REVIEW ARTICLE



Solid-State Electrolytes for Lithium-Ion Batteries: Fundamentals, Challenges and Perspectives

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Abstract

With the rapid popularization and development of lithium-ion batteries, associated safety issues caused by the use of flammable organic electrolytes have drawn increasing attention. To address this, solid-state electrolytes have become the focus of research for both scientific and industrial communities due to high safety and energy density. Despite these promising prospects, however, solid-state electrolytes face several formidable obstacles that hinder commercialization, including insufficient lithium-ion conduction and surge transfer impedance at the interface between solid-state electrolytes and electrodes. Based on this, this review will provide an introduction into typical lithium-ion conductors involving inorganic, organic and inorganic–organic hybrid electrolytes as well as the mechanisms of lithium-ion conduction and corresponding factors affecting performance. Furthermore, this review will comprehensively discuss emerging and advanced characterization techniques and propose underlying strategies to enhance ionic conduction along with future development trends.

Keywords Solid-state electrolyte · Lithium-ion transport · Structure tuning · Interface engineering

1 Introduction

The large-scale application of electric vehicles (EVs) is considered to be a viable method to reduce CO_2 emissions and alleviate greenhouse effects, and in the past few years, the use of EVs has become more accepted by the general

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public. However, with the rapid adoption of EVs, safety issues as well as other associated issues in the use of lithiumion batteries (LIBs) as a power source are becoming more pronounced and will require further research and development [1-3]. In general, a commercial LIB is composed of an anode, a non-aqueous electrolyte, a separator and a cathode and the use of non-aqueous electrolytes possesses serious safety issues due to the flammability and easy ignition of the organic solvents used in these electrolytes (i.e., propylene carbonate, ethylene carbonate, ethylene carbonate) [4-6]. In addition, the growth of Li dendrites in non-aqueous LIBs is unavoidable and can increase the possibility of short-circuiting and limit application in EVs [7–9]. Furthermore, these issues can further be exacerbated in cases of improper use, leading to fires and possible explosions. Moreover, the use of conventional non-aqueous electrolytes also limits the practicality of high-voltage cathode materials to achieve high specific energy batteries [10-12]. Based on these negative attributes, the use of nonflammable solid-state electrolytes to replace non-aqueous electrolytes is considered to be an appealing strategy [13, 14]. Solid-state electrolytes generally include inorganic Li-ion conductors, polymer electrolytes and organic-inorganic hybrid composites [15], and several essential requirements are necessary for their application in LIBs. These include high ionic conductivity (> 10^{-4} S

 cm^{-1}), negligible electronic conductivity, feasible electrochemical stability windows as well as favorable transfer impedance between interfaces [16–18]. Unfortunately, these requirements are difficult to achieve individually or together. For example, although single -ion conductors can avoid concentration polarization effects, their practical application is plagued by low ionic conductivity at room temperature and poor interfacial contact with electrodes. Moreover, the stability of solid-state electrolytes against anode and cathode materials also needs to be taken into consideration. And overall, numerous challenges need to be addressed before commercialization and will require intensive investigation into the fundamentals of solid-state electrolytes. Based on this, extensive research has been carried out recently among industrial and scientific communities to speed up innovation in this field [19-22], and although achievements have been made, the large-scale application of solid-state electrolytes still faces numerous problems ranging from fundamental understanding to industrial manufacturing [23–25]. For example, the mechanism of Li-ion transport remains controversial and has been fiercely debated by researchers for decades, especially around issues associated with Li-ion transport across the interface of anodes, solid-state electrolytes and cathodes, and is a major bottleneck in the practical application of all-solid-state batteries [26]. In addition, aside from the development of advanced characterization techniques, more visual results and direct evidence are required to promote the understanding of fundamentals [27-29]. Based on all of this and to guide future development, this review into solid-state electrolytes will provide the fundamentals of Li-ion transport as well as promising enhancement strategies along with future perspectives and proposed research directions.

2 Overviews of Typical Li-Ion Conductors

2.1 Inorganic Li-Ion Conductors

2.1.1 Oxide-Type Li-Ion Conductors

2.1.1.1 Perovskite Conductors Perovskite (ABO₃)-type Liion conductors and structurally related variants have recently attracted extensive attention based on optimal bulk Li⁺ conductivities among various solid electrolytes with numerous studies reporting varying ionic conductivities through the replacement of A and B sites with different ions in which ionic conductivities can reach 10^{-3} S cm⁻¹ ($x \approx 0.11$) after optimization. The general formula of perovskite-type lithium lanthanum titanate solid-state electrolytes can be expressed as Li_{3x}La_{(2/3)-x}TiO₃ (LLTO) with the value of *x* ranging from 0.07 to 0.13 and the activation energy ranging from 0.3 to 0.4 eV (Fig. 1). Here, the high ionic conductiv-

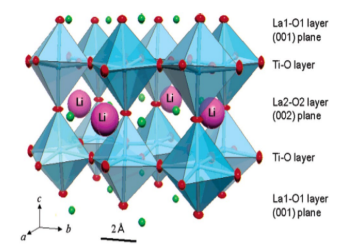


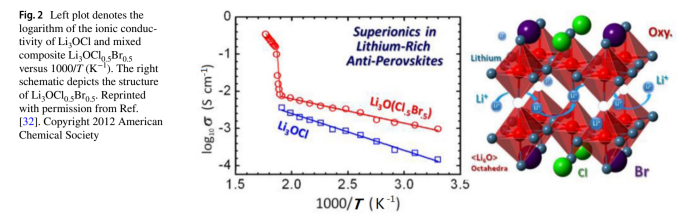
Fig. 1 Crystal structure of $La_{0.62}Li_{0.16}TiO_{3.0}$. Pink, green and blue spheres represent Li, La and Ti atoms, respectively, red ellipsoids represent oxygen ions, and TiO₆ octahedrons are depicted as blue squares. Reprinted with permission from Ref. [31]. Copyright 2005 American Chemical Society

ity of LLTOs is a result of the large concentration of A-site vacancies with researchers reporting that cation deficiencies at A sites as formed by 4 adjacent TiO_6 octahedra can facilitate the migration of Li cations through bottlenecks [30].

In general, different synthesis methods and heterogeneous lattice vacancies can result in different perovskite-like compounds with diverse crystal structures, including cubic, hexagonal, orthorhombic and tetragonal. And to advance all-solid-state electrolytes, it is essential to probe the ionic conduction mechanism and crystal structure of LLTOs. For example, Bohnke et al. [30] confirmed that the ionic conduction mechanism of LLTOs incorporated the tilting and/or rotation of BO₆ octahedra as promoted by either rising temperatures or increasing vacancies in the structure. Yashima et al. [31] also studied the migration path of the Li-ion conductor La_{0.62}Li_{0.16}TiO₃ (Fig. 1) and found that Li cations can migrate from the 2c site on the (002) La-deficient layer to the 2c-4f-2c or 2c-2d-2c tie line at 77 K to enhance conductivity.

Overall, perovskite-type conductors are considered to be promising candidates as all-solid-state electrolytes based on their appropriate electrochemical windows and acceptable ionic conductivities; however, the intercalation of Li-ions and limited electrodes greatly restrict practical application.

2.1.1.2 Anti-Perovskite Conductors In 2012, Zhao et al. [32] proposed a new strain of anti-perovskites (LiRAPs) that were "electronically inverted" as compared with standard perovskite-type electrolytes and reported that this novel class of electrolytes can be referred to as anti-perovskites because although they share similar structures with standard perovskite-type electrolytes, the ions of the anti-perovskites in the corresponding lattice sites can undergo electronic



inversion. For example, A (monovalent) and B (divalent) can serve as metallic cations in typical ABX₃ perovskite structures with strongly electronegative C anions, whereas A, B and X all possess inverted charges in LiRAP. In addition, B sites located in the octahedral center of LiRAP are occupied by chalcogens^{2–} (O^{2–}, S^{2–}), and any halogens[–] (F[–], Cl[–], Br[–], I[–]) can act as dodecahedral centers (A sites). Here, these researchers reported that the ionic conductivity of LiRAP can reach greater than 10^{–3} S cm^{–1} at ambient temperatures with activation energy of 0.2–0.3 eV and that these LiRAPs possessed good stability even under high temperatures.

Moreover, researchers have reported that chemical substitution can facilely manipulate Li⁺ conduction in antiperovskites. For example, Zhang et al. [33] reported that dodecahedral sites can be supplanted by larger anions (Br⁻, I⁻) to allow for the replacement of Li with divalent cations (Mg²⁺, Ca²⁺, Ba²⁺) at the Li₆O octahedral center and that the depletion of LiA (samples: Li_{2.95}OCl_{0.95}, Li_{2.90}OCl_{0.90}, Li_{2.85}OCl_{0.85} and Li_{2.80}OCl_{0.80}) can optimize superionic conductivity, in which the conductivity of Li₃OCl_{0.5}Br_{0.5} at room temperature can jump to 6.05×10^{-3} S cm⁻¹. In addition, thermal treatments (i.e., annealing, pressing, heating/ cooling) can also affect conductivity in which the ionic conductivity of Li₃OCl, Li₃OBr and Li₃OCl_{0.5}Br_{0.5} (Fig. 2) at ambient temperatures can be significantly enhanced by 2 orders of magnitude after annealed in vacuum for 24*h* [33].

The mechanism of Li^+ transport in LiRAP has also been investigated by Zhang et al. [33], in which phase diagrams and electronic structure calculations proved that antiperovskites (Li₃OCl, Li₃OBr and their mixed composite Li₃OCl_{0.5}Br_{0.5}) were thermodynamically metastable but electrochemically stable. Furthermore, ab initio molecular dynamic simulations were conducted and revealed that increased structural disorders and Li vacancies can lead to enhanced Li⁺ diffusion. Despite these results, however, the mechanism of Li-rich anti-perovskites remains a muchdebated topic and further studies are needed to address this issue. As another anti-perovskite, $\text{Li}_{3-2x}M_x$ HalO (M stands for divalent cations such as Mg²⁺, Ca²⁺ or Ba²⁺, Hal stands for halides such as Cl⁻, I⁻ or a mixture) was first reported by Braga et al. [34] as a novel class of glassy electrolytes based on LiRAP that can exhibit ultra-fast ionic conduction (25 mS cm⁻¹ at 25 °C). And based on the merits of these novel glassy electrolytes (lightness, nonflammability, nonpolluting, thermal and electrochemical stability, etc.), they are expected to be promising in future applications in which higher valence dopants can result in higher conductivities.

2.1.1.3 NASICON **Conductors** As another oxygen type of Li conductors, NASICON-type conductors $(Na_{1+r}Zr_2P_{3-r}Si_rO_{12})$ are promising fast ion conductors with high conductivity that are derived from $NaZr_2(PO_4)_3$ after the replacement of P with Si due to the poor ionic conductivity of P. Here, researchers have reported that the enhanced conductivity of $Na_{1+x}Zr_2P_{3-x}Si_xO_{12}$ can be attributed to 3 reasons, including the increased mobility of Na⁺ ions in the structure, the higher density of the sintered pellets and the more favorable environment for Na⁺ ion migration through the 3D structure with enlarged tunnel sizes [35]. Of these NASICON-type conductors, iso-structural $LiM_2(PO_4)_3$ (M = Zr, Ti, Hf, Ge or Sn) was first reported in 1977 in which the skeletons consisting of MO₆ octahedra and PO₄ tetrahedra sharing oxygen atoms (Fig. 3) [36]. These $LiM_2(PO_4)_3$ series can further be divided into $LiM_2(PO_4)_3$ (M = Ti, Ge) with rhombohedral symmetry and $LiM_2(PO_4)_3$ (M = Zr, Hf, or Sn) with a triclinic phase and lower symmetry. Here, alkaline ions in the rhombohedral phase are likely to occupy two crystallographic sites in which M1 sites are surrounded by 6 oxygens and M2 sites are positioned between 2 adjacent M1 positions with tenfold oxygen coordination. As for the triclinic phase, Li cations are driven by structural distortions in a fourfold coordination at the intermediate M12 sites. In addition, researchers have reported that the conductive performance of LiTi₂(PO₄)₃ outshines other $LiM_2(PO_4)_3$ (M = Zr, Ge or Hf) materials because of its optimal lattice size for conducting Li⁺ ions. Moreover,

