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Synthesis of Antiperovskite Solid Electrolytes: Comparing Li₃SI, Na₃SI, and Ag₃SI

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Supporting Information

ABSTRACT: Prior calculations have predicted that chalcohalide antiperovskites may exhibit enhanced ionic mobility compared to oxyhalide antiperovskites as solid-state electrolytes. Here, the synthesis of Ag-, Li-, and Na-based chalcohalide antiperovskites is investigated using first-principles calculations and *in situ* synchrotron X-ray diffraction. These techniques demonstrate that the formation of Ag₃SI is facilitated by the adoption of a common body centered cubic packing of S²⁻ and I⁻ in the reactants and products at elevated temperatures, with additional stabilization achieved by the formation of a solid solution of the anions. The absence of these two features appears to hinder the formation of the analogous Li and Na antiperovskites.

he development of all-solid-state batteries has attracted increasing attention in recent years. The adoption of solid-state electrolytes (SSEs) in place of conventional organic liquid electrolytes is projected to enable the use of high voltage cathode materials and high capacity metallic anodes. Solid electrolytes also greatly enhance battery safety. 1,2 The advantages of solid-state rechargeable batteries have motivated the search for inorganic solid-state ionic conductors. Several categories of SSEs with ionic conductivity comparable to that of liquids have been identified including glass-ceramic-type lithium sulfides, NASICON-type phosphates, garnet-type $\text{Li}_7\text{La}_3\text{Zr}_2\text{O}_{12}$, and antiperovskites (A₃OX, with A = Li, Na; X = Cl, Br) etc.³ However, the practical application of SSEs still remains challenging. 4,5 Some of these challenges can be traced to the need for higher ionic conductivity $(>10^{-2} \text{ S})$ cm⁻¹), better chemical stability against electrodes, a wide electrochemical window, and mechanical resilience.6-

The antiperovskite (AP) family has been known as ionic conductors for more than half a century. ^{9,10} In recent years, Libased APs, such as Li₃OCl, Li₃OBr, and Li₃OCl_{0.5}Br_{0.5}, have attracted considerable attention as potential candidates for solid-state batteries because of their good electrochemical stability against Li metal, high ionic conductivity (above 10⁻³ S cm⁻¹ at room temperature), low activation enthalpy, negligible electronic conductivity, good cyclability, moderate weight, and low cost. ¹¹ Na-based antiperovskites have also been reported as potential solid electrolytes for use in sodium batteries. ^{12,13} Nevertheless, additional improvements to the performance of antiperovskite SSEs are desirable.

In 2019, a correlation was reported between the ion migration barriers in antiperovskites and their degree of lattice distortion, which could be tuned through substitutions on the anion sublattice: for example, substituting O²⁻ by S²⁻ or Cl⁻/Br⁻ by I⁻.¹⁴ In this work, Na₃SI and Li₃SI were predicted as new (or "hypothetical") antiperovskites with the potential to exhibit higher mobility for Na⁺ and Li⁺, respectively. Nevertheless, the larger distortions of the antiperovskite lattice

resulting from such substitutions were predicted to come at a cost to stability: 14 Na $_3$ SI and Li $_3$ SI are predicted to be unstable relative to metal chalcogenide and halide reagents at low temperature, but entropic contributions are expected to stabilize them upon heating (Figure 1). 15 Similar to Li $_3$ OBr, 16 these materials could potentially be realized as kinetically stabilized phases at ambient temperature, following synthesis at, and quenching from, the higher temperatures at which they are stable.

