

www.acsmaterialsletters.org

🔤 😳 💽

Mechanochemical Synthesis and Structure of Lithium Tetrahaloaluminates, LiAIX₄ (X = Cl, Br, I): A Family of Li-Ion Conducting Ternary Halides

Nicolás Flores-González, Nicolò Minafra, Georg Dewald, Hazel Reardon, Ronald I. Smith, Stefan Adams, Wolfgang G. Zeier, and Duncan H. Gregory*



ACCESS

Cite This: ACS Materials Lett. 2021, 3, 652-657



Article Recommendations

S Supporting Information

ABSTRACT: State-of-the-art oxides and sulfides with high Liion conductivity and good electrochemical stability are among the most promising candidates for solid-state electrolytes in secondary batteries. Yet emerging halides offer promising alternatives because of their intrinsic low Li⁺ migration energy barriers, high electrochemical oxidative stability, and beneficial mechanical properties. Mechanochemical synthesis has enabled the characterization of LiAlX₄ compounds to be extended and the iodide, LiAlI₄, to be synthesized for the first time (monoclinic P2₁/c, Z = 4; a = 8.0846(1) Å; b = 7.4369(1) Å; c = 14.8890(2) Å; β = 93.0457(8)°). Of the tetrahaloaluminates, LiAlBr₄ exhibited the highest ionic conductivity at room temperature (0.033 mS cm⁻¹), while LiAlCl₄ showed a

III Metrics & More



conductivity of 0.17 mS cm⁻¹ at 333 K, coupled with the highest thermal and oxidative stability. Modeling of the diffusion pathways suggests that the Li-ion transport mechanism in each tetrahaloaluminate is closely related and mediated by both halide polarizability and concerted complex anion motions.

eplacing flammable liquid electrolytes in conventional Li-ion batteries (LIBs) with solid-state alternatives could lead to a breakthrough in battery safety and longevity. Moreover, otherwise inaccessible high energy density cells (using Li-metal anodes and high-voltage cathodes) could become a reality by employing thermodynamically stable solid-state electrolytes (SSEs) in all-solid-state batteries (SSBs).¹ Inorganic SSEs with sufficiently high ionic conductivity and chemical/electrochemical stability are almost within reach. Among them, oxide- and sulfide-based materials have been the main focus of research because of the remarkable ionic conductivity that can be achieved at room temperature. However, oxides lack mechanical strength and require high processing temperatures, whereas both the narrow electrochemical windows and limited stability of sulfides have proved challenging to their adoption as SSEs.^{2,3} Among alternatives, the complex halides, $\hat{Li}_3 M^{III} X_6$ ($M^{III} = Sc, Y, In$, La, Ho, Er; X = halogen) and $Li_{3-x}M_{1-x}Zr_xCl_6$ (M^{III} = Y, Er), are raising interest with appreciable ionic conductivity at room temperature, low activation energies for Li⁺ migration and wide electrochemical windows.⁴⁻¹¹ As originally discovered in the

1990s, the synthesis of $\text{Li}_3 M^{\text{III}} X_6$ requires several steps including high temperature annealing.¹² By comparison, ternary lithium-light element halides, including LiAlCl₄, have become well-known over the past four decades on account of remarkable ionic conductivity in the solution and molten states. With liquid LiAlCl₄·6SO₄ showing Li⁺ ionic conductivity of >0.10 S cm⁻¹ at room temperature,¹³ such halides have attracted renewed scrutiny recently,^{14,15} while in solution, LiAlX₄ (X = Cl, Br) can surpass the Li-ion conductivity of the ubiquitous electrolyte, LiPF₆.¹⁶ By contrast, the structure and conductivity of lithium tetrahaloaluminates in the solid state have scarcely been studied and only very recently has solid LiAlCl₄ been identified as one of several promising halides for

Received:January 22, 2021Accepted:April 15, 2021Published:April 20, 2021





electrolytes in SSBs.¹⁷ In fact, the ionic conductivity of monoclinic LiAlCl₄ was first reported by Weppner and Huggins using DC polarization measurements in 1977.¹⁸ At 298 K, single crystals of LiAlCl4 were reported with a conductivity of 1.2×10^{-6} S cm⁻¹, increasing to 1.4×10^{-4} S cm⁻¹ at 413 K (just below the melting temperature). The feasibility of using LiAlCl₄ as a SSE was first demonstrated 15 years later by deploying it in a $\text{Li}_x \text{TiS}_{2(s)}/\text{LiAlCl}_{4(s)}/$ $Li_{1-x}CoO_{2(s)}$ solid-state cell (0 < x < 0.45) at 373 K.¹⁹ The cell exhibited an open-circuit potential of 2.1 V in the charged state. Moreover, it showed excellent discharge characteristics at current densities up to 0.1 mA cm⁻² with minimal capacity loss over 100 charge-discharge cycles. Despite this promising result, there has been a lack of studies since. The corresponding bromide is uncharacterized, while the iodide has never been isolated. Consequently, the conductivity of either is unknown in the solid state.

