Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}] – a layered lithium superionic conductor
Li$_{0.6}$[Li$_{0.2}$Sn$_{0.8}$S$_2$] – a layered lithium superionic conductor†

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One of the key challenges of energy research is finding solid electrolytes with high lithium conductivities comparable to those of liquid electrolytes. In this context, developing new structural families of potential Li$^+$ ion conductors and identifying structural descriptors for fast Li$^+$ ion conduction to occur is key to expand the scope of viable Li$^+$ ion conductors. Here, we report that the layered material Li$_{0.6}$[Li$_{0.2}$Sn$_{0.8}$S$_2$] shows a Li$^+$ ion conductivity comparable to the currently best lithium superionic conductors (LISICONs).

Li$_{0.6}$[Li$_{0.2}$Sn$_{0.8}$S$_2$] is composed of layers comprising edge-sharing Li/SnS$_6$ octahedra, interleaved with both tetrahedrally and octahedrally coordinated Li$^+$ ions. Pulsed field gradient (PFG) NMR studies on powder samples show intragrain (bulk) diffusion coefficients $D_{\text{NMR}}$ on the order of 10$^{-11}$ m$^2$ s$^{-1}$ at room temperature, which corresponds to a conductivity $\sigma_{\text{NMR}}$ of 9.3 $\times$ 10$^{-3}$ S cm$^{-1}$ assuming the Nernst–Einstein equation, thus putting Li$_{0.6}$[Li$_{0.2}$Sn$_{0.8}$S$_2$] en par with the best Li solid electrolytes reported to date. This is in agreement with impedance spectroscopy on powder pellets, showing a conductivity of 1.5 $\times$ 10$^{-2}$ S cm$^{-1}$. Direct current galvanostatic polarization/depolarization measurements on such samples show negligible electronic contributions (less than 10$^{-9}$ S cm$^{-1}$) but indicate significant contact resistance (d.c. conductivity in a reversible cell is 1.2 $\times$ 10$^{-4}$ S cm$^{-1}$ at 298 K). Our results suggest that the partial occupation of interlayer Li$^+$ positions in this layered material is beneficial for its transport properties, which together with tetrahedrally coordinated Li sites provides facile Li$^+$ ion diffusion pathways in the intergallery space between the covalent Sn(Li)S$_2$ layers. This work therefore points to a generic design principle for new layered Li$^+$ ion conductors based on the controlled depletion of Li$^+$ ions in the interlayer space.

**Broader context**

Fast solid lithium conductors with negligible electronic contribution are materials of enormous scientific and technological significance. Solid electrolytes enabling rapid ion transport are key ingredients of all-solid-state devices such as batteries, supercapacitors, electrochemical sensors and actuators. Especially in the field of Li-based batteries, appropriate electrolytes with high Li ion conductivities are key ingredients, given the fact that conventional liquid electrolytes suffer from severe safety risks. Therefore, discovering new solid electrolytes composed of earth-abundant elements is in high demand. Here, we describe the first simple ternary ultrafast solid electrolyte and point to a new design principle according to which the implementation of vacant Li sites can drastically enhance the Li ion mobility by orders of magnitude.

**Introduction**

Efficient battery systems for mobile applications, particularly in portable devices, are currently in high demand. To date, rechargeable Li$^+$ ion batteries are leading in performance but suffer from safety concerns augmented by the use of liquid electrolytes. Current lines of research are therefore directed towards the development of non-flammable, non-toxic solid Li electrolytes for use in all-solid-state Li$^+$ ion batteries with similar power densities. In view of the delicate nature of the solid–solid contacts they are particularly relevant for thin film battery applications. Furthermore, solid electrolytes are relevant for various other applications such as electrochemical capacitors, electrochemical sensors and electrochemical actuators.

Since its discovery in the 1970s, lithium nitride (Li$_3$N) and its doped variants have been the best Li$^+$ ion conductors ($\sigma_{\text{298K}}$ = 6 $\times$ 10$^{-3}$ S cm$^{-1}$). However, due to its high reactivity and low
electrochemical decomposition potential, Li2N is not suitable for commercial applications. Other Li+ ion conducting materials with high conductivities are e.g. oxide perovskites (Li0.3Li0.7TiO3; \( \sigma_{298K} \approx 1 \times 10^{-3} \) S cm\(^{-1}\)), garnet-type systems such as Li2La3Zr2O12 (\( \sigma_{298K} \approx 3 \times 10^{-4} \) S cm\(^{-1}\)), sulfides such as Li2SnS4 (\( \sigma_{298K} \approx 7 \times 10^{-5} \) S cm\(^{-1}\)) and the 70Li2S-30P2S5 glass ceramic (\( \sigma_{298K} \approx 3.2 \times 10^{-3} \) S cm\(^{-1}\)) as well as mixed chalcogenide glassy materials (Li2S–SiS2–Li3PO4; \( L - \omega = 0.75 \)). One of the most prominent classes of crystalline Li electrolytes are the so-called lithium superionic conductors (LISICONs), which evolved from the prototype Li2+xZn1-xGeO4 with \( \sigma_{298K} \approx 1 \times 10^{-7} \) S cm\(^{-1}\). The highest Li+ ion conductivities in this class of materials (up to \( 10^{-3} \) S cm\(^{-1}\)) are found in the isotopic thio-LISICON solid solutions with composition \( Li_{1-x}Ge_{1-x}P_{2}S_{12} \) (\( x = 0.75 \)).

Recently, Li+ ion conductivity has been significantly improved in the structurally related tetragonal solid solution with composition \( Li_{11-x}M_{2-x}P_{1+x}S_{12} (M = Si, Ge, Sn) \), which displays Li+ ion conductivities on the order of \( 10^{-2} \) S cm\(^{-1}\) at room temperature for \( Li_{10}GeP_{2}S_{12} \) and \( Li_{10}SiP_{2}S_{12} \).

The crystal structure of the layered lithium intercalated tin sulfide \( Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}] \) shows Li distributed both in and between Li/Sn-ordered honeycomb sulfide layers.\(^{11,12} \) Brant \textit{et al.}\(^{11,12} \) reported \( Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}] \) to be a fast Li+ ion conductor with \( \sigma_{298K} = 1.5 \times 10^{-5} \) S cm\(^{-1}\), environmental stability under ambient conditions and high thermal stability up to 750 °C. \( Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}] \) crystallizes in the Na2IrO3 structure type (space group \( C2/c \)), an ordered variant of the layered \( ZnFeO_{2} \) structure. The structure of \( Na_{2}IrO_{3} \) differs from that of the \( Na_{2}FeO_{2} \) (which consists of alternating layers of Na and Fe octahedrally coordinated by oxygen) in having a mixed Na/Fe layer that orders in a honeycomb fashion. Both of these structures are related to the Delafosse structure, which has linear, rather than octahedral, coordination of the interlayer cation. Note that these materials are structurally related to the widely used electrode material LiCoO2.\(^{15,16} \) We previously reported a related compound with a stoichiometry of \( Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}] \) to exist in the ternary Li–Sn–S phase diagram, however the details of its crystal structure remained unsolved.\(^{11} \) Aqueous exfoliation of the layered \( Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}] \) into single layer nanosheets is extremely facile.

For easier comparison of the two materials \( Li_{0.6}[Li_{0.3}Sn_{0.6}S_{2}] \) and \( Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}] \) we use the general formula \( Li_{1-x}Li_{x}Sn_{1-x}S_{2} \), where \( Li_{1-x} \) corresponds to the interlayer Li atoms and \( Li_{x} \), \( Sn_{1-x} \) to the covalent layers. This representation displays that for each Li atom located in the layers, three times as many Li+ ions must be found in between the layers. The lithium content within the tin sulfide layer decreases from Li:Sn = 1:2 in \( Li_{0.3}Sn_{0.6}S_{2} \) (\( x = 0.33 \); \( Li_{0.3}Sn_{0.7}S_{2} \)) to 1:4 in \( Li_{0.5}Sn_{0.5}S_{2} \) (\( x = 0.2 \)). Similarly, the Li content in the interlayer gallery decreases from full occupation in \( Li_{0.3}Sn_{0.6}S_{2} \) to only 60% occupancy in \( Li_{0.5}Sn_{0.5}S_{2} \). With respect to these interlayer Li+ ions, \( Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}] \) can be viewed as a lithium-depleted version. Here the Li+ deficiency is compensated by Sn2+ excess, while in various other layer compounds discussed in the literature the interlayer cation deficiency is compensated by electron holes, corresponding to an oxidation of framework species.\(^{17,18} \) Here we show that the ionic conductivity of \( Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}] \) is as high as \( 10^{-5} \) S cm\(^{-1}\) at room temperature (determined by PFG NMR), which is more than two orders of magnitude higher than the ionic conductivity of \( Li_{0.6}[Li_{0.3}Sn_{0.7}S_{2}] \) on cold pressed powders. This is most likely related to the reduced amount of Li+ ions in between the layers, in agreement with a recent report concerning Na+ ion conductivity in \( Na_{2}FeO_{2} \) related materials.\(^{19} \) Our findings therefore strengthen the argument that interlayer cation depleted layered materials can be excellent ion conductors with Li+ ion conductivities \textit{en par} with those of 3D thio-LISICON materials. The high ionic conductivity of \( 10^{-3} \) S cm\(^{-1}\) and the very low electronic contribution (< \( 10^{-9} \) S cm\(^{-1}\)) places \( Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}] \) among the best solid Li+ electrolytes known to date.

**Experimental section**

Polycrystalline \( Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}] \) was synthesized in three different procedures at 700–750 °C, with three different starting reagents. Synthesis 1 was reported by Kuhn \textit{et al.},\(^{13} \) synthesis 2 was performed with \( Li_{2}CO_{3}, SnS_{2} \) and S, and synthesis 3 with \( Li_{2}S, SnS_{2} \) and S. For synthesis 1 an amorphous phase was found in the NMR spectra which could be avoided by using syntheses 2 or 3. Further details are described in the ESI.\(^{1} \) Note that without quenching – for synthesis 2 and 3 – we did not find any evidence for the existence of other ternary compounds in addition to \( Li_{0.6}[Li_{0.3}Sn_{0.7}S_{2}] \) and \( Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}] \). Other compositions prepared by our method resulted in mixed phases. Small single crystals were obtained by subliming \( Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}] \) at 750 °C in a vacuum-sealed ampoule for several days.

Single-crystal X-ray diffraction (SXRD) was conducted on a Bruker APEX II diffractometer with Mo-K\(_{α}\) radiation at 100 K. Unit cell refinement and data integration was performed with the Bruker APEX II software and CELL_NOW. The crystal structure was refined using the full-matrix least-squares method on \( F^{2} \), implemented through SHELXL-2013 and WinGX.

SEM/EDX measurements were performed with a scanning electron microscope (SEM; Vega TS 5130 MM, Tescan) using a Si/Li detector (Oxford). ICP-AES was done with a Vista Pro ICP-AES spectrometer.

\( Li \) and \( ^{11} \)Sn solid-state (ss) NMR spectra were obtained on a Bruker Avance III 400 MHz instrument (\( B_{0} = 9.4 \) T) at Larmor frequencies of 400 MHz, 58.88 MHz and 149.12 MHz, respectively.\(^{7} \)\( Li \) T1 relaxation time measurements at various temperatures were performed to determine the activation energies. Further details are described in the ESI.\(^{1} \)\( ^{7} \)Li Pulsed Field Gradient (PFG) NMR measurements were performed on a Bruker Avance-III 400 MHz instrument (magnetic field of 9.4 T, and \( ^{7} \)Li Larmor frequency of 155.56 MHz), equipped with a diff60 single gradient diffusion probe. The probe allows for pulse field gradients \( g \) of up to 30 T m\(^{-1}\) and variable temperature measurements up to +150 °C. The diffusion measurements were accomplished using a Stimulated Echo pulse sequence.\(^{20} \) The echo attenuation curves \( S(g,\delta,\Delta) \) were processed using the Stejskal–Tanner equation,\(^{21} \)

\[ S(g,\delta,\Delta) = E(-g^{2}\delta^{2}D\Delta/(\Delta - \delta/3)) \]
where $\gamma = 1.398 \times 10^8 \text{ Hz}^{-1}$ is the $^7\text{Li}$ gyromagnetic ratio, $g$ is the strength of the pulse field gradient of duration $\delta$, $D$ is the effective diffusion coefficient, and $\Delta$ is the time interval between the field gradients that defines the diffusion time scale. Diffusion constants near room temperature, where diffusion is slow, were obtained by varying the gradient strength between 0.1 and 30 T m$^{-1}$ at a fixed $\delta = 1.0$ ms, in a range of $\Delta = 10$–100 ms. At higher temperatures, when diffraction effects in an attenuation curve become apparent, the diffusion measurements were performed by varying $\delta$ in a range of 0.25–4.0 ms at a fixed gradient strength $g$ of 900 G, and a fixed $\Delta$ of 20 ms.

For electrical conductivity measurements two different configurations were used. In each of them, the thoroughly ground powder was filled into a Teflon die with an inner diameter of 4 mm. The powder was compressed by two stamps (4 mm) and they were fixed in a uniaxial press during the measurements. The density of the cold-pressed powders is estimated to be 75%.

In configuration 1, in order to avoid any undesired reaction between lithium and the stamp to ensure a good contact, a 0.1 mm thick gold foil was placed between the plunger and the powder at both sites. Configuration 1 is described therefore as $\text{Au}[[\text{Li}_{0.6}[\text{Li}_{0.2}\text{Sn}_{0.8}\text{S}_2]]\text{Au}$. For configuration 2, an additional small layer of LiAl alloy was placed between the sample and the gold foil at both sides ($\text{Au}[[\text{LiAl}]\text{Li}_{0.6}[\text{Li}_{0.2}\text{Sn}_{0.8}\text{S}_2]]\text{LiAl}[\text{Au}]$). The alloy acts as a lithium reservoir and allows for both ionic and electronic charge transfer – thereby, concentration polarization effects on the electrodes can be avoided. For each configuration, DC galvanostatic polarization/depolarization measurements were carried out with a Keithley 2604B source-meter, while impedance spectra were analyzed using the commercial software Z-View by Scribner Assoc. The density of the cold-pressed powders is estimated to be 75%.

**Results and discussion**

The crystal structure of Li$_{0.6}$[Li$_{0.2}$Sn$_{0.8}$S$_2$] was solved by single crystal X-ray diffraction in the monoclinic space group $C2/m$ (no. 12) with $a = 19.217(7)$ Å, $b = 3.6996(14)$ Å, $c = 6.529(2)$ Å, and $\beta = 109.056(4)^\circ$ to a $wR_2$ value of 6.6% and a $R_1 (F_o > 2\sigma(F_o))$ value of 2.4%. The structure is displayed in Fig. 1, the crystallographic details are listed in Table 1, the atomic coordinates and the anisotropic displacement parameters are given in Tables S1 and S2 (see ESI†). We have confirmed with powder X-ray diffraction that the solution of the single crystal data largely corresponds to the structure of the bulk phase. Slight differences in the powder X-ray pattern compared to the single crystal structure are observed in the 2θ range between 7 and 12°, which is diagnostic of different concentrations of stacking faults in the polycrystalline powder compared to the single crystal, as also observed by J. A. Brant et al.$^{14}$ for Li[Li$_{0.33}$Sn$_{0.67}$S$_2$]. The observed monoclinic superstructure arises from the rhombohedral subcell through cation ordering within the layers, which is commonly observed in materials of this type.$^{38-41}$ The chemical formula was confirmed using SEM/EDX and ICP-AES (Fig. S2, Tables S3 and S4, ESI†). SS NMR was performed to confirm the $+4$ oxidation state of Sn (Fig. S3–S6, see ESI†).

