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# Tuning lithium-yttrium chloride local structure through coordination control and mixing during synthesis<sup>†</sup>

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Lithium halide-based solid electrolytes have high  $Li^+$  conductivity and can be synthesized through lowtemperature aqueous solution routes. While  $Li_3InCl_6$  can be readily synthesized through dehydration, other analogous materials, such as  $Li_3YCl_6$ , cannot. This difference may be due to differences in  $H_2O$ coordination strength, which leads to partial hydrolysis to form YOCl. In this work, we followed and compared  $Li_3YCl_6$  synthesis using three different methods using *in situ* neutron diffraction. The data revealed that forming an ammonium halide complex intermediate is essential in synthesizing  $Li_3YCl_6$  from an aqueous solution. In carefully examining the  $Li_3YCl_6$  products, we found that changes in local structure follow on to drive significant differences in ionic transport and  $Li^+$  diffusivity as determined through diffusion NMR measurements. These changes were ascribed to the change in the correlative transport of  $Li^+$ . This work provides insight into the reaction mechanisms involved in synthesizing halide solid electrolytes and highlights the importance of considering their synthetic and processing conditions to optimize their performance in all-solid-state batteries.

# 1. Introduction

The demand for denser, safer, and more portable energy storage has led to significant interest in all-solid-state batteries (ASSBs) and solid electrolytes.<sup>1</sup> Solid Electrolytes enable the replacement of conventional flammable organic liquid electrolytes<sup>2</sup> and span many material classes including polymers,<sup>3</sup> oxides,<sup>4</sup> sulfides,<sup>5</sup> hydrides,<sup>6</sup> and more recently, halides.<sup>7</sup> Halide solid electrolytes have shown promise due to their high ionic conductivity, high voltage cathode compatibility, and adaptability in synthesis and processing routes.<sup>8</sup>

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Compositions such as  $Li_3InCl_6$  have garnered interest due to their compatibility with aqueous and organic solvent-based synthesis.<sup>9,10</sup> Direct synthesis of other  $A_3MX_6$  (A = Li, M = Sc, Y, Ho, Yb, Er, *etc.*, X = F, Cl, Br, I) compositions in water or other solvents is limited due to the hydrolysis of  $MCl_3 \cdot nH_2O$ intermediates upon heating. This decomposition route leads to the formation of MOCl and LiCl impurities. However, recent reports have shown that it is possible to synthesize these other compositions in water using ammonium halide coordination reagents.<sup>11,12</sup>

This report provides a detailed examination of the formation pathway for  $Li_3YCl_6$  and establishes synthesis-structure property correlations to guide the optimization of halide solid electrolyte-based ASSBs. Large-scale synthesis and optimization of battery materials is not trivial<sup>13,14</sup> and often requires process monitoring. A key enabler of large-scale chemical production is an intimate understanding of the reaction mechanism during synthesis.<sup>15</sup> In addition, many solid electrolyte properties are greatly influenced by the chosen synthesis route.<sup>16</sup> The formation mechanism of halide solid electrolytes through these novel solvent-based routes is poorly understood. Previous work suggested that the addition of ammonium halides resulted in the formation of a complex halide intermediate of the form  $(NH_4)_3MX_6$ , eqn (1).<sup>17,18</sup> This intermediate, shown in Fig. 1, is suspected to provide an alternative

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Fig. 1 Representative crystal structures of the  $(NH_4)_3YCl_6$  and anhydrous Li<sub>3</sub>YCl<sub>6</sub> phases. Octahedrals represent  $YCl_6^{3-}$  polyanions.

reaction route away from the oxyhalide formation side reaction (eqn (2)) during the dehydration and the subsequent annealing steps at 350-550 °C.

$$\operatorname{YCl}_{3} \cdot n\operatorname{H}_{2}\operatorname{O} + 3\operatorname{NH}_{4}\operatorname{Cl} \xrightarrow{-n\operatorname{H}_{2}\operatorname{O}} (\operatorname{NH}_{4})_{3}\operatorname{YCl}_{6}$$
(1)

$$\operatorname{YCl}_3 \cdot n\operatorname{H}_2\operatorname{O} \xrightarrow{\Delta} \operatorname{YOCl} + (n-1)\operatorname{H}_2\operatorname{O} + 2\operatorname{HCl}$$
(2)

We utilized in situ neutron powder diffraction (NPD) to track the formation of Li<sub>3</sub>YCl<sub>6</sub> (LYC) across multiple synthesis routes. NPD is well suited for probing reactions at relevant mass scales due to its high penetrative power and sensitivity to Li.<sup>19,20</sup> In addition, neutrons are far more sensitive to protons and deuterons groups within NH4Cl and complex halide intermediates than X-ray diffraction. Deuterated precursors (D<sub>2</sub>O and ND<sub>4</sub>Cl) were used for NPD experiments to minimize the incoherent scattering background of protons. We confirmed the formation of (ND<sub>4</sub>)<sub>3</sub>YCl<sub>6</sub> during the aqueous synthesis of LYC and its decomposition. We show that dehydration of the complex before reaching high temperatures is critical even in the wet chemical synthesis route to avoid the formation of LiCl and YOCl impurities. We observe in multiple reaction mechanisms that the LYC formation and then melting at higher temperatures is crucial to driving proper mixing of precursors and intermediates, agreeing with a previous report by Asano et al. where LYC was thermally stable up to 450 °C.<sup>21</sup> To our knowledge, this is the first instance of providing structural characterization after melting the solid electrolyte phase using in situ diffraction techniques.<sup>22</sup> Melting of the LYC phase could play an essential role in separator fabrication through melt infiltration or casting methods.<sup>23</sup> We characterize the influence of the LYC synthesis route on the ionic and electronic conductivity and observe the significant role of grain boundaries in the total ionic conductivity of LYC solid electrolytes that undergo annealing.