Inspired by the recent performance of halides in the solid state and by the historically promising properties of halide salts, we were motivated to investigate the structures, stabilities and electrochemical properties of the solid lithium tetrahaloa-luminates, LiAlX₄, (X = Cl, Br, I) systematically. Here, we demonstrate how mechanochemistry can be employed to synthesize the tetrahaloaluminates in one step, without heating. This is critical for synthesizing powders of the low melting point iodide, which we could not isolate thermally. The high purity, bulk samples so-obtained have enabled us to determine the structures of the halides and to make evaluations of their Li⁺ conductivity. Both thermal and electrochemical oxidative stability have also been determined. Some preliminary hypotheses for cation conductivity mechanisms in the lithium tetrahaloaluminates can be proposed on the basis of the data.

High-purity LiAlX₄ (X = Cl, Br, I) powders were synthesized by milling the respective component binary halides in an inert atmosphere (Tables S1 and S2 and Figures S1–S8). Although lab-based powder X-ray diffraction (PXD) provided basic crystal structure models of the halides (Figures S9–11), we undertook synchrotron PXD (SPXD) and time-of-flight (ToF) powder neutron diffraction (PND) experiments to locate the Li positions accurately and to determine anisotropic thermal displacement parameters, allowing full characterization of the underlying structural chemistry of the LiAlX₄ materials. Figure 1 shows the profile fits for LiAlI₄, from structure refinement against PND and SPXD data, respectively. The respective plots



Figure 1. Room-temperature profile fits from Rietveld refinement of the structure of LiAlI₄ against: (a) ToF PND data ($\langle 2\theta =$ 92.59° \rangle detector bank; Polaris, ISIS),^{22,23} and (b) SPXD data ($\lambda =$ 0.56466 Å; X04SA, PSI). Experimental (black), calculated (red), and difference profiles (blue) are shown; vertical markers indicate Bragg reflection positions for LiAlI₄ (green) and LiI (purple), respectively.

for the chloride and bromide analogues can be found in the Supporting Information (Figures S12, S13, and S15).

Crystallographic data from these refinements are collated in Tables S3-S18. Our room temperature diffraction data confirmed that LiAlBr₄ and LiAlI₄ are isostructural to the chloride analogue (monoclinic, space group $P2_1/c$). The structural models for the bromide and iodide were further assessed by means of the Global Instability Index (GII),^{20,21} which corroborated their plausibility with low values of 0.10 and 0.07, respectively. The crystal structure of the haloaluminates can be described as a slightly distorted hcp X^{-} sublattice within which Li^{+} and Al^{3+} occupy octahedral and tetrahedral interstices, respectively. The extended structure can be considered to be constructed from distorted LiX₆ octahedra and AlX₄ tetrahedra. Two LiX₆ octahedra link across a common edge to form "Li₂X₁₀ dimers". Each Li-X dimer is connected to four others by 2 axial and 2 equatorial vertices in a "trans" confirmation that creates stepped or buckled layers that propagate in all three dimensions. Meanwhile, each AlX₄ tetrahedron is connected to one Li-X dimer via two edges and two other dimers by one vertex each (Figure 2a and b).



Figure 2. Crystal structure of mechanochemically synthesized LiAlI₄ ($P2_1/c$) projected along the *b*-axis as visualized with VESTA²⁶ (a) showing a polyhedral representation of the extended structure and the linking of Li-X "dimers", (b) showing the linkage between an LiX₆ octahedron and neighboring AlX₄ tetrahedra, and (c) highlighting the positions of the Li⁺ cations with respect to the isolated haloaluminate anions: Li (4*e*, green spheres), Al (4*e*, light blue spheres), and I (4*e*, purple).