Unfortunately, preliminary attempts to synthesize Na₃SI and Li₃SI have not been successful thus far. In contrast, the Ag₃SI antiperovskite is a well-known fast ionic conductor. ^{17–20} At least four different polymorphs of Ag₃SI, namely, α , α^* , β , and γ , have been observed at different temperatures. ^{9,10,18,21} These polymorphs differ in their ordering of S^{2–}/I[–] and the location of Ag⁺ within interspaces between anions. The S^{2–}/I[–] disordered α -Ag₃SI is a superionic phase (ionic conductivity: ~1.5 S cm⁻¹ above 520 K). α^* -Ag₃SI, which differs from α -Ag₃SI in the positions of the Ag ions, has an ionic conductivity of ~0.13 S cm⁻¹ at ambient temperature. In contrast, the ionic conductivity of S^{2–}/I[–]ordered β -Ag₃SI is approximately two orders of magnitude lower than that of α^* -Ag₃SI. ²²

Similar to Na_3SI and Li_3SI , Ag_3SI is predicted by DFT calculations to be unstable with respect to AgI and Ag_2S at low temperatures as shown in Figures 1 and S1. However, their stabilities increase as temperature rises. Given the similar free-energy behavior across these antiperovskites, the difference in our ability to synthesize them is surprising. If the formation mechanism associated with the synthesis of Ag_3SI could be

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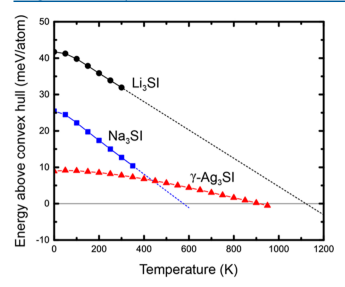


Figure 1. Energy above the convex hull, ΔG (given by $\Delta G = G_{A_3SI} - (G_{A_2S} + G_{AI})$, A = Li, Na, and Ag) as a function of temperature. (A description of the computational methodology can be found in the Supporting Information.) The simulation results for Li₃SI and Na₃SI are from ref 15.

revealed, then this knowledge might be exploited in the synthesis of Na_3SI and Li_3SI .

Here, *in situ* synchrotron X-ray diffraction is used to carefully map out the phase evolution during the formation of Ag_3SI as a function of temperature. Analogous experiments are performed on Na_3SI and Li_3SI as well. A comparison of reaction pathways for these three antiperovskites reveals that the formation of Ag_3SI occurs via a mechanism that is not accessible for Na_3SI and Li_3SI . In particular, the adoption of a common body centered cubic (bcc) packing of anions (S^{2-} or I^-) in the reactants at elevated temperatures facilitates the formation of the antiperovskite structure with the same bcc anion packing. *In situ* X-ray diffraction data, as shown in Figure 2a, reveal a series of phase evolutions during the heating of the 1:1 ratio mixture of AgI and Ag_2S , which are vacuum sealed in a quartz tube, from room temperature up to 600 °C. The annealing profile was chosen based on previous syntheses. 17,18

The synthesis process can be categorized into three stages (Figure 2b-d). At room temperature, AgI crystallizes in the β polymorph. This polymorph adopts space group P63mc, with hexagonal close packing (hcp) of I⁻. Ag₂S crystallizes in the α polymorph with space group $P2_1/c_1$, which has a distorted body centered cubic (bcc) packing of $S^{2-23,24}$ In the first stage, shown in Figure 2b, AgI and Ag₂S undergo phase transitions at ~175 °C and ~200 °C into α -AgI (S.G. Im-3m) and β -Ag₂S (S.G. Im-3m) respectively. 9,25 The polymorphs of the two binaries possess the same body centered cubic (bcc) packing of anions (I^-, S^{2-}) , which also is the anion packing sequence for Ag₃SI antiperovskites (Figure 2e, Stage 1). Following these phase transitions, the diffraction peaks of α -AgI and β -Ag₂S shift to lower angles for temperatures up to 475 °C, as shown in Figure 2c. The linear correlations of the binaries' lattice parameters with temperature in this regime are consistent with thermal expansion (Figure S4).

In the second stage, corresponding to temperatures from 475 to 600 °C, broad Bragg peaks grow between the two binaries, indicating a continuous merging into an $Ag_xS_yI_{2-y}$ (2 $\leq x \leq 4$, $0 \leq y \leq 2$) solid-solution, which possesses the same bcc anion packing. After holding at 600 °C for 2.5 h, a set of symmetric diffraction peaks, indexed with $Im\overline{3}m$ symmetry, is observed. Further changes in peak positions and shapes are negligible, indicating that the distribution of S^{2-} and I^- in the $Ag_xS_yI_{2-y}$ solid solution has become homogeneous. Therefore, at the end of stage 2, an Ag_3SI phase with a random distribution of I^- and S^{2-} is formed.

Finally, in the subsequent cooling process (stage 3), the Ag₃SI phase maintains $Im\bar{3}m$ symmetry until the temperature reaches 210 °C. Another cubic phase with an extra hkl peak at 2.36 degrees starts to evolve as shown in Figure 2d. This new cubic phase can be indexed with a lower symmetry (Pm-3m), which has an extra ordering along the [111] direction. This extra ordering results from the rearrangement of S²⁻ and I⁻. As shown in Figure 2e, stage 3, the center of the cubic lattice (0.5, 0.5, 0.5) is occupied by I⁻, while the corners (0, 0, 0) are occupied by S²⁻. Thus, the S/I ordered β -Ag₃SI is gradually transformed from the S/I disordered α -Ag₃SI at 200 °C. Nonetheless, after holding at 200 °C for 2 h, the α to β phase transition is still not complete. Upon lowering the temperature to 90 °C, new sharp diffraction peaks appeared at 2.85, 3.29,

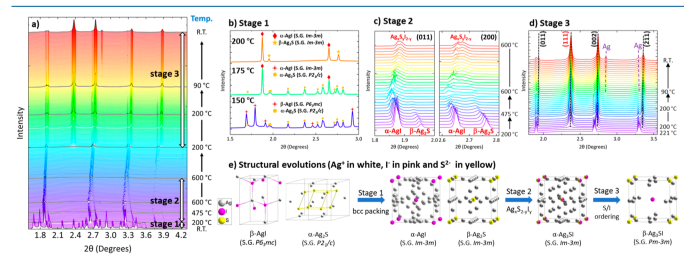


Figure 2. In situ synchrotron X-ray diffraction (sXRD) data collected during the synthesis of Ag_3SI with temperatures on the right side (a) and selected sXRD data for three major stages (b-d) with the representation of structural evolutions in each stage below (e).

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and 4.65 degrees, which correspond to the diffraction peaks of Ag metal. The metallic silver may be the reduction product of excess Ag⁺ by carbon, which is coated on the inner wall of quartz tube. Excess Ag⁺ could originate from the evaporation of I at high temperatures. Filling the quartz tube with inert gas instead of vacuum sealing is expected to reduce Ag metal formation.

The final products obtained at room temperature consist of dominating β -Ag₃SI, minor α^* -Ag₃SI, and other byproducts, including Ag metal and the two binary reactants (Figure S5). It has been reported that α -Ag₃SI could dissociate itself into the binaries or transform to α^* -Ag₃SI during cooling. The instability of α -Ag₃SI compared with other polymorphs at low temperature agrees with the computed Gibbs free energies shown in Figure S2. In particular, the free energy of α -Ag₃SI is predicted to be larger than that of β -Ag₃SI below 314 K. Similarly, the formation energy of α -Ag₃SI relative to AgI and Ag₂S is predicted to become increasingly positive at low temperatures. (Figure 1)