#### 2. Experimental

#### 2.1 Synthesis

The aqueous ammonium coordinated (AC) and ammonium free (AF) synthesis of LYC was prepared by weighing out 34 mol% excess of anhydrous YCl<sub>3</sub> (Sigma Aldrich 99.99%, 451363), stoichiometric amounts of LiCl (Sigma Aldrich 99%, 793620) and NH<sub>4</sub>Cl (Sigma Aldrich 99.5%, A4514) in a 1:3:3 molar ratio in a glovebox (1:3:0 for AF synthesis). The mixture was then transferred to a round bottom flask and dissolved in DI water, sonicated for 15 minutes, and excess water was evaporated *via* rotovap. The sample was then dried overnight at 150 °C under vacuum to remove excess water. The white powder was ground in an agate mortar and pestle, pelletized, placed in an alumina combustion boat, and annealed at 500 °C (heating rate 5 °C min<sup>-1</sup>) for 5 h under argon flow (30 mL min<sup>-1</sup>) in a glovebox.

The mechanochemical (MC) synthesis of Li<sub>3</sub>YCl<sub>6</sub> was conducted in a Fritsch Pulvirisette 7 planetary ball mill. Here, the LiCl and anhydrous YCl<sub>3</sub> were loaded into zirconia milling jars (45 mL volume) with Y-stabilized zirconia milling media (5 mm diameter) inside a glovebox. The media-to-powder ratio was 30:1. The mixture was milled at 510 rpm for 48 cycles with 15 min of milling and a 15 min pausing step to allow the sample to cool; after the initial 48 cycles, the jar was taken into a glovebox, and the sample was scraped off the walls of the jar to prevent the formation of hot zones during milling, this was followed by another 48 cycles under the same conditions. The milled powder was pelletized and annealed at equivalent conditions to the AC and AF samples. For clarity, this sample is referred to as the mechanochemical annealed (MCA) sample, and the unannealed sample is referred to as the MC sample.

#### 2.2 Characterization

*Ex situ* NPD patterns were collected on POWGEN (BL-11A) at 1.5 Å wavelength. The samples were loaded into vanadium cans under an argon atmosphere in a glovebox and sealed with copper gaskets and aluminum lids. The patterns were measured at 300 K. The samples were absorption corrected after data collection. Rietveld refinements against the neutron diffraction data were performed using GSAS II.<sup>24</sup> The following parameters were refined: (1) background using a Chebyshev polynomial with 10 coefficients, (2) scale factor, (3) sample-induced peak broadening, (4) lattice parameters, (5) atomic positions, (6) isotropic thermal displacement parameters, and (7) occupancies.

The loss of ammonium salt was confirmed using infrared spectroscopy using an Agilent Cary 650 spectrometer equipped with an attenuated total reflection accessory (Specac Ltd). Before measurement, the samples were protected from moisture by sealing them in the diamond crystal holder in a glovebox.

**2.2.1 Impedance spectroscopy.** The solid electrolyte samples were pelletized at 5.9 t  $cm^{-2}$  for 1 min in a 12.7 mm die set with carbon-coated aluminum foil on either side to be

used as blocking electrodes for the ionic conductivity measurements. The measurements were performed by electrochemical impedance spectroscopy (EIS) at OCV, applying an excitation amplitude of 10 mV in a frequency range between 7 MHz and 100 mHz with 6 points per decade using a VSP potentiostat (Biologic).

**2.2.2 DSC and TGA.** TA instruments TGA Q500 and DSC Q100 were used to measure thermogravimetry and differential scanning calorimetry (TGA and DSC), respectively. The samples were loaded into aluminum pans and heated to 500 °C at a heating rate of 5 °C min<sup>-1</sup> under a nitrogen flow of 50 mL min<sup>-1</sup>.

2.2.3 In situ neutron diffraction. To follow the product distribution of a larger scale reaction, large pellets ( $\sim$ 3–5 grams) of the post-vacuum dried AC, AF, and MC samples were loaded into separate quartz baskets along with quartz wool. The basket was placed into the AGES gas flow system at POWGEN (BL-11A) diffractometer at the Spallation Neutron Source (SNS) at Oak Ridge National Laboratory.25,26 An initial room temperature scan was taken and then ramped to 500 °C at a rate of 5 °C min<sup>-1</sup> under flowing Ar, with 1 h holds at 100, 200, 300, 400, and 500 °C for ND data collection. Samples were cooled and monitored in event mode to enable the re-binning of data after the experiment. ND data were collected at a 1.5 Å center wavelength that covered a d-space range of 0.7 to 8 Å. Refinements of all the neutron-diffraction data were conducted using the Rietveld method in GSAS-II.<sup>24</sup> For the neutron-diffraction spectra, the background was fitted using log-interpolated polynomials.

**2.2.4 NMR spectroscopy.** 1D <sup>7</sup>Li static and diffusion NMR measurements were performed on a 300 MHz NMR spectrometer equipped with a DOTY Z-spec PFG NMR probe. The stimulated echo pulse sequence was used to collect the <sup>7</sup>Li signal by varying the gradient strength up to the gradient probe with a maximum gradient of 900 G cm<sup>-1</sup>. The diffusion coefficients were measured with varying temperatures (from 25 °C to 88 °C) and gradient pulse separation ( $\Delta$ ) from 40 ms to 150 ms with gradient pulse length ( $\delta$ ) of 4 ms. The diffusion coefficients were calculated by using the Stejkal-Tanner equation.<sup>27</sup> 1D <sup>7</sup>Li static NMR spectra (single pulse experiment with 90° pulse length of 12.6 µs) were collected with 32 scans and 10 s recycling delay and fitted using DMFIT software.<sup>28 7</sup>Li chemical shifts were externally referenced to 1 M LiTFSI solution.