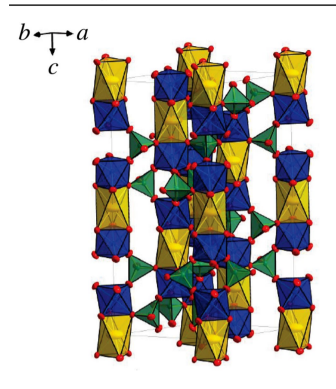


Fig. 3 Unit cell of $\text{LiTi}_2(\text{PO}_4)_3$. Li⁺, Ti⁴⁺ and P⁵⁺ occupy yellow octahedra, blue octahedra and green tetrahedra, respectively. Reprinted with permission from Ref. [36]. Copyright 2017 American Chemical Society

researchers have also reported that trivalent cations such as Al³⁺, Sc³⁺, Ga³⁺, Fe³⁺, In³⁺ and Cr³⁺ replacing partial Ti⁴⁺ cations in $Li_{1+r}R_rTi_{2-r}(PO_4)_3$ can enhance conductivity in which Li_{1.3}Al_{0.3}Ti_{1.7}(PO₄)₃ (LATP) can gain an enhanced ionic conductivity of 7×10^{-4} S cm⁻¹. Furthermore, the ionic radius of M4+ cations can influence the lattice constants of $LiM_2(PO_4)_3$ (M = Ge, Hf, or Zr) as well to further contribute to the tunnel size for Li^+ diffusion (Fig. 4) [37]. Augmenting densification, increasing the mobile ion content in the structure and reducing activation energy derived from the incorporation of trivalent ions can also enhance ionic conductivity in NASICON-type conductors [35]. In another study, Arbi et al. [37] also investigated the influence of cation miscibility in the optimization of the transport properties of $\text{Li}_{1+x}\text{Ti}_{2-x}\text{Sc}_x(\text{PO}_4)_3$ ($0 \le x \le 0.5$) series and reported that in Sc-poor samples, NASICON phases with rhombohedral symmetry were characterized by using XRD patterns whereas in Sc-rich structures, the existence of secondary TiO₂, LiScO₂ and LiScP₂O₇ phases was confirmed, which were found to suppress the incorporation of Li in the NASICON phase. Furthermore, Aono et al. [35] proposed another strategy to enhance conductivity by introducing secondary Li compounds including Li₂O, Li₃PO₄ and Li₃BO₃ to serve as fluxes to promote crystallization and form conductive components at the grain boundary in which

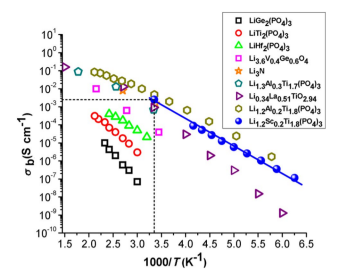


Fig. 4 Ionic conductivity of NASICON-type Li-ion solid electrolytes. Reprinted with permission from Ref. [37]. Copyright 2017 American Chemical Society

the addition of 20% Li_2O into $LiTi_2(PO_4)_3$ leads to a constant conductivity of 5×10^{-4} S cm⁻¹.

2.1.1.4 Garnet-Type Conductors In 2003, a novel family of Li-ion electrolytes possessing a garnet-like structure and a nominal composition of $\text{Li}_5\text{La}_3\text{M}_2\text{O}_{12}$ (M=Nb, Ta) was proposed by Thangadurai et al. [38] that has since attracted wide interest due to unique crystal structures, high conductivities and wide electrochemical windows, in which compared with LiPON, $\text{Li}_9\text{AlSiO}_8$ and Li-β-alumina, the ionic conductivity of these garnet-type conductors was higher and the bulk conductivity of both niobium and tantalum compounds could reach a similar order of magnitude (~10⁻⁶ S cm⁻¹ at 25 °C) with activation energy of 0.43 eV and 0.56 eV, respectively. In addition, the Ta members ($\text{Li}_5\text{La}_3\text{Ta}_2\text{O}_{12}$) were also chemically stable in molten Li and moisture.

Of these garnet-type conductors, orthosilicate garnets possess the formula $A_3^{II}B_2^{III}(SiO_4)_3$ in which A and B stand for 8- and 6-coordinated cation sites. And in this structure, separated SiO₄ tetrahedra are bonded to each other through ionic bonds between interstitial B cations and by replacing silicon with other elements, various complex oxides have been revealed to crystallize in the garnet-like conductors with a general formula of $A_3B_5O_{12}$ (A=Ca, Mg, Y or Ln=La, or rare-earth elements; B=Al, Fe, Ga, Ge, Mn, Ni, V), resulting in a 3D structure made up of BO₄ tetrahedra and BO₆ octahedra sharing corners with each other. For example, in the crystal structure of Li₅La₃M₂O₁₂ (M=Nb, Ta) (Fig. 5), 8- and 6- coordination sites are occupied by La and Ta ions with Li-ions on both the octahedral and tetrahedral sites as confirmed by neutron diffraction investigations

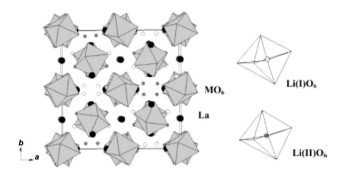


Fig. 5 Crystal structure of $Li_5La_3M_2O_{12}(M=Nb, Ta)$. Reprinted with permission from Ref. [42]. Copyright 2005 John Wiley and Sons

[38] in which $Li(I)O_6$ was much more distorted than Li(II) O_6 . In addition, researchers have also reported that 6 LiO₆ octahedra and 2 vacant sites of Li-ions surrounding MO₆ octahedra can result in high Li-ion conduction [39]. Thangadurai et al. [40] also enhanced the ionic conductivity of garnet-like Li₅La₃Nb₂O₁₂ through the partial substitution of K⁺ for La³⁺ and In³⁺ for Nb⁵⁺. These researchers also reported that high sintering temperatures can decrease grain boundary resistances and create large grain size particles to fashion microstructures and enhance conductivity. Moreover, Thangadurai et al. [41, 42] in subsequent studies generated original garnet-like conductors through the replacement of La with alkaline earth elements and reported that ionic conductivity reduced as the ionic radius of alkaline earth ions decreased in which among various conductors synthesized, Ba compound $Li_6BaLa_2Ta_2O_{12}$ showed optimal ionic conduction performances $(4 \times 10^{-5} \text{ S cm}^{-1} \text{ at room temperature})$ with activation energy of 0.40 eV). These researchers also reported that the ionic conductivity of parent compound Li₅La₃Nb₂O₁₂ can be enhanced by Y- and Li-codoping to create a novel Li-rich Li_{5+2x}La₃Nb_{2-x}Y_xO₁₂ structure in

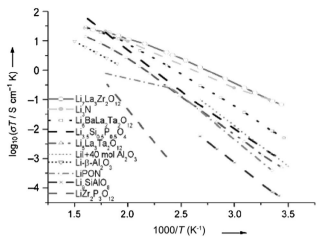
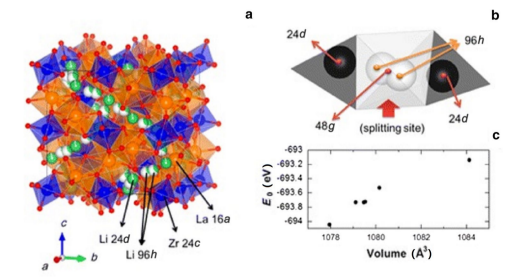


Fig. 7 Comparison of the total conductivity of Li₇La₃Zr₂O₁₂ and other Li-ion conductors. Reprinted with permission from Ref. [44]. Copyright 2007 John Wiley and Sons

which trivalent ions such as Y^{3+} surpassed other dopants such as M^{2+} (alkaline earth ions), In^{3+} and Zr^{4+} [43].

Cubic Li₇La₃Zr₂O₁₂ (LLZO) is another garnet-like structure (Fig. 6) that was first demonstrated by Murugan et al. [44] and is a promising all-solid-state ion conductor that possesses a higher total conductivity (3×10^{-4} S cm⁻¹) than that of any previously reported Li garnet-like materials (Fig. 7). Here, researchers report that in cubic LLZO, Li-ions are distributed over the octahedral 48*g*, tetrahedral 24*d* Li(1) and 96*h* Li(2) sites [45]. Another tetragonal-type LLZO polymorph with low-temperature tetragonal symmetry was also reported; however, the well-ordered arrangement of Li-ions at the octahedral 16*f* and 32*g* sites and tetrahedral 8*a* sites resulted in a much lower conductivity value of 10^{-6} S cm⁻¹ [46, 47]. As a result, many studies have been devoted to the stabilization of cubic structures in garnet-type conductors

Fig. 6 Crystal structure of cubic LLZO. Li-ions distributed over the octahedral 48*g*, tetrahedral 24*d* Li(1) and 96*h* Li(2) sites. Reprinted with permission from Ref. [45]. Copyright 2013 American Chemical Society



either through the increase in Li vacancy concentrations or the reduction in the Li content. For example, Rangasamy et al. [48] demonstrated that the addition of Al as a substitute for Li can generate more Li vacancies and investigated the optimal ratio of Al to Li to form desirable LLZOs.

The Li-ion diffusion mechanism of garnet-type conductors is also controversial; however, better understandings of Li⁺ transport mechanisms require advanced characterization techniques to determine the exact location of Li-ions. For example, Thangadurai et al. [49] utilized the bond valence sum method to explore the characteristic structure of Li₅La₃M₂O₁₂ and confirmed possible ion transport pathways through a minimization procedure of bond valence mismatch in which the obtained results revealed that Liions exclusively occupied octahedral sites. However, Cussen et al. [50] later disputed this finding and proposed that Li-ions in the framework of Li₅La₃M₂O₁₂ (M=Ta, Nb) were located in both octahedral and tetrahedral sites in which Li⁺ in the octahedra sites was a mobile species whereas Li⁺ in the tetrahedral sites was considered to be trapped and immobile. This conclusion was also reached by Wüllen et al. [51] employing advanced NMR strategies. Furthermore, researchers have also reported the significance of final annealing temperatures because the mobility of Li-ions is sensitive to Li-ions migration. For example, annealing with high temperatures can facilitate more Li-ions in octahedral sites to improve ionic conductivity. As for the migration mechanism of LLZOs, this is typically determined by the imposed restrictions of occupied site-to-site interatomic isolation allowing only 48g/96h-24d-48g/96h and the temporal stay of Li^+ at the 24d site which can promote ion mobility and reconfigure surrounding Li neighbors [45]. In addition, simulation methods are useful tools to gain deeper understandings of migration mechanisms in which Xu et al. [52] use the nudged elastic band (NEB) method to find the minimum energy path for Li diffusion in $Li_{3+x}La_3M_2O_{12}(x=0,$ 2, 4).

2.1.2 Sulfide-Type Li-ion Conductors

2.1.2.1 Thio-LISICONs The thio-LiCICONs family has been proposed as a sulfide-type Li-ion conductor due to its enhanced electrochemical performance as compared with LISICON family conductors and can be described through the nominal composition of $\text{Li}_{4-x}\text{M}_{1-y}\text{M}_{y}\text{S}_{4}$ (M=Si, Ge, and M'=P, Al, Zn, Ga) [53, 54]. And among these conductors, the ionic conduction of $\text{Li}_{4-x}\text{Ge}_{1-x}\text{P}_x\text{S}_4$ (2.2×10⁻³ S cm⁻¹ at 25 °C) showed superior performances than any other members. Researchers have also reported that ionic conduction performances are mainly dependent on the radius and polarizability of constituted ions in which the replacement of O by S can diminish interactions between Li⁺ in the sub-

lattice and increase the concentration of mobile Li⁺, which explains the higher conductivity of S-doped LISICONs as compared with their oxide analogues.

Kamaya et al. [55] also developed a novel crystalline sulfide Li-ion electrolyte (Li₁₀GeP₂S₁₂) with outstanding electrochemical properties (exceedingly high conductivity of 1.2×10^{-2} S cm⁻¹ and a wide potential window) and other advantages and reported that this material can be applied in LIBs with LiCoO₂ and Li-In alloy as the cathode and the anode, respectively, to achieve desirable performances. Here, the researchers determined the crystal skeleton of Li₁₀GeP₂S₁₂ using XRD and synchrotron radiation X-ray data associated with the ab initio method to reveal a 3D framework with LiS_6 octahedra and $(Ge_{0.5}P_{0.5})S_4$ tetrahedra forming 1D chains connected by a common edge, which are connected at the corners with PS₄ tetrahedra. In addition, neutron Rietveld analysis was used to further confirm Li occupation and the Li content. Du et al. [56] also reported that tetrahedrally coordinated Li1 (16h) and Li3 (8f) sites were responsible for ionic conduction and formed a 1D tetrahedral chain along the c direction whereas the octahedrally coordinated Li2 sites were not available for Li⁺ diffusion, suggesting that Li⁺ diffusion paths along the *c*-axis were better for Li⁺ diffusion than that in the *ab* plane (reflecting only weak anisotropy of Li⁺ diffusion) with the thermodynamic stability being studied by using ab initio methods. Furthermore, the strong Coulombic connection between mobile ions can prompt Li-ion transport in a string-like manner in LGPS with low activation energy barriers and extremely high ionic conductivities [57].

2.1.2.2 LGPS Family Since being first proposed in 2011, $Li_{10}GeP_2S_{12}$ (LGPS) has attracted wide interest due to an incredible ionic conductivity that is comparable to organic liquid electrolytes (Fig. 8) [55]. However, the unstable interface of LGPS needs to be addressed [58] as well as the low abundance and high cost of Ge in the electrolyte. Based on this, the exploration of new LGPS-type materials is necessary to advance the application of LGPS conductors in which the LGPS family can be divided into Ge-free and Ge-doped electrolytes ($Li_{10}GeP_2S_{12}$ and Li_7GePS_8) [59].

To address the high cost of Ge, researchers have attempted to substitute Ge⁴⁺ with homologous Si⁴⁺ or Sn⁴⁺, resulting in materials such as Li₁₀SnP₂S₁₂ [60], Li_{10+δ}(Sn_ySi_{1-y})_{1+δ}P_{2-δ}S₁₂ [61], Li₁₁Si₂PS₁₂ [62], Li_{9.54}Si_{1.74}P_{1.44}S_{11.7}Cl_{0.3} [63]. And overall, researchers report that these Si- and Sn-doped materials exhibited lower conductivities than parent LGPS in which Si-doping, to some extent, can narrow tunnel sizes for Li migration and lower conductivity. As for the reduction in conductivity in Sn analogues, Zeier et al. [64] in their study clearly found that Sn⁴⁺ substitution can render the bottleneck of Li⁺ diffusion along

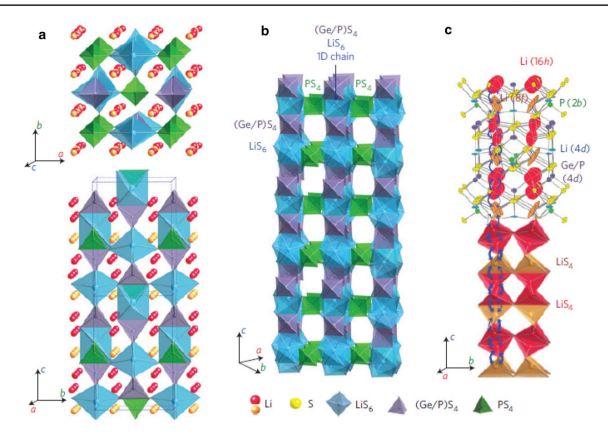
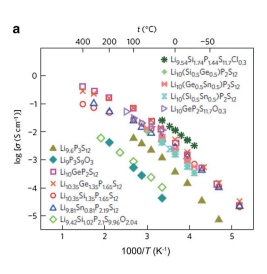


Fig.8 a Framework structure and Li-ion environment of $Li_{10}GeP_2S_{12}$. **b** Framework of $Li_{10}GeP_2S_{12}$. **c** Conduction pathway of Li-ions in $Li_{10}GeP_2S_{12}$. Reprinted with permission from Ref. [55]. Copyright 2011 Nature Publishing Group

the z-direction, which induces tougher activation barriers. And of these substituted LGPSs, Kato et al. [63] reported that Li_{9.54}Si_{1.74}P_{1.44}S_{11.7}Cl_{0.3} possessed an exceedingly high conductivity of up to 25 mS cm⁻¹, which was the highest ever reported for Li conductors (specific data demonstrated in Fig. 9a) and double that of the archetype LGPS [63]. Here, the researchers used the anisotropic thermal displacement of Li (Fig. 9b) and nuclear density distribution (Fig. 9c) to verify 3D Li migration channels with 1D channels along the c-axis and 2D channels in the ab plane and suggested that this 3D conduction pathway in the LGPS-type conductor can contribute to optimal conductivities. In addition, these researchers also reported good chemical stability of the LGPS-type conductor, especially at 100 °C, and attributed this to the lack of elemental diffusion and small interfacial resistance. Moreover, Li_{9.6}P₃S₁₂ also possesses a similar LGPS structure and has been reported to also possess excellent electrochemical stability and longer lifespans under extreme cell operation conditions. Recently, Sun et al. [61] also successfully obtained a Sn-Si derivative of LGPStype conductors $[Li_{10+\delta}(Sn_{v}Si_{1-v})_{1+\delta}P_{2-\delta}S_{12}]$ in the form of a Li₃PS₄-Li₄SnS₄-Li₄SiS₄ quasi-ternary system in which the researchers reported that by adjusting the ratio of Sn/Si and M^{4+} (Sn⁴⁺ and Si⁴⁺)/P⁵⁺ in the [M1/P1] (4*d*) site, the dual doping of the original LGPS with Sn and Si can improve ionic conduction, resulting in a $\text{Li}_{10.35}(\text{Sn}_{0.27}\text{Si}_{1.08})\text{P}_{1.65}\text{S}_{12}$ achieving an ionic conductivity of 1.1×10^{-2} S cm⁻¹ that is extremely close to that of the archetype LGPS (1.2×10^{-2} S cm⁻¹).

2.1.2.3 Argyrodites Argyrodites are characterized by high ionic conductivities and good cation mobility, and fast Liion conducting argyrodites such as Li_6PS_5X (X = Cl, Br, I) are representative members of this family. Here, the addition of halides into argyrodites can lead to conductivities in the range of 10^{-2} – 10^{-3} S cm⁻¹ in which Deiseroth et al. [65] reported that Li₆PS₅X can be formed by mixing Li₂S, P₂S₅ and LiX under 550 °C and that Li₆PS₅Cl and Li₆PS₅Br showed better conductivity than Li₆PS₅I. As for the crystal structure of Li₆PS₅X (Fig. 10a), the framework of the lattice is based on a cubic Laves topology and PS_4^{3-} anions are located in the middle of 4b sites whereas the remaining sulfur atoms take up 4a and 4c sites [66]. Here, despite the replacement of sulfur by halogens, the sulfur of PS_4^{3-} is not substituted and halogens take up 4aor 4c sites to display a face-centered structure in which the remaining tetrahedral interstices (24g and 48h Wyckoff sites) are occupied by Li-ions. In addition, position 24g



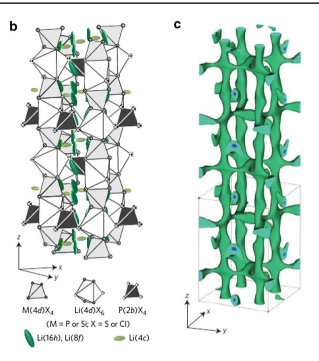


Fig.9 Ionic conductivity and framework of $Li_{9,54}Si_{1.74}P_{1.44}S_{11.7}Cl_{0.3}$. **a** Arrhenius conductivity plots for the LGPS family, $Li_{9,6}P_3S_{12}$ and $Li_{9,54}Si_{1.74}P_{1.44}S_{11.7}Cl_{0.3}$. **b** Framework structure of

can serve as the transition state for jumps between 2 48h sites and 12 48h sites can constitute a cage structure by surrounding each 4c site (Fig. 10b).

As for Li diffusion caused by these partially occupied positions, this can be attributed to 3 different jump processes, including next-neighbor jumps (48h-24g-48h), intra-cage jumps (48h-48h jumps within the cage) and inter-cage jumps (48h-48h jumps between cages) [66]. And among these processes, inter-cage jumps are considered to play a dominate role in macroscopic long-range ion transport whereas intra-cage jumps with low jump rates can constrain diffusion (Fig. 9c, d) [23, 67]. In addition, the replacement of sulfur with halogen can generate vacancies, which is associated with disorder on the S^{2-}/X^{-} sublattice and can enhance local Li-ion diffusion. For example, Deiseroth et al. [65] reported that Cl^{-1} and Br^{-1} can take up 4a and 4c sites in Li_6PS_5X to allow for the significant disordering of halogen ions, resulting in high conductivities for Li₆PS₅Cl and Li₆PS₅Br. Alternatively, I^{-1} can barely takes up 4*a* sites and I-containing argyrodites lacking the disorder to display higher activation barriers for conduction. In addition, Kraf et al. [66] reported the effect of lattice dynamics on ionic motion and concluded that the migration barrier of moving cations decreases with softer lattices.

2.1.2.4 Other New Thiophosphates Aside from the above thiophosphates, several novel sulfide-based Li-ion conduc-

 $Li_{9,54}Si_{1.74}P_{1.44}S_{11.7}Cl_{0.3}$. **c** Nuclear distributions of Li atoms in $Li_{9.54}Si_{1.74}P_{1.44}S_{11.7}Cl_{0.3}$ at ambient temperature. Reprinted with permission from Ref. [63]. Copyright 2016 Nature Publishing Group

tors with different features have also been recently reported in which $\text{Li}_4\text{PS}_4\text{I}$ as a superionic conductor is one representative example. Here, Sicolo et al. [68] synthesized $\text{Li}_4\text{PS}_4\text{I}$ using a solvent-based method in which $\text{Li}_4\text{PS}_4\text{I}$ presented a new structure that was composed of insular PS_4^{-3-} tetrahedra aligned by layers vertical to the *c*-axis and Li-ions were dispersed over 5 partially occupied sites and diffused in a 3D migration pathway. And according to topo-structural analyses, the Li-ion conductivity of the $\text{Li}_4\text{PS}_4\text{I}$ was ~ 1.2×10^{-4} S cm⁻¹ and based on recent calculations, a higher conductivity > 10^{-2} S cm⁻¹ is attainable [68].

Minafra et al. [69] also synthesized superionic Li-argyrodite $\text{Li}_{6+x}\text{P}_{1-x}\text{Si}_x\text{S}_5\text{Br}$ through aliovalent doping into argyrodites and through X-ray, neutron diffraction and electrochemical impedance spectroscopy, reported that the $\text{Li}_{6+x}\text{P}_{1-x}\text{Si}_x\text{S}_5\text{Br}$ exhibited both structural changes and a threefold enhancement in ionic conductivity as compared with non-doped argyrodites. Here, the researchers suggested that the expansion of unit cells can be explained by the substitution of P⁵⁺ with larger Si⁴⁺ and the inclusion of additional Li⁺, with the increasing Li content leading to increasing occupancy of Li⁺ on the transition states and therefore improved transport properties.

More recently, theoretical studies have also revealed that $Li_{1+2x}Zn_{1-x}PS_4$ (LZPS) as characterized by a body-centered-cubic structure can provide high Li-ion conductivity in which Richards et al. [70] employed AIMD and DFT to study the LZPS and reported that $Li_{1+2x}Zn_{1-x}PS_4$ with higher

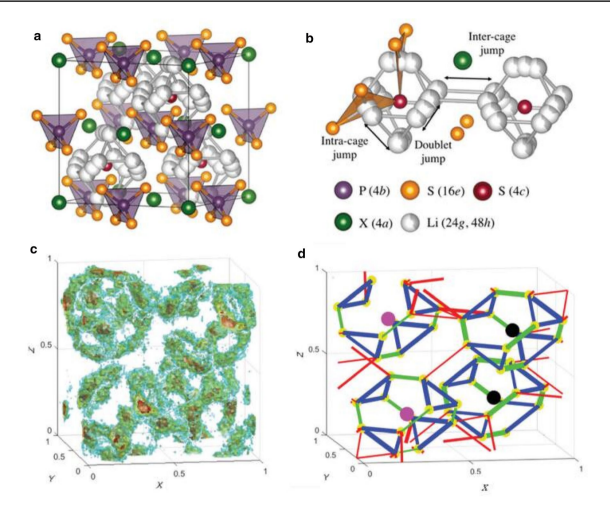
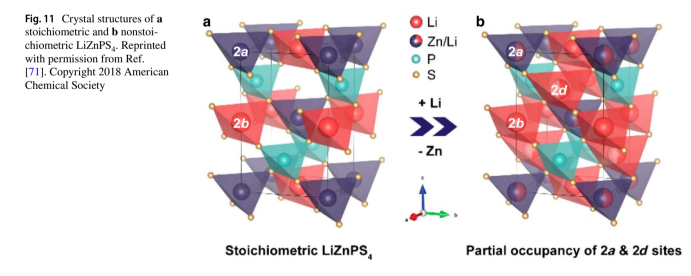


Fig. 10 a Crystal structure of $\text{Li}_6\text{PS}_5\text{X}$. b Li positions and possible jump routes. c MD simulation of Li-ion density in $\text{Li}_6\text{PS}_5\text{Cl}$. d MD results of jump statistics of $\text{Li}_6\text{PS}_5\text{Cl}$. Reprinted with permission from Ref. [23]. Copyright 2018 Royal Society of Chemistry



x can exhibit lower Li migration barriers. Here, the structure of the parent phase stoichiometric LiZnPS₄ (Fig. 11a) is composed of PS_4^{3-} and ZnS_4^{6-} tetrahedra in which Li takes

over 2*b* sites and Zn distributes over 2*a* sites. As for the crystal structure of $\text{Li}_{1+2x}\text{Zn}_{1-x}\text{PS}_4$ (x>0) (Fig. 11b), these researchers reported that excess Li-ions can share 2*a* sites

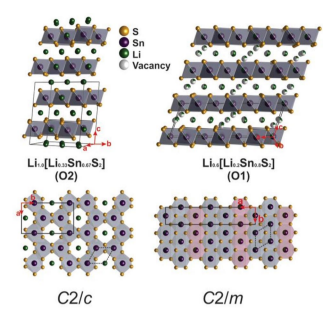


Fig. 12 Left: Crystal structure of $\text{Li}[\text{Li}_{0.33}\text{Sn}_{0.67}\text{S}_2]$ with a full Sn/Li order. Right: crystal structure of $\text{Li}_{0.6}[\text{Li}_{0.2}\text{Sn}_{0.8}\text{S}_2]$ with partially substituted Sn positions and an O1 type stacking. Reprinted with permission from Ref. [75]. Copyright 2016 Royal Society of Chemistry

with Zn and occupy 2*d* sites that are charge compensated due to the replacement of Zn²⁺ in the 2*a* sites by Li⁺. Kaup et al. [71] also synthesized a well-crystallized LZPS with a high *x* value of 0.35 and reported that the resulting Li_{1.7}Zn_{0.65}PS₄ can exhibit high conductivities of > 10⁻⁴ S cm⁻¹, which the researchers attributed to the Li occupation over 2*d* interstitial sites and decreased Li-ion migration barriers [72]. Despite these performances, however, the conductivities of these experimental Li_{1+2x}Zn_{1-x}PS₄ samples were all lower than those obtained through theoretical calculations, which may be due to the amorphous content and highly metastable defect composition. In addition, studies have revealed that Li_{2.5}Zn_{0.25}PS₄ with a glassy matrix can provide conductivities up to 8×10^{-4} S cm⁻¹.

2.1.2.5 Layered Sulfides Recently, a series of fast Li-ion conductors based on $\text{Li}_{3x}[\text{Li}_x\text{Sn}_{1-x}\text{S}_2]$ has been identified in which $\text{Li}[\text{Li}_{0.33}\text{Sn}_{0.67}\text{S}_2]$ (Li: Sn = 1: 2) and $\text{Li}_{0.6}[\text{Li}_{0.2}\text{Sn}_{0.8}\text{S}_2]$ (Li: Sn = 1: 4) are two outstanding examples that can exhibit different structures and ion conductivities. Here, both layered sulfides can be synthesized by using several approaches such as a facile wet chemistry approach for both [73], a high-temperature solid-state approach for Li_2SnS_3 (x = 0.33) [74] and other methods for $\text{Li}_2\text{Sn}_2\text{S}_5$ (x = 0.2) [75]. And in the case of the Li-rich prototype Li_2SnS_3 (in *C2/c*), Brant et al. [74] reported the crystallization of Li_2SnS_3 in a layered A₂BO₃ structure with a NaCl-like network (Fig. 12), in which the framework was built up by Li/Sn-ordered sulfide layers with Sn ions being arranged in a honeycomb fashion

 Table 1
 Ionic conductivities of representative crystalline inorganic solid Li-ion conductors at room temperature

Electrolyte	Ionic conductivity (S cm ⁻¹)	References
$\overline{\text{Li}_{3x}\text{La}_{(2/3)-x}\text{TiO}_3}$	10 ⁻³	[30, 31]
LiRAP	10^{-3}	[32]
LiM ₂ (PO ₄) ₃	$10^{-3} - 10^{-4}$	[35–37]
Li ₅ La ₃ M ₂ O ₁₂	$10^{-4} - 10^{-6}$	[38–50, 52]
$Li_{4-x}M_{1-y}M_{y}S_{4}$	$10^{-2} - 10^{-3}$	[53–57]
LGPS	$1.1 \times 10^{-2} - 10^{-4}$	[58, 59, 61–64]
Li ₆ PS ₅ X	$10^{-2} - 10^{-3}$	[23, 65–67]
$Li_{1+2x}Zn_{1-x}PS_4$	~10 ⁻⁴	[70–72]
$\operatorname{Li}_{3x}[\operatorname{Li}_{x}\operatorname{Sn}_{1-x}\operatorname{S}_{2}]$	~10 ⁻⁵	[73–75]

and Li-ions being sandwiched between the layers, showed an ionic conductivity of 1.5×10^{-5} S cm⁻¹ at room temperature and 1.6×10^{-3} S cm⁻¹ at 100 °C. As for the Li-depleted version, Holzmann et al. [75] reported that the structure of $Li_2Sn_2S_5$ (in C2/m) was similar to the parent compound Li₂SnS₃ whereas the Li content decreased to 60% in the interlayer gallery (Fig. 12). Here, these researchers reported that the Li depopulation resulted in an increase in the interlayer distance due to charge compensation by additional Sn in the honeycomb layers as well as the enhancement of Li mobility in the interlayer gallery, which contributed the most to high ionic conductivities. And based on impedance spectroscopy, these researchers reported that the activation energy of Li₂Sn₂S₅ was 0.17 eV, which was lower than that of Li_2SnS_3 (0.59 eV) and attributed this to the different Li diffusion pathways in which for Li₂Sn₂S₅, there was an addition diffusion pathway of hops between tetrahedral (T) and face-sharing octahedral (O) Li sites as compared with the pure O-O trajectories in Li₂Sn₂S₅. And as a result, Li₂Sn₂S₅ is the best Li solid electrolyte currently based on a bulk conductivity of $\sigma_{\rm NMR} = 9.3 \times 10^{-3} \, {\rm S \, cm^{-1}}$ by using data taken from field gradient (PFG) NMR which is consistent with a conductivity of 1.5×10^{-2} S cm⁻¹ as measured by impedance spectroscopy. Furthermore, in the case of singlecrystalline Li₂Sn₂S₅ samples, even higher conductivities are possible along the *ab* plane based on structural anisotropy (Table 1).

2.2 Polymer Electrolyte-Based Li-ion Conductors

All-solid-state electrolytes consist of inorganic solid electrolytes (ISEs), solid polymer electrolytes (SPEs) and organic–inorganic hybrid electrolytes [25]. And although this review primarily presents typical organic Li-ion conductors (oxide-based or sulfide-based materials), it is still necessary to provide a brief overview of solid polymer electrolytes (SPEs) and organic–inorganic hybrid materials due to their wide application in Li batteries.

2.2.1 Polyether-Based SPEs

In 1973, Wright et al. [76] first reported the successful synthesis of crystalline alkali metal salt complexes of poly(ethylene oxide) (PEO) with ionic conduction. And since 1978, SPEs have been considered as candidates for application in electrochemical devices due to a combination of solid-state electrochemical properties and facile processability inherent to plastics [77]. SPEs are solid solutions of alkali metal salts in polymer matrixes without the presence of organic liquid solvents. These electrolytes possess many advantages over liquid electrolytes, including the lack of leakage, low flammability, absolute safety, great flexibility and facile processability [78, 79]. And of the many SPEs, PEO is a semi-crystalline polymer that can enable interconnection between Li⁺ ions and oxygen atoms in the PEO matrix [80]. Here, it is widely acknowledged that the mobile and disordered Li transport of PEO SPEs can be ascribed to the continuous segmental motion of the PEO chain to transfer Li⁺ ions to the surrounding chains and create new sites [80, 81]. However, the motion of PEO chains can be negatively affected by decreased temperatures and aggravated crystallinity, thus restricting practical application [25]. To address this, researchers have reported that the potential solution to promote the segmental movement of SPEs is to reduce the presence of crystalline domains because amorphous phases are mainly accountable for high ionic conductivity in polymer materials rather than crystalline domains [79].

In addition, researchers have also reported that the suitable glass transition temperature (T_g) of materials is a major factor for high ionic conductivity because the movement of amorphous polymer chain segments is continuous only above T_g , which is crucial to facilitate the transport of ions [82]. And although elevated temperatures can to some extent expand the amorphous region and reinforce interfacial stability with electrodes [83, 84], high temperatures over 80 °C can result in growth of Li dendrites and limited electrochemical windows [85]. Therefore, ideal polymer electrolytes should possess a low T_g to maintain conductivity even at room temperature and preserve good ionic conductivity [86].

Furthermore, efforts have been made to inhibit crystallization, which has major implications in ionic conductivity in which the most common method is the addition of liquid plasticizers into PEO-LiX matrixes. However, this method can also reduce the intrinsic nature of polymer electrolytes due to solid-state deconfiguration and less compatibility with electrodes. Researchers have also suggested the addition of ceramic powder into PEO-type polymer electrolytes to enhance mechanical and interfacial properties and, most importantly, low-temperature conductivity [87–90]. Moreover, researchers have also reported that smaller nanosized ceramic powders (TiO₂, SiO₂, Al₂O₃ [91] and MgO [92]) can better retard crystallization. In addition, blend-based polymers [93–96] and copolymerization with other polymers [85, 97, 98] have also been proposed as effective means to develop amorphous polymer matrixes to reduce the possibility of increasing crystallization. Other polyethers such as PPO [99, 100] and nitrile-based polymer electrolytes such as PAN [101–103] have also have been reported. However, due to the limited space here, these will not be elaborated upon.

2.2.2 Other Organic SPEs

2.2.2.1 Polycarbonate-Based SPEs PEO-based SPEs were first mentioned by Wright et al. [76] and have recently experienced a renaissance. And although great efforts have been paid to compensate for the drawbacks of PEO-based solid electrolytes (extremely low ionic conductivity of 10^{-8} S cm⁻¹), tight coordination structures enabling strong coupling between Li-ion diffusion and segmental movements of polymer chains also need to be modified. Based on this, Koinuma et al. [104] synthesized aliphatic polycarbonates as novel hosts for SPEs through alternate copolymerization with carbon dioxide and epoxide. In addition, numerous aliphatic polycarbonate-based SPEs have been developed, such as PEC (polyethylene carbonate), PPC (polypropylene carbonate), PTMC (polytrimethylene carbonate). [105].

PEC can also be incorporated into alternating copolymers made up of carbon dioxide and epoxide in which a chain of (-O-(C=O)-O-) can be distinguished from the structure (Fig. 13) [106]. Based on this, Tominaga et al. [107] synthesized a novel composite of PEC and lithium bis(trifluoromethanesulfonyl)imide (LiTFSI) with appropriate ionic conductivity and high Li transference numbers in which the researchers reported that the ionic conductivity of PEC/LiTFSI became higher with increasing salt concentrations due to the imide anions from LiTFSI, which can cause plasticizing effects and enhance the segmental movement of the polymer chains. And as a result, the highest ionic conductivity obtained for the PEC/LiTFSI polymer electrolyte was 0.47 mS cm⁻¹ at ambient temperatures and the Li-ion transference number was much larger than that of a PEO-based counterpart (0.4 and 0.1 for PEC and PEO, respectively) [108].

PPC is another biodegradable copolymer of propylene oxide and carbon dioxide that has attracted considerable interest [109–111], and the application of PPC-based SPEs in batteries is mainly focused on pristine PPCs and its copolymer SPEs. In addition, cellulose is widely considered to be one of the most ample and renewable resources bearing excellent properties, including outstanding thermal stability, chemical stability, environmental benignity and a high dielectric constant [112–115]. And based on the properties of these 2 materials, Cui et al. [116] developed a novel rigid-soft and high-performance polymer electrolyte

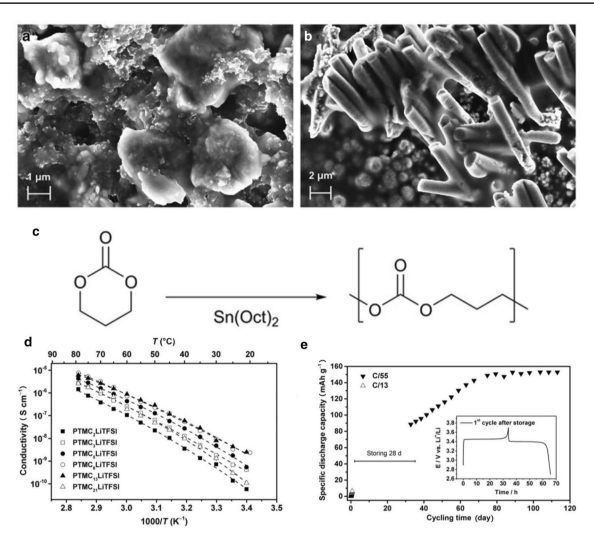


Fig. 13 SEM images of PTMC-LiTFSI materials with a LiFePO_4 composite electrode and b 3D Cu-nanopillar substrate. c Synthesis of PTMC. d Ionic conductivity of PTMC/LiTFSI materials as a function of temperature. e Diagram of the specific discharge capacities of

with nonwoven cellulose serving as the backbone and PPC transporting Li-ions in which the resulting safetyreinforced cellulose-PPC-LiTFSI SPE exhibited outstanding ionic conductivities, an appropriate electrochemical window and acceptable interfacial stability. Furthermore, LiFePO₄/Li and LiFe_{0.2}Mn_{0.8}PO₄/Li batteries using this cellulose-PPC-LiTFSI SPE showed extraordinary rate capabilities and cycling stability. These researchers also found an interesting phenomenon in which if a piece of the well-functioning cell was cut away, the remaining cell remained functional and could still light up a red LED lamp. Moreover, the safety of the cells using the cellulose-PPC-LiTFSI SPE also increased to unprecedented levels, providing a new avenue for the synthesis of high-voltage, ambient-temperature SPEs for LIBs.

a LiFePO₄/PTMC5LiTFSI/Li cell cycled at C/13 and C/55. Insert is the charge/discharge curve for the first cycle. Reprinted with permission from Ref. [117]. Copyright 2014 Elsevier

Moreover, PTMC is also a new host material for SPEs that can be synthesized through the ring-opening polymerization of cyclic monomers (Fig. 13), through which a high-molecular-weight polymer serving as a base material for electrolytes can be obtained. Here, Sun et al. [117] reported that a LiFeO₄-based LIB using a homogenous mixture of PTMC and LiTSI can exhibit high discharge capacities and reach a specific plateau of 153 mAh g⁻¹. These researchers also reported that the ionic conductivity of this mechanically robust and flexible PTMC-based electrolyte can reach 10^{-7} S cm⁻¹ at elevated temperatures and that it possessed good electrochemical stability, demonstrating the potential of PTMC in different battery applications.