Alternatively, considering only complex $[AlX_4]^-$ anions and Li⁺ cations, then the latter can be seen to occupy space within "pseudo-layers" between the isolated haloaluminate tetrahedra (Figure 2c).

The unit cell expands linearly in all three dimensions as Cl^- (167 pm) is replaced by Br^- (182 pm) and I^- (206 pm),²⁴ and there is a concomitant increase in both the average Li–X and Al–X bond lengths. The monoclinic distortion of the cell decreases very slightly with increasing halide radius (Figure S16). Other than these expected differences in cell volume, our refinements hinted at differences between the structural

Гable 1. Thermal Properties ar	d Transport Data o	of Mechanochemically	Synthesized Lithium	Tetrahaloaluminates
--------------------------------	--------------------	----------------------	---------------------	---------------------

Material	Melting point (K)	Decomposition onset temperature (K)	$\sigma_{\rm RT} \times 10^5 \ ({\rm S \ cm^{-1}})$	$\sigma_0 \times 10^{-5} \ ({\rm K \ S \ cm^{-1}})$	$E_{\rm a}~({\rm eV})$
LiAlCl ₄	420.0 [419, ¹⁸ 422 ¹⁶]	~643	$2.9(2)$ [2.1^{25}]	8.6(7)	$0.473(2) [0.47,^{18} 0.50(6)^{25}]$
LiAlBr ₄	464.9 [466 ¹⁶]	~593	3.3(2)	2.5(4)	0.437(4)
$LiAlI_4$	509.0	~593	1.2(1)	0.61(7)	0.429(3)

models of the mechanochemically synthesized lithium tetrahaloaluminates and the previously reported structures of (thermally synthesized) LiAlCl₄ and LiAlBr₄ (although the latter structure was previously determined only from single crystal data at 100 K).¹⁶ First, attempts were made to refine the occupancy of the Li site in each halide, and although for X =Br, this did not vary from 100%, for X = Cl and I, respectively, values of 91(4)% and 94(2)% were obtained with slight reductions in R-factors. More interestingly, if the Li occupancy of the normally vacant i2 interstitial position (0.236, 0.014, 0.792) identified by SoftBV in the conduction mechanism (see Table S20) was simultaneously refined, then occupancies of 0.84(3) and 0.16(3) were obtained for the normal and interstitial Li sites in LiAlCl₄ when the respective thermal parameters were fixed. By contrast, no evidence for antisite mixing (Li-Al disorder) was obtained for any of the halides. Although these results are not conclusive, they do tend to support observations from solid-state NMR spectroscopy of a partially occupied interstitial site in LiAlCl₄²⁵ and provide a rationale for the Li⁺ diffusion mechanism elucidated by BVSE and MD analyses for the haloaluminates (see below). It will be interesting to see whether local structural approaches, such as pair distribution function (PDF) analysis can provide further information regarding the links between defect structure and Li-ion motion.

The thermal stabilities of LiAlX₄ were studied by simultaneous thermogravimetric-differential thermal analysis (TG-DTA). For all samples, the TG profiles are typical of thermal decomposition with volatile decomposition products (Figures S17–19). The decomposition is preceded by melting in each case (the melting points, as determined by the respective DTA peak onsets, are summarized in Table 1). The melting points of the tetrahaloaluminates increase from X = Clthrough Br to I, with the new iodide, LiAlI₄, melting at ~509 K (and decomposing from ~593 K). TG-DTA also confirmed the absence of unreacted AlX₃ in the synthesized LiAlX₄ materials with no AlX₃ melting transitions visible in the DTA data (e.g., AlI₃ melts at 461.43 K).²⁷

The ionic transport properties of mechanochemically synthesized LiAlX₄ were analyzed by variable temperature electrochemical impedance spectroscopy (EIS) with blocking Au electrodes. Figure 3a shows the room temperature Nyquist plots. For LiAlCl₄, the spectrum was fitted with an equivalent circuit consisting of one parallel constant phase element (CPE)/resistor in series with a CPE representing the behavior of the electrolyte and blocking electrodes, respectively. The capacitance of the CPE/resistor is 2.5×10^{-11} F cm⁻² with an α -value of 0.99, which together indicate a predominant bulk contribution to the impedance response.²⁸ Conversely, the LiAlBr₄ and LiAlI₄ spectra were fitted with an equivalent circuit consisting of two parallel constant phase elements (CPE)/ resistors in series with a further CPE. The capacitances of the high-frequency CPE/resistor elements are 2.4×10^{-11} (LiAlBr₄) and 1.7 \times 10⁻¹¹ F cm⁻² (LiAlI₄) with α -values of 0.96 and 0.91, respectively, which represent the ideality of the CPE and confirm a bulk-process.