# 3. Results and discussion

#### 3.1 Synthesis and phase evolution

Summary plots of *in situ* neutron diffraction measurements are presented in Fig. 2, along with phase evolution and thermal analyses. We compared three reaction routes for LYC production at relatively large scales (3–5 grams): ammonium coordinated (AC), ammonium free (AF), and dry mechanochemical (MC). This sample size aligns with a typical laboratory-scale synthesis procedure. For the two aqueous routes, the precursors (LiCl and YCl<sub>3</sub>, with and without ammonium chloride) were dissolved in water and dried in a vacuum at 80 °C before measurement. Note that the deuteron will contribute to coherent scattering, making it possible to observe the formation of complex intermediates during the phase evolution. The reactions were conducted by heating the dried precursors under flowing Ar with a temperature ramp rate set to 5 °C min<sup>-1</sup> and 1 hour holds at 100, 200, 300, 400, and 500 °C before cooling to room temperature (~30 °C).

After comparing the three ND contour plots, the first observation reveals that the AC route (Fig. 2a) is more complex than the others. At 50 °C, YCl<sub>3</sub> and ND<sub>4</sub>Cl precursors are present (identified by peaks at 2.76 and 1.58 Å, respectively), and the diffraction intensities begin to fade at 100 °C (Fig. 2a). We also initially observe both YCl<sub>3</sub> and YCl<sub>3</sub>·6H<sub>2</sub>O (peaks at 2.74 and 3.91 Å respectively), as well as small amounts of ND<sub>4</sub>Cl and LiCl (1.93 and 2.96 Å respectively). Rietveld refinement of the patterns within the contour plot quantifies the phase evolution (Fig. 2b). As the guidelines point out, these precursors are consumed and form the  $(ND_4)_3YCl_6$  complex intermediate, AYC, at 100 °C (identified by the peak at 3.83 Å). The AYC complex assumes a C2/c structure of the form  $Cs_3BiCl_6$ , where  $YCl_6^{3-1}$ octahedrals are indeed formed and stabilized.<sup>29</sup> Unlike the final product, trigonal LYC ( $P\bar{3}m1$ ), the cations are not well ordered into rows, as shown in Fig. 1. This reaction is accompanied by an intermediate phase(s), which we call LYC-I. We note here that this intermediate is only seen during the AC route. The formation of AYC is seen in the thermal analysis (Fig. 2b), which displays an endothermic peak at 100 °C (Peak 1) and significant weight loss corresponding to  $D_2O$  loss. The AYC complex is not long-lived, being consumed at 200 °C and giving way to more intense LYC-I peaks and YCl<sub>3</sub> formation (Fig. 2a). DSC shows an endothermic process at 175 °C (Peak 2) that appears to be a convolution of the AYC decomposition and LYC-I formation. Between 200 and 500 °C, the phase fraction of LYC-I increases and YCl<sub>3</sub> decreases, ending with LYC-I melting, also congruent with an endothermic feature at 475 °C (Peak 4). Within this range, we also observed an endothermic process with significant mass loss between 330 and 375 °C (Peak 3), which correlates with ND<sub>4</sub>Cl decomposition.<sup>30</sup> Given that LYC has known polymorphs and no additional phase was observed in the reaction, we believe LYC-I to be a metastable polymorph.

*Ex situ* infrared spectroscopy in Fig. S4<sup>†</sup> confirms the presence of  $H_2O$  with  $NH_4^+$  cations with strong vibrations at 3100–3200 cm<sup>-1</sup> (N–H stretching) and 1420 cm<sup>-1</sup> (N–H scissoring). These bands are absent after annealing at 350 °C and the O–H bands are absent after annealing at 500 °C. The resulting LYC phase that crystallized from all *in situ* melts was the target trigonal LYC phase ( $P\bar{3}m1$ ). We note that while the YCl<sub>3</sub> impurity appears significant in the AC sample, it is likely the result of the LYC melt dripping through the quartz basket's porous bottom. The *ex situ* samples do not show this impurity in similarly large phase fractions, as discussed later.

The AF sample contour plot (Fig. 2c) begins with LiCl and  $YCl_3{\cdot}6D_2O$  phases being present with no anhydrous  $YCl_3$ 

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**Fig. 2** Visualization of the synthesis pathways using *in situ* neutron diffraction of the AC (a), AF (c), and MC (e), along with their respective phase evolutions with thermal analysis spectra from TGA and DSC (b, d, and f). The contour plots are during heating from 47 to 500 °C only. The initial NPD pattern is the precursors after dissolution and vacuum drying (or mixing) prior to ramping, and the final NPD pattern is the RT product obtained after cooling. Endothermic peaks are numbered for textual clarity (see Fig. S1–3† for high-resolution images).

phase detected (Fig. 2d). The DSC shows no feature at 100 °C associated with intermediate complexes (Fig. 2d); however, during heating to 175 °C the DSC curve shows a large endothermic process alongside substantial mass loss (Peak 1). These results correlate with YCl<sub>3</sub> hydrate consumption and LYC formation in Fig. 2d. The LYC formed here is the targeted trigonal phase and is associated with the second more minor endothermic process in the DSC (230 °C, Peak 2 in DSC). The consumption of the YCl<sub>3</sub> hydrate also results in the formation of the YOCl impurity phase that is formed around 350 °C (Peak 3).<sup>31</sup> This fact demonstrates that the synthesis routes result in differing intermediate phases and reaction kinetics and may allow for trapping novel metastable phases. LYC continues to increase until 500 °C when it melts, leaving insoluble YOCl and LiCl behind. DSC shows LYC melting as a smaller endothermic peak at 475 °C (Peak 4) commensurate with this route's less efficient LYC formation. After cooling, LYC reforms while YOCl and LiCl impurities remain, allowing us to conclude that ammonium chloride is required to protect the YCl<sub>3</sub> precursor from hydrolysis during heating.