2.2.2.2 Polysiloxane-Based SPEs In general, the development of oligo(ethylene oxide) polymers is considered to be

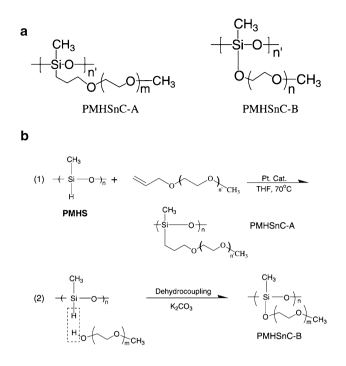


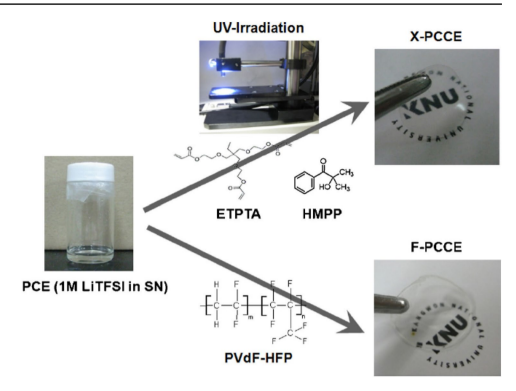
Fig. 14 a Chemical structures of mono-comb poly(siloxane-g-ethylene oxide). **b** Synthesis of mono-comb poly(siloxane-g-ethylene oxide). Reprinted with permission from Ref. [121]. Copyright 2003 Elsevier

a promising means to obtain amorphous polymers that are preferred for ideal SPEs and various polymer backbones have been studied [118]. Among these, polysiloxanes possess a very flexible backbone that can act as a potential host for polymers due to its low glass transition temperature, large free volume and good electrochemical stability [119] in which various organic groups can be connected to the silicon including ethylene oxide with side chains (chemical structure and synthesis of mono-comb poly(siloxane-gethylene oxide)). Despite the comparable conductivity for practical use, polysiloxane-based SPEs have been reported to possess low mechanical strength; however, this can be ameliorated through cross-linking or networking with polymers [25, 120]. In addition, several studies have investigated the ionic conductivity of polysiloxane-based SPEs based on the number and molecular weight of oligo(ethylene oxide) substituents and researchers have also developed comblike poly(siloxane-g-ethylene oxide) as backbones for SPEs (Fig. 14). For example, Oh et al. [121] synthesized a LiNi_{0.8}Co_{0.2}O₂/SPEs/Li cell that exhibited exceptional cycling performances with negligible capacity decline, suggesting that polysiloxane-based SPEs can serve as potential candidates in large-scale battery applications. Fonseca et al. [122] also explored poly(dimethyl siloxane-co-ethylene oxide) (P(DMS-co-EO)) copolymers, and through AC impedance analysis they reported that the maximum ionic conductivity of the copolymer film with 5 wt% LiClO₄ can reach $2.6 \times 10^{-4} \text{ S cm}^{-1}$ at room temperature and that the electrochemical stability window was wide, suggesting that P(DMS-co-EO)/LiClO₄ electrolytes are promising in electrochemical devices.

Furthermore, Hooper et al. [123] proposed in 2011 the possibility that disubstituted polysiloxanes are superior to monosubstituted analogues in terms of electrochemical performance and synthesized a series of double-comb polysiloxane polymers with a growing number of ethylene oxide units in the side chains and reported improved ionic conduction performances as compared with mono-comb polysiloxanes.

2.2.2.3 Plastic Crystal-Based SPEs Novel approaches revolving around the core idea of dissolving salts into highly flexible plastic crystals have been adopted by many researchers [124, 125]. This is because plastic crystals are a class of materials with disordered structures and lattice defects resulting from rotatory motions of included globular-shaped cations and anions [125, 126], allowing for excellent properties such as greater plasticity and enhanced ionic conduction as compared with normal rigid crystals [127]. And among LIB electrolytes, nitrile materials have attracted great attention due to their high thermodynamic stability as well as their extraordinary electrochemical oxidation resistance [4] as a result of the strong interaction between nitrile functional groups ($-C \equiv N$) and Li⁺ ions that can enhance the ionic conduction of nitrile-based SPEs [83]. Based on these factors, nitrile-based succinonitrile (N \equiv C–CH₂–CH₂–C \equiv N, SN) is a highly polar plastic-crystalline molecule that can dissolve various types of Li salts to achieve great ionic conductivity (two orders of magnitude above polymers) [125]. In addition, SN can maintain its plastic-crystalline phase in temperatures ranging from -40 °C to almost 60 °C (the melting point) [128].

Nevertheless, SN cannot be used as a self-standing film copolymer because it is extremely plastic and prone to deformation [25] and therefore is usually used as a versatile additive to obtain plastic crystal composite electrolytes (PCCEs) with enhanced ionic conductivity and robust mechanical properties. For example, Maier et al. [129] reported the feasibility of SN as an ionic conductivity enhancer in 3 polymer electrolyte systems, including PEO-LiTFSI, P(VdF-HFP)-LiTFSI and P(VdF-HFP)-LiBETI (LiBETI: lithium bisperfluoroethylsulfonylimide) in which the researchers attributed the enhanced ionic conduction to the strong solubility of SN for Li salts and the decreased crystallinity of the polymer. Lee et al. [127] also proposed a simple method to synthesize self-standing PCCEs through integration with UV (ultraviolet)-cured ethoxylated trimethylolpropane triacrylate (ETPTA) and SN-LiTFSI in which a transparent, nonsticky, and self-standing X-PCCE was synthesized **Fig. 15** UV curing process and images of the PCE, the X-PCCE and the F-PCCE. Reprinted with permission from Ref. [127]. Copyright 2011 Elsevier



after the trifunctional groups on an ETPTA monomer were cross-linked under exposure to UV lights for 20 s (Fig. 15). In addition, these researchers also synthesized F-PCCE through the integration of the conventional PCCE with polyvinylidenefluoride-co-hexafluoropropylene (PVdF-HFP) and reported that the resulting F-PCCE provided high ionic conductivity and unprecedentedly robust mechanical strength. Lee et al. [130] further fabricated a PCCE with a semi-interpenetrating polymer network (semi-IPN) as a matrix and a plastic crystal electrolyte (PCE, 1M lithium bis-trifluoromethanesulfonimide in succinonitrile) and reported unprecedentedly improved bendability and good ionic conduction, providing a novel method to fine-tune the electrochemical and physicochemical properties of PCCEs through anomalous UV-curable semi-IPN architectures as polymer matrixes.

2.3 Organic–Inorganic Hybrid SPEs

Organic–inorganic hybrid SPEs, including inert ceramic fillers (Al₂O₃, TiO₂, SiO₂, etc.), ferroelectric ceramic fillers (LiNbO₃, PbTiO₃ and BaTiO₃), clay and fast ion conductors [131–136], have also been reported to be promising in LIBs. In organic–inorganic hybrid SPEs, inorganic materials can be dispersed into polymer matrixes to effectively promote Li-ion conduction, create robust mechanical strength and provide more stable interfaces in SPEs [137–139]. The general idea of dispersing fine ceramic materials into ionic conductors has been prevalent since 1973 in which Liang et al. [140] proposed the feasibility of incorporating Al₂O₃

particles into Li salts to enhance Li⁺ conduction. Here, the two-phase coexistence of ceramic fillers and ion conductive polymers has attracted considerable interest after observations of enhanced conductivity and interfacial stability as compared with ceramic-free systems. In addition, a series of studies have reported that the addition of ceramic fillers can inhibit the crystallization kinetics of PEO polymer chains, extend the amorphous regions and promote specific surface interactions [87, 131, 141]. Furthermore, Sun et al. [132] in their study measured the conductivity of composites made up of different ferroelectric ceramics (BaTiO₃, PbTiO₃ and LiNbO₃) and Li salts (LiClO₄, LiBF₄, LiPF₆, LiCF₃SO₃ and $LiN(CF_3SO_2)_2$) (Fig. 16) and rationalized that the improved conductivity was based on the association tendencies of anions with Li⁺ and the spontaneous polarization of fillers from their particular crystal frameworks, resulting in substantially reduced interfacial resistances between electrolytes and electrodes. Murugan et al. [142] also proposed a novel polymer electrolyte of poly(ethylene) glycol (PEO) complexed with SrBi₄Ti₄O₁₅ fillers and LiClO₄ and used the citrate gel technique to form the same ferroelectric fillers with particle sizes that were much smaller than that of the conventional method and reported that the reduction in the particle size of fillers can lead to better ionic conduction.

In addition, Croce et al. [141, 143] utilized inert ceramic fillers (SiO₂, Al₂O₃, TiO₂, etc.) to develop nanosized PEObased polymer electrolytes with improved properties and reported that among the inert materials, TiO₂ was the most promising in the improvement in ionic conduction due to the fragile connection between Li⁺ ions and polymer chains

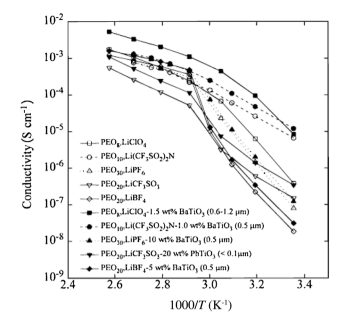


Fig. 16 Conductivity of composites consisting of ferroelectric ceramics, Li salts and PEO. Conductivity values were evaluated during the first heating scan by using impedance spectroscopy measurements. Reprinted with permission from Ref. [132]. Copyright 2000 The Electrochemical Society

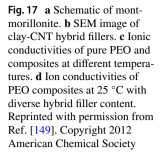
[25]. Lin et al. [144] also found that the reduction in TiO_2 grain size can increase the ionic conductivity of nanosized TiO₂-modified composite polymer electrolytes (CPEs) and that there was a great dichotomy in the ionic conduction between PEO-10% LiClO₄-5% TiO₂ (3.7 nm) and pure PEO-10% LiClO₄ in which the value of the former was an order of magnitude larger than that of the latter. Furthermore, these researchers also reported that transference numbers of 0.51 and 0.21 were achieved for the TiO₂-modified CPE and pure PEO-LiClO₄. Researchers have also reported that the interaction between TiO2-doped Li salts and PEO chains can promote structural modifications and endow Li-ions with favorable migration pathways [139] and that the high mobility of charge carriers can be regarded as the result of the interactions of polymers in the amorphous region even at ambient temperatures.