Figure 3. (a) Room-temperature Nyquist plots of LiAlX₄ (X = Cl, Br, I) normalized to the thickness of the pellets, showing the impedance responses (open circles) and fits (solid lines). (b) Arrhenius plots of conductivity values obtained from temperature-dependent impedance spectroscopy.

At lower frequencies, impedance contributions are observed with capacitances of 2.5×10^{-9} (X = Br) and 8.1×10^{-7} F cm⁻² (X = I). This can be attributed to a surface layer, which may be formed on thermal decomposition during gold coating, for example.^{6,28}

Arrhenius behavior was noted for all samples in the LiAlX₄ series across a temperature range of 233–333 K (Figure 3b). Room-temperature total ionic conductivities ($\sigma_{\rm RT}$) and the parameters extracted from linear fits of the Arrhenius plots (σ_0 , E_{a}) are summarized in Table 1. We found that the room temperature Li⁺ conductivity first increases but then subsequently decreases when switching from X = Cl through Br to I. The results corroborate the premise that both the exponential prefactor, σ_0 , and the activation energy for Li⁺ diffusion decrease with increasing anion polarizability, although the differences in the E_a values for LiAlBr₄ and LiAlI₄ are not statistically significant.^{29,30} In this regard, it should be noted that a reduction in $E_{\rm a}$ does not always lead to improved ionic conductivity, since E_{a} and σ_{0} are correlated in line with the Meyer–Neldel rule.^{31,32} By considering both parameters in Table 1, it can be appreciated why LiAlI₄ might have the lowest ionic conductivity within the haloaluminate series. It should be noted that the ionic conductivity for LiAlCl₄ measured here is 1 order of magnitude higher than that reported by Weppner and Huggins.¹⁸ Although our mechanochemical syntheses are quicker and less energyintensive to that recently reported for LiAlCl₄ by Tanibata et al., the EIS data do further support the premise that ball milling positively influences the ionic conductivity in the lithium halides (likely facilitating defect formation compared to thermal synthesis methods, which are evidently not viable for X = I).²⁵ Indeed, EIS measurements performed on a pellet of the LiAlCl₄ sample that was subsequently annealed at 373 K for 14 h yielded a lower room temperature conductivity (of $1.2(2) \times 10^{-5}$ S cm⁻¹) than that of the cold-pressed mechanochemically synthesized chloride.

The comprehensive structural models obtained from neutron and synchrotron diffraction allowed the probable Li^+ diffusion pathways in the haloaluminates to be established via bond-valence site energy (BVSE) analysis.²¹ In this class of

materials, our analysis shows that the ionic conductivity is governed by the presence of intrinsic tetrahedral and octahedral interstitial sites. The BVSE map of LiAlI_4 is presented as an example in Figure 4a, and shows that a



Figure 4. (a) BVSE map showing Li⁺ migration pathways in a (100) projection of the LiAlI₄ structure, as visualized with VESTA.²⁶ The highest isosurface level of 0.64 eV over the global minimum is shown in yellow. Red dots indicate octahedral Li⁺ lattice sites and yellow spheres indicate tetrahedral/octahedral interstitial sites. (b) BVSE model of migration barriers for LiAlI₄ derived from Rietveld refinements against SPXD and PND data. The relative site energy is zero for Li⁺ lattice sites.

bottleneck for 2D conduction involves Li⁺ hopping from its normal octahedral 4*e* lattice position to an adjacent tetrahedral 4*e* site ("i7" in the BVSE map notation in Figure 4b). Alternatively, 3D conduction requires hops from/to interstitial octahedral 4*e* lattice sites ("i3" \rightarrow "i2"). Qualitatively, the BVSE models of the migration barriers in the LiAlX₄ series depict very similar energy landscapes (while noting that the overall activation energies for LiAlBr₄ and LiAlI₄ are higher than the experimental values due to the level of accuracy of SoftBV, Figures 4b, S20, and S21). These energy profiles indicate that the conduction pathways do not change significantly with the halide and that the observed differences in ionic conductivity cannot be rationalized only via a static, crystal chemistry treatment.

Empirical molecular dynamics simulations for a 768 atom 4 \times 4 \times 2 supercell of LiAlCl₄ over the temperature range 250–400 K over 1500–18000 ps harmonize to the experimentally observed conductivity (Figure S22). A more detailed analysis shows that the 2D Li⁺ motion in the *y*–*z* plane is coupled to dynamic anion disorder (librations) that in the experimental study may be facilitated by the mechanochemical synthesis. Details are given in the Supporting Information (Figures S23 and S24). Given the role of polyanion motions in other cation conductors, the combination of experimental and computational data encourages further investigations of the role of

defects and dynamic anion effects in LiAlX₄ materials and how such effects might be tuned.

In view of the promising ionic transport behavior, preliminary linear sweep voltammetry (LSV) experiments were conducted to determine the oxidative stability of LiAlX₄. InLi and a carbon + SSE composite were employed as the counter and working electrodes, respectively. The oxidative stability limits were defined by the onset potentials (E_{onset}) and calculated via linear fitting of the nonfaradaic and faradaic region.³³ The room temperature voltammograms are shown in Figure 5.



Figure 5. Room temperature linear sweep voltammogram (0.1 mV s⁻¹) of InLilLiAlX₄|LiAlX₄ + C cells. Dashed lines indicate linear fits of the faradaic region. The bottom *x*-axis shows the values of the voltage versus In/InLi, the top *x*-axis shows the corresponding values of voltage versus Li⁺/Li.³⁴

We determined that oxidation of LiAlX₄ materials starts at 3.8 (4.4), 2.8 (3.4), and 2.0 (2.6) V versus In/InLi (vs Li⁺/Li) for X = Cl, Br, and I, respectively. Density functional theory (DFT) calculations predicted the electrochemical windows of LiAlX₄ to be either $1.7-4.5 \text{ V}^{35}$ or $1.54-4.45 \text{ V}^{7}$ for X = Cl and $1.8-3.9 \text{ V}^{35}$ (vs Li⁺/Li), for X = Br. Above the oxidation limits, LiAlX₄ are predicted to produce AlX₃ and X₂, while below the reduction limits, LiAlX₄ are predicted to form Al and LiX, indicating that lithium tetrachloro- and tetrabromoaluminates are not stable against Li metal, as is true for the liquid electrolyte, LiAlCl₄·3SO₂.³⁶ These results indicate the feasibility of combining LiAlCl4 with high voltage cathode materials, while the bromide and iodide analogues would be better suited to use with lower potential electrodes, such as sulfide-based cathodes, for potential cell applications. Equally, use of LiAlBr₄ and LiAlI₄ with high voltage cathodes might be enabled with an appropriate coating of the positive electrode to prevent SSE decomposition.^{37,38} Further investigations into various half- and full-cell architectures and their performance are currently underway and will be reported elsewhere.

In summary, we have shown that high purity lithium tetrahaloaluminate powders can be easily synthesized by mechanochemical methods. This approach is extremely effective in preparing bulk quantities of LiAlX₄ including the new iodide, LiAlI₄, which could not be synthesized by thermal

methods. Synchrotron and neutron diffraction have shown not only that the bromide and iodide are isostructural to the chloride analogue but that Li vacancies and interstitials are likely prevailing features of mechanically synthesized tetrahaloaluminates. Each material exhibits appreciable Li-ionic conductivity, good thermal stability and reasonable stability to oxidation, such that the chloride, especially, might be employed with high voltage cathodes. Moreover, Li⁺ conductivity and stability can likely be improved still further by tuning the microstructure, composition and defect chemistry of the haloaluminates through doping, substitution and compositing. The LiAlX₄ family suggests these and other polyanion halides offer considerable promise as new classes of cation conductor and as candidates for testing systematically as SSEs.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsmaterialslett.1c00055.