The structure of Li$_{0.6}$[Li$_{0.2}$Sn$_{0.8}$S$_2$] is related to the structure of its parent compound Li[Li$_{0.33}$Sn$_{0.67}$S$_2$]. Both compounds (Li[Li$_{0.33}$Sn$_{0.67}$S$_2$] in $C2/c$ and Li$_{0.6}$[Li$_{0.2}$Sn$_{0.8}$S$_2$] in $C2/m$) are...
composed of layers of edge-sharing tin sulfide octahedra which face in the same direction, and Li\(^+\) ions that are sandwiched between the layers (see Fig. 1). The Sn atoms are arranged in a hexagonal net; part of the Sn is substituted by Li in both structures and (partial) ordering of the Sn atom in the layer causes a monoclinic supercell. In Li\([\text{Li}_{0.33}\text{Sn}_{0.67}\text{S}_2]\) the Sn atoms are arranged in a honeycomb order with Li occupying the voids of the honeycomb. In \(\text{Li}_6\text{[Li}_{0.6}\text{Sn}_{0.4}\text{S}_2]\), however, the reason for the monoclinic supercell is preferential occupancy of two different Sn positions. This preferential occupancy causes an enlargement of the \(a\) axis compared to the rhombohedral subcell by a factor of \(3\sqrt{3}\). Differences in intralayer ordering were also confirmed by ssNRM (see ESIF). Upon Li\(^+\) ion depopulation of the interlayer Li sites, compensated by increasing the amount of Sn in the honeycomb layers, the interlayer distance \(d\) slightly increases from 6.11 Å in \(\text{Li}[\text{Li}_{0.33}\text{Sn}_{0.67}\text{S}_2]\) to 6.17 Å in \(\text{Li}_6[\text{Li}_{0.6}\text{Sn}_{0.4}\text{S}_2]\) due to decreased electrostatic attraction forces between the layers, as clearly seen in the PXRD patterns (Fig. S1, ESIF).

Both \(\text{Li}[\text{Li}_{0.33}\text{Sn}_{0.67}\text{S}_2]\) and \(\text{Li}_6[\text{Li}_{0.6}\text{Sn}_{0.4}\text{S}_2]\) display O-type stacking of the layers, where the interlayer atoms are octahedrally coordinated by sulfur atoms, which is similar to many other honeycomb type compounds.\(^{22-27}\) In \(\text{Li}_6[\text{Li}_{0.6}\text{Sn}_{0.4}\text{S}_2]\), however, only 77% of the interlayer lithium atoms present occupy those octahedral sites; the other 23% occupy tetrahedral sites that are closer to the covalent layers. We assume that due to the unoccupied positions in the interlayer gallery and the slightly increased distance between the layers, the occupation of both octahedral and tetrahedral sites is possible. The distance between Li in octahedral and tetrahedral positions is about \(2.3\) Å, which makes it possible, in principle, that lithium occupies adjacent tetrahedral and octahedral sites, according to the Shannon radius of Li\(^+\) (0.59 for tetrahedral and 0.76 for octahedral coordination).\(^{28}\) Nevertheless, it seems more likely that due to the vacant sites either a tetrahedral or an octahedral position is occupied in a particular local region of the structure. This is also supported by the fact that due to the Li depletion, out of all possible octahedral and tetrahedral interlayer Li positions in sum only 38% are occupied.