Fig. 2e and f show MC reaction route evolution starting with planetary milled LiCl and YCl<sub>3</sub> for 24 h. This reaction proceeds through the most direct route, with LYC being already formed with unreacted LiCl at 68 and 32 wt%, respectively (Fig. 2f). While  $YCl_3$  was not initially detected, we suspect it was present as an amorphous phase, given its immediate appearance with heating. During the temperature ramp, the LiCl phase decreases, and after 175 °C, an endothermic process (Peak 1) is observed and the YCl<sub>3</sub> phase decreases. The latter effect correlated to the small exothermic peak ~275 °C (Peak 2) in Fig. 2f. Again, LYC melts at 475 °C (Peak 3 in the DSC) with residual YCl<sub>3</sub> being present (9 wt%) after cooldown. The LYC phase observed throughout this route was trigonal; however, it started from a nanocrystalline subproduct that has a significant amount of Y2-Y3 site disorder in line with previous reports.32,33

We followed the formation of  $(ND_4)_3YCl_6$  (AYC) by mixing  $ND_4Cl$  and  $YCl_3$  in  $D_2O$  using *in situ* NPD (Fig. 3). While solving the crystal structure is outside the scope of this report, we highlight a few findings. The temperature-dependent stack

plot in Fig. 3a shows the initial sample containing anhydrous  $YCl_3$  and  $ND_4Cl$  (peaks at 2.76 and 3.94 Å, respectively). As the temperature reaches 100 °C, the  $ND_4Cl$  peaks vanish, and the AYC peaks form, which are stable up to 250 °C. The thermal analysis shows residual dehydration and AYC formation are likely convoluted between Peaks 1 and 2 in the DSC. During the AC reaction, AYC was observed during the dehydration, and the peaks vanished as the temperature reached 175 °C (Fig. 2b). This suggests that the reaction to form LYC from AYC through the replacement of ammonia with Li<sup>+</sup> is thermodynamically and kinetically favorable.

This conclusion is reinforced by the fact that Meyer *et al.* proposed a decomposition of  $(ND_4)_3$ [YCl<sub>6</sub>] back into YCl<sub>3</sub> and NH<sub>4</sub>Cl at around 425 °C, which we do not see evidence for in the AC route.<sup>34</sup> We also do not observe the presence of large amounts of LiCl during the AC phase evolution prior to forming Li<sub>3</sub>YCl<sub>6</sub>, which could signal the existence of a distinct  $(NH_4)_3$ [YCl<sub>6</sub>]·3LiCl composite structure.<sup>11</sup> We note that when comparing the thermal analysis of all the aqueous processes (Fig. 2b, d and f), an approximate weight loss of 30% from room temperature to 200 °C is seen, corresponding to D<sub>2</sub>O



Fig. 3 (a) *in situ* NPD data showing AYC formation heating from ~45 to 250 °C from a dehydrated ND<sub>4</sub>Cl + YCl<sub>3</sub> mixture and (b) TGA/DSC of the same precursor. Diffraction tick marks were obtained by phase identification match to  $Cs_3BiCl_6$ . Endothermic peaks are labeled for textual clarity.

being removed from the structure. Ammonium chloride allows water to be removed from YCl<sub>3</sub>·6H<sub>2</sub>O without hydrolysis to YOCl at lower temperatures, presumably by shifting the structure from coordination with water to a hexahedral coordination with Cl<sup>-</sup>. The last two endothermic peaks (3 and 4) in Fig. 3b do not seem to directly correlate to any observed changes in the in situ NPD data. Some possible explanations for this can be gleaned from a similar process observed in the other samples. For the MCA sample, we observe a similar small endothermic peak coinciding with the crystallization of  $YCl_3$  (Peak 1), which could coincide with the  $YCl_3$  observed in the 150 and 250 °C NPD data. Previous work observed a phase transition of NH<sub>4</sub>Cl from an  $\alpha$  to a  $\beta$  high-temperature phase at 180 °C; however, we do not see the characteristic strong reflection for this phase in the NPD data (the expected peak at 3.298 Å is absent).<sup>34</sup> The final small endothermic process at ~250 °C (Peak 4 in Fig. 3b) could be the consumption of residual YCl<sub>3</sub> as observed in the AF reaction route. Lastly, YOCl formation is not observed in the NPD because the sample stayed below the crystallization temperature (~330 °C).

#### 3.2 Synthesis-structure-property correlation

Recent literature reviews have established the significant relationship between the synthesis route and solid electrolyte properties ranging from the room temperature ionic conductivity to the electrochemical stability window.<sup>16,35</sup> Here, we summarize observations related to atomic structure from ND (lattice parameters), particle morphology, and bulk and grain boundary conductivity. In Fig. 4 we display the Rietveld refinement results for NPD data collected on each of the three products within sealed vanadium cans at room temperature. We note here that the samples always have approximately 5% chloride impurity, whether it was YCl<sub>3</sub> or LiCl. In this study, the impurity was stochastic; however, we didn't apply significant effort in resolving the causal relationship other than annealing drives YCl<sub>3</sub> impurity formation. When comparing the lattice parameters of the MC and MCA samples obtained from NPD, we find that upon annealing, the lattice volume contracts from 662.13 to 659.02  $\pm$  0.02 Å<sup>3</sup>, and the lattice parameters a and c contract from 11.235 to 11.220  $\pm$  0.001 Å and 6.0573 to 6.0493  $\pm$  0.0006 Å, respectively. The AC product has a similar lattice volume to the MCA sample, 660 Å<sup>3</sup>. It is possible for the differences in lattice parameters to be driven by Y2-Y3 site disorder, where zero site disorder would correspond to 100% occupation of the Y2 site.<sup>21,33</sup> Upon milling, the MC sample starts with 39% site disorder, and after annealing, this is reduced to 22% site disorder for the MCA sample. This trend of decreasing site disorder as a function of annealing agrees with previous reports.33

The degree of disorder can be qualitatively evaluated through <sup>7</sup>Li static NMR. As shown in Fig. 6, the MC sample exhibits two distinct features: a narrow and a broad <sup>7</sup>Li peak. The narrow Gaussian peak itself is a superposition of a quadrupole central transition and a partially narrowed response, which is attributed to  $Li^+$  ions undergoing rapid hopping. At the same time, the latter broad satellite quadrupolar feature

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Fig. 4 (a) Refined crystal structure of tetragonal LYC product highlighting the Y2 and Y3 site disorder along with Rietveld refinement of the (b) LYC-AC (c) LYC-MC (d) LYC-MCA products' measured ND data. See Fig. S5–7† for high-resolution images.

corresponds to those occupying crystallographic inequivalent sites.<sup>36</sup> The disorder in the latter is featured by distribution in quadrupolar parameters, resulting in a rounded line shape without well-defined divergences. These narrow and broad peaks converge at elevated temperatures due to rapid Li<sup>+</sup> motion (see Fig. S8<sup>†</sup>).

In the annealed MCA sample, two distinct <sup>7</sup>Li features have been identified. The peak associated with Li<sup>+</sup> ions situated in inequivalent crystallographic sites exhibits a well-defined quadrupole powder pattern, consistent with the diffraction result that annealing induces ordering in the Li<sup>+</sup> sublattice. On the contrary, the static spectrum of the solution-synthesized AC product reveals three distinct <sup>7</sup>Li features. A single narrow peak is ascribed to the fast-moving Li<sup>+</sup> ions, while two well-defined quadrupolar features are associated with Li<sup>+</sup> ions sited in different inequivalent crystallographic sites. This highlights the more heterogeneous structure of sample AC compared to the MCA sample. In both samples, the quadrupolar features narrow with temperature, which can be attributed to increased Li<sup>+</sup> motion. The representative deconvolution of static spectra is shown in ESI Fig. S9.<sup>†</sup>

We used electrochemical impedance spectroscopy (EIS) to investigate the impact of the synthesis route on the ionic conductivity and activation energy. The activation energy was obtained by the Arrhenius relationship for solid-state ionic transport in eqn (3),

$$\sigma_{\rm ionic}T = \sigma_0 \, \exp\!\left(-\frac{E_{\rm a}}{k_{\rm B}T}\right) \tag{3}$$

where  $\sigma_0$ ,  $k_{\rm B}$ , and T are the temperature-independent Arrhenius prefactor, the Boltzmann constant, and the absolute temperature, respectively.  $\sigma_0$  represents the frequency of lithium hopping events at a standard carrier concentration within the solid ionic conductor.<sup>37,38</sup> Fig. 5a shows representative Nyquist plots showing low ionic resistance and the presence of grain boundaries. For the LYC AC sample, the room temperature lattice ionic conductivity is  $0.57 \pm 0.05 \text{ mS cm}^{-1}$ , while the grain boundary is 0.043 mS cm<sup>-1</sup>, more than an order of magnitude lower. For the LYC MCA sample, we see a much smaller difference in the two processes,  $0.42 \pm 0.03$  mS  $cm^{-1}$  for the lattice vs. 0.21 ± 0.01 mS  $cm^{-1}$  for the grain boundary. For the MC product, the grains are neglectable, showing 0.47  $\pm$  0.01 mS cm<sup>-1</sup> for lattice conductivity. Fig. 5b compares the Arrhenius behaviour for the total conductivity of the LYC products, showing that the products have similar values conductivity activation energies of  $\sim 0.35 \pm 0.01$  eV. However, deconvoluting the bulk and grain boundary shows that the activation energy for the lattice conductivity for AC, MC, and MCA are 0.22, 0.15, and 0.17 eV, respectively. This result suggests that the Y distribution between the Y2-Y3 sites strongly influences ionic conduction. The change (a 13% increase in the activation energy) is likely due to the 17% reduction in site disorder observed in the NPD data. This observation is in line with previous reports regarding the effect of annealing time on the mechanochemically synthesized Li3YCl6, where it was observed that the conductivity decreased as the milled sample was annealed for more extended periods of time ranging from 1 minute to 1 hour.<sup>33</sup> That said, we also observed an increase of 25% in the grain boundary activation energy, which increased from 0.28 to 0.39 eV between the MC-derived product and the AF product. This shows that microstructure is equally important in determining the ionic transport properties of LYC and must be optimized during synthesis or processing.

PFG NMR was utilized to measure  $\text{Li}^+$  diffusivity,  $D_{\text{Li}^+}$ , as functions of temperature and diffusion time (*i.e.*, gradient pulse



Fig. 5 (a) Representative Nyquist plots of each sample at 35 °C, along with representative fit of the Nyquist plot and equivalent circuit for the LYC MCA sample at 35 °C. (b) Arrhenius plot comparison for the Li<sub>3</sub>YCl<sub>6</sub> M, MCA, and AC samples' lattice conductivity.

separation), as displayed in Fig. 7. It is worth noting that, firstly, Li<sup>+</sup> diffusion in MCA is slower compared to AC and MC, which is consistent with conductivity results. Secondly, The  $D_{\text{Li}^+}$  of AC and MC are found to be comparable. All  $D_{Li^+}$  exhibit increase with temperature, demonstrating Arrhenius behavior;  $E_a = 0.43$ eV for AC and MC, and 0.33 eV for MCA. The dependence of  $D_{Li^+}$ on  $\Delta$  shows a decreasing trend for all samples (Fig. 7b). In free isotropic diffusion, D is independent of  $\Delta$ . The results here indicate that free diffusion is restricted roughly to the length scale of the PFG experiment, given by  $\sim (Dt)^{1/2}$ , which is on the order of 1 µm and is similar to the average particle size of 800 nm as determined by electron micrographs.

We estimated the correlated migration of the Li<sup>+</sup> ions using the inverse Haven ratio  $(H_{\rm R}^{-1})$ , which is the ratio of the measured conductivity and the theoretically derived value obtained by a tracer diffusion coefficient and the Nernst-Einstein relationship.



Fig. 7 Diffusion coefficients of Li<sup>+</sup> of AC (black circle), MC (red triangle) and MCA (blue square) measured as a function of (a) temperature from 25 to 88 °C and (b) gradient pulse separation ( $\Delta$ ) from 40 ms to 150 ms.

$$\sigma_{\rm diff} = \frac{ne^2}{k_{\rm B}T} D_{\rm Li^+} \tag{4}$$

where *n* is the concentration of the mobile ion. We assumed Li<sup>+</sup> motion is from the more mobile one of the two lithium sites. When  $H_{\rm R}^{-1}$  is greater than unity, correlated charge motion is present. For the sake of comparison, we hold *n* constant for each sample at  $6.8 \times 10^{-3}$  Li<sup>+</sup> Å<sup>-3</sup>; thus, the values in Table 1 have convolutions of correlated motion and mobile Li<sup>+</sup> concentration. The table shows that the solution and mechanochemical-synthesized products have higher correlated

**Table 1** Inverse Haven ratio  $(H_R^{-1})$  of Li<sub>3</sub>YCl<sub>6</sub> products

Product	${H_{\mathrm{R}}}^{-1}$ at 25 °C	${H_{\mathrm{R}}}^{-1}$ at 70 °C
AC	2.69	2.55
MC	1.80	1.47
MCA	0.35	0.62



Fig. 6 <sup>7</sup>Li static NMR spectra of MC, MCA, and AC samples and the deconvolution fits.

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migration than the annealed sample. These results are similar in value to those obtained for doped and undoped  $\text{Li}_3\text{ScCl}_{6}^{39}$  $H_{\text{R}}^{-1}$  decreases with temperature, suggesting that more Li<sup>+</sup>ions become mobile or more correlated jumps occur during self-diffusion. We note that  $H_{\text{R}}^{-1}$  is significantly suppressed for the MCA sample, being below unity, suggesting that grain boundary influences affect the conductivity measurement. That said, given the contraction of the lattice due to annealing and the well-defined quadrupole satellite peaks in the static <sup>7</sup>Li NMR (Fig. 6), a significant portion of the Li<sup>+</sup> is in welldefined sites, suggesting that the mobile Li<sup>+</sup> concentration is less than the other two samples. These results highlight the correlations between the synthesis procedure and the microstructure of the product.

#### 4. Conclusions

Our work compares the reaction mechanism across multiple synthetic routes, showing that LYC formation differs in terms of intermediate phases and kinetics. Aqueous ammonium coordinated routes are a promising strategy for optimizing not only the performance of halide solid electrolytes but also the processing and, eventually, large-scale production. We show the importance of forming complex halide intermediates necessary to synthesize Li<sub>3</sub>YCl<sub>6</sub> through aqueous routes. Control of this intermediate is the key to enabling process control for large batch reactor reaction engineering for halide solid electrolytes. The LYC products were also found to differ significantly in their ionic transport. These differences are shown to be driven by changes in local microstructure, which affected the correlated Li<sup>+</sup> transport or the concentration of mobile Li<sup>+</sup>. We find annealing of LYC induces grain boundary formation regardless of the synthesis route. For the LYC properties to be optimized for use in either halide-based or bilayer-based ASSBs, careful consideration must be made as to the ideal synthesis route to use for both processing and optimization performance. Potential future directions include optimizing processing parameters to enable microstructures with smaller grain boundary resistance and favorable particle size distributions, as well as exploring novel liquid-phase techniques compatible with higher and lower valence transition metal chlorides such as ZrCl<sub>2</sub> and HfCl<sub>2</sub>.

#### Conflicts of interest

There are no conflicts to declare.

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# References

- 1 J. Janek and W. G. Zeier, A Solid Future for Battery Development, *Nat. Energy*, 2016, 1(9), 16141, DOI: 10.1038/ nenergy.2016.141.
- 2 J. Wang, R. Chen, L. Yang, M. Zan, P. Chen, Y. Li, W. Li, H. Yu, X. Yu, X. Huang, L. Chen and H. Li, Raising the Intrinsic Safety of Layered Oxide Cathodes by Surface Re-Lithiation with LLZTO Garnet–Type Solid Electrolytes, *Adv. Mater.*, 2022, 34(19), 2200655, DOI: 10.1002/ adma.202200655.
- 3 L. Yue, J. Ma, J. Zhang, J. Zhao, S. Dong, Z. Liu, G. Cui and L. Chen, All Solid-State Polymer Electrolytes for High-Performance Lithium Ion Batteries, *Energy Storage Mater.*, 2016, 5, 139–164, DOI: 10.1016/j.ensm.2016.07.003.
- 4 M. Jia, N. Zhao, H. Huo and X. Guo, Comprehensive Investigation into Garnet Electrolytes Toward Application-Oriented Solid Lithium Batteries, *Electrochem. Energy Rev.*, 2020, 3(4), 656–689, DOI: 10.1007/s41918-020-00076-1.
- 5 X. Bai, T. Yu, Z. Ren, S. Gong, R. Yang and C. Zhao, Key Issues and Emerging Trends in Sulfide All Solid State Lithium Battery, *Energy Storage Mater.*, 2022, **51**, 527–549, DOI: **10.1016/j.ensm.2022.07.006**.
- 6 Y. Pang, Y. Liu, J. Yang, S. Zheng and C. Wang, Hydrides for Solid-State Batteries: A Review, *Mater. Today Nano*, 2022, 18, 100194, DOI: 10.1016/j.mtnano.2022.100194.
- 7 H. Kwak, S. Wang, J. Park, Y. Liu, K. T. Kim, Y. Choi, Y. Mo and Y. S. Jung, Emerging Halide Superionic Conductors for All-Solid-State Batteries: Design, Synthesis, and Practical Applications, ACS Energy Lett., 2022, 7(5), 1776–1805, DOI: 10.1021/acsenergylett.2c00438.
- 8 C. Wang, J. Liang, J. T. Kim and X. Sun, Prospects of Halide-Based All-Solid-State Batteries: From Material Design to Practical Application, *Sci. Adv.*, 2022, **8**(36), eadc9516, DOI: **10.1126/sciadv.adc9516**.
- 9 X. Li, J. Liang, N. Chen, J. Luo, K. R. Adair, C. Wang, M. N. Banis, T. Sham, L. Zhang, S. Zhao, S. Lu, H. Huang, R. Li and X. Sun, Water–Mediated Synthesis of a Superionic

Halide Solid Electrolyte, *Angew. Chem., Int. Ed.*, 2019, 58(46), 16427–16432, DOI: 10.1002/anie.201909805.

- 10 X. Luo, D. Cai, X. Wang, X. Xia, C. Gu and J. Tu, A Novel Ethanol-Mediated Synthesis of Superionic Halide Electrolytes for High-Voltage All-Solid-State Lithium–Metal Batteries, *ACS Appl. Mater. Interfaces*, 2022, **14**(26), 29844– 29855, DOI: **10.1021/acsami.2c06216**.
- 11 C. Wang, J. Liang, J. Luo, J. Liu, X. Li, F. Zhao, R. Li, H. Huang, S. Zhao, L. Zhang, J. Wang and X. Sun, A Universal Wet-Chemistry Synthesis of Solid-State Halide Electrolytes for All-Solid-State Lithium-Metal Batteries, *Sci. Adv.*, 2021, 7(37), eabh1896, DOI: 10.1126/ sciadv.abh1896.
- 12 X. Shi, Z. Zeng, H. Zhang, B. Huang, M. Sun, H. H. Wong, Q. Lu, W. Luo, Y. Huang, Y. Du and C. Yan, Gram–Scale Synthesis of Nanosized Li <sub>3</sub> HoBr <sub>6</sub> Solid Electrolyte for All– Solid–State Li–Se Battery, *Small Methods*, 2021, 5(11), 2101002, DOI: 10.1002/smtd.202101002.
- 13 M. R. Palacin, Battery Materials Design Essentials, *Acc. Mater. Res.*, 2021, 2(5), 319–326, DOI: 10.1021/accountsmr.1c00026.
- 14 R. Jain, A. S. Lakhnot, K. Bhimani, S. Sharma, V. Mahajani, R. A. Panchal, M. Kamble, F. Han, C. Wang and N. Koratkar, Nanostructuring versus Microstructuring in Battery Electrodes, *Nat. Rev. Mater.*, 2022, 7(9), 736–746, DOI: 10.1038/s41578-022-00454-9.
- 15 V. Sebastian, M. Arruebo and J. Santamaria, Reaction Engineering Strategies for the Production of Inorganic Nanomaterials, *Small*, 2014, **10**(5), 835–853, DOI: **10.1002**/ **smll.201301641**.
- 16 A. Banik, T. Famprikis, M. Ghidiu, S. Ohno, M. A. Kraft and W. G. Zeier, On the Underestimated Influence of Synthetic Conditions in Solid Ionic Conductors, *Chem. Sci.*, 2021, 12(18), 6238–6263, DOI: 10.1039/D0SC06553F.
- 17 M. Simon and G. Meyer, The Oxidation of Tantalum with Ammonium Chloride as an Example of a Novel Route to Early Transitional Metal-Nitrogen Cluster Compounds. Synthesis and Crystal Structure of (NH<sub>4</sub>) <sub>6</sub> [Ta <sub>5</sub> (NH) <sub>4</sub> Cl <sub>17</sub>], *J. Chem. Soc., Chem. Commun.*, 1993, (5), 460–461, DOI: 10.1039/C39930000460.
- 18 G. Meyer, E. Garcia and J. D. Corbett, The Ammonium Chloride Route to Anhydrous Rare Earth Chlorides-The Example of Ycl<sub>3</sub>, in *Inorganic Syntheses*, ed. H. R. Allcock, John Wiley & Sons, Inc., Hoboken, NJ, USA, 2007, pp. 146–150. DOI: 10.1002/9780470132562.ch<sub>3</sub>5.
- 19 R. L. Sacci, T. H. Bennett, A. R. Drews, V. Anandan, M. J. Kirkham, L. L. Daemen and J. Nanda, Phase Evolution during Lithium–Indium Halide Superionic Conductor Dehydration, *J. Mater. Chem. A*, 2021, 9(2), 990– 996, DOI: 10.1039/D0TA10012A.
- 20 R. L. Sacci, R. D. McAuliffe, T. F. Malkowski, N. Kidder, X. C. Chen, A. Huq, M. Kirkham, B. L. Armstrong, L. L. Daemen and G. M. Veith, La <sub>2</sub> Zr <sub>2</sub> O <sub>7</sub> Nanoparticle-Mediated Synthesis of Porous Al-Doped Li <sub>7</sub> La <sub>3</sub> Zr <sub>2</sub> O <sub>12</sub> Garnet, *Inorg. Chem.*, 2021, 60(13), 10012–10021, DOI: 10.1021/acs.inorgchem.1c01300.

- 21 T. Asano, A. Sakai, S. Ouchi, M. Sakaida, A. Miyazaki and S. Hasegawa, Solid Halide Electrolytes with High Lithium-Ion Conductivity for Application in 4 V Class Bulk-Type All-Solid-State Batteries, *Adv. Mater.*, 2018, **30**(44), 1803075, DOI: **10.1002/adma.201803075**.
- 22 Y. Wu, S. Wang, H. Li, L. Chen and F. Wu, Progress in Thermal Stability of all-solid-state-Li-ion-batteries, *InfoMat*, 2021, **3**(8), 827-853, DOI: **10.1002/inf2.12224**.
- 23 Y. Xiao, K. Turcheniuk, A. Narla, A.-Y. Song, X. Ren, A. Magasinski, A. Jain, S. Huang, H. Lee and G. Yushin, Electrolyte Melt Infiltration for Scalable Manufacturing of Inorganic All-Solid-State Lithium-Ion Batteries, *Nat. Mater.*, 2021, 20(7), 984–990, DOI: 10.1038/s41563-021-00943-2.
- 24 B. H. Toby and R. B. Von Dreele, *GSAS-II*: The Genesis of a Modern Open-Source All Purpose Crystallography Software Package, *J. Appl. Crystallogr.*, 2013, 46(2), 544–549, DOI: 10.1107/S0021889813003531.
- 25 M. Kirkham, L. Heroux, M. Ruiz-Rodriguez and A. Huq, AGES: Automated Gas Environment System for *in Situ* Neutron Powder Diffraction, *Rev. Sci. Instrum.*, 2018, **89**(9), 092904, DOI: **10.1063/1.5031432**.
- 26 A. Huq, M. Kirkham, P. F. Peterson, J. P. Hodges, P. S. Whitfield, K. Page, T. Húgle, E. B. Iverson, A. Parizzi and G. Rennich, POWGEN: Rebuild of a Third-Generation Powder Diffractometer at the Spallation Neutron Source, *J. Appl. Crystallogr.*, 2019, 52(5), 1189–1201, DOI: 10.1107/ S160057671901121X.
- 27 E. O. Stejskal and J. E. Tanner, Spin Diffusion Measurements: Spin Echoes in the Presence of a Time-Dependent Field Gradient, *J. Chem. Phys.*, 1965, **42**(1), 288– 292.
- 28 D. Massiot, F. Fayon, M. Capron, I. King, S. Le Calvé, B. Alonso, J.-O. Durand, Z. Gan and G. Hoatson, Modelling One- and Two-Dimensional Solid-State NMR Spectra, *Magn. Reson. Chem.*, 2002, 40, 70–76, DOI: 10.1002/mrc.984.
- 29 H. Mattfeld and G. Meyer, Ternäre Halogenide Vom Typ A3MX6. I A3YCl6 (A = K, NH4, Rb, Cs): Synthese, Strukturen, Thermisches Verhalten. Über Einige Analoge Chloride Der Lanthanide, *Z. Anorg. Allg. Chem.*, 1992, 618(12), 13–17, DOI: 10.1002/zaac.19926180103.
- 30 G. Yudoyono, N. Ichzan, V. Zharvan, R. Daniyati, H. Santoso, B. Indarto, Y. H. Pramono and M. Zainuri, Darminto. Effect of Calcination Temperature on the Photocatalytic Activity of TiO2 Powders Prepared by Co-Precipitation of TiCl3; Semarang, Indonesia, 2016, p. 020099. DOI: 10.1063/1.4945553.
- 31 P. K. Todd, M. J. McDermott, C. L. Rom, A. A. Corrao, J. J. Denney, S. S. Dwaraknath, P. G. Khalifah, K. A. Persson and J. R. Neilson, Selectivity in Yttrium Manganese Oxide Synthesis via Local Chemical Potentials in Hyperdimensional Phase Space, *J. Am. Chem. Soc.*, 2021, 143(37), 15185–15194, DOI: 10.1021/jacs.1c06229.
- 32 M. Gombotz and H. M. R. Wilkening, Fast Li Ion Dynamics in the Mechanosynthesized Nanostructured Form of the Solid Electrolyte Li<sub>3</sub>YBr<sub>6</sub>, *ACS Sustainable*

Inorganic Chemistry Frontiers

*Chem. Eng.*, 2021, **9**(2), 743–755, DOI: **10.1021**/ **acssuschemeng.0c06694**.

- 33 R. Schlem, S. Muy, N. Prinz, A. Banik, Y. Shao-Horn, M. Zobel and W. G. Zeier, Mechanochemical Synthesis: A Tool to Tune Cation Site Disorder and Ionic Transport Properties of  $\text{Li}_3\text{MCl}_6$  (M = Y, Er) Superionic Conductors, *Adv. Energy Mater.*, 2020, **10**(6), 1903719, DOI: **10.1002**/ **aenm.201903719**.
- 34 G. Meyer and P. Ax, An Analysis of the Ammonium Chloride Route to Anhydrous Rare-Earth Metal Chlorides, *Mater. Res. Bull.*, 1982, 17(11), 1447–1455, DOI: 10.1016/ 0025-5408(82)90231-8.
- 35 Y. W. Choi, C. M. Araujo and R. Lizárraga, Amorphisation-Induced Electrochemical Stability of Solid-Electrolytes in Li-Metal Batteries: The Case of Li 3 ClO, *J. Power Sources*, 2022, 521, 230916, DOI: 10.1016/j. jpowsour.2021.230916.

- 36 V. Epp, Ö. Gün, H.-J. Deiseroth and M. Wilkening, Long-Range Li+ Dynamics in the Lithium Argyrodite Li7PSe6 as Probed by Rotating-Frame Spin–Lattice Relaxation NMR, *Phys. Chem. Chem. Phys.*, 2013, **15**, 7123–7132, DOI: **10.1039/C3CP44379E**.
- 37 V. Jagannadham, How Do We Introduce the Arrhenius Pre-Exponential Factor (A) to Graduate Students?, *Creat. Educ.*, 2010, 01(02), 128–129, DOI: 10.4236/ce.2010.12019.
- 38 Y. Gao, N. Li, Y. Wu, W. Yang and S. Bo, Rethinking the Design of Ionic Conductors Using Meyer-Neldel-Conductivity Plot, *Adv. Energy Mater.*, 2021, **11**(13), 2100325, DOI: **10.1002/aenm.202100325**.
- 39 R. Li, P. Lu, X. Liang, L. Liu, M. Avdeev, Z. Deng, S. Li, K. Xu, J. Feng, R. Si, F. Wu, Z. Zhang and Y.-S. Hu, Superionic Conductivity Invoked by Enhanced Correlation Migration in Lithium Halides Solid Electrolytes, *ACS Energy Lett.*, 2024, 9, 1043–1052, DOI: 10.1021/acsenergylett.3c02496.