Experimental procedures, characterization techniques, crystallographic information, TG-DTA profiles, BVSE analysis, and details of MD simulations (PDF)

AUTHOR INFORMATION

Corresponding Author

Duncan H. Gregory – School of Chemistry, University of Glasgow, Glasgow G12 8QQ, U.K.; o orcid.org/0000-0002-4585-3280; Email: duncan.gregory@glasgow.ac.uk

Authors

- Nicolás Flores-González School of Chemistry, University of Glasgow, Glasgow G12 8QQ, U.K.
- Nicolò Minafra Institute for Inorganic and Analytical Chemistry, University of Münster, 48149 Münster, Germany
- **Georg Dewald** Institute of Physical Chemistry, Justus-Liebig-University Giessen, D-35392 Giessen, Germany
- Hazel Reardon School of Chemistry, University of Glasgow, Glasgow G12 8QQ, U.K.
- Ronald I. Smith ISIS Pulsed Neutron and Muon Source, STFC Rutherford Appleton Laboratory, Didcot, Oxfordshire OX11 0QX, U.K.; orcid.org/0000-0002-4990-1307
- Stefan Adams Department of Materials Science and Engineering, National University of Singapore, 117575, Singapore; orcid.org/0000-0003-0710-135X
- Wolfgang G. Zeier Institute for Inorganic and Analytical Chemistry, University of Münster, 48149 Münster, Germany; orcid.org/0000-0001-7749-5089

Complete contact information is available at: https://pubs.acs.org/10.1021/acsmaterialslett.1c00055

Author Contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

The authors acknowledge the Advanced Human Capital Program of the National Commission for Scientific and Technological Research (CONICYT/Becas Chile/No. 72170338) for a PhD scholarship for N.F.G., the Royal Society of Chemistry for Researcher Mobility Grant M19-8459 and the EPSRC for associated funding under grant EP/ N001982/1. The authors also thank the UK Science and Technology Facilities Council (STFC) and the Material Science Beamline X04SA at the Paul Scherrer Institute (PSI) for the award of ISIS Xpress access and MESQUIK access measurements, respectively and Dr. Nicola Casati at PSI for collecting the latter data. N.M. and W.Z. are grateful for support by the Deutsche Forschungsgemeinschaft (DFG) under grant number ZE 1010/4-1.

REFERENCES

(1) Janek, J.; Zeier, W. G. A solid future for battery development. *Nat. Energy* **2016**, *1*, 16141.

(2) Han, F.; Zhu, Y.; He, X.; Mo, Y.; Wang, C. Electrochemical Stability of $Li_{10}GeP_2S_{12}$ and $Li_7La_3Zr_2O_{12}$ Solid Electrolytes. *Adv. Energy Mater.* **2016**, *6*, 1501590.

(3) Han, X.; Gong, Y.; Fu, K.; He, X.; Hitz, G. T.; Dai, J.; Pearse, A.; Liu, B.; Wang, H.; Rubloff, G.; Mo, Y.; Thangadurai, V.; Wachsman, E. D.; Hu, L. Negating interfacial impedance in garnet-based solidstate Li metal batteries. *Nat. Mater.* **2017**, *16*, 572–579.

(4) Asano, T.; Sakai, A.; Ouchi, S.; Sakaida, M.; Miyazaki, A.; Hasegawa, S. Solid Halide Electrolytes with High Lithium-Ion Conductivity for Application in 4 V Class Bulk-Type All-Solid-State Batteries. *Adv. Mater.* **2018**, *30*, 1803075.

(5) Schlem, R.; Muy, S.; Prinz, N.; Banik, A.; Shao-Horn, Y.; Zobel, M.; Zeier, W. G. Mechanochemical Synthesis: A Tool to Tune Cation Site Disorder and Ionic Transport Properties of Li_3MCl_6 (M = Y, Er) Superionic Conductors. *Adv. Energy Mater.* **2020**, *10*, 1903719.

(6) Schlem, R.; Bernges, T.; Li, C.; Kraft, M. A.; Minafra, N.; Zeier, W. G. Lattice Dynamical Approach for Finding the Lithium Superionic Conductor Li₃ErI₆. ACS Appl. Energy Mater. **2020**, *3*, 3684–3691.

(7) Wang, S.; Bai, Q.; Nolan, A. M.; Liu, Y.; Gong, S.; Sun, Q.; Mo, Y. Lithium Chlorides and Bromides as Promising Solid-State Chemistries for Fast Ion Conductors with Good Electrochemical Stability. *Angew. Chem., Int. Ed.* **2019**, *58*, 8039–8043.

(8) Li, X.; Liang, J.; Chen, N.; Luo, J.; Adair, K. R.; Wang, C.; Banis, M. N.; Sham, T.-K.; Zhang, L.; Zhao, S.; Lu, S.; Huang, H.; Li, R.; Sun, X. Water-Mediated Synthesis of a Superionic Halide Solid Electrolyte. *Angew. Chem., Int. Ed.* **2019**, *58*, 16427–16432.