Montmorillonite (MMT) as a layered clay with a lamellar shape has attracted intense interest from researchers in the development of polymer–clay electrolytes due to the fact that lamellar elements can exhibit high in-plane strength, high aspect ratio and stiffness [145]. For example, Zhang et al. [146] reported that PEO/MMT/LiTFSI solid polymer electrolytes demonstrated reinforced Li-ion conductivities in which an optimal electrolyte composed of 10 wt % MMT reached a conductivity of 3.22×10^{-4} S cm⁻¹ at 60 °C and a corresponding solid-state battery displayed good cycling stability and an acceptable discharge capacity of 643 mAh g⁻¹ under the same conditions. Carbon nanotubes (CNTs) are widely thought to be ideal polymer matrixes to reinforce mechanical properties through high aspect ratios, good strengths and modulus. In addition, CNTs possess rich electron clouds that can create strong affinity with Li-ions and facilitate Li salt dissociation as well as ion migration through the framework [147, 148]. However, the electrical shorting of CNTs greatly restricts practical production. As an example, Tang et al. [149] successfully assembled hybrid 3D nanofillers with 1D CNTs and 2D MMT clay platelets (Fig. 17) and reported that the Li-ion conductivity and tensile strength of PEO-LiClO₄ electrolytes with the synthesized nanofiller were greatly improved, demonstrating the feasibility of embedding conducting–insulating hybrid materials in the synthesis of desirable SPEs.

Fast ionic conductors also possess many extraordinary merits, such as the lack of solution leakage, environmental benignity, nonflammability, high thermal stability, good electrochemical properties. [150]. Because of this, Murugan et al. [83] synthesized cubic Li₇La₃Zr₂O₁₂ (LLZO) with a garnet-like structure and Lee et al. [151] synthesized organic-inorganic membranes consisting of PEO and the diverse tetragonal LLZO content in which through AC impedance measurements, 52.5% LLZO was confirmed to be able to provide an optimal ionic conductivity of 4.42×10^{-4} S cm⁻¹ at 55 °C. Here, these researchers suggested that the hybridization of inorganic and organic materials can create synergetic effects to improve ionic conductivity in which the charge/discharge test of Li/composite membrane/ $LiNi_{0.6}Co_{0.2}M_{0.2}O_2$ was more satisfactory than that of PEOonly materials and a counterpart material with 52.5% Al₂O₃, demonstrating the viability of all-solid organic-inorganic hybrid composite membranes. Despite this, further studies into these hybrids should be devoted to the improvement in cycling performances for practical battery systems.

3 Factors Influencing the Ionic Conductivity of Solid-State Electrolytes

The ionic conductivity of solid-state electrolytes is a key parameter to determine electrochemical performance; however, few possess acceptable intrinsic conductivity. Based on this, great efforts have been put into the enhancement of ionic conductivity in solid-state electrolytes. Here, the understanding of the mechanisms and influences of ionic conductivity in solid-state electrolytes is the key, and therefore, this section will introduce relevant research based on 3 aspects of solid-state electrolytes, including structure, composition and the interphase.



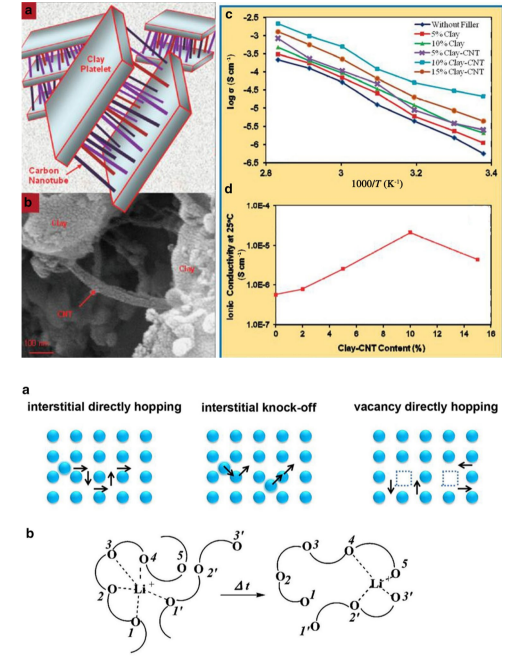


Fig. 18 Schematic of Li-ion transport in a crystalline and b polyether media. Reprinted with permission from Ref. [152]. Copyright 2004 American Chemical Society

3.1 Structure

The mechanism and conductivity of ionic transport are closely related to crystal structure and vary greatly for different systems. In general, there are 3 Li⁺ diffusion mechanisms in inorganic solid electrolytes, including interstitial direct hopping, interstitial knock-off and vacancy direct hopping (Fig. 18) [153] and a thorough comprehension of ionic transport is imperative to remold and design optimal solid electrolytes. For example, Ceder et al. [72] reported that frameworks with a body-centered cubic (bcc) anion arrangement that possess lower activation barriers are superior to

other anion packing models (fcc, hcp) and can achieve optimal ion transfer rates. Overall, the conductivity of ions in crystalline structures follows the Arrhenius law as follows:

$$\sigma_{\rm DC}T = A_{\sigma}\exp(-E_{\sigma}/k_{\rm B}T) \tag{1}$$

However, for structures that are not in a long-range order (*i.e.*, glass, polymers or crystalline materials with randomly occupied defects), the conductivity of ions does not follow the linear Arrhenius law and can be better represented by the Vogel–Fulcher–Tammann (VFT) or Williams–Landel–Ferry (WLF) equations as follows [154]:

$$\sigma_{\rm DC}T \propto A_{\sigma} \exp[-E_{\sigma}/k_{\rm B}(T-T_0)] \tag{2}$$

in which in crystalline inorganic solid electrolytes, defect sites may weakly impact each other, whereas for amorphous systems with many defects, the correlation between mobile ions and the interaction between Li⁺ and framework ions should be taken into consideration [155]. And until now, no explicit theories into ion diffusion behaviors in disordered systems have been proposed; however, a few amorphous solid electrolytes have been reported to show higher ionic conductivities than crystalline or rigid solid electrolytes in laboratory settings in which the improvements may be due to the vast defects in the amorphous materials. In addition, researchers have also reported that the mixing of crystalline materials with amorphous glass can also enhance conductivity, which may be attributed to greater defects around the crystalline region [26].

As for SPEs, it is generally accepted that ions move mainly in the amorphous phase [156] in which the motion of ions in polymer systems is closely related to the structural relaxation of polymeric chains. For example, Ratner et al. [157] studied Li-ions coordinated with polar groups in polymer chains (Fig. 18) and reported that accompanying the continuous peristalsis of segments, polar groups that are solvated by Li-ions can be replaced to create the incessant movement of Li-ions, demonstrating that the number and stability of polar groups as well as the chain strength and peristalsis of the polymer host all affect the conductivity of SPEs.

As for inorganic-organic composite electrolytes, multiphase structures can increase the complexity of the ion transport process in which both amorphous and crystalline components can provide access to ion transport to diversify pathways and the structure and distribution of different phases can make the process even more viable. Researchers have also reported that the introduction of nanosized inorganic components to polymers can enhance mechanical strength and ionic conductivity in which numerous researchers have reported the occurrence of Liion hopping near polyether-filler interfaces [158, 159]. Moreover, Wieczorek et al. [160] used structural characterizations obtained from FT-IR, NMR, DSC and EDX methods to propose an ionic transport mechanism in composite polymeric systems in which they demonstrated that the formation of various complexes near polyether-filler interfaces as a result of acid-base reactions is responsible for conductivity enhancements. And based on NMR, ac conductivity spectra and powder X-ray patterns results, Liu et al. [161] proposed a suppositional ionic conductivity mechanism for highly crystalline polymer electrolytes in which polymer molecules remained stationary whereas Li-ions hopped between adjacent units driven by defects and cavities in the crystalline phase, suggesting that larger

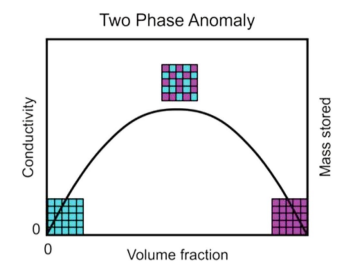


Fig. 19 Qualitative diagram of the enhanced conductivity and higher storage capacity of two-phase composites. Reprinted with permission from Ref. [166]. Copyright 2007 Elsevier

crystallite sizes with more abundant defects can enhance ionic transport.

3.2 Composition

Initial research into ion transport systems was mainly focused on bulk properties; however, researchers soon realized that boundary layers are fundamentally connected with significant impact on ionic conduction properties in which Liang et al. reported in 1973 that two-phase LiI-Al₂O₃ possessed unusually high ionic conductivity in comparison with single-phase LiI. Since this study, serious research has been conducted into these solid electrolytes (referred to as "composite electrolytes" or "heterogeneous electrolytes") with Wagner et al. making significant contributions. Here, it is commonly believed that the improved conductivity of these solid electrolytes arises from the interfacial zones in which different compositions can bring about different interfaces with different conductive mechanisms [162]. Wagner et al. [163] further proposed that enhanced conductivity may originate from multiple reasons, including space-charge layers (SCLs), more intensive dislocation densities and the formation of new phases. Here, the SCL theory can be further referred to in publications by Maier et al. [162, 164, 165] and is universally approved and accepted (Fig. 19).

3.3 Dimension

The interface represents a discontinuity of the structure and usually arises simultaneously with variations in charge carrier concentration. In addition, interfacial regions can expand by reducing particle sizes or dimensions (i.e.,

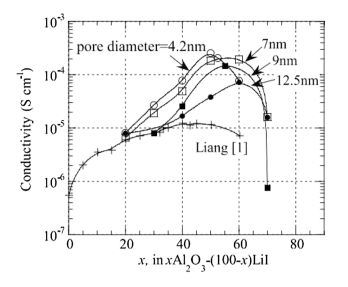


Fig. 20 Composition and pore size correlations with dc conductivity for Al_2O_3 -LiI composites. Reprinted with permission from Ref. [167]. Copyright 2004 Elsevier

nanocrystallization) and in comparison with bulk materials, nanosized solid electrolytes possess much larger interfacial regions to allow for significantly enhanced ionic conductivity. Maekawa et al. [167, 168] also conducted research into the influence of Al₂O₃ pore sizes in composite electrolytes to improve ion conductivity and reported that conductivity can be greatly improved by 10-50-folds with the reduction in particle size. Here, the researchers suggested that this enhancement originated from overlapping space charges and the formation of new components in the interfacial region. In addition, Maier et al. [169] also proposed the possibility of the generation of metastable subsystems in the interface, and more detailed discussions can be found in their published literature. Overall, due to the introduction of enormous interfaces, the ionic conductivity of solid electrolytes has been significantly advanced through nanocrystallization,

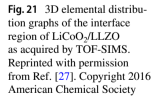
and therefore, this "nanoionics" is vital for future research (Fig. 20).

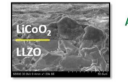
4 Advanced Characterization Techniques for the Study of Solid-State Electrolytes

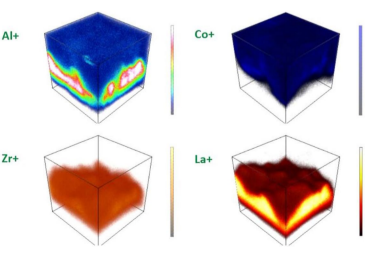
In addition to traditional characterization techniques such as X-ray diffraction (XRD), energy dispersive X-ray detection (EDX), X-ray photoelectron spectroscopy (XPS), scanning electron microscopy (SEM), transmission electron microscopy (TEM), up-to-date and advanced techniques have emerged recently in the analysis and detection of solid-state electrolytes. Based on this, this section will introduce some of the latest techniques reported.

4.1 Time-of-Flight Secondary Ion Mass Spectroscopy

The interphase between electrode materials and solid-state electrolytes plays a definitive role in overall performance. Therefore, meticulous surface characterization by using powerful tools to confirm chemical changes is imperative. Here, time-of-flight secondary ion mass spectroscopy (TOF-SIMS) is a powerful characterization technology that possesses the ability to probe the composition, spatial distribution and structure of samples in which vast quantities of information can be obtained within a short period of time through high-quality spectra and direct chemical imaging. As a result, the high sensitivity and adaptability of TOF-SIMS have aroused increasing interest in application for material characterization. For example, Park et al. [27] studied the interface between LiCoO₂ and LLZO. Through the use of TOF-SIMS, they were able to obtain a 3D highresolution elemental distribution mapping in which colored dots represented various elements with ionic concentrations shown based on color scales (Fig. 21). And as a result, these







researchers were able to observe that at the $LiCoO_2/LLZO$ interface, Co diffused across the interface from $LiCoO_2$ into LLZO whereas Zr/La in LLZO diffused into $LiCoO_2$. In addition, these researchers also unexpectedly observed the results for Al, with Al diffusing over long distances and even at the end of $LiCoO_2$. These researchers also used XRD and TEM to confirm that this leach out of Al in LLZO can further cause the transformation of cubic LLZO into a tetragonal phase. Overall, due to its nondestructive testing and high sensitivity, TOF-SIMS will become more versatile in the field of electrochemistry.