On the basis of above in situ diffraction studies, the key step to form the Ag₃SI antiperovskite phase is the phase transformations of AgI and Ag2S binaries into polymorphs having bcc anion packing. This facilitates the formation of the $Ag_{x}S_{y}I_{2-y}$ solid solution, which has a similar anion packing. The $\alpha - \beta$ phase transition around 200 °C is an important step to obtain the S²⁻/I⁻ ordered β -Ag₃SI structure at room temperature. The minimum temperature for these two steps is approximately 200 °C, which is much lower than the common synthesis temperature for Ag₃SI. Moreover, the present DFT calculations shown in Figure S1 predict that α -Ag₃SI becomes stable at ~80 °C relative to the AgI and Ag₂S reactants. In addition, since the ramping rate for the in situ synthesis was relatively fast, the high onset temperature observed for the merging process in stage 2 may be an artifact of the heating rate. To test this hypothesis, ex situ synthesis is attempted at 220 °C. As a comparison, another synthesis is conducted at 550 °C followed by holding at 220 °C. Highresolution X-ray diffraction data of the two products are shown in Figure S6. Rietveld refinements confirm that pure β -Ag₃SI can be obtained after holding the reaction temperature at 550 °C for 2 days followed by another 2 days at 220 °C. Interestingly, 70 wt % of β -Ag₃SI is obtained after annealing at 220 °C for 4 days. Lattice parameters of two β -Ag₃SI samples are almost identical: 4.9009 (4) Å versus 4.9031(1) Å. Therefore, the merging of bcc anion packing binaries can occur at a much lower temperature than that observed in the in situ measurement (475 °C).

In situ experiments are conducted to investigate the possible formation of Na₃SI and Li₃SI. As shown in Figure 3, both reactants, NaI and Na2S, retain face centered cubic packing up to 650 °C. The Rietveld fit confirms two cubic binaries at 650 °C (Figure S7). Furthermore, their lattice parameters exhibit linear correlations with temperature, indicating that only thermal expansion has occurred. In situ studies on the synthesis of Li₃SI show the same result (Figure S8). Phase transitions to bcc anion packing of the initial binaries does not occur in the case of sodium and lithium compounds. The reason could be the low stability of the bcc anion packing of S2- and I- with occupations of Na+ or Li+ among the interspaces. According to previous literature, 25 in contrast to sodium and lithium iodides and sulfides, the bcc anion packing could be stabilized by the orbital mixing between Ag+ (d orbitals) and I⁻, S²⁻ (p orbitals) in silver analogues. Further

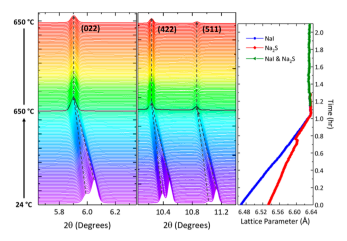


Figure 3. In situ sXRD data for the annealing of Na_2S/NaI mixture with temperature program labeled on the left. Changes in lattice parameters were plotted on the right.

increasing the synthesis temperature could be a strategy to bypass these barriers. However, any such strategy must address the consequences of evaporation of NaI and LiI, which was found to be severe very shortly after melting, as proved by the rapid mass loss measured in TGA/DSC measurements shown in Figure S9. We found that these vapors were extremely reactive against the quartz container used in these syntheses, even when coated with carbon, precluding any reaction between the two alkali metal binaries.

In conclusion, the stability and formation mechanism of Ag_3SI antiperovskite is investigated using *in situ* synchrotron X-ray diffraction and first-principles calculations. Three major stages are involved in the formation of the final S^{2-}/I^- ordered Ag_3SI polymorph. The formation of Ag_3SI is facilitated by phase transitions of the two reactants, Ag_2S and AgI, to structures having the same bcc anion packing. The merging of these two anion bcc packing binaries into Ag_3SI antiperovskite can initiate at as low as 220 °C. However, in the case of Na_2SI and Li_3SI , the adoption of a common bcc anion packing in the binary reactants does not occur below their melting points. This hinders the facile formation of the respective ternary antiperovskites.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.inorgchem.0c01705.

Experimental details of *in situ* and *ex situ* syntheses, computational methods, additional diffraction data for phase identification and Rietveld fits, *in situ* diffraction data for the synthesis of Li₃SI, TGA, and DSC data, details of additional figures and tables (PDF)

Rietveld fits of synchrotron data of ex situ synthesized Ag₃SI (condition 1) (TXT)

Rietveld fits of synchrotron data of ex situ synthesized Ag₃SI (condition 2) (TXT)

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Notes

The authors declare no competing financial interest.

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