(9) Xu, Z.; Chen, X.; Liu, K.; Chen, R.; Zeng, X.; Zhu, H. Influence of Anion Charge on Li Ion Diffusion in a New Solid-State Electrolyte, Li₃LaI₆. *Chem. Mater.* **2019**, *31*, 7425–7433.

(10) Liu, Y.; Wang, S.; Nolan, A. M.; Ling, C.; Mo, Y. Tailoring the Cation Lattice for Chloride Lithium-Ion Conductors. *Adv. Energy Mater.* **2020**, *10*, 2002356.

(11) Park, K.-H.; Kaup, K.; Assoud, A.; Zhang, Q.; Wu, X.; Nazar, L. F. High-Voltage Superionic Halide Solid Electrolytes for All-Solid-State Li-Ion Batteries. *ACS Energy Lett.* **2020**, *5*, 533–539.

(12) Steiner, H.-J.; Lutz, H. D. Neue schnelle Ionenleiter vom Typ $M_3^{I}M^{III}Cl_6$ (M^{I} = Li, Na, Ag; M^{III} = In, Y). Z. Anorg. Allg. Chem. **1992**, 613, 26–30.

(13) Foster, D. L.; Kuo, H. C.; Schlaikjer, C. R.; Dey, A. N. New Highly Conductive Inorganic Electrolytes: The Liquid Solvates of the Alkali and Alkaline Earth Metal Tetrachloroaluminates. *J. Electrochem. Soc.* **1988**, *135*, 2682–2686.

(14) Grundish, N.; Amos, C.; Goodenough, J. B. Communication-Characterization of $LiAlCl_4$:xSO₂ Inorganic Liquid Li⁺ Electrolyte. J. Electrochem. Soc. **2018**, 165, A1694–A1696.

(15) Ramar, V.; Pszolla, C.; Rapp, M.; Borck, M.; Zinck, L. Non-flammable Inorganic Liquid Electrolyte Lithium-Ion Batteries. *J. Electrochem. Soc.* **2020**, *167*, 070521.

(16) Scholz, F.; Unkrig, W.; Eiden, P.; Schmidt, M. A.; Garsuch, A.; Krossing, I. Synthesis, Spectroscopic Characterization, Crystal Structures, Energetics, and Thermal Stabilities of $Li[AIX_4]$ (X = Cl,

Br): Investigation and Performance of Their Electrolyte Solutions. *Eur. J. Inorg. Chem.* **2015**, 2015, 3128–3138.

(17) Kahle, L.; Marcolongo, A.; Marzari, N. High-throughput computational screening for solid-state Li-ion conductors. *Energy Environ. Sci.* **2020**, *13*, 928–948.

(18) Weppner, W.; Huggins, R. A. Ionic Conductivity of Solid and Liquid LiAlCl₄. J. Electrochem. Soc. **1977**, *124*, 35–38.

(19) Plichta, E. J.; Behl, W. K.; Vujic, D.; Chang, W. H. S.; Schleich, D. M. The Rechargeable Li_xTiS₂/LiAlCl₄/Li_{1-x}CoO₂ Solid-State Cell. *J. Electrochem. Soc.* **1992**, *139*, 1509–1513.

(20) Salinas-Sanchez, A.; Garcia-Muñoz, J. L.; Rodriguez-Carvajal, J.; Saez-Puche, R.; Martinez, J. L. Structural characterization of R_2BaCuO_5 (R = Y, Lu, Yb, Tm, Er, Ho, Dy, Gd, Eu and Sm) oxides by X-ray and neutron diffraction. *J. Solid State Chem.* **1992**, *100*, 201– 211.

(21) Chen, H.; Wong, L.; Adams, S. SoftBV - a software tool for screening the materials genome of inorganic fast ion conductors. *Acta Crystallogr., Sect. B: Struct. Sci., Cryst. Eng. Mater.* **2019**, *75*, 18–33.

(22) Smith, R. I.; Hull, S.; Tucker, M. G.; Playford, H. Y.; McPhail, D. J.; Waller, S. P.; Norberg, S. T. The upgraded Polaris powder diffractometer at the ISIS neutron source. *Rev. Sci. Instrum.* **2019**, *90*, 115101.

(23) Flores-González, N. Insights into the structure of mechanochemically-synthesised LiAlX₄ (X = Cl, Br, I) by powder neutron diffraction. *STFC ISIS Neutron and Muon Source Data Journal* **2018**, RB1890322.

