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Design strategies for liquid-phase synthesis of sodium-based quaternary solid-state electrolytes†

Saeed Ahmadi Vaselabadi.* Brynn Benham and Colin A. Wolden

The cost-effective and scalable production of solid-state electrolytes is essential for advancing all-solidstate sodium batteries as a safer, lower-cost alternative to Li-ion batteries. Herein, we explored liquidphase synthesis methods for quaternary chalcogenide solid-state electrolytes (SSEs) with the formula $Na_{11}Sn_2PnCh_{12}$ (Pn = P, Sb, Ch = S, Se). We outline the key design principles, including reagent selection, solvent chemistry, and thermal treatment, while elucidating reaction mechanisms through systematic studies using XRD, Raman, and FTIR spectroscopy. We demonstrate that acetonitrile, an aprotic solvent with high polarity, enables the synthesis of Na₁₁Sn₂PS₁₂ via a simple solution process followed by heat-treatment, achieving high ionic conductivity ($\sigma_{Na^+} = 0.2$ mS cm⁻¹). The antimony analog, Na₁₁Sn₂SbS₁₂, can be produced in an aqueous solution facilitated by reactive bisulfide intermediates, yielding a highly crystalline chalcogenide at low temperatures with excellent transport properties ($\sigma_{\text{Na}^+} = 0.4 \text{ mS cm}^{-1}$). For selenide analogs (e.g. $\text{Na}_{11}\text{Sn}_2\text{SbSe}_{12}$), alkahest amine—thiol solvents facilitate synthesis from elemental and binary precursors, marking the first report of its kind. Additionally, we identified $Na_{11}Sn_2SbS_{12}$ as the most promising candidate for utilization in sodium all-solid-state batteries owing to its wide electrochemical stability and strong compatibility with Na-Sn alloy anodes.

Introduction

Solid-state sodium-ion batteries are attractive candidates to facilitate renewable energy integration due to sodium's abundance, low cost, and reduced environmental impact relative to their high-performance lithium counterparts. Sodium's relatively low atomic weight and its similar electrochemical potential make it a promising target for "post-Li" chemistry. Moreover, substitution of solid-state electrolytes for conventional organic liquid electrolytes can enhance energy density and specific capacity of the cell while mitigating flammability safety concerns. Sodium-based chalcogenides are promising candidates to realize sodium all-solid-state batteries owing to their high ionic conductivity ($>0.1 \text{ mS cm}^{-1}$), favorable interfacial contact with electrodes based on their deformability, and mild processing requirements.1-5

Synthesis of cubic Na₃PS₄ with high ionic conductivity (0.2 mS cm⁻¹) ignited interest in Na-based multinary chalcogenides. Since then, extensive research has led to the development of a variety of ternary and quaternary compounds such as Na_3PnCh_4 (Pn = P, Sb, As, Ch = S, Se), ⁷⁻¹³ $Na_{11}Sn_2$ -PnCh₁₂, ¹⁴⁻¹⁶ Na_{3-x}Pn_{1-x}W_xS₄, ¹⁷ etc. Quaternary chalