO <sub>4</sub> , Co <sub>3</sub> O <sub>4</sub> , LiCl	
LATP	LCO	Co <sub>3</sub> O <sub>4</sub> , CoAl <sub>2</sub> O <sub>4</sub> , Co <sub>2</sub> TiO <sub>4</sub> , and Li <sub>3</sub> PO <sub>4</sub>	262
LLZO	LCO	$La_2O_3$ , $Li_6Zr_2O_7$ , $Li_5CoO_4$	232
Li <sub>3</sub> ScCl <sub>6</sub>	LCO	LiCl, Co <sub>3</sub> O <sub>4</sub> , LiClO <sub>4</sub> , Sc <sub>2</sub> O <sub>3</sub>	232

critically impacts the overall performance and durability of these batteries. In typical solid-state Li-metal batteries, the cathode is often a composite, consisting of cathode active materials and ISEs. Achieving effective contact between these components frequently necessitates high-temperature cosintering. However, this process can induce thermal decomposition or Li loss from the active materials,263 as well as promote undesirable elemental diffusion at the cathode/solid electrolyte interface, further exacerbating interfacial instability. The challenges associated with the cathode/LLZO interface can be broadly categorized into three key types: mechanical, chemical, and electrochemical instability. These issues are critical as they contribute to increased interfacial resistance, which hinders ion transport and degrades battery performance. To address these challenges, various strategies have been proposed. Cheng et al. have made significant advancements in this area by developing LLZO-based, castable, and flexible sheet electrolytes aimed at improving interfacial contact (Fig. 17b1).61,264 In addition, their work highlights the use of a buffer layer composed of lowmelting-point Li<sub>3</sub>BO<sub>3</sub> to enhance interfacial contact between LCO particles and the LLZO substrate. Upon heating, Li<sub>3</sub>BO<sub>3</sub> melts and forms a Li-ion-conducting liquid phase, which infiltrates cracks and voids, thereby improving the connectivity and stability at the interface (Fig. 17b2).73 Such innovative approaches hold promise for overcoming the interfacial challenges in solid-state Li-metal batteries and should be further explored in future studies.

The integration of oxide-based cathodes in sulfide-based ASSBs presents challenges to their applications: (1) decomposition of sulfide electrolyte at high voltage; (2) interface reactions and detrimental decomposition products; (3) formation of space-charge layer; and (4) structural degradation of oxide cathodes. Transition-metal sulfide cathodes are the preferred cathode type because they provide strong interfacial compatibility with sulfide-type solid electrolytes while also

conductivity providing superior ionic and capacity. 12,48,109,110 The use of a single material as a cathode, such as amorphous niobium polysulfides (a-NbSx, x = 3, 4, 5), has drawbacks for ASSBs applications owing to a-NbSx's weak electronic and ionic conductivity, which affects battery cycle performance. The usage of carbon-based materials may increase the cathode material's electrical conductivity while protecting the cathode with sulfide electrolyte layers improves the solid-solid interface contact issue. Xie and co-workers109 investigated the performance of ASSBs using a-NbS<sub>4.5</sub>/20% Super P@15% Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub> cathodes produced via ball milling and annealing. Bilayer electrolytes consisted of Li<sub>10</sub>GeP<sub>2</sub>S<sub>12</sub> (6.2 ×  $10^{-3} \text{ S cm}^{-1}$ ) and 75% Li<sub>2</sub>S-24% P<sub>2</sub>S<sub>5</sub>-1% P<sub>2</sub>O<sub>5</sub> (1.54 × 10<sup>-3</sup> S cm<sup>-1</sup>). The results demonstrate an initial discharge capacity of 975.7 mA h  $g^{-1}$  at 0.1 A  $g^{-1}$  and a reversible capacity of 464.8 mA h  $g^{-1}$  after 2500 cycles at 0.5 A  $g^{-1}$ . After 40 cycles of EIS measurement, the battery cathode containing a-NbS<sub>4.5</sub>/20% Super P@15%  $\text{Li}_7\text{P}_3\text{S}_{11}$  had lower  $R_e$  and  $R_{ct}$  values than the a-NbS<sub>4.5</sub> and a-NbS<sub>4.5</sub>/20% Super P cathodes. It was concluded that using Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub> layer and Super P in the cathode might increase electrochemical performance as well as the electrolyteelectrode interface. Jiang et al.110 used the solvothermal technique to synthesize the NiS-CNT@Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub> nanocomposite (260 °C, 1 h). LCO-NiS-CNT@Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub> composite cathodes were produced by combining 2.5, 5.0, and 7.5 wt% NiS-CNT@Li<sub>7</sub>P<sub>3</sub>-S<sub>11</sub> with LCO. Another composite cathode consists of LCO-NiS-CNT@Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub> and Li<sub>10</sub>GeP<sub>2</sub>S<sub>12</sub> in a 7:3 ratio. The solid electrolyte utilized is a combination of Li<sub>10</sub>GeP<sub>2</sub>S<sub>12</sub> and 75% Li<sub>2</sub>S-24%  $P_2S_5-1$ %  $P_2O_5$ . In the  $Li_{10}GeP_2S_{12}/NiS-CNT@Li_7P_3S_{11}$ nanocomposite/Li<sub>10</sub>GeP<sub>2</sub>S<sub>12</sub> symmetric cell, NiS-CNT@Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub> achieves a higher ionic conductivity of  $1.5 \times 10^{-4} \, \mathrm{S \ cm^{-1}}$  than LCO, which is  $4.5 \times 10^{-5}$  S cm<sup>-1</sup>. In ASSBs with LCO-5%NiS-CNT@Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub> cathode has a capacity of 118.6 mA h  $g^{-1}$ , which is higher than the pristine LCO cathode's capacity of 115.1 mA h  $g^{-1}$  at 0.1C after 10 cycles. At 0.5C, LCO-5%NiS-

CNT@Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub> achieves a reversible capacity of 99.6 mA h g<sup>-1</sup> after 100 cycles, compared to pure LCO, which has a capacity of 81.9 mA h g<sup>-1</sup> (Fig. 17c).

Scheld et al.77 used a blacklight flash lamp sintering approach using an Xe flash lamp to fabricate an LCO/LLZO composite cathode in 20 seconds, rivaling the speed of RTP and laser sintering methods. The three used sintering modes low temperature (LT), medium temperature (MT), and high temperature (HT)-revealed that the HT mode induced cracks in the composite cathode, led to the decomposition of the LCO phase and resulted in the development of secondary phases such as Li<sub>2</sub>CoZrO<sub>4</sub> (Fig. 17d1-d3). Composite cathodes sintered

at medium-temperature conditions exhibited many linkages between LLZO and LCO particles. The upper cross-section exhibited cracks resulting from thermal shock induced by rapid and strong heating and cooling conditions. Nevertheless, cracks were only observable on the LCO surface, since this region experienced the most substantial heat contrast relative to the lower area (Fig. 17d4 and d5). In LLZO composites, the migration of Co-ions from LCO and LLZO must be prevented. Raman spectroscopy may be used to assess the contamination of LLZO by Co (Fig. 17d6). The contamination level was around 120 µm, lower than RTP sintering. 213 The short sintering duration resulted in reduced Co ion migration into LLZO, or maybe

### Coating process of interface layers

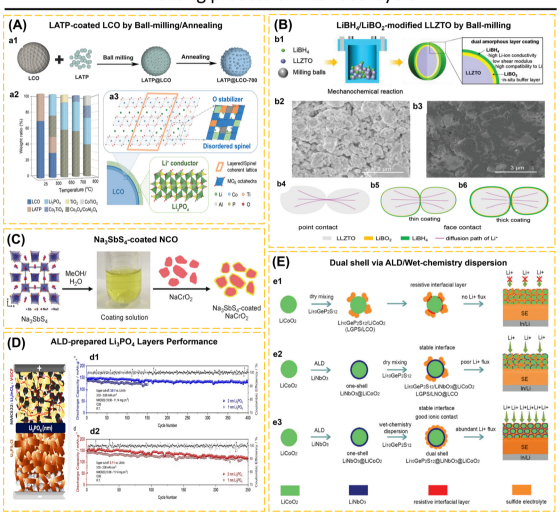


Fig. 18 (A) (a1) Schematic illustrations of the surface engineering of LCO. (a2) Phase constitutions of reaction products of LiCoO<sub>2</sub> and LATP heattreated at different temperatures. (a3) Schematic illustration of the surface layer growth mechanism. The spinel phase is structurally more stable than the layered phase with weaker oxygen-anion oxidizing ability at high voltages. Reproduced with permission from ref. 262. Copyright 2020, Wiley-VCH. (B) (b1) Schematic of the preparation of LiBH<sub>4</sub>-modified LLZTO. SEM images of pellets of (b2) pristine LLZTO and (b3) LLZTO-4LiBH<sub>4</sub>. Schematic illustration of contact and the Li<sup>+</sup> diffusion path between particles in the pellets of (b4) pristine LLZTO sample, with (b5) low and (b6) high LiBH<sub>4</sub> content. Reproduced with permission from ref. 265. Copyright 2021, Wiley-VCH. (C) Schematic illustrations of Na<sub>3</sub>SbS<sub>4</sub>-coated NCO. Reproduced with permission from ref. 271. Copyright 2016, Wiley-VCH. (D) Long-term cycling performance of cells (NMC622: Li₃InCl<sub>6</sub>:  $VGCF|Li_{3}PO_{4}-coated\ Li_{6}PS_{5}Cl\ |\ Li_{6}PS_{5}Cl\ |\ Li_{0.5}In)\ with\ varied\ surface\ coating\ thicknesses\ at\ (d1)\ 3.6\ V\ and\ (d2)\ 3.7\ V\ cutoff\ voltages.\ Reproduced\ description and the produced of the produced o$ with permission from ref. 272. Copyright 2022, American Chemical Society. (E) Schematic diagram of ASSLIBs with various interfacial nanostructures. (e1) LiCoO<sub>2</sub> directly mixed with Li<sub>10</sub>GeP<sub>2</sub>S<sub>12</sub> without interfacial design for ASSLIBs. (e2) A one-shell LiNbO<sub>3</sub>@LiCoO<sub>2</sub> cathode for ASSLIBs. (e3) A dual shell LGPS@LNO@LCO for ASSLIBs. Reproduced with permission from ref. 270. Copyright 2019, Wiley-VCH.

none at all. The findings suggest that this sintering approach is viable for commercializing the manufacturing of composite cathodes by enhancing the solid-solid contact between the cathode and ISE, although further advancements in the production of LLZO composites are necessary to minimize surface cracks.

Koç  $et~al.^{219}$  evaluated the stability of a core-shell-like cathode composite of ISEs-coated NMC622 using ISEs based on lithium thiophosphates (β-Li<sub>3</sub>PS<sub>4</sub>), argyrodite (Li<sub>6</sub>PS<sub>5</sub>Cl), and halide (Li<sub>3</sub>InCl<sub>6</sub>). In cases of ISEs using β-Li<sub>3</sub>PS<sub>4</sub> and Li<sub>6</sub>PS<sub>5</sub>Cl, the battery cells were arranged as follows: NMC622/ISE/ VGCF|ISE|Li<sub>0.5</sub>In/ISE. Specifically, Li<sub>3</sub>InCl<sub>6</sub> was arranged as follows: NMC622/Li<sub>3</sub>InCl<sub>6</sub>/VGCF|Li<sub>6</sub>PS<sub>5</sub>Cl|Li<sub>0.5</sub>In/Li<sub>6</sub>PS<sub>5</sub>Cl. The NCM/Li<sub>6</sub>PS<sub>5</sub>Cl and NCM622/β-Li<sub>3</sub>PS<sub>4</sub> cathode composites exhibited capacity retention rates of 97% and 90%, respectively, but the NCM622/Li<sub>3</sub>InCl<sub>6</sub> cathode composite dropped to 50%. Galvanostatic profile testing of the composite cathode (NCM622/Li<sub>3</sub>InCl<sub>6</sub>) in a three-electrode cell configuration revealed a reduction in electrochemical performance, corroborating findings from two-electrode configuration cell testing, attributed to the highly resistive positive electrode interface and the incompatibility between the Li<sub>3</sub>InCl<sub>6</sub>/Li<sub>6</sub>PS<sub>5</sub>Cl and Li<sub>3</sub>InCl<sub>6</sub>/ Li<sub>3</sub>PS<sub>4</sub> interfaces. The EIS analysis reveals a considerable rise in positive electrode resistance until the 20th cycle.

3.2.3. Coating process of the interface layers. The interface layer is applied to suppress side reactions at the electrode-ISEs interface, enhance poor interface contact, and improve the performance stability of ASSBs. 1,97,107,108,262,265-267 The laver is applied to the interface using an in situ approach (direct synthesis on the electrode surface) or direct coating (pressing method). Both have been implemented in the case of the anode. In the case of the cathode, to obtain complete interface contact with ISEs: (1) a composite cathode is fabricated by mixing the CAM with ISE (catholyte) in a specific ratio;75,101,110 (2) the coating material is directly pressed or coated onto the prepared cathode surface; 66,76,266 (3) the coating is synthesized in situ on the cathode surface;97 (4) CAM is coated with metal oxide, followed by densification or other fabrication methods to produce the electrode;1 (5) coated CAM mixed with ISE (catholyte) to produce a composite cathode. 7,80

The charge-discharge process will often induce volume fluctuations in CAM, resulting in the loss of contact force among CAM particles and leading to the formation of cracks in the composite cathode materials. The inherent characteristics of the ISE surface that are incompatible with certain cathodes, intact with the cathode-ISE interface reactions capable of generating unstable products, worsen the interface contact,268 and ultimately degrade battery performance. CAM protection layers and artificial interface layers will serve to enhance the structural and interfacial stability of the electrolyte-electrode and mitigate unstable side reactions during charging and discharging.262,269 The parameters for CAM coating materials and their interaction processes with cathode and electrolyte materials have been clarified.105 The choice of coating type and processes in the design of ASSBs must be meticulous to reduce performance deterioration.268 If CAM coating is the selected approach, the compression pressure during densification

should not result in the CAM coating breaking and possibly contacting the ISE directly. Alternatively, the composite cathode may be densified first, followed by the compression or synthesis of the interface layer above the composite cathode layer. The variation in construction will influence battery performance. ISE-coated cathodes offer superior advantages over the CAM and ISE mixing methods due to a more uniform partial distribution between CAM and ISE, which minimizes voids between particles, thereby enhancing particle contact and increasing ionic conduction in the cathode and electrodes. <sup>270,271</sup>

Here we will explain the method of preparing coated CAM and coating the electrode-ISE interface. Several articles have described the method for depositing the interface layer including ALD, <sup>106,108,269,270,272</sup> mechanical mixing, <sup>262,265</sup> solution-processed coating, <sup>273</sup> direct pressing process, <sup>274</sup> magnetic sputtering, <sup>1</sup> spin coating, <sup>98,266</sup> *in situ* synthesis. <sup>97</sup> The electrode-electrolyte interface material applied can be a metal oxide such as SnO<sub>2</sub>, <sup>107</sup> ZrO<sub>2</sub>, <sup>108</sup> LiNbO<sub>3</sub>, <sup>273,275</sup> Li<sub>3</sub>PO<sub>4</sub>, <sup>106,272,273</sup> LiTaO<sub>3</sub>, <sup>269</sup> Li<sub>4</sub>Ti<sub>5</sub>O<sub>12</sub>, <sup>273</sup> graphite, <sup>91</sup> Li<sub>3</sub>N-LiF, <sup>97</sup> ZnO-LiF, <sup>1</sup> and ISEs. Here, we will explain the progress of ISE application not only as electrolyte and catholyte but also as an artificial interface layer. Until now, ISE has been applied as an interface layer, for example LATP, <sup>262</sup> Na<sub>3</sub>SbS<sub>4</sub>, <sup>271</sup> Li<sub>7.5</sub>La<sub>3</sub>Zr<sub>1.5</sub>Co<sub>0.5</sub>O<sub>12</sub>, <sup>76</sup> Li<sub>3</sub>OCl, <sup>98</sup> Li<sub>6</sub>PS<sub>5</sub>Cl, <sup>22,42,67,75</sup> and Li<sub>3</sub>YCl<sub>6</sub>. <sup>57</sup>

3.2.3.1. Mechanical mixing technique. Wang et al. <sup>262</sup> applied a simple mechanical mixing technique to coat the surface of the LCO electrode with LATP at high-temperature conditions. The LATP and LCO powders were blended until homogeneous in a mixing machine. The composite samples were calcined at 400–900 °C. The adjustments of ball-milling and calcination conditions influenced the thickness and uniformity of LATP on the LCO surface (Fig. 18a1). The deposition of the LATP layer on the LCO surface leads to the formation of a passivation layer consists of spinel phases (Co<sub>3</sub>O<sub>4</sub>, CoAl<sub>2</sub>O<sub>4</sub>, and Co<sub>2</sub>TiO<sub>4</sub>) and Li<sub>3</sub>PO<sub>4</sub> (Fig. 18a2 and a3), which exhibit stable properties at high voltages, suppress the decomposition reactions, enhance Li-ion transport, and extend the voltage range to 4.6 V. LATP-coated LCO heated at 700 °C has a capacity retention of 88.3% at 25 °C and 72.9% at 45 °C after 100 cycles.

Garnet-type LLZO is often manufactured using hightemperature sintering to optimize ionic conductivity; however, this process leads to inadequate solid-solid contact with the electrode and increased brittleness. Gao et al.265 addressed the issue by doping LiBH4 by the ball-milling technique. The interaction between LLZO and the LiBH4 layer results in the formation of a LiBO2 passivation layer, which stabilizes the LLZO-LiBH<sub>4</sub> interface (Fig. 18b1). The cold pressing method resulted in a reduction in open porosity from 26.4% (LLZO) to 14.4% (LLZO-4LiBH<sub>4</sub>) (Fig. 18b2 and b3). This accomplishment elucidates that the coating outcomes function as fillers (enhancing the density of LLZO), adhesives (improving solidsolid contact and reducing interfacial resistance), and connections (efficient ionic conductors) (Fig. 18b4-b6). Ionic conductivity improved from 4.17 imes 10<sup>-9</sup> S cm<sup>-1</sup> (LLZO) to 8.02 imes10<sup>-5</sup> S cm<sup>-1</sup>. Hydroborate serves as an alternate option for a coating layer to enhance oxide-based ion-selective electrodes (ISEs).

Solution-processed coating. The technique offers a straightforward preparation procedure for coating, although the challenges of this method are the control of coating concentration, thickness, and homogeneity of the coating. Liu et al.273 synthesized solution-based LiNbO3/Li3PO4/Li4Ti5O12coated LiNi<sub>0.5</sub>Mn<sub>1.5</sub>O<sub>4</sub> (LNMO). LNMO and the precursors of Li<sub>3</sub>PO<sub>4</sub> coating (LiOH, NH<sub>4</sub>H<sub>2</sub>PO<sub>4</sub>) were synthesized in aqueous solution and subsequently given heat treatment to produce Li<sub>3</sub>PO<sub>4</sub>-coated LNMO. HRTEM examination verified that the coating layer effectively encapsulated LNMO with a thickness of 10-20 nm. The cathode composite comprising 8 wt% LiNbO<sub>3</sub>coated LNMO and Li<sub>7</sub>PS<sub>5</sub>Cl (70:30) exhibited an initial discharge capacity of 115 mA h  $\mathrm{g}^{-1}$  and a reversible discharge capacity of 80 mA h g<sup>-1</sup> after 20 cycles. Banerjee et al.<sup>271</sup> deposited a Na<sub>3</sub>SbS<sub>4</sub> layer onto the NaCrO<sub>2</sub> surface in a methanol solvent followed by low-temperature treatment (Fig. 18c). Na<sub>3</sub>SbS<sub>4</sub> layers (13 wt%) applied on NaCrO<sub>2</sub> ensured optimal ionic contact and strong adherence. High-Resolution Transmission Electron Microscopy (HRTEM) verified that the thickness of the Na<sub>3</sub>SbS<sub>4</sub> layers on NaCrO<sub>2</sub> was about 200 nm. NASICON could serve as a protective layer by using a 10 nm LAGP buffer layer (with a conductivity of 0.41 mS cm<sup>-1</sup>) as a solid electrolyte interface coating that covers the sulfide electrolyte infiltrating NCM622 cathode.18 The LAGP layer formed on the NCM622 surface using the sol-gel method may diminish interfacial side reactions while enhancing Li-ion transport. The dip-coating approach is then used, wherein the LGPS/NMP solution is permeated into the LAGP-coated NCM622 cathode sheet to improve solid-solid contact. NCM622, infused with LGPS and LAGP buffer layers on ASSLBs, had an initial discharge capacity of 141.5 mA h  $g^{-1}$  at 0.05C and maintained stability for 100 cycles at 0.1C.

3.2.3.3. Direct pressing process. The Li<sub>3</sub>N-based interface layer was applied on the Li/Li<sub>2</sub>ZrCl<sub>6</sub> (LZC) interface. Commercial Li<sub>3</sub>N (c-Li<sub>3</sub>N) was converted into the α-phase and β-phase using ball-milling and annealing treatment, respectively. The EIS study of  $\beta$ -Li<sub>3</sub>N reveals an ionic conductivity of 2.46  $\times$ 10<sup>-4</sup> S cm<sup>-1</sup>, and its deformability characteristics allow production of a pellet by direct pressing with the solid electrolyte. The conductivity of LZC with and without the β-Li<sub>3</sub>N layer is  $2.69 \times 10^{-4} \text{ S cm}^{-1}$  and  $2.95 \times 10^{-4} \text{ S cm}^{-1}$ , respectively, indicating that the β-Li<sub>3</sub>N layer is stable on LZC. The stability of lithium plating/stripping in symmetrical Li|LZC|Li cells at a current density of 0.1 mA cm<sup>-2</sup> exhibits an initial overpotential of 250 mV, which expands to around 1 V after 80 hours. The symmetrical Li $|\beta$ -Li<sub>3</sub>N/LZC/ $\beta$ -Li<sub>3</sub>N|Li cells exhibit an initial overpotential of 100 mV. After reaching 300 hours of operation with an overpotential of around 50 mV. The Li<sub>3</sub>N layer effectively decreases interfacial resistance, enhances Li<sup>+</sup> diffusion, and preserves the stability of the plating/stripping cycle.<sup>274</sup>

3.2.3.4. Atomic layer deposition. For halide ISE, alongside the use of metal alloys to mitigate the spontaneous interaction between halides and ISEs, the implementation of sulfide-based anode interface layers (i.e. Li<sub>6</sub>PS<sub>5</sub>Cl) presents an alternative strategy. 46,199,212 Li<sub>6</sub>PS<sub>5</sub>Cl in contact with lithium will create SEI layers (Li<sub>2</sub>S, Li<sub>3</sub>P, and P<sub>2</sub>S<sub>5</sub>) that restrict the interaction between

the halide ion-conducting electrolyte and lithium. The contact established between the halide and sulfide electrolytes is significant. Koç et al.272 elucidated that the SXRD investigation of the Li<sub>3</sub>InCl<sub>6</sub>/Li<sub>6</sub>PS<sub>5</sub>Cl interface revealed the decomposition products, LiCl. The assembly of dual ISEs using Li<sub>3</sub>InCl<sub>6</sub> and Li<sub>6</sub>PS<sub>5</sub>Cl facilitates interfacial processes that impede the efficiency of ASSBs. A possible approach involves interface engineering using a protective layer, such as Li<sub>3</sub>PO<sub>4</sub>. Koc and coworkers used the ALD process as an outcome. ALD has difficulties in modifying layer thickness because of the rough surface of Li<sub>6</sub>PS<sub>5</sub>Cl. Nonetheless, the benefit of ALD is found in its ability to produce ultra-uniform and ultra-thin structures on a wide scale without damaging the basic crystal structure of the coated material.269 The thickness of the applied layer is determined by the number of cycles, 17 ALD cycles for a 2 nm thick Li<sub>3</sub>PO<sub>4</sub>. In the cell configuration NCM622(coated)/Li<sub>3</sub>InCl<sub>6</sub>/ VGCF|Li<sub>3</sub>InCl<sub>6</sub>/Li<sub>3</sub>PO<sub>4</sub>-coated Li<sub>6</sub>PS<sub>5</sub>Cl|Li<sub>0.5</sub>In/Li<sub>6</sub>PS<sub>5</sub>Cl, the cell with a 2 nm thick Li<sub>3</sub>PO<sub>4</sub> layer exhibits 92.3% capacity retention after 400 cycles, while the cell with a 1 nm thick Li<sub>3</sub>PO<sub>4</sub> layer demonstrates 88% retention after 151 cycles. The voltage of 3.7 V exhibited comparable retention rates of 79.9% and 80.1% after 250 cycles for Li<sub>3</sub>PO<sub>4</sub> films of 2 nm and 1 nm thickness, respectively (Fig. 18d1 and d2). Raising the cut-off voltage to 3.8 V results in diminished performance. The application of a Li<sub>3</sub>PO<sub>4</sub> layer with a thickness of up to 10 nm exhibited no electrochemical activity.

Wang et al.270 integrated atomic layer deposition and wetchemical techniques to create a dual shell design for lithium cobalt oxide cathodes. The direct interaction between LCO and LGPS electrolyte promotes the formation of a high-resistivity CEI layer (Fig. 18e1); therefore, the application of LiNbO<sub>3</sub>based layer material via the ALD process results in a one-shell LiNbO<sub>3</sub>@LCO structure. LiNbO<sub>3</sub> serves as an interfacial layer and stabilizes the LGPS-LCO contact. The LiNbO3@LCO/LGPS composite, synthesized by the dry mixing process, exhibits reduced aggregation; nevertheless, the ionic conductivity of the combination requires enhancement (Fig. 18e2). The second layer, LGPS, is formed on LiNbO3@LCO via a wet-chemistry dispersion technique, resulting in the dual LGPS@LiNbO3@LCO. This approach effectively produces LiNbO<sub>3</sub>@LCO particles uniformly coated with LGPS (Fig. 18e3). The GITT study indicates that the Li<sup>+</sup> diffusion efficiency of the dual-shell design is four times higher than that of the singleshell design. ASSBs with a single-shell LiNbO3@LCO cathode exhibited a specific capacity of 20 mA h g<sup>-1</sup> at 1C, while the dual-shell LGPS@LiNbO3@LCO electrode demonstrated a capacity of 87.7 mA h g<sup>-1</sup> at the same rate. The advancement of dual-shell LGPS@LiNbO3@LCO using ALD/Wet-chemical techniques is expected to provide promising outcomes for high-performance ASSBs.

3.2.3.5. In situ synthesis process. According to Kato et al., 276 lithium superionic conductors such as Li<sub>9.6</sub>P<sub>3</sub>S<sub>12</sub> and Li<sub>9.54</sub>-Si<sub>1.74</sub>P<sub>1.44</sub>S<sub>11.7</sub>Cl<sub>0.3</sub> have LGPS-like crystal structures, with Li<sub>9.54</sub>Si<sub>1.74</sub>P<sub>1.44</sub>S<sub>11.7</sub>Cl<sub>0.3</sub> having the highest conductivity of 25 mS cm<sup>-1</sup>, which is twice the LGPS conductivity value (12) mS cm<sup>-1</sup>) reported by Kamaya et al.<sup>277</sup> In ASSBs, the coulombic efficiency of cells with solid electrolytes based on Li<sub>9.6</sub>P<sub>3</sub>S<sub>12</sub>,

# Coating process of interface layers

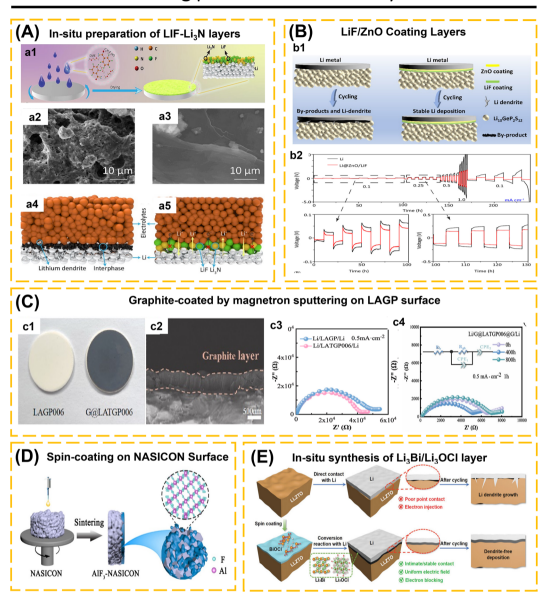


Fig. 19 (A) (a1) The Li@LiF-Li<sub>3</sub>N anode is *in situ* formed *via* a manipulated reaction of C<sub>7</sub>H<sub>2</sub>F<sub>5</sub>NO with lithium metal, realizing highly stable all-solid-state lithium batteries. Surface SEM images of Li anodes obtained from (a2) LCO|LGPS|Li and (a3) LCO|LGPS|Li@LiF-Li<sub>3</sub>N batteries after 20 cycles at 0.1C. Lithium deposition behaviors of (a4) Li/electrolytes and (a5) Li@LiF-Li<sub>3</sub>N/electrolytes. Reproduced with permission from ref. 97. Copyright 2023, Elsevier. (B) (b1) Proposed interfacial evolution of Li and Li@ZnO/LiF negative electrodes with LGPS after deposition. (b2) Rate capability of the symmetric cells of Li@ZnO/LiF|LGPS|Li@ZnO/LiF, Li|LGPS|Li at 0.1, 0.25, 0.5, 1.0 mA cm<sup>-2</sup>. Reproduced with permission from ref. 1. Copyright 2023, Wiley-VCH. (C) (c1) Optical images of the polished LATGP006 pellet and the LATGP006 pellet with a graphite layer. (c2) SEM images of the LATGP006 pellet with a graphite layer: cross-sectional image. (c3) EIS spectra of the Li/LAGP/Li and Li/LATGP006/Li cells after cycling at 0.5 mA cm<sup>-2</sup>. (c4) Nyquist plots of the symmetric Li/G@LATGP006@G/Li cell before and after cycling at current densities of 0.5 mA cm<sup>-2</sup>. Reproduced with permission from ref. 91. Copyright 2023, Elsevier. (D) (d1) Schematic illustration of the synthesis process of AlF<sub>3</sub>-NASICON pellets. (d2) SEM images of the cross-sections of Na/AlF<sub>3</sub>-NASICON interfaces. Insets are the corresponding digital images showing the wetting behaviors of molten Na on NASICON and AlF<sub>3</sub>-NASICON, respectively. Reproduced with permission from ref. 266. Copyright 2020, Elsevier. (E) Schematic diagrams of the fabrication of alloy@antiperovskite hybrid layer by *in situ* conversion reaction between garnet and Li metal, and the comparison of dendrite suppression capability for the pristine and modified Li/garnet interfaces. Reproduced with permission from ref. 98. Copyright 2023, Wiley-VCH.

 ${\rm Li_{10}GeP_2S_{12}}$ , and  ${\rm Li_{9.54}Si_{1.74}P_{1.44}S_{11.7}Cl_{0.3}}$  is 90%, 61%, and 39%, respectively. The loss in efficiency is caused by a certain amount of lithium being consumed during the reaction, which forms an interfacial layer on the electrode/SE interface. To prevent the

formation of lithium dendrites, an electron-insulating layer for the lithium anode must be introduced. Wu *et al.*<sup>97</sup> developed an anode interface layer based on LiF–Li<sub>3</sub>N (Fig. 19a1). Li@LiF–Li<sub>3</sub>N was synthesized by an *in situ* reaction of C<sub>7</sub>H<sub>2</sub>F<sub>5</sub>NO in DMF

solvent. The C<sub>7</sub>H<sub>2</sub>F<sub>5</sub>NO-DMF solution was dropped on lithium foil using spin coating methods and dried at 60 °C for 2 hours to produce Li@LiF-Li3N electrodes. Stability performance analysis of Li/Li<sub>10</sub>GeP<sub>2</sub>S<sub>12</sub>/Li symmetric cell shows the overpotential reaches 2 V after 403 hours a 0.1 mA cm<sup>-2</sup>/0.1 mA h cm<sup>-2</sup>. The Li@LiF-Li3N/Li10GeP2S12/Li@LiF-Li3N symmetric cell can achieve stability of more than 9000 hours with an overpotential of 0.39 V. SEM was used to observe the anode morphology of Li and Li@LiF-Li3N anodes from all-solid-state batteries after 20 cycles (Fig. 19a2 and a3). The anode without a protective coating looks to have holes on its surface, as well as side reactions that damage the electrode/electrolyte contact. The anode's LiF-Li<sub>3</sub>N layer ensures robust surface integrity. Li<sub>3</sub>N, as a good ionic conductor, may prevent lithium side reactions that produce sulfide, but LiF, with its poor electronic conductivity and high interfacial energy, can reduce dendrite formations. The combination of both allows the electrode-electrolyte interface to remain stable while also providing superior electrochemical performance stability for ASSBs (Fig. 19a4 and a5).

3.2.3.6. Sputtering technique. Chang et al. applied a double layer of ZnO/LiF to prevent side reactions at the Li-LGPS metal contact. Magnetic sputtering provides benefits over ALD and CVD due to its high deposition rate, favorable environmental

impact, ease of control, and high production efficiency on an industrial scale. ZnO was sprayed on the surface of the lithium metal disc using magnetic sputtering (RH450) at 50 W, 0.5 Pa, and a rotation speed of 20 rpm for 1 hour. Then, LiF in the same way for 6 hours to generate Li@ZnO/LiF. The lithiophilic ZnO layer limits direct contact between LGPS and lithium anode, while the lithiophobic LiF layer prevents lithium dendrite growth (Fig. 19b1). The Li@ZnO/LiF|LGPS|Li@ZnO/LiF symmetric cell demonstrates stability after 2000 hours of lithium plating/striping cycling, with a small overpotential of 200 mV at 0.1 mA cm<sup>-2</sup> for 1 hour. Fig. 19b2 illustrates tiny overpotentials of 15, 153, 564, 1431, and 198 mV for current densities of 0.1, 0.25, 0.5, 1.0, and 0.1 mA cm $^{-2}$ . The conclusion is that the ZnO/LiF coating may improve the stability of the Li/ LGPS interface.

Graphite exhibits good conductivity, superior mechanical properties, and electrochemical stability when used as the Li-LATGP interface layers. The graphite layer uniformly covers the LATGP surface with a thickness of 0.5 μm (Fig. 19c1 and c2). The Nyquist plots of LAGP, LAGTP006, and G@LAGTP006 in a symmetrical cell at 0.5 mA cm<sup>-2</sup> indicate that the impedance of G@LAGTP006 is inferior to that of LAGP and LAGTP006 (Fig. 19c3 and c4). The graphite layer effectively enhances the

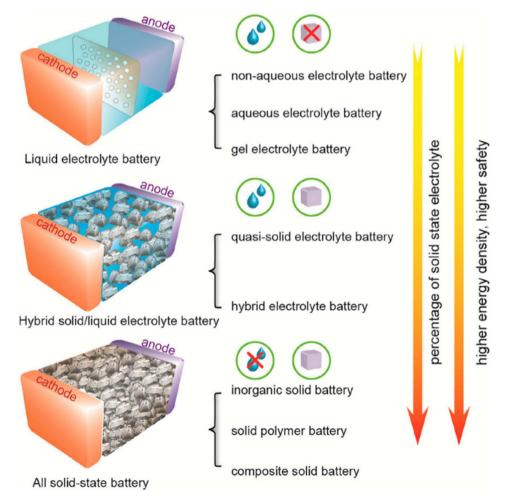


Fig. 20 Classification of batteries based on the use of liquid electrolyte. Reproduced with permission from ref. 278. Copyright 2020, American Chemical Society.

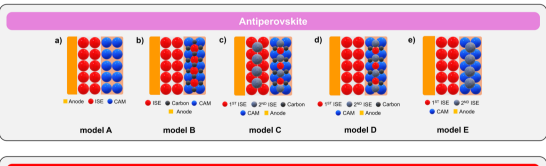
Table 6 Comparison of the main components of ASSSIBs and ASSLIBs

Materials	ASSSIBs	ASSLIBs
Cathode active material	NaTi <sub>2</sub> (PO <sub>4</sub> ) <sub>3</sub>	LFP
	$Na_3V_2(PO_4)_3$	NCA
	$Na_{0.67}Mn_{0.47}Ni_{0.33}Ti_{0.2}O_2$	LMO
	$Na[Ni_{1/3}Fe_{1/3}Mn_{1/3}]O_2$	LCO
	$\mathrm{TiS}_2$	LTO
	$Na_3V_2(PO_4)_2O_2F$	NCM811
	$Na_{0.85}Mn_{0.5}Ni_{0.4}Fe_{0.1}O_2$	NCM622
	$NaNi_{0.68}Mn_{0.22}Co_{0.1}O_{2}$	NCM83
	$Na_3V_2(PO_4)_2F_3$	NCM85
	$NaMO_2$ (M = Fe, Ni, Mn, Co)	NCM90
	· · · · · · · · · · · · · · · · · · ·	$TiS_2$
Catholyte	Na-based ISEs	Li-based ISEs
Conductive material	Carbon black, carbon nanofiber, MWCNT, RGO, etc.	Carbon black, carbon nanofiber, MWCNT, RGO, etc.
Cathode interface layer	_	Metal oxide-based layer
•		Oxide-based ISEs
		Halide-based ISEs
Electrolyte	Na-based ISEs	Li-based ISEs
Anode interface layer	Metal oxide	Sulfide-based ISEs
•	Metal halide	Halide-based ISEs
		Metal halide-based layer
		Metal oxide-based layer
		Graphite-based layer
		Nitride-based layer
Anode	Na	Li
	Na-Sn	Li–In
	Na-SiO <sub>2</sub>	Li–Si
	Hard carbon	Graphite

stability of the Li-LAGTP006 contact. The initial discharge of the Li/LAGTP006/LFP cell reaches 142.1 mA h  $\rm g^{-1}$ , exhibiting a retention of 17.6% after 100 cycles. The Li/G@LAGTP/LFP cell has a retention rate of 87.8%. The graphite interface layer

enhances cycling performance by preventing damage at the interface during charge-discharge cycles.<sup>91</sup>

3.2.3.7. Spin-coating with heat-treatment process. Miao et al.  $^{266}$  synthesized an AlF<sub>3</sub>-based protective layer at the Na/NASICON interface by a spin-coating technique. The AlF<sub>3</sub>



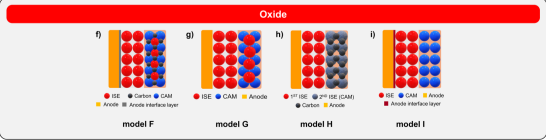
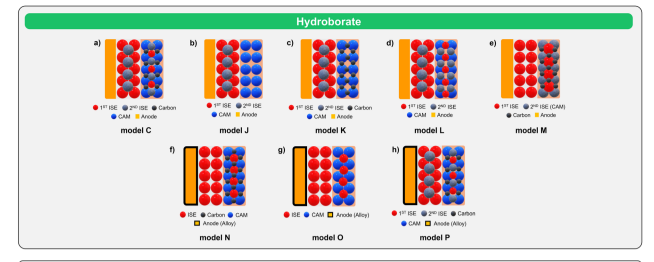


Fig. 21 Current models of ASSBs assembly for antiperovskite-based ISEs: (a) model A, (b) model B, (c) model C, (d) model D, (e) model E, and oxide-based ISEs: (f) model F, (g) model G, (h) model I.



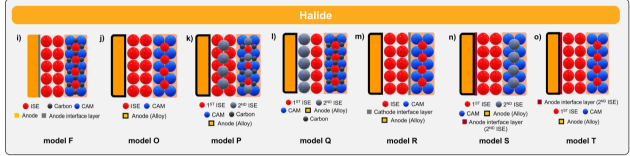


Fig. 22 Current models of ASSBs assembly for hydroborate-based ISEs: (a) model C, (b) model J, (c) model K, (d) model L, (e) model M, (f) model N, (g) model O, (h) model P, and halide-based ISEs: (i) model F, (j) model O, (k) model P, (l) model Q, (m) model R, (n) model S, (o) model T.

solution was prepared from Al(NO<sub>3</sub>)<sub>3</sub>.9H<sub>2</sub>O and NH<sub>4</sub>F. The NASICON pellet was coated with 60 µL of AlF<sub>3</sub> at 4000 rpm for 30 seconds, then sintered at 450 °C for 5 hours (Fig. 19d). The AlF<sub>3</sub> solution enhanced interfacial contact, exhibiting a conductivity of  $2.415 \times 10^{-4} \text{ S cm}^{-1}$  at RT, and filled the gaps in the NASI-CON to inhibit dendrite growth. Under high voltage conditions, the AlF3-coated NASICON exhibited cycle stability, while the uncoated NASICON demonstrated significant voltage drop and short circuit. Furthermore, the propensity of AlF<sub>3</sub> to react with Na during the first phase of activation led to the formation of stable SEI layers, which reduces electron distribution and increases the dendrite resistance ability. This significantly enhances the cycle stability of solid-state batteries. Bi et al.98 explained the formation of a hybrid layer consisting of Li<sub>3</sub>Bi alloy nanoparticles embedded inside an antiperovskite Li<sub>3</sub>OCl matrix (Li<sub>3</sub>Bi@Li<sub>3</sub>OCl) in situ at the interface of Li|Li<sub>6.75</sub>La<sub>3</sub>-Zr<sub>1.75</sub>Ta<sub>0.25</sub>O<sub>12</sub> (Fig. 19e). BiOCl is deposited onto the Li<sub>6.75</sub>La<sub>3</sub>-Zr<sub>1,75</sub>Ta<sub>0,25</sub>O<sub>12</sub> surface using spin-coating and undergoes a spontaneous reaction with Li, resulting in the formation of a hybrid Li<sub>3</sub>Bi@Li<sub>3</sub>OCl layer. The density of states (DOS) indicates that Li<sub>3</sub>OCl, possessing a bandgap of 5.06 eV, can inhibit electron tunneling from Li to LLZTO and serve as a good ionic conductor at the interface. On the other hand, Li<sub>3</sub>Bi alloy, with a bandgap of 0.44 eV, functions as an electron conductor, with Li<sub>3</sub>Bi uniformly dispersed within the Li<sub>3</sub>OCl matrix, thereby restricting electron conduction and preventing electron

movement through the Li<sub>3</sub>OCl matrix. This functions to inhibit dendritic growth. Li3B exhibits superior lithiophilic capabilities due to its reduced surface energy in comparison to Li<sub>3</sub>OCl. The Li<sub>3</sub>Bi@Li<sub>3</sub>OCl hybrid layer effectively decreased the interface resistance from 349  $\Omega$  cm<sup>2</sup> to 27  $\Omega$  cm<sup>2</sup>. In conclusion, BiOCl can spontaneously react with Li to form a Li3Bi@Li3OCl hybrid layer at the Li-LLZTO interface and exhibits good lithiophilicity, ionic conductivity, and electron-blocking properties. This also indicates that antiperovskite can act as an interfacial layer to improve the incompatibility of garnet to Li metal.

# Current models of ASSBs and multi-phase ISEs-based ASSBs

#### 4.1. Current models of all-solid-state lithium/sodium-ion batteries

The solid-state battery configuration model is an innovative improvement over liquid electrolyte-based batteries. It replaces the liquid electrolyte and separator with a solid electrolyte, such as ISE, which performs the functions of both.200 Chen et al.278 clarified the classification of batteries based on the presence or absence of LE. In general, solid-state batteries consist of an anode (Li or Na metal), ISE, and a cathode (containing cathode active materials such as LCO, LTO, NMC, and LFP). This setup forms a straightforward interface system between the anode/ISE

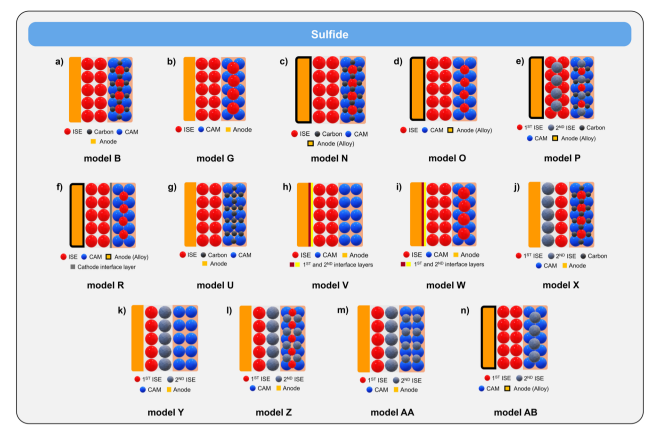


Fig. 23 Current models of ASSBs assembly for sulfide-based ISEs: (a) model B, (b) model G, (c) model N, (d) model O, (e) model P, (f) model R, (g) model U, (h) model V, (i) model W, (j) model X, (k) model Y, (l) model Z, (m) model AA, (n) model AB

and ISE/cathode, as shown in Fig. 20. This section has summarized some published model designs of lithium/sodiumbased ASSBs. First, it is necessary to know the general configuration of both systems. Table 6 shows the comparison of components in both systems based on the information gathered. Some solid electrolytes may exhibit incompatibility with either the anode or the cathode or both. In conventional solidstate batteries, a certain quantity of liquid electrolytes is frequently applied to improve the wetting properties of the interface between the ISE and cathode. 91,125,158,159,163,279 In order to eliminate the demand for flammable organic materials in solid-state batteries, the liquid electrolyte can be replaced with a solid interface layer made of metal oxide, or ISEs. A solid interface layer is typically applied to facilitate solid-solid contact interactions in ASSBs. 91,97,270 Additionally, a composite cathode consisting of CAM, conductive carbon, and catholyte (ISEs) can be applied to enhance resistance and reduce side reactions at the ISE/cathode interface.77,103,109,110,165,196,269 This will complicate the interface system in ASSBs such as CAMcarbon, CAM-catholyte, carbon-catholyte, CAM-cathode interface layer, carbon-cathode interface layer, catholyte-cathode interface layer, ISE/cathode interface layer, ISE-anode interface layer, and anode-anode interface layer. In Fig. 21-23 present the existing 28 ASSBs models and categorize them based on ISEs type. (1) Antiperovskite: this group of ISEs remains stable when

in contact with a lithium anode, even without the presence of an interface layer. The cathode typically consists of CAM, ISEs, and conductive carbons. (2) Sulfide: ASSBs that use sulfide electrolytes often employ Li/Na metal or alloy as the anode. Several models display an interface layer on either the anode, cathode, or both. The composition of the cathode varies. Hybrid sulfidebased electrolytes may be used either as a homogenous mixture or as stacked layers of electrolyte. (3) Oxide: this ISE not only serves as an electrolyte and a catholyte, but also functions as a CAM based on the model presented. Further development, single-class ASSBs in NASICON class can be collected, with the possibility of the material having the same or different chemical composition. This ISE exhibits stability when exposed to both the anode and cathode. In some cases, an interface layer may be added. (4) Halide: electrolytes exhibit such high reactivity towards lithium anodes that the use of anode interface layers or alloys is increasingly prevalent. For some cases with halide electrolyte, the utilization of interface layers is applied even though anodes (alloys) or composite cathodes have been used. (5) Hydroborate: in the ASSBs models do not include the usage of anode and cathode interface layers. This ISE has excellent stability when used with metal anodes, alloy anodes, and composite cathodes. Tables 7 and 8 summarize example configurations of each current ASSB model along with a comparison of their respective electrochemical performances.

Table 7 Performance comparison of Na-based ISEs in ASSSIBs

	Solid electrolyte	Anode	ASSBs model	Capacity	Potential window	Capacity retention	Ref.
$NaTi_2(PO_4)_3 + Carbon$	$\rm Na_{3.16}Zr_{1.84}Y_{0.16}Si_2PO_{12}$	Na	Н	Reversible capacity of	1.5–2.8 V vs. Na/Na <sup>+</sup>	28.3% capacity retention	78
		;	;	56.6 mA h g <sup>-1</sup> at C/5	+	after 950 cycles at C/5	
$ m Na_3V_2(PO_4)_3 + carbon$ black + PVDF	UHS-NZSP	Na	Ħ	Initial discharge capacity of 102 mA h $\mathrm{g}^{-1}$ (NVP loading of $\sim$ 2 mg cm $^{-2}$ )	2.0-4.0 V vs. Na/Na	82% capacity retention after 850 cycles	39
$\mathrm{NaTi}_2(\mathrm{PO}_4)_3 + \mathrm{RGO}$	$NLZSP-NBO_3$	Na	н	Initial discharge capacity of 95.4 mA h $g^{-1}$	1.5–3.0 V $\nu s$ . Na/Na $^+$	91.4% capacity retention after 200 cycles at 1C	41
${\rm Na_{2/3}Ni_{1/3}Mn_{2/3}O_2}$	NZSP	Na	A	Initial discharge capacity of $92.5 \text{ mA h g}^{-1}$ at $0.1C$	2.0–4.0 V υs. Na/Na <sup>+</sup>	I	229
$TiS_2 + Na_3SbS_4$	$\mathrm{Na_3SbS_4}$	Na	Ð	Initial charge capacity of $115.9 \text{ mA h g}^{-1}$ at $0.1\text{C}$	1.2–2.4 V vs. Na/Na <sup>+</sup>	I	104
TiS <sub>2</sub> + Ca-doped Na <sub>3</sub> PS <sub>4</sub>	${ m Ca-doped~Na_3PS_4}$	Na-Sn alloy	Ь	2nd charge capacity of 198 mA h g <sup>-1</sup> at 0.06C	1.0–2.5 V vs. Na/Na <sup>+</sup>	91% capacity retention after 100 cycles	101
$Na_3V_2(PO_4)_3$ + carbon + t- $Na_3PS_{3.8}Cl_{0.2}$	$\text{t-Na}_{3}\text{PS}_{3.8}\text{Cl}_{0.2}$	Na	M	Initial discharge capacity of 100 mA h $g^{-1}$ at 10 mA $g^{-1}$	2.5–3.8 V νs. Na/Na <sup>+</sup>		63
$Na[Ni_{1/3}Fe_{1/3}Mn_{1/3}]O_2 + Na_3B_{24}H_{23}-5Na_2B_{12}H_{12} + carbon black$	${ m Na_3B_{24}H_{23}}{ m -}5{ m Na_2B_{12}H_{12}}$	Na	U	Initial discharge of $51.2 \text{ mA h g}^{-1}$ at $0.1\text{C}$	2.0–4.2 V vs. Na/Na <sup>+</sup>	87% capacity retention after 50 cycles at 0.1C at 25 °C	70
${ m TiS}_2$	$Na_{3}B_{24}H_{23} - 5Na_{2}B_{12}H_{12}$	Na	'n	Initial discharge of $171.4 \text{ mA h g}^{-1} \text{ at } 0.1\text{C}$	1.5–2.5 V vs. Na/Na <sup>+</sup>	$\sim$ 150 mA h g $^{-1}$ after 100 cycles at 0.1C	20
TiS <sub>2</sub> + acetylene black + PEG	$NaB_3H_8 \cdot xNH_3 \otimes NaB_3H_8$ (x < 1)	Na	X	Initial discharge of $237.2 \text{ mA h g}^{-1}$ at 1C	$1.40-2.35~\mathrm{V}~\nu\mathrm{s}.~\mathrm{Na/Na^+}$	90.1% capacity retention after 240 cycles at 1C	20
${ m Na_3V_2(PO_4)_2O_2F} + { m Na_4B_36H_{34}-7Na_2B_{12}H_{12}} + { m carbon\ black}$	${ m Na_4B_36H_{34}}^{-7{ m Na}_2B_{12}H_{12}}$	Na	M	Initial discharge of 71.2 mA h $g^{-1}$ at 0.1C	2.5–4.5 V vs. Na/Na <sup>+</sup>	85.8% capacity retention at 25 °C after 100 cycles	69
${ m TiS_2 + Na_2B_{20}H_{18} - } \ 4{ m Na_2B_{12}H_{12}}$	$\rm Na_2B_{20}H_{18}{\rm ^-}4Na_2B_{12}H_{12}$	Na	ı	Initial discharge of $64.8 \text{ mA h g}^{-1}$	1.5–2.7 V vs. Na/Na <sup>+</sup>	82% capacity retention after 100 cycles at 0.1C	51
${ m TiS_2 + Na_4B_{20}H_{18} - } \ 3Na_2B_{12}H_{12}$	$\rm Na_4B_{20}H_{18}{\rm -}3Na_2B_{12}H_{12}$	Na	L	Initial discharge of $38.1 \text{ mA h g}^{-1}$ at $0.1\text{C}$	1.2–2.7 V vs. Na/Na <sup>+</sup>	92% capacity retention after 50 cycles at 0.1C	9
$\begin{array}{l} \text{NMNFO} + \text{CB} + 0.62 \\ \text{[HCF-Sm]} \cdot 0.38 \text{[LCF-Ta]} \end{array}$	0.62[HCF-Sm] 0.38[LCF-Ta]	Na-Sn alloy	Ъ	Initial discharge of 117.9 mA h $\mathrm{g}^{-1}$	2.3-4.0 V vs. Na/Na <sup>+</sup>	88.3% after 100 cycles at 0.2C (25 $^{\circ}$ C)	30
$NMNFO + CB + 0.57$ $[HCF-La] \cdot 0.43[LCF-Ta]$	0.57[HCF-La] 0.43[LCF-Ta]	Na-Sn alloy	Ъ	Initial discharge of 95.1 mA h $\mathrm{g}^{-1}$	2.3-4.0 V vs. Na/Na <sup>+</sup>	91.0% after 100 cycles at 0.2C (25 °C)	30

Table 8 Performance comparison of Li-based ISEs in ASSLIBs

Li <sub>3</sub> BO <sub>3</sub> -coated LiCoO <sub>2</sub>	•	Anode	model	Capacity	window	Capacity retention	Ref.
	Al-doped Li <sub>7</sub> La <sub>3</sub> Zr <sub>2</sub> O <sub>12</sub>	Li (Au layer)	I	Initial discharge capacity of 128 mA h g <sup>-1</sup>	2.8-4.3 V	87% after 30 cycles	73
LCO/Ta, Al-doped LLZO	Ta, Al-doped LLZO	In	Ŋ	at 0.2C Initial discharge capacity of 60 mA h ${ m g}^{-1}$	2.8-3.6 V	ı	77
LFP + LAGP + carbon black	(Li <sub>6.45</sub> Al <sub>0.05</sub> La <sub>3</sub> Zl <sub>1.6</sub> 1a <sub>0.4</sub> O <sub>12)</sub> Te-doped LAGP	Li (graphite layer)	ĹΉ	Initial discharge of 154.4 mA h $\rm g^{-1}$ at 0.1	2.5-4.0 V vs.	135.6 mA h $g^{-1}$ (87.8% capacity	91
Linbo <sub>3</sub> @NCA +	$\rm Li_7P_{2.9}Ge_{0.05}S_{10.75}O_{0.1}$	Li-In	0	Initial discharge capacity of	2.1-3.7 V vs.	89% capacity retention after 50	65
$\text{Li}_7 P_{2.9}^{2} \text{Ge}_{0.05}^{2}_{10.75}^{2} \text{O}_{0.1}$ $\text{LiNbO}_3 \text{@NCA} + \text{Li}_7 P_3 \text{S}_{11}$	$\mathrm{Li}_7\mathrm{P}_3\mathrm{S}_{11}$	Li-In	0	Ibs.9 mA n g at 0.064 mA cm , KI Initial discharge capacity of	2.1-3.7 V vs.	cycles 61.57% capacity retention after 50	65
ГСО	$\begin{array}{l} Li_{10}GeP_2S_{12} + 75\%\ Li_2S - \\ 24\%\ P_2S_5 - 1\%\ P_2O_5 \end{array}$	Li	¥	118.4 mA h g $^{-}$ at 0.064 mA cm $^{-}$ , KT Initial discharge capacity of $\sim$ 124 mA h g $^{-1}$ at 0.1C	Li/Li 3.0–4.2 V νs. Li/Li <sup>+</sup>	cycles $115.1 \text{ mA h g}^{-1}$ after 10 cycles at 0.1C	110
70 wt% LCO + 30 wt% (5%		Li	×	Initial discharge capacity of	3.0-4.2 V vs.	93.4 mA h g <sup>-+</sup> after 50 cycles at 0.1C $118.6$ mA h g <sup>-1</sup> after 10 cycles at $\frac{1}{2}$	110
NIO-CINI (d/L/7F3O11)	$24\% V_2 V_5 = 1\% V_2 V_5$			~124 IIIA II g at 0.10	רו/ רו	114.3 mA h g <sup>-1</sup> (91.9% capacity retention) after 50 evoles at 0.10	
70 wt% LCO + 30 wt%	$\text{Li}_{10}\text{GeP}_2\text{S}_{12} + 75\% \text{ Li}_2\text{S} - 340\% \text{ B S} - 10\% \text{ D O}$	Li	×	Initial discharge capacity of $\frac{1}{1251}$ and $\frac{1}{10000}$	3.0–4.2 V vs.	118.5 mA h $g^{-1}$ after 25 cycles at	110
$100 \text{ Cer}_{2}$ $212 \text{ 80 wt% (LCO + 5% NiS-CNIT%) I is p. c.) + 20 urth 200 \text{ cm}$	$24\% \text{ F}_2\text{S}_5 - 1\% \text{ F}_2\text{O}_5$ $\text{Li}_{10}\text{GeP}_2\text{S}_{12} + 75\% \text{ Li}_2\text{S} - 24\% \text{ D} \text{ C}_2$	Li	Z	135.1 MA II $g$ at 0.05C Initial discharge capacity of 125.1 mA b $g^{-1}$ at 0.05C	3.0-4.2 V vs.	$124 \text{ mA h g}^{-1}$ after 25 cycles at	110
$ ext{Li}_{10} ext{GeP}_2 ext{S}_{12}$	24/01295 - 1/01295			100:1 III B at 0:000	17/17		
LiNi <sub>0.7</sub> Mn <sub>0.15</sub> Co <sub>0.15</sub> O <sub>2</sub> + Li <sub>3</sub> PS <sub>4</sub> -2.3LiBH <sub>4</sub> + conductive carbon	$\mathrm{Li}_3\mathrm{PS}_4$ –2.3 $\mathrm{LiBH}_4$	Li-In	Ъ	Initial discharge of 177.5 mA h $\rm g^{-1}$ at 0.05C	2.4–3.7 V νs. Li/Li <sup>+</sup>	I	71
LCO + SbCl <sub>3</sub> -doped LGPS + VGCF + n(MMA/nBA)	SbCl <sub>3</sub> -doped LGPS	Li	В	Initial discharge capacity of 125 6 mA h o <sup>-1</sup>	I	86.3% capacity retention at 0.1C (60 °C) after 250 evoles	49
NCM 811 +  Li <sub>10.2</sub> P <sub>2.95</sub> Sb <sub>0.05</sub> S <sub>12</sub> I + vapor- orown nanofiber carbon	$Li_{10.2}P_{2.95}Sb_{0.05}S_{12}I$	Li	В	Initial discharge of 147 mA h g <sup>-1</sup> at 0.1C	2.5–4.25 V νs. Li/Li <sup>+</sup>	141 mA h g <sup>-1</sup> , 96.6% capacity retention (200 cycles)	111
NCM 811 + Li <sub>10</sub> P <sub>3</sub> S <sub>12</sub> I + vapor-grown nanofiber	$\mathrm{Li}_{10}\mathrm{P}_{3}\mathrm{S}_{12}\mathrm{I}$	Li	В	Initial discharge of 126 mA h $\rm g^{-1}$ at 0.1C	2.5–4.25 V vs. Li/Li <sup>+</sup>	95 mA h $\mathrm{g}^{-1}$ , 75.4% capacity retention (200 cycles)	111
TiS <sub>2</sub> + LGSbS-04	LGSbS-04	Li-In	0	Initial discharge of 221 mA h $\rm g^{-1}$	0.78–2.08 V vs. Li/Li	$157 \text{ mA h g}^{-1}$ after $44 \text{ cycles}$	25
LCO + LGPS (LLZCO coating laver)	LGPS	Li-In	В	Initial discharge of 121.0 mA h $\mathrm{g}^{-1}$ at 0.2C	1.9–3.6 V vs. Li/Li <sup>+</sup>	96% capacity retention after 100 cycles	92
TCO + TGPS	LGPS	Li-In	0	Initial discharge of 73.1 mA h $\mathrm{g}^{-1}$ at 0.2C	1.9–3.6 V vs. Li/Li <sup>+</sup>	59% capacity retention after 100 cycles	9/
$\rm NCM622 + Li_{10}SnP2S_{12}$	$\mathrm{Li}_{10}\mathrm{SnP2S}_{12}$	Li-In	0	Initial discharge of 122.9 mA h $\mathrm{g}^{-1}$ at 0.1C	2.4–3.7 V νs. Li/Li <sup>+</sup>	}	99

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Table 8 (Contd.)

Cathode	Solid electrolyte	Anode	ASSBs model	Capacity	Potential window	Capacity retention	Ref.
$NCM622 + Li_3InCl_6$	$\mathrm{Li}_{10}\mathrm{SnP2S}_{12}$	Li-In	AB	Initial discharge of 176.1 mA h $\mathrm{g}^{-1}$ at	2.4–3.7 V vs.	$106.2 \text{ mA h g}^{-1}$ after $100 \text{ cycles}$	99
(catholyc/coating layer) LCO + LGPS	LGPS	Li (Li <sub>3</sub> N–LiF layer)	W	Initial discharge of 132 mA h $\mathrm{g}^{-1}$ at 0.1C	3.0–4.2 V vs. Li/Li <sup>+</sup>	93.9% capacity retention after 50 cycles	97
				Initial discharge 95 mA h $\rm g^{-1}$ at 1C	ļ	93.5% capacity retention after 500 cycles at 1C	
TCO + TGPS	LGPS	Li	ß	Initial discharge of 126.5 mA h $\mathrm{g}^{-1}$ at	3.0–4.2 V vs.	14.6 mA h g <sup>-1</sup> after 20 cycles at 0.10 (14.5% consoits retention)	26
$LiNbO_{3}\text{-coated LCO} + LGPS - LGPS-Al_{2}O_{3} \ (90:10)$	$LGPS-Al_2O_3\left(90:10\right)$	Li-In	0	Initial discharge of 111 mA h g <sup>-1</sup> at 0.2C (unavroced)	1.9–3.6 V vs.	90% capacity retention after 50 carles at 0.20 (imaxposed)	80
				Initial discharge 108 mA h $g^{-1}$ at 0.2C (exposed at $-30$ °C, 1 h)	25. 11/11	cycles at 0.20 (unexposed) 92.6% capacity retention after 50 cycles at 0.20 (exposed at a dew point of -30 °C. 1 h)	
LiNbO <sub>3</sub> -coated LCO + L-LSiPSCI-1.0	L-LSiPSCI-1.0	Li-In	0	Initial discharge of 123 mA h $\rm g^{-1}$ at 0.2C	1.9–3.6 V νs. Li/Li <sup>+</sup>	>97% capacity retention after 100 cycles at 0.2C	7
LiNbO <sub>3</sub> -coated LCO + LGPS	RGPS	Li (ZnO-LiF layer)	>	5th -discharge capacity of 80 mA h $\mathrm{g}^{-1}$ at 0.10	3.0–4.2 V vs.	97% capacity retention after 500 cycles at 0.1C	П
$\begin{array}{c} NCM811 + Li_6PS_5Cl + super  Li_6PS_5Cl \\ P \end{array}$	${ m Li_6PS_5Cl}$	Li-In	z	Initial discharge of 192.4 mA h ${ m g}^{-1}$	1.9–3.6 V vs. vs. Li/Li <sup>+</sup>		89
$LCO + Li_6PS_5Cl$	$\mathrm{Li_6PS_5Cl} + \mathrm{Li_{5.6}PS_{4.6}I_{1.4}}$	Li	AA	Initial discharge of 128.8 mA h $\rm g^{-1}$ at 0.1C	2.5–4.2 V vs. Li/Li <sup>+</sup>	79.6% capacity after 50 cycles	209
LFP + PVDF + Super P	$\mathrm{Li}_{6}\mathrm{PS}_{5}\mathrm{F}_{0.5}\mathrm{Cl}_{0.5}$	Li	Ω	Initial discharge of 122 mA h $\rm g^{-1}$ at 0.2C	2.5–4.0 V νs. Li/Li <sup>+</sup>	Discharge capacity above 105 mA h g <sup>-1</sup> after 50 cycles	16
LTO + PVDF + Super P	${ m Li_6PS_5F_{0.75}I_{0.25}}$	Li	Ω	Initial discharge of 140 mA h $\mathrm{g}^{-1}$	$1.0-2.8 \text{ V } \nu s.$ Li/Li <sup>+</sup>	91.5% capacity retention after 50 cycles	92
${ m TiS_2} + { m Li_2B_{12}H_{12}} - { m 5Li_2B_{10}H_{10}} - { m 5Li_2B_{10}H_{10}} - { m 6Li_2B_{10}}$	${ m Li}_2{ m B}_{12}{ m H}_{12}$ – ${ m 5Li}_2{ m B}_{10}{ m H}_{10}$ – ${ m 6LiBH}_4$	$\mathrm{In_{1.3}Li_{0.3}}$	0	Initial discharge of 147.8 mA h $\rm g^{-1}$	2.0–2.7 V νs. Li/Li <sup>+</sup>	80% capacity retention after 120 cycles	52
NCM811 + $\text{Li}_3(\text{CB}_{11}\text{H}_{12})_2(\text{CB}_9\text{H}_{10})$ + carbon fibers	${ m Li}_3({ m CB}_{11}{ m H}_{12})_2({ m CB}_9{ m H}_{10})$	Li–In	Ъ	Initial discharge of $\sim$ 145 mA h g $^{-1}$ at C/10, RT	1.9–3.6 V νs. Li/Li <sup>+</sup>	$54\%$ capacity retention after 2000 cycles at C/2, 60 $^{\circ}\mathrm{C}$	59
$NCM811 + Li_3(CB_1H_{12})_2(CB_9H_{10}) + Carbon fibers$	${\rm Li}_3({\rm CB}_{11}{\rm H}_{12})_2({\rm CB}_9{\rm H}_{10})$	Graphite	Z	Initial discharge of $\sim$ 125 mA h g $^{-1}$ at C/10, RT	2.4–4.1 V νs. Li/Li <sup>+</sup>	75% capacity retention after 350 cycles at C/2, RT	59
$\mathrm{Fe}_2(\mathrm{MoO}_4)_3$	${ m Li}_2{ m OHBr}$	Li	A	Initial discharge of ${\sim}30~\mu\text{Ah}~\text{cm}^{-2}~\mu\text{m}^{-1}$	2.7 to 3.2 V	$\sim$ 35 $\mu$ Ah cm <sup>-2</sup> $\mu$ m <sup>-1</sup> after 60 cycles	74
$\rm LFP + Li_3 InCl_6$	$({ m Li}_2{ m OH})_{0.99}{ m K}_{0.01}{ m Cl}$	Li	ы	Initial discharge of $\sim 120 \text{ mA h g}^{-1}$ at 80 mA g <sup>-1</sup>	2.9–3.8 V vs. Li/Li <sup>+</sup>	96.1% capacity retention after 150 cycles at 80 mA $g^{-1}$	72
$LFP + LLZTO + Super-P + Li_{3-r}(OH_r)Cl_{0.9}F_{0.1}$	$LLZTO + Li_{3-x}(OH_x)Cl_{0.9}F_{0.1}$	Li	O	Initial discharge of 83.6 mA h g <sup>-1</sup> at 0.05C, 90 °C	2.7-4.0 V vs. Li/Li <sup>+</sup>	70.1% capacity retention after 50 cycles	24
$LFP + LLZTO + Super-P + Li_{3-x}(OH_x)Cl_{0.9}F_{0.1}$	$\mathrm{Li}_{3-x}(\mathrm{OH}_x)\mathrm{Cl}_{0.9}\mathrm{F}_{0.1}$	Li	Q	Initial discharge of 70.8 mA h $\rm g^{-1}$ at 0.05C, 90 $^{\circ}\mathrm{C}$	2.7-4.0 V νs. Li/Li <sup>+</sup>	62.5% capacity retention after 50 cycles	24

Table 8 (Contd.)

Cathode	Solid electrolyte	Anode	ASSBs model	Capacity	Potential window	Capacity retention	Ref.
NCM622 + Li <sub>3</sub> SI + Super P	Li <sub>3</sub> SI	In	В	Initial discharge of 96.8 mA h g <sup>-1</sup> at	I	50% capacity retention after 50 cycles at 0.04C	32
$VCI_3 + LIC + AB$	LIC	Li (Li <sub>6</sub> PS <sub>5</sub> Cl layer)	Ĺτι	Initial discharge capacity of $\sim$ 90 mA h $\sigma^{-1}$ at 6C	2.45–3.25 V	85.7% retention after 200 cycles at	29
$NCM811 + Li_{2.73}Ho_{1.09}Cl_6$ (1.1. PO., coating layer)	$\mathrm{Li}_{2.73}\mathrm{Ho}_{1.09}\mathrm{Cl}_{6}$	Li-In	В	Initial charge capacity of 209 mA h g <sup>-1</sup> at 0.1C (25 °C)	1.9–3.7 V νs. Li/Li <sup>+</sup>	Specific capacity of 125.5 mA h $\rm g^{-1}$ after 180 eveles (25 °C)	26
LCO + LIC	LIC	Li–In (Li <sub>6</sub> PS <sub>5</sub> Cl laver)	L	Initial discharge of 102.9 mA h $g^{-1}$ at 1C	2.7–4.2 V vs. Li/Li <sup>+</sup>	89.1% retention after 150 cycles at	42
NCM83 + Li-Y-Cl	$BM\text{-}SmCl_3\cdot 0.5Li_2ZrCl_6$	Li-In (Li-Y-Cl electrolyte laver)	S	Initial discharge capacity of $\sim$ 135 mA h g <sup>-1</sup> at 0.5C	2.2–3.7 V vs. Li/Li <sup>+</sup>	85% retention after 600 cycles	2
NCM88 + Li-LaCeZrHfTa- Cl + AB	$Li-LaCeZrHfTa-Cl+Li_6PS_5Cl$	Li-In alloy	$\circ$	Initial discharge capacity of 217.2 mA h $g^{-1}$ at 0.1C, 25 °C	2.5–4.3 V νs. Li/Li <sup>+</sup>	80% retention after 3000 cycles at 4C	54
80 wt% NCM85 + 20 wt% Li <sub>2</sub> $_6$ Zr <sub>0</sub> $_4$ Ho <sub>0</sub> $_6$ Cl <sub>6</sub>	$\mathrm{Li}_{2.6}\mathrm{Zr}_{0.4}\mathrm{Ho}_{0.6}\mathrm{Cl}_{6}$	In/Li–In (Li <sub>6</sub> PS <sub>5</sub> Cl coating laver)	H	Initial discharge of 180 mA h g <sup>-1</sup> at 0.5C	2.5–4.3 V νs. Li/Li <sup>+</sup>	96.8% retention after 180 cycles at 0.5C	75
80 wt% NCM85 + 20 wt% $Li_{2.6}Zr_{0.4}Lu_{0.6}Cl_6$	$\mathrm{Li}_{2.6}\mathrm{Zr}_{0.4}\mathrm{Lu}_{0.6}\mathrm{Cl}_{6}$	In/Li-In (Li <sub>6</sub> PS <sub>5</sub> Cl coating layer)	Т	Initial discharge of 202 mA h $\rm g^{-1}$ at 0.5C	$2.5-4.6 \text{ V} \nu s.$ $\text{Li/Li}^+$	88.9% retention after 180 cycles at 0.5C	75
80 wt% NCM85 + 20 wt% $\text{Li}_2  _{6} \text{Zr}_0  _{4} \text{Lu}_0  _{6} \text{Cl}_6$	$\mathrm{Li}_{2.6}\mathrm{Zr}_{0.4}\mathrm{Lu}_{0.6}\mathrm{Cl}_{6}$	Li <sub>0.7</sub> Si	0	Initial discharge of 16.1 mA h $\rm cm^{-2}$ at 0.5 mA $\rm cm^{-2}$	1.8–4.3 V νs. Li/Li <sup>+</sup>	I	75
$NCM83 + Li_3ZrCl_4O_{1.5}$	$\mathrm{Li}_3\mathrm{ZrCl}_4\mathrm{O}_{1.5}$	Li–In (Li <sub>3</sub> YCl <sub>6</sub> coating laver)	T	Initial discharge of $\sim$ 135 mA h g $^{-1}$ at 1C	1.9–3.7 V νs. Li/Li <sup>+</sup>	90.1% retention after 300 cycles at 1C	57
$\rm LCO + Li_3ZrCl_4O_{1.5}$	${ m Li}_3{ m ZrCl}_4{ m O}_{1.5}$	Li–In (Li <sub>3</sub> YCl <sub>6</sub> coating laver)	T	Initial discharge of $\sim 115 \text{ mA h g}^{-1} \text{ at}$ 0.5C	1.9–3.7 V νs. Li/Li <sup>+</sup>	90% retention after 240 cycles at 0.5C	57
NCM811 + LSZC	TSZC	Li-In (Li <sub>6</sub> PS <sub>5</sub> Cl coating laver)	T	al discharge of $\sim$ 200 mA h $\sim$ 150 mA h g <sup>-1</sup> at 1C	2.8–4.3 V νs. Li/Li <sup>+</sup>	82.7% retention after 1000 cycles at 1C, 30 °C	22
NCM90 + LSZC	TSZC	Li–In (Li <sub>6</sub> PS <sub>5</sub> Cl coating laver)	T	Initial discharge of 219.4 mA h g <sup>-1</sup> at 0.1C, 200.7 mA h g <sup>-1</sup> at 0.5C	2.8–4.3 V νs. Li/Li <sup>+</sup>	78.1% retention after >200 cycles at 0.5C, 30 °C	22
LCO + LZTC	LZFC	Li–In foil (Li <sub>6</sub> PS <sub>5</sub> Cl coating layer)	T	Initial discharge capacity of 129.9 mA h $g^{-1}$ at 0.05C	$2.3-3.6 \text{ V } \nu s.$ Li-In/Li <sup>+</sup>	Discharge capacity of 76.3 mA h g <sup>-1</sup> after 200 cycles	46
NCM811 + LZTC	LZTC	Li–In foil (Li <sub>6</sub> PS <sub>5</sub> Cl coating layer)	Т	Initial discharge capacity of $187.8 \text{ mA h g}^{-1}$ at $0.05C$	$2.2-3.7 \text{ V } \nu \text{S}.$ $\text{Li-In/Li}^+$	85.7% retention after 600 cycles	46
$\mathrm{LCO} + \mathrm{Li}_2\mathrm{ZrCl}_6$	$\mathrm{Li}_2\mathrm{ZrCl}_6$	Li–In foil (Li <sub>6</sub> PS <sub>5</sub> Cl coating layer)	Т	Initial discharge capacity of 137 mA h $\mathrm{g}^{-1}$	1.9–3.6 V νs. Li–In/Li	Discharge capacity of $114~{\rm mA~h~g}^{-1}$ after $100~{\rm cycles}$ at $0.5{\rm C}$	212
$NCM811 + Li_2ZrCl_6$	${ m Li}_2{ m ZrCl}_6$	Li–In foil (Li <sub>6</sub> PS <sub>5</sub> Cl coating layer)	Т	Initial discharge capacity of 181 mA h $\rm g^{-1}$	$2.2-3.8 \text{ V } \nu s.$ Li-In/Li <sup>+</sup>	Discharge capacity of $149 \mathrm{\ mAh\ g}^{-1}$ after 200 cycles at 1C	212
${ m LCO} + { m Li}_{2.25} { m Hf}_{0.75} { m Fe}_{0.25} { m Cl}_6$	Li <sub>2.25</sub> Hf <sub>0.75</sub> Fe <sub>0.25</sub> Cl <sub>6</sub>	Li–In foil (Li <sub>6</sub> PS <sub>5</sub> Cl coating layer)	L	Initial discharge capacity of 113.4 mA h $\rm g^{-1}$	2.5–4.2 V νs. Li–In/Li	88% retention after 100 cycles	199

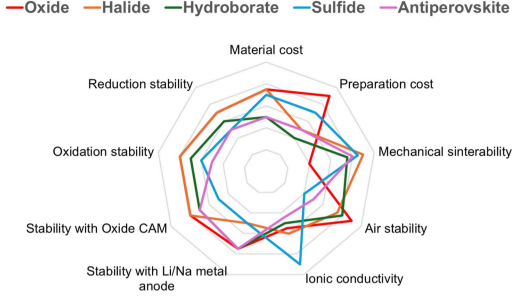


Fig. 24 Comparison of the properties of Na/Li-based ISEs.

#### 4.2. Multi-phase ISEs-based ASSBs

The development and application of ISEs-based ASSBs in recent decades have yielded favorable outcomes across various fields, demonstrating energy densities comparable to liquid electrolyte-based batteries and exhibiting promising safety levels.216,274 The stability of ISE with respect to electrode materials is critical, as it influences battery performance. The stability of the ISE/cathode interface has been addressed through the development of composite cathodes with an ISEbased catholyte and/or a cathode interface layer. 51,65,70,110 The stability of the ISE-anode interface is addressed through the utilization of Li/Na alloy and/or an anode interface layer. 30,42,71,75,101 The recent advancement indicates that ISE can be utilized as an anode-cathode interface layer. This expands the potential applications of ISEs, which can function in three distinct roles within ASSBs: as an electrolyte, catholyte, and interface layer at both anode and cathode sites.

The ongoing innovation has resulted in the advancement of multi-phase ISEs-based ASSBs, characterized by the incorporation of multiple types of ISEs in the latest ASSB assembly model. The application of a single ISE as an electrolyte encounters numerous challenges, including ionic conductivity, chemical and electrochemical stability, and potential window limitations. Hybrid ISEs combinations may serve as an alternate approach, with reported examples including hybrid LGPS-LPS (Li<sub>10</sub>GeP<sub>2</sub>- $S_{12}$  + 75%  $Li_2S$  - 24%  $P_2S_5$  - 1%  $P_2O_5$ ), 110 LPS-hydroborate (Li<sub>3</sub>PS<sub>4</sub>-2.3LiBH<sub>4</sub>),<sup>71</sup> double-argyrodite (Li<sub>6</sub>PS<sub>5</sub>Cl  $\text{Li}_{5.6}\text{PS}_{4.6}\text{I}_{1.4}$ ),<sup>209</sup> double-hydroborate ( $\text{Li}_3(\text{CB}_{11}\text{H}_{12})_2(\text{CB}_9\text{H}_{10})$ ),<sup>59</sup> garnet-antiperovskite (LLZTO + Li<sub>3-x</sub>(OH<sub>x</sub>)Cl<sub>0.9</sub>F<sub>0.1</sub>),<sup>24</sup> double- $(BM-SmCl_3 \cdot 0.5Li_2ZrCl_6)$ ,<sup>2</sup> halide-argyrodite LaCeZrHfTaCl + Li<sub>6</sub>PS<sub>5</sub>Cl).<sup>54</sup> The hybrid combination of ISEs at the catholyte and electrolyte sites includes catholyte (halide)-ISE (antiperovskite),72 catholyte (garnet)-ISE (antiperovskite),24 catholyte (halide)-ISE (halide),2 catholyte (halide)-ISE (halidesulfide),54 catholyte (halide)-ISE (sulfide),66 catholyte (sulfide)-

hybrid ISE (sulfide-sulfide),110 and catholyte (sulfide)-ISE (halide). 198,200 The integration of ISEs at both the electrolyte and interface layers includes ISE (halide)-anode interface layer (sulfide),22 ISE (halide)-anode interface layer (halide),57 ISE (sulfide)-cathode interface layer (garnet),76 and ISE (sulfide)cathode interface layer (halide).66 The wide scope of ISE types developed requires more investigation into the integration of multiple ISE types and the analysis of hybrid ISE interfaces to advance the next generation of ASSBs.

#### Dynamic database of solid-state electrolyte (DDSE)

Given the significant progress of various ISE classes in recent decades, it is imperative for researchers to have an ISE performance database. This database serves as a valuable resource for comparing and referencing current findings with existing ones. It also serves as a motivation for researchers to carry on their commitment to innovation in ISE research. Yang et al.280 developed a database referred to as the DDSE, which utilizes Machine Learning techniques. The program facilitates researchers in assessing the performance of ISEs discovered and allows for comparison with existing datasets. The DDSE database includes data from many ISEs that include both mono- and divalent cations such as Li<sup>+</sup>, Na<sup>+</sup>, K<sup>+</sup>, Ag<sup>+</sup>, Ca<sup>2+</sup>, Mg<sup>2+</sup>, and Zn<sup>2+</sup>, as well as anions including halide, hydride, and sulfide. This database will undoubtedly be invaluable for the progress of the next generations of ISEs and SSBs.

### Conclusion and future perspectives

In this review, we summarize recent advances in ISEs in terms of new chemical compounds, doping types, crystal structures, preparation methods, electrochemical behavior, chemical

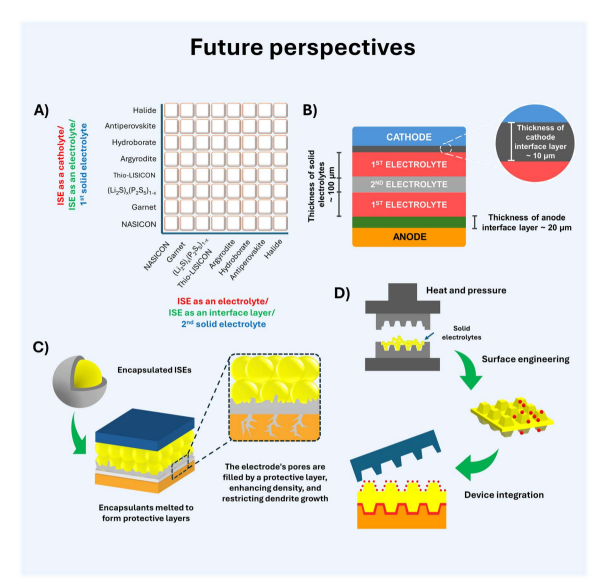


Fig. 25 Schematic representation of the future perspectives on ASSBs based on inorganic solid electrolytes: (A) multi-phase ISEs in ASSBs, (B) effect of layer thickness on ASSBs performance, (C) encapsulated electrolytes, and (D) lock and key concept.

reactivity, and alternative strategies to alleviate the limitations of ISEs. As demonstrated, halides exhibit relatively wide stability windows, highlighting the importance of further investigation into their structures and characteristics to fulfill the performance requirements of high-voltage electrodes. In addition, their ionic conductivity is comparable to other ISEs. Oxide electrolytes typically exhibit excellent chemical stability; nevertheless, their stiffness compromises the contact with the electrode materials, necessitating the use of liquid electrolytes in some instances. Regarding garnet, engineering the cathode-LLZO interface is critical to improving the overall battery performance and longevity. Moreover, efforts should be directed toward scaling up the production of LLZO-based sheet electrolytes. NASICON is an ISE class with various benefits, including good mechanical properties, non-toxic, air stability, and wide electrochemical window. NASICON's further development is going to focus on low-temperature synthesis and

controlling impurities concentration during synthesis, as well as addressing poor interface contact with the electrode materials. Sulfide has the highest conductivity among ISEs and good mechanical deformability; however, sulfides are vulnerable to atmospheric moisture, leading to hydrolysis and a consequent reduction in conductivity. Antiperovskite is a new class of ISE that warrants additional investigation due to its good mechanical properties and ease of production, but it must be handled carefully since it is less stable to moisture and has poor ionic conductivity when compared to other ISEs. Finally, the hydroborate class has considerable promise due to its straightforward preparation, good chemical stability, and wide potential window for high-voltage electrodes. To boost the ionic conductivity, the chemical structure of the hydroborate group should be improved. ISE innovation has advanced rapidly providing promising features for high-performance ASSBs.

Review

Fig. 24 compares five classes of ISEs as considerations for the sustainable development of high-performance ASSBs.

Based on our analysis, the strategies, and recommendations to encourage the widespread application of ISEs and ASSBs are described as follows and presented in Fig. 25:

- (1) The incorporation of multiphase ISE into ASSBs. The ISEs group has the potential to function as an electrolyte, an interface layer, and a catholyte. Considering that there is a lack of knowledge about the combination of two different types of ISEs in the same class or different classes, it is necessary to conduct further investigation on this topic. When it comes to the application of ISEs in three different positions in ASSBs, we demonstrate 64 different combination arrangements between ISEs that can serve as a guide. These arrangements include (a) ISE as a catholyte-ISE as an electrolyte; (b) ISE as an electrolyte-ISE as an interface layer; and (c) utilization of the hybrid inorganic solid electrolytes.
- (2) According to point No. 1, it is critical to investigate the impact of hybrid inorganic solid-state electrolyte thickness on ASSB device performance. In the case of hybrid ISEs, we divide them into two concepts: first, hybrid ISEs are manufactured by mixing two types of ISE in a specific ratio until they reach a homogeneous equilibrium and are implemented as a unified ISE electrolyte; and second, hybrid ISEs manufactured by stacking two layers of different ISEs while paying attention to their respective thickness ratios. We will focus on the second point in this section; the use of this concept in ASSBs necessitates further research, particularly when stacking the combination into several thin layers.
- (3) Continuing from point No. 2, we are interested in the influence of the thickness of each component layer in an ASSBs system consisting of a composite cathode, a cathode interface layer, a solid electrolyte, an anode interface layer, and an anode. Developing thin-film batteries with a high energy density offers a huge opportunity. An interface layer is an alternate option for solid electrolytes that are reactive towards the anode and cathode in order to minimize side reactions. Furthermore, the solid electrolyte's reactivity on both sides varies, necessitating research into regulating the thickness of the interface layer. The application of interface layers at insufficient levels is likely to have a negative impact on ASSB performance and production costs.
- (4) Develop ISE derivative compounds from the present ISE group. Each class of ISE has benefits and drawbacks that can be attributed to the type and coordination of cations and anions, respectively. As a result, using modern computational methods to design the chemical structure is critical for developing nextgeneration ISE electrolytes.
- (5) Utilization of the encapsulation concept for solid electrolytes with poor air stability. How does this concept perform for ASSBs? Sulfide, antiperovskite, and other solid electrolytes are covered with encapsulants that exhibit strong ionic conductivity and low electron conductivity. The aim is to simplify the preparation and incorporation of electrolytes into ASSBs devices. The encapsulant (coating) will melt after a certain period of time when the encapsulated electrolyte is incorporated into the ASSB device, therefore releasing ISEs. The

molten encapsulant will fill the gaps between ISE particles concurrently to improve the interaction between them. At the electrolyte-electrode interface, the molten encapsulant also generates a protective layer to control interfacial side reactions. Melted encapsulant is supposed to be an electronic insulator and an ionic conductor to improve ionic transport and suppress dendrite growth.

- (6) Utilization of modified textured surfaces. Enhancing the contact area between two solid surfaces will enhance the transfer of mobile ions. The sophistication of computing systems can be utilized to design textured surfaces for electrodes and solid electrolytes. The touch-plane concept is derived from the mechanism of enzymes interacting with biological substrates via the "lock and key" principle. Enzymes exhibit the best performance when they encounter a substrate surface that precisely fits their structure. By using this principle, the contact surfaces of the two substances, the electrode, and solid electrolyte, are deliberately modified to have a textured structure that is coherent and compatible with each other. Furthermore, one of the surfaces is coated with a substance that enhances the adhesion between the two surfaces and serves as an interface laver.
- (7) Comparative analysis of the effectiveness of developing hybrid ISEs vs. using electrolyte additives. Both exhibit the same objectives for ISEs, including reducing sintering temperature, inhibiting impurity growth, enhancing grain connectivity, decreasing grain boundary resistance, improving ionic conductivity, increasing density, rectifying mechanical issues, and enhancing chemical and moisture stability. Materials in ISE classes (NASICON, halide, sulfide, antiperovskite, hydroborate, etc.) and electrolyte additives such as oxides, nitrides, carbides, and solid polymer electrolytes can be alternatives and tailored to match the specific substrate properties and affordability for fabricating high-performance ISEs.

#### List of formulas

Cu-doped NZSP	$Na_{3.12}Zr_{1.94}Cu_{0.06}Si_{2}PO_{12}$
LAGP	$Li_{1.5}Al_{0.5}Ge_{1.5}(PO_4)_3$
LATP	$\text{Li}_{1+x}\text{Al}_x\text{Ti}_{2-x}(\text{PO}_4)_3$
LCO	${\rm LiCoO_2}$
LFP	${ m LiFePO_4}$
LGLZO	$\mathrm{Li}_{6.25}\mathrm{Ga}_{0.25}\mathrm{La}_{3}\mathrm{Zr}_{2}\mathrm{O}_{12}$
LGPS	$\mathrm{Li_{10}GeP_{2}S_{12}}$
LIC	Li <sub>3</sub> InCl <sub>6</sub>
LLZC	$\mathrm{Li}_{2.5}\mathrm{Lu}_{0.5}\mathrm{Zr}_{0.5}\mathrm{Cl}_{6}$
LLZO	$\text{Li}_7\text{La}_3\text{Zr}_2\text{O}_{12}$
LLZCO	$\text{Li}_{7.5}\text{La}_{3}\text{Zr}_{1.5}\text{Co}_{0.5}\text{O}_{12}$
LLZTO	${\rm Li_{6.5}La_{3}Zr_{1.5}Ta_{0.5}O_{12}}$
LLZWO	${\rm Li_{6.3}La_3Zr_{1.65}W_{0.35}O_{12}}$
LNMO	$\mathrm{LiNi}_{0.5}\mathrm{Mn}_{1.5}\mathrm{O}_4$
LSZC	$\text{Li}_{2.375}\text{Sc}_{0.375}\text{Zr}_{0.625}\text{Cl}_{6}$
LSiPSCl	$\text{Li}_{10.02}\text{Si}_{1.47}\text{P}_{1.56}\text{S}_{11.7}\text{Cl}_{0.3}$
LTO	$\text{Li}_4\text{Ti}_5\text{O}_{12}$
LYB	$\mathrm{Li_{3}YBr_{6}}$
LYC	$\mathrm{Li}_{3}\mathrm{YCl}_{6}$

AB

Mg-doped NZSP-	$Na_{3.456}Zr_{1.872}Mg_{0.128}Si_{2.2}P_{0.8}O_{12}$
0.128	
NCM622	$\text{LiNi}_{0.6}\text{Co}_{0.2}\text{Mn}_{0.2}\text{O}_2$
NCM811	$\text{LiNi}_{0.8}\text{Co}_{0.1}\text{Mn}_{0.1}\text{O}_2$
NCM83	$LiNi_{0.83}Co_{0.11}Mn_{0.06}O_2$
NCM85	$LiNi_{0.85}Co_{0.1}Mn_{0.05}O_2$
NCM88	$LiNi_{0.88}Co_{0.09}Mn_{0.03}O_2$
NCM90	$LiNi_{0.9}Co_{0.05}Mn_{0.05}O_2$
NLZSP	$Na_{3.3}La_{0.3}Zr_{1.7}Si_2PO_{12}$
NMNFO	$Na_{0.85}Mn_{0.5}Ni_{0.4}Fe_{0.1}O_2$
NZSP	$Na_3Zr_2Si_2PO_{12}$
NZSP-ATO	Na <sub>3</sub> Zr <sub>2</sub> Si <sub>2</sub> PO <sub>12</sub> with 5 wt% antimony ti
	oxide
NLZSP-NBO <sub>3</sub>	Na <sub>3.3</sub> La <sub>0.3</sub> Zr <sub>1.7</sub> Si <sub>2</sub> PO <sub>12</sub> with 3 wt%
	$Na_2B_4O_7$
NVPF	$Na_3V_2(PO_4)_2F_3$
SbCl <sub>3</sub> -doped LGPS	$Li_{9.88}GeP_{1.96}Sb_{0.04}S_{11.88}Cl_{0.12}$
Te-doped LAGP	$\text{Li}_{1.5}\text{Al}_{0.5}\text{Te}_{0.06}\text{Ge}_{1.44}(\text{PO}_4)_3$

#### List of abbreviations

acetylene black

ACN	acetonitrile
AGG	abnormal grain growth
AIMD	Ab initio molecular dynamics
ALD	atomic layer deposition
ASR	area-specific resistance
ASSBs	all-solid-state batteries
ASSLIBs	all-solid-state lithium-ion batteries
ASSSIBs	all-solid-state sodium-ion batteries
BVEL	bond valence energy landscape
BVSE	bond-valence site energy
CAD	computer-aided design
CAM	cathode active materials
CB	carbon black
CCP	cubic close-packed
CEI	cathode interphase layer
CI-NEB	climbing-image nudged elastic band
CNT	carbon nanotube
CSP	cold sintering process

CVD chemical vapor deposition DDSE dynamic database of solid-state electrolyte DFT density functional theory **DME** 1,2-dimethoxyethane activation energy Ea **EDS** energy-dispersive X-ray spectroscopy EIS electrochemical impedance spectroscopy **GITT** galvanostatic intermittent titration technique high-coordination frameworks HCF HCF-Sm high coordination frameworks-Na<sub>0.75</sub>Sm<sub>1.75</sub>Cl<sub>6</sub> HCF-La high coordination frameworks-Na<sub>0.75</sub>La<sub>1.75</sub>Cl<sub>6</sub>

**HCP** hexagonal close-packed high-pressure low-temperature **HPLT** hard and soft acids and bases **HSAB ISEs** inorganic solid electrolytes LCF low-coordination frameworks

LCF-Ta low coordination frameworks-NaTaCl<sub>6</sub>

LCF-Zr	low coordination frameworks-Na <sub>2</sub> ZrCl <sub>6</sub>
LCF-Hf	low coordination frameworks-Na <sub>2</sub> HfCl <sub>6</sub>
LIBs	lithium-ion batteries
LiRAP	Li-rich antiperovskite
LISICON	lithium superionic conductor
LPS	lithium thiophosphates
MAS NMR	magic angle spinning nuclear magnetic

MEM maximum entropy method MSD mean squared displacement **MWCNT** multi-walled carbon nanotube NASICON natrium superionic conductor

resonance

NMP N-methylpyrrolidone NPD neutron powder diffraction PDF pair distribution function

p(MMA/nBA) poly(methyl methacrylate)/n-butyl acrylate

grain boundary resistance RTroom temperature RTP rapid thermal processing SEI solid interphase layer **SEM** scanning electron microscopy sodium-ion batteries **SIBs** 

**SSBs** solid-state batteries SPS spark plasma sintering THF tetrahydrofuran

UHS ultrafast high-temperature sintering

**XRD** X-ray diffraction

#### Data availability

No primary research results, software or code have been included and no new data were generated or analysed as part of this review.

#### Conflicts of interest

The authors declare no competing financial interest.

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