(24) Shannon, R. D. Revised effective ionic radii and systematic studies of interatomic distances in halides and chalcogenides. *Acta Crystallogr., Sect. A: Cryst. Phys., Diffr., Theor. Gen. Crystallogr.* **1976**, 32, 751–767.

(25) Tanibata, N.; Takimoto, S.; Nakano, K.; Takeda, H.; Nakayama, M.; Sumi, H. Metastable Chloride Solid Electrolyte with High Formability for Rechargeable All-Solid-State Lithium Metal Batteries. *ACS Materials Lett.* **2020**, *2*, 880–886.

(26) Momma, K.; Izumi, F. VESTA 3 for three-dimensional visualization of crystal, volumetric and morphology data. *J. Appl. Crystallogr.* **2011**, *44*, 1272–1276.

(27) Haynes, W. M. CRC Handbook of Chemistry and Physics, 97th ed.; CRC Press, 2016.

(28) Irvine, J. T. S.; Sinclair, D. C.; West, A. R. Electroceramics: Characterization by Impedance Spectroscopy. *Adv. Mater.* **1990**, *2*, 132–138.

(29) Kraft, M. A.; Culver, S. P.; Calderon, M.; Böcher, F.; Krauskopf, T.; Senyshyn, A.; Dietrich, C.; Zevalkink, A.; Janek, J.; Zeier, W. G. Influence of Lattice Polarizability on the Ionic Conductivity in the Lithium Superionic Argyrodites Li_6PS_5X (X = Cl, Br, I). *J. Am. Chem. Soc.* **2017**, *139*, 10909–10918.

(30) Muy, S.; Bachman, J. C.; Chang, H.-H.; Giordano, L.; Maglia, F.; Lupart, S.; Lamp, P.; Zeier, W. G.; Shao-Horn, Y. Lithium Conductivity and Meyer-Neldel Rule in Li₃PO₄-Li₃VO₄-Li₄GeO₄ Lithium Superionic Conductors. *Chem. Mater.* **2018**, *30*, 5573–5582.

(31) Meyer, W.; Neldel, H. Relation between the energy constant and the quantity constant in the conductivity-temperature formula of oxide semiconductors. *Z. Technol. Phys.* **1937**, 588.

(32) Muy, S.; Schlem, R.; Shao-Horn, Y.; Zeier, W. G. Phonon-Ion Interactions: Designing Ion Mobility Based on Lattice Dynamics. *Adv. Energy Mater.* **2020**, 2002787.

(33) Asakura, R.; Duchêne, L.; Kühnel, R.-S.; Remhof, A.; Hagemann, H.; Battaglia, C. Electrochemical Oxidative Stability of Hydroborate-Based Solid-State Electrolytes. *ACS Appl. Energy Mater.* **2019**, *2*, 6924–6930.

(34) Yu, C.; van Eijck, L.; Ganapathy, S.; Wagemaker, M. Synthesis, structure and electrochemical performance of the argyrodite $\text{Li}_6\text{PS}_5\text{Cl}$ solid electrolyte for Li-ion solid state batteries. *Electrochim. Acta* **2016**, *215*, 93–99.

(35) Richards, W. D.; Miara, L. J.; Wang, Y.; Kim, J. C.; Ceder, G. Interface Stability in Solid-State Batteries. *Chem. Mater.* **2016**, *28* (1), 266–273.

(36) Park, C. W.; Oh, S. M. Performances of Li/Li_xCoO₂ cells in LiAlCl₄:3SO₂ electrolyte. *J. Power Sources* **1997**, *68*, 338–343.

(37) Culver, S. P.; Koerver, R.; Zeier, W. G.; Janek, J. On the Functionality of Coatings for Cathode Active Materials in Thiophosphate-Based All-Solid-State Batteries. *Adv. Energy Mater.* **2019**, *9*, 1900626.

(38) Zhang, W.; Sun, Y.; Deng, H.; Ma, J.; Zeng, Y.; Zhu, Z.; Lv, Z.; Xia, H.; Ge, X.; Cao, S.; Xiao, Y.; Xi, S.; Du, Y.; Cao, A.; Chen, X. Dielectric Polarization in Inverse Spinel-Structured Mg₂TiO₄ Coating to Suppress Oxygen Evolution of Li-Rich Cathode Materials. *Adv. Mater.* **2020**, *32*, 2000496.