To add evidence to the distribution of the Li positions, we performed \(^6\)Li ssNMR studies on both \(\text{Li}[\text{Li}_{0.33}\text{Sn}_{0.67}\text{S}_2]\) and \(\text{Li}_6[\text{Li}_{0.6}\text{Sn}_{0.4}\text{S}_2]\). The NMR spectra of both samples exhibit narrow, well-resolved signals suggesting phase pure samples with high crystallinity. The \(^6\)Li NMR spectrum of \(\text{Li}[\text{Li}_{0.33}\text{Sn}_{0.67}\text{S}_2]\) shows two signals at \(-0.2\) ppm and \(1.5\) ppm with an integral ratio of around 75 : 25 (Fig. 2, upper panel). This agrees well with the two different types of Li – 75% (Li1 + Li3) between the layers (see Table S1, ESIF) and Fig. 2, lower panel). There are five types of lithium atoms in the unit cell; two (Li1 + Li2) are within the layer and three (Li3–Li5) reside in the interlayer gallery, where Li3 and Li4 occupy octahedral positions and Li5 occupies a tetrahedral position. The labelling of each atom position is displayed in Fig. 2. In agreement with \(\text{Li}[\text{Li}_{0.33}\text{Sn}_{0.67}\text{S}_2]\) the signal of intra-layer Li is likely downfield shifted due to the Sn–S environment, hence the intralayer lithium atoms, Li1 and Li2, can be assigned to the signal at 2.2 ppm. Furthermore, we assume that Li5, which is tetrahedrally coordinated, behaves similar to the intralayer Li due to the closeness to the tin sulfide layers. The two octahedrally coordinated Li\(^+\) ions Li3 and Li4 can be assigned to the signal at 1.1 ppm. From the crystal structure we then would expect an integral ratio of 42 : 58. This is almost what we see in the NMR spectrum (47 : 53). Note that single crystal XRD data were taken at room temperature. Fig. 2 (top left) displays the diffusion coefficient \(D_{\text{NMR}}\) as a function of the diffusion time \(\Delta\). The longer \(\Delta\), the higher is the likelihood of crossing a grain boundary. Note that the mean free path for diffusion is on the order of \(\sim 1\) μm for diffusion times in the ns range. SEM analysis of the powder samples show that grain sizes vary between 100 μm and less than 1 μm (see Fig. S2, ESIF). This indicates that grain boundaries can affect Li\(^+\) ion diffusion at all diffusion times \(\Delta\) measured here. Note that the values of the room temperature diffusion coefficient \(D_{\text{NMR}}\) are on the order of \(10^{-11}\) m\(^2\) s\(^{-1}\), which is notably higher than the ones reported for LGPS (\(\sim 10^{-12}\) m\(^2\) s\(^{-1}\)).\(^{29}\) We converted \(D_{\text{NMR}}\) to conductivity \(\sigma_{\text{NMR}}\) using the extended Nernst–Einstein relation \(\sigma_{\text{NMR}} = (D_{\text{NMR}}nZ^2e^2) (k_BT)^{-1}\) (see ESIF). We find
that for the shortest diffusion time the conductivity reaches a value of \( \sigma_{\text{NMR}} = 9.3 \times 10^{-3} \text{ S cm}^{-1} \). Note that for this analysis we only included interlayer lithium as participants of ion transport (see TOPOS calculations below). The overall \(^7\text{Li} \) NMR signal in the static sample consists of two unresolved components with substantially different spin–spin relaxation times \( T_2 \). Only the part of the signal with the longer \( T_2 \) contributes to the diffusion attenuation curve. That part corresponds to the lithium in the interstitial space and has been consecutively used in all further calculations related to the conductivity (see Fig. S10, ESI†). This was further confirmed by 2D exchange experiments – below ~330 K we do not see any exchange between intra and interlayer lithium. The grain boundary conductivity of \( 9 \times 10^{-4} \text{ S cm}^{-1} \) (\( \Delta t = 100 \text{ ms} \)) assessed by PFG NMR exceeds the value measured with d.c. measurements, \( 1.2 \times 10^{-4} \text{ S cm}^{-1} \).

At an elevated temperature of 407 K, \( D_{\text{NMR}} = 2.6 \times 10^{-10} \text{ m}^2 \text{s}^{-1} \) is obtained which corresponds to \( \sigma_{\text{NMR}} = 6.7 \times 10^{-2} \text{ S cm}^{-1} \). PFG NMR measurements at high temperatures displayed scattering effects that likely arise from the anisotropic lithium movement within the crystals. These problems have been previously observed in different anisotropic materials.\(^{30,31}\)

Owing to the scattering effects at high temperatures, we could not determine activation energies with PFG NMR. Further, the preparation of dense samples by sintering suitable for precise impedance spectroscopy measurements was unsuccessful. Therefore, we performed \(^7\text{Li} \) \( T_1 \) relaxation time measurements in a region of slow molecular motion at varying temperatures from 320–440 K to determine activation energies (see Fig. 3).\(^{32,33}\) We find an activation energy of \( E_{\text{act}} = 0.17 \text{ eV} \) that is comparable to values for the currently best LISICONs such as \( \text{Li}_{10}\text{GeP}_2\text{S}_12 \) (0.21 eV) and \( \text{Li}_1\text{Si}_{13}\text{PS}_{12} \) (0.19 eV).\(^{34}\) Nevertheless, different mechanisms can be present simultaneously and we cannot determine which mechanism contributes most.