4.2 Solid-State NMR Techniques

Solid-state nuclear magnetic resonance spectroscopy (SS NMR) is a critical and requisite characterization method for the detection of atomic level local structures and ion motions of specific species. In addition, various NMR techniques including pulsed-field gradient (PFG) NMR, triple-quantum MAS (3QMAS), magic angle spinning (MAS) NMR, spin-lattice (SLR) NMR, 3D exchange (2D EXSY) NMR and spin alignment echo (SAE) NMR have been employed and more often are combined to trace structural evolutions, ion diffusion pathways and ion conductor dynamics. This is because NMR techniques are nondestructive, non-contact and possess a wide range of applications toward not only crystalline but also amorphous materials. For example, Arbi et al. [170] explored $Li_{1+x}Al_xM_{2-x}(PO_4)_3$ materials (M = Ti, Ge) using MAS-NMR to ascertain ion locations and Li-ion diffusion pathways in which ²⁷Al and ³¹P MAS-NMR as well as the spectra of ⁷Li (Fig. 22) were investigated. Here, the researchers reported that the spectra were observed to be centered at ~0 ppm, which can be attributed to Li^+ at M1 and M3 sites with further fitting results of the spectra revealing that the quadrupole interactions at the M3 sites were significantly lower than that at the M1 sites, demonstrating enhanced Li⁺ mobility at the M3 sites. In another study, Wang et al. [171] investigated the doped garnet-type solid electrolyte Li₇La₃Zr₂O₁₂ to reveal the influence of aliovalent substitution on ionic conductivity using highresolution NMR analysis and found that the Li⁺ ions had 3 sites in the garnet-type electrolyte, including 24d (tetrahedral sites), 48g and 96h (octahedral sites). Here, the researchers carried out 2D chemical exchange experiments using NMR to obtain a 2D ⁶Li-⁶Li exchange spectrum of $Li_{7-2x-3y}Al_{y}La_{3}Zr_{2-x}W_{x}O_{12}$ (Fig. 23) in which the crosspeak at $\omega_1 = 0.8$, $\omega_2 = 1.6$ ppm revealed the close relationship between the 96h and 24d sites, supporting Li migration along 24d-96h-48g-96h-24d. Liang et al. [172] also used diverse SS NMR techniques to investigate the origins of the optimal Li-ion conductivity in Li₁₀GeP₂S₁₂ (LGPS) solid electrolytes and found that Li-ions in LGPS systems can diffuse along 1D tunnels simultaneously with 2D plane pathways, which

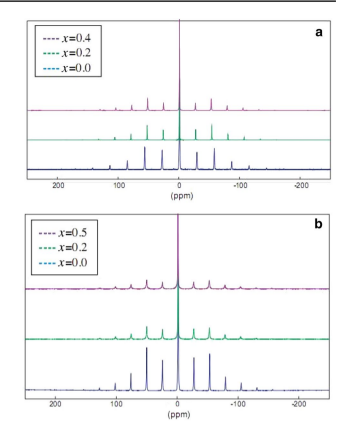


Fig. 22 ⁷Li MAS-NMR spectra of **a** LATP and **b** LAGP samples. Reprinted with permission from Ref. [170]. Copyright 2015 Elsevier

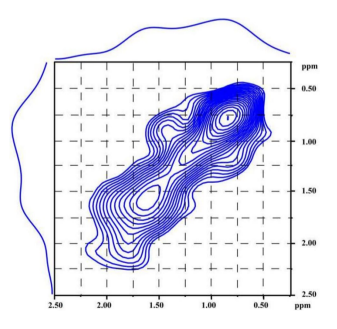


Fig. 23 ${}^{6}\text{Li}{}^{-6}\text{Li}$ exchange spectra of $\text{Li}_{7-2x-3y}\text{Al}_{y}\text{La}_{3}\text{Zr}_{2-x}W_{x}O_{12}$. Reprinted with permission from Ref. [171]. Copyright 2015 American Chemical Society

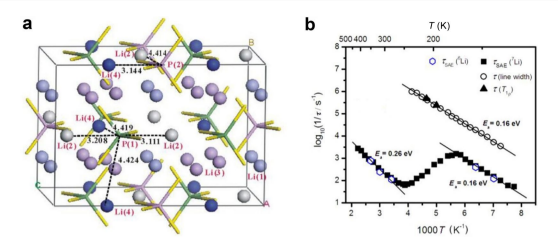


Fig. 24 a Crystal structure diagram of LGPS. b ⁷Li $1/\tau$ of LGPS, ⁷Li spin-lock SLR NMR and the line width. Reprinted with permission from Ref. [172]. Copyright 2015 American Chemical Society

corresponded to activation energy of 0.16 eV and 0.26 eV, respectively. In addition, these researchers obtained the relationship curve of $1/\tau_{SAE}$ versus the reciprocal of temperature for LGPS (Fig. 24) in which τ_{SAE} is a parameter denoting the correlation time of Li⁺ migration based on Eq. (3) and found that τ_{SAE} experienced an initial decline followed by an upsurge and subsequent decline with rising temperatures, suggesting the existence of 2 different processes for Li-ion diffusion in LGPS systems. Furthermore, based on the 2 gradients of the Arrhenius curve in the low- and high-temperature areas, activation energy of 0.16 eV and 0.26 eV can be obtained. For a more detailed reference, Xiang et al. [173] provided a detailed overview of NMR techniques utilized in the solid electrolyte field.

$$S_2(t_{\rm m}, t_{\rm p}) = \left\{ A \exp\left(-\frac{t_{\rm m}}{\tau_{\rm SAE}}\right) + B \right\}^{\gamma_{\rm c}} \exp\left(-\frac{t_{\rm m}}{T_{\rm 1Q}}\right) \quad (3)$$

4.3 Magnetic Resonance Imaging

Magnetic resonance imaging (MRI) is an up-and-coming technology in the fields of catalysis [174, 175], polymers [176], corrosion [177], fuel cells [178], etc, due to its high reliability and nondestructive testing. And recently, studies using MRI have been emerging in the area of batteries to provide both quantitative and spatial information. For example, Chien et al. [177] used 2D and 3D ⁷Li MRI to study Li distribution in the solid electrolyte $Li_{10}GeP_2S_{12}$ and found Li deficiency at the interface and enhanced heterogeneity of Li distribution through repeated charge and discharge in which quantitative 3D ⁷Li MRI images of $Li_{10}GeP_2S_{12}$ and PEO-coated $Li_{10}GeP_2S_{12}$ pellets after cycling (Fig. 25) showed significant Li depletion on the top and bottom layers

with Li depletion being more severe at the top whereas none was observed in the bulk. In addition, these researchers analyzed the scattering of histograms and found that the Li content for both top and bottom layers was heterogeneous in the unmodified sample whereas with the modification of the PEO film at the interface, Li loss and the heterogeneity distribution of Li were dramatically reduced. Romanenko et al. [28] also conducted in situ MRI studies of organic ionic plastic crystals (OIPCs) in a solid-state battery with a focus on Li⁺ transport at the interface and proposed the relationship between the ¹H MRI signal and Li-ion content/ distribution through the use of a novel SPRITE (singlepoint ramped imaging with T₁ enhancement) technique in which the displacement of the front electrolyte, a fast grain boundary transfer and Li⁺ exhaustion at the interface were detected.

5 Promising Strategies to Enhance Li-Ion Transport

5.1 Structural Tuning

5.1.1 Tuning lattice Volume Through Substitution

For crystalline electrolytes, incremental lattice volumes can result in higher Li-ion conduction and lower activation energy in which lattice volumes can be changed through 2 effective approaches including substitution and mechanical strain. In terms of substitution, increased ionic conductivity can be achieved through iso-valent substitution in a given structural framework to determine lattice volumes and bottleneck sizes for Li-ion diffusion. For example, the conductivity of NASICON-type $\text{LiM}_2(\text{PO}_4)_3$ (M=Ge, Zr or Ti) can be enhanced through elemental substitution to **Fig. 25 a, b** ⁷Li 3D MRI images of cycled $\text{Li}_{10}\text{GeP}_2\text{S}_{12}$ pellets with and without PEO coating. **c** Plot of normalized ⁷Li densities versus the layer index. **d, e** Bar graphs of normalized Li counts at different depths corresponding to **a, b.** Reprinted with permission from Ref. [179]. Copyright 2018 American Chemical Society

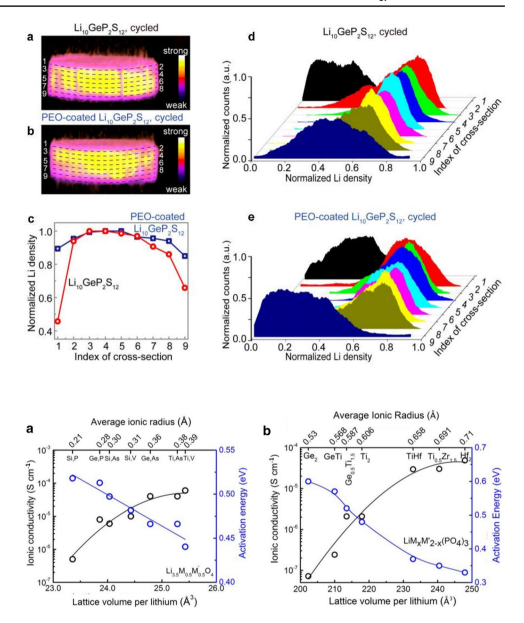


Fig. 26 Li-ion conductivity of **a** LISICON-like $\text{Li}_{3,5}\text{M}_{0,5}\text{M}'_{0,5}\text{O}_4$ and **b** NASICON-like $\text{LiM}_x\text{M}'_{1-x}(\text{PO}_4)_3$ as a function of lattice volume per Li atom. Reprinted with permission from Ref. [180]. Copyright 2016 American Chemical Society

generate $\text{Li}_{1+x}M_x\text{Ti}_{2-x}(\text{PO}_4)_3$ (M = Al, Cr, Ga, Fe, Sc, In, Lu, Y or La) (Fig. 26) [181]. In addition, aliovalent substitution of O^{2-} with S^{2-} in the sulfide electrolyte Li_3PS_4 can form the thio-LISICON compound $\text{Li}_{10}\text{GeP}_2\text{S}_{12}$ (LGPS) and improve Li-ion conductivity through the enlargement of ion transport channels [182]. As for perovskite-type structures with a formula of ABO₃ (A = La, Sr, or Ca; B = Al or Ti), substitutions with larger size A-site rare-earth metal ions (Sm < Nd < Pr < La) can also enlarge lattice volumes and improve Li-ion conductivity and lower activation energy [183]. Furthermore, enhanced Li-ion conductivity induced by substitution was also reported for LISICON-type electrolytes [184].

Aside from substitution, mechanical strain can also alter lattice volume through tensile or compressive strain on crystalline conductors in which according to DFT studies on cubic Li₇La₃Zr₂O₁₂ (cLLZO) [185] and LGPS [186], isotropic compressive strain can significantly lower Li-ion conductivity whereas tensile strain showed no effects. This suggests that the further increase in lattice volume has little effect on the reduction in Li-ion activation energies when transiting bottleneck points or the lattice volume of these 2 conductors is close to optimization. Recent studies have also demonstrated that the tailoring of Li conductivity is attainable through the generation of strain from lattice mismatches in the substrates of thin-film growth of Li-ion conductors with Li_{0.33}La_{0.56}TiO₃ thin films grown on NdGaO₃ (NGO) being a good example [187, 188] in which the distinct strain imposed by NGO along the aand *b*-axes directions can lead to the anisotropy of the ionic conductivity. Despite these results, however, more efforts are needed to deeper probe into the implications

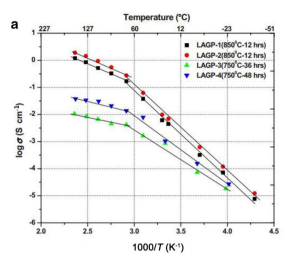
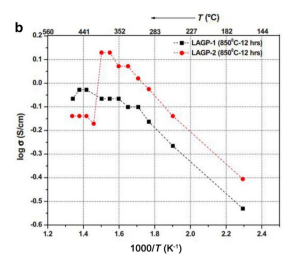


Fig. 27 a Arrhenius plots of the conductivity of LAGP specimens crystallized at diverse temperatures and times. **b** Arrhenius plots of the conductivity of LAGP specimens crystallized in the high-temper-

of mechanical strain on Li-ion conduction with considerations of the negligible variations in ionic conductivity.

5.1.2 Decreasing Grain Boundary Resistances Through the Control of the Space-Charge Effect

Decreasing grain boundary resistance is another attractive approach to increase Li-ion conductivity, and researchers have reported that significant decreases can be achieved through the incorporation of other phases at the grain boundary as caused by the space-charge effect [189]. Here, 2 separate processes have been adopted to create new phases at interfacial boundaries in which the first involves additive fillers. This is because additive fillers can play an important role in enhancing conductivity in which in addition to conventional ionic transport in liquid electrolytes, Li-ion adsorption on filler particles can lead to increased vacancy concentrations of Li-ions in space-charge regions, contributing to higher conductivities. For example, Cao et al. [190] reported that polymer electrolytes can show high ionic conductivity due to well-dispersed filler nano-TiO₂ particles in the conductor. And based on C-rate and AC impedance analysis, these researchers proposed that the interfacial resistance of the cells can be decreased due to ionic mobility in the vicinity of TiO₂ nanoparticle boundaries. Furthermore, more favorable Li-ionic conduction pathways can be created through Lewis acid-base interactions or through the formation of favorable transport processes closely connected to TiO₂ nanoparticles [136, 191]. For example, Kumazaki et al. [192] reported that the grain boundary resistance of garnet-structured Li₇La₃Zr₂O₁₂ can be lowered through the incorporation of Si and Al and attributed this to the interface of amorphous



ature region. Reprinted with permission from Ref. [196]. Copyright 2009 The Electrochemical Society

Li-Al-Si-O as well as to the nanocrystalline LiAlSiO₄ that can improve Li-ion migration between $Li_7La_3Zr_2O_{12}$ grains. Moreover, the addition of B_2O_3 , TiO₂ and ZrO₂ has also been reported to be able to promote the rearrangement of grains to lead to better connections during the crystal-lization process [193–195].

The second process to create new phases at interfacial boundaries is a sintering process that does not require fillers and can provide conductive phases to enhance Liion conductivity. For example, Kumar et al. [196, 197] reported that during the synthesis of LAGP, the formation of Li₂O and AlPO₄ at the grain boundaries can result in space-charge-mediated effects on conductivity in which the effects displayed in the LAGP specimens were attributed to their impact on activation energy. Here, the researchers reported that the formation of AlPO₄ at ~70 °C showed small space-charge effects as reflected in the Arrhenius plot whereas the formation of Li₂O at ~400 °C contributed to significant drops in conductivity (Fig. 27), allowing the formation of Li₂O:Li⁺ complexes to display enhancements in ionic conductivity. And aside from synthesis temperatures, these researchers also reported that the concentration of dielectric phases can also influence space-charge effects [196].

Moreover, ambient conditions and preparation methods can also affect the performance of grain boundaries. For example, Kali et al. [198] proposed that spark plasma sintering can impart desirable properties such as fast densification, minimization of coarse particles and fine grain-to-grain bonding in which Li₇La₃Zr₂O₁₂ sintered in Ar atmosphere displayed lower Li-ion conductivity due to fewer grain boundaries as compared with that prepared in air [199].

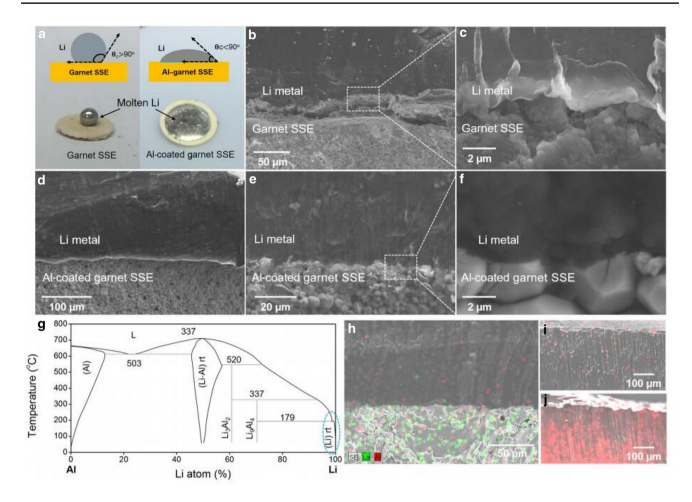


Fig. 28 a-f Wetting properties and appearances of interfaces between bare and Al-coated LLZO and Li. g Phase diagram of Li-Al. h Elemental mapping of the cross section. i, j Elemental mapping of the top area of Li. Reprinted with permission from Ref. [200]. Copyright 2017 Science

5.2 Interface Engineering

Different from common batteries or semisolid batteries, solid-state batteries are composed of solid materials in which a prominent solid-solid interface barrier occurs between the electrolyte and electrodes, making ion diffusion sluggish. In addition, the gaps between the electrolyte and electrodes can also impair battery cycling ability or render them inoperable. Therefore, sufficient contact and the more important issues of good electrochemical compatibility and fast interfacial ionic transfer between electrodes and electrolytes are the key to the future application of solid-state electrolytes. Based on this, this section will present related studies on this topic.

5.2.1 Interface Between the Electrolyte and the Anode

Garnet-type $Li_7La_3Zr_2O_{12}$ (LLZO) solid electrolytes are promising materials for next-generation batteries due to remarkable intrinsic conductivity and stability toward Li anodes. However, a poor interfacial wetting ability limits further application as a result of insufficient physical contact. To address this, Hu et al. [200] proposed the physical deposition of an Al layer as thick as 20 nm using Ebeam evaporation to form a Li-Al alloy at the interface between LLZO and Li (Fig. 28). Here, the researchers reported that the as-formed artificial alloy layer can enable the wetting of the LLZO surface to allow the interface resistance to be significantly reduced from 950 to 75 Ω cm². In addition, these researchers also reported that this electrode design showed outstanding electrochemical performances for Liion, Li-sulfur and Li-oxygen batteries. Furthermore, Hu et al. [19] proposed another strategy to address the interfacial challenges of garnet-type solid electrolytes through the deposition of an ultrathin layer of Al₂O₃ onto the surface of the garnet electrolyte using atomic layer deposition in which greatly reduced conductivity resistances from 1710 to 1 Ω cm² at room temperature were achieved. As a result, a full cell with Li2FeMn3O8 (LFMO) as the cathode, garnet $Li_7La_{2.75}Ca_{0.25}Zr_{1.75}Nb_{0.25}O_{12}$ (LLCZN) as the electrolyte and Li as the anode were tested to have significant

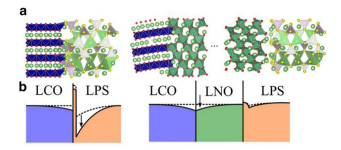


Fig. 29 a Schematic of the atomic structure in equilibrium and \mathbf{b} Li concentration at the initial stage of charging at the interface. Reprinted with permission from Ref. [210]. Copyright 2014 American Chemical Society

improvements, which the researchers attributed to the deposited Al_2O_3 layer that can accelerate Li^+ transfer, prevent electrolyte decomposition and strengthen contact between the electrolyte and Li metal.

Li dendrites can also form along the grain boundaries in $Li_7La_3Zr_2O_{12}$ (LLZ) solid electrolytes as detected by TEM, EELS and MAS NMR, resulting in the short-circuiting of batteries. To address this, Tsai et al. [201] proposed that the reason for this dendrite growth can be the inhomogeneous contact at the interface through carefully polishing and modifying the surface with thin Au layers. Here, the researchers reported that as a result, surface contact was greatly enhanced and resistances reduced, with growth of Li dendrites being prevented at the same current densities. However, these researchers also reported that Li dendrites still appeared at higher current densities, indicating that more comprehensive and in-depth research is required. Furthermore, many new reports have been published to address interfacial engineering on LLZO [27, 202–207].

Sulfide electrolytes can also be made into all-solid-state batteries and have also attracted much attention recently [208, 209]. For example, Yao et al. [208] studied the intimate contact at the solid electrolyte–cathode interface using an in situ liquid-phase method and suggested that the $\text{Li}_7\text{P}_3\text{S}_{11}$ electrolyte particles were directly anchored onto cobalt sulfide nanosheets to lead to excellent cycling and rate performances.

5.2.2 Interface Between the Electrolyte and the Cathode

The prominent issue of interfaces between electrolytes and cathodes is instability as caused by mutual diffusion, side reaction, etc., as well as the space-charge layer effect. And similar to the solutions of anode interfaces, the introduction of buffer layers into interfaces is also effective. For example, Haruyama et al. [210] built up a buffer layer of LiNbO₃ (LNO) between an oxide cathode (LiCoO₂ (LCO)) and a sulfide electrolyte (β -Li₃PS₄ (LPS)) in an all-solid-state

Li-ion conductor and through theoretical calculations of the 3 interfaces (Fig. 29), elucidated the matching structures with atomic levels and electronic properties in the interfacial regions. Here, the researchers found that the introduction of LNO layers can alleviate structural disorder at the interface, balance Li distribution and suppress growth of SCLs to boost Li conduction.

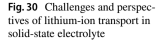
Interfacial reactions of solid electrolytes with anode and cathode materials can also affect performance [211]. For example, Auvergniot et al. [212] studied the interfacial phenomenon of argyrodite $\text{Li}_6\text{PS}_5\text{Cl}$ toward LiCoO_2 , $\text{LiNi}_{1/3}\text{Co}_{1/3}\text{Mn}_{1/3}\text{O}_2$ and LiMn_2O_4 electrodes using Auger electron spectroscopy and X-ray photoelectron spectroscopy and found that the electrolyte can be oxidized into sulfur, lithium polysulfides, P_2S_x , phosphates and LiCl at the interface. Wenzel et al. [213, 214] also investigated the interfacial zone between $\text{Li}_7\text{P}_3\text{S}_{11}$ and Li metal using in situ photoelectron spectroscopy and discovered the existence of a nongrowing resistive interphase in contact with Li metal analogous to an SEI film. Similar approaches were also taken to study the interface between LLTO and Li electrodes and showed the reduction in Ti ions and Ti metal.

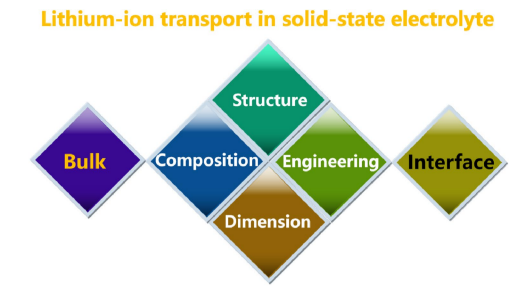
Overall, the introduction of buffer layers between solid electrolytes and electrodes is an effective method to control interfacial resistance. However, these artificial layers can also increase conduction resistance at the interface. Therefore, search for novel interfacial materials and synthesis of thinner deposition layers is a major direction for future development [215].

6 Conclusions and Perspectives

With the rapid progress of solid-state Li-based batteries in recent years, numerous efforts have been devoted among academic and industrial communities to enhance the performance of solid-state electrolytes. However, compared with commercial liquid electrolytes, numerous challenges need to be addressed before large-scale application. These challenges include methods to enhance Li-ion transport in the bulk, decrease grain boundary resistance at the interface between solid electrolytes and electrodes and enhance the stability of the structure and the interface of solid-state electrolytes during long term cycling (Fig. 30).

Currently, although some material systems with relatively high conductivities as Li-ion conductors have been reported, more efforts are required. And through numerous investigations, researchers have found that anion arrangements can determine the intrinsic ionic conductivity of materials in which the body-centered cubic (bbc) anion framework is proposed to be the most desirable for fast ion diffusion and low activation energy (*E*a). In addition, substitution with optimal-sized ions for a given material has extensively





proven to be effective in the enhancement of conductivity for LISICON, NASICON, perovskite and other families by orders of magnitude in which activation energy can be reduced through expanded lattice volumes and enlarged bottleneck sizes. Aside from ionic sizes and lattice volumes, the introduction of amorphous glass into crystalline electrolytes can also enhance conductivity through so-called "composite electrolytes" or "heterogeneous electrolytes". Moreover, more intensive defects and space-charge layer effects can also contribute to improved conductivity.

Aside from Li diffusion in the bulk of solid-state electrolytes, interfacial properties also play a crucial role in the performance of all-solid-state electrolytes in which for practical applications of all-solid-state batteries, the interface between solid-state electrolytes and electrodes needs to be stable and electrochemically compatible to minimize interfacial resistances. In addition, different from liquid electrolytes, the contact at solid/solid interfaces is often insufficient. To address this, many sustainable solutions have been demonstrated to improve battery performance. For example, various surface engineering treatments can be carried out to deposit thin layers of metals, oxides and semiconductors to modify the electrolyte/electrode interface. Moreover, the polishing of electrolyte surfaces and the exerting of appropriate pressures can also be effective. More intimate contact and much smaller resistance can also be obtained through the introduction of additional soft polymer interlayers. In addition, the design and fabrication of artificial buffer layers into electrolyte/electrode interfaces can enable intimate contact to suppress side reactions and reduce transfer resistance.

Overall, future studies into solid-state electrolytes should focus on the design of multi-component nanoscale interfaces and the minimization of surface coating thicknesses. In addition, the understanding of Li-ion transport in the bulk and the interface of solid-state electrolytes through advanced characterization techniques is fundamental to the future advancement of solid-state electrolytes and can provide new perspectives into the design of high-capacity solid-state batteries.

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