Fig. 4 summarizes the main findings of the electrical conductivity measurements (both d.c. galvanostatic polarization/depolarization and a.c. impedance spectroscopy), which were carried out at room temperature (298 K) and – due to hydration of the material in air – within an argon filled glove box using two different experimental setups with ion-blocking and non-blocking electrodes, respectively (Au|Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}]|Au and Au|LiAl|Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}]|LiAl|Au). In the first case (Fig. 4, panel a), the voltage change after switching on a d.c. current of 1 nA (at approximately \( t = 70 \text{ min} \)) is characteristic of a slow polarization process due to the accumulation of mobile charge carriers at the sample/electrode interface. Note that turning off the d.c. current (at \( t = 400 \text{ min} \)) results in an equally slow depolarization process. As in this case ion-blocking Au electrodes were used, this clearly indicates that the majority of mobile charge carriers in \( \text{Li}_{0.6}[\text{Li}_{0.2}\text{Sn}_{0.8}\text{S}_{2}] \) are ions. From Fig. 4a one can estimate the electronic transference number of \( \text{Li}_{0.6}[\text{Li}_{0.2}\text{Sn}_{0.8}\text{S}_{2}] \) to be less than \( 10^{-5} \). Further details for the calculation of the transference number are described in the ESI†; the refined evaluation shifts the upper limit of the transference number to \( 10^{-6} \) (see Fig. S11, ESI†). In addition, an impedance spectrum acquired in this configuration (ion blocking electrodes) is shown in Fig. S12 (ESI†); a Warburg behavior can be clearly recognized in the low frequency range. The high ionic conductivity is confirmed by the experiments which were performed in the non-blocking Au|LiAl|Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}]|LiAl|Au configuration (Fig. 4, panel b). In this case, the response of the voltage on the application of a d.c. current of 5 \( \mu \text{A} \) is instantaneous and corresponds to an effective Li\(^+\) ion conductivity of the sample of \( 1.2 \times 10^{-4} \text{ S cm}^{-1} \). In this configuration, also impedance spectroscopy measurements were carried out and a representative complex impedance spectrum is shown in Fig. 4c. Here, the spectrum is rather distorted and seems to be composed of at least two contributions. This is not surprising as the sample consists of cold-pressed powders, in which contributions from the bulk, grain boundaries, electrodes and current constrictions (between adjacent particles) can be present. It is noteworthy that, in the low frequency range, the intercept of
the spectrum with the real axis ($\sim 12.500 \, \Omega$) is in good agreement with the resistance value that one can extract from the voltage change after the application of the d.c. current (in Fig. 4b) the change of voltage corresponds to a resistance of $11600 \, \Omega$). By fitting the high frequency part of the spectrum with a resistance $R$ in parallel to a constant phase element $Q$ with which the capacitance can be obtained as $C = (R^{1-n}Q)^{1/2n}$ with $n$ being an additional fitting parameter) a capacitance value of $2.6 \times 10^{-10} \, F$ is obtained, which corresponds to an apparent relative dielectric constant of 4200. Since $Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}]$ is not expected to possess such a large dielectric constant (e.g. it is not a ferroelectric material), this large capacitance cannot be ascribed to the bulk properties of the material, but rather seems to stem from the grain-to-grain contacts. This is supported also by the fact that the extrapolation of the spectrum towards frequencies exceeding $1 \, MHz$ results in a finite intercept at about $130 \, \Omega$ (see inset in Fig. 4c) which corresponds most likely to the bulk properties. The inset shows the finite intercept of the extrapolation of the spectrum towards frequencies exceeding $1 \, MHz$ at around $130 \, \Omega$ (see also Fig. S13 in the ESI†) indicating a bulk conductivity that is two orders of magnitudes greater. The intercept of the fitted semicircle in the low frequency range most likely belongs to grain boundary resistance. The suppressed semicircle indicates the presence of a second contact resistance, which is not fitted here. Unlike the high frequency intercept, the low frequency intercept is sensitive to the pressure applied to the powder.

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ions are assumed to be able to jump between different sites if open channels between two sites exist. A channel is considered to be accessible for lithium motion if the sum of the radii of a lithium and a framework atom (here sulfur) does not exceed the channel radius by more than 10–15%. For Li[Li_{0.33}Sn_{0.67}S_{2}] we find that Li migrates through the covalent Li/SnS_{2} layers, along the [101] direction (see Fig. S8, ESI†). Following the calculated pathways, lithium migration is only possible passing through unoccupied tetrahedral sites. The calculation for Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}] is more difficult as TOPOS is not able to handle mixed occupancies. Therefore, we chose four different configurations: (i) all Sn/Li positions are fully occupied by tin and all interlayer positions are fully occupied by lithium, (ii) all Sn2/Li2 positions are occupied by tin and Sn1/Li1 positions as well as interlayer positions are occupied by lithium, (iii) all Sn1/Li1 positions are occupied by tin and Sn2/Li2 positions as well as interlayer positions are occupied by lithium, (iv) all tin and all lithium positions are fully occupied by lithium. All these configurations can be seen as borderline cases which only partially reflect the single-crystal X-ray structure. (Note, however, that charge balance is not considered by TOPOS and the calculations are purely geometrical in nature). Interestingly, the migration pathway analysis leads to chains along b in all four cases (see Fig. S9, ESI†). This can arise from the fact that the anion framework (here sulfur), which greatly influences the Li motion, is the same in all calculations. In contrast to Li[Li_{0.33}Sn_{0.67}S_{2}] though, the migration pathway does not cross the covalent layers. The calculated migration pathway of lithium is marked by white dotted lines in the electron density maps (Fig. 5, top). These calculations lend further evidence that lithium movement is only possible via tetrahedrally coordinated sites. There is no hopping from octahedral to octahedral voids in both structures but only hopping from octahedral to tetrahedral sites and vice versa.

Conclusions

We have shown that Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}] is a fast Li⁺ ion conductor with a bulk conductivity of ~10⁻² S cm⁻¹ at room temperature, exceeding electronic contributions by at least 7 orders of magnitude. This places Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}] among the best solid Li electrolyte materials known to date. While being compositionally simpler, the new superionic conductor Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}] is also structurally distinct from the known LGPS solid electrolytes and can be viewed formally as a Li sulfide-depleted version of Li[Li_{0.33}Sn_{0.67}S_{2}], a recently reported Li⁺ ion conductor with a much lower conductivity. By analyzing the structure–property relationships in the two compounds by means of single crystal X-ray diffraction and ssNMR we demonstrate that removal of Li⁺ ions from the interlayer gallery significantly improves the ionic conductivity. Clearly, such partial occupancy slightly decreases the number of mobile Li⁺ ions while at the same time it facilitates Li⁺ ion motion by increasing the mobility of the Li⁺ ions. Therefore, besides a distinctly different Sn/Li order in the mixed Li/SnS₂ layers, the Li⁺ ions in Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}] have more space to move, due to the slightly larger layer distance in Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}], but also to the lower occupancy of the Li sites located in between the layers. Importantly, we show that interlayer Li does not only occupy the central octahedral but also tetrahedral sites, which is critical for facile ion movement as it opens up a Li trajectory involving face-sharing tetrahedral and tetrahedral voids with lower activation energies for Li hopping. In Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}], however, Li⁺ ions are only found in the octahedral positions which are fully occupied, rendering movement of those Li⁺ ions more difficult. Our study therefore proves that the partial occupancy of interlayer cation sites in layered NaCl-analogues and Delafossite-related structure types can significantly enhance ionic conductivity as recently also seen in Na₃₋ₓSbx₋₄Sc₀₋₄SbNaO₋₄. This effect can be very important for further studies on those materials since many Delafossite-related compounds have been reported to also exist in cation depleted versions. In the non-optimized cold-pressed samples, the transport measurements indicate high bulk conductivities (~10⁻² S cm⁻¹) along with significant contact resistances (~10⁻⁴ S cm⁻¹). Given the structural anisotropy of Li_{0.6}[Li_{0.2}Sn_{0.8}S_{2}], even higher conductivity can be expected for a single crystalline sample along the ab plane. Further studies on large enough single crystals and densified sintered samples are required for a deeper analysis. As far as the applicability of the new solid electrolyte is concerned, it will be key to combine it with electrodes of moderate Li chemical potential, or to find ways of kinetic passivation. Studies along those lines are currently ongoing in our lab.

Acknowledgements

We thank Marie-Luise Schreiber for ICP-AES measurements. Financial support by the Max Planck Society, Nanosystems Initiative Munich (NIM), Center for Nanoscience (CeNS) and Fonds der Chemischen Industrie is gratefully acknowledged. Work at Princeton was supported by the NSF-sponsored MRSEC at Princeton University, grant DMR-1005438. LMS gratefully
acknowledge support from the Minerva fast track fellowship by the Max Planck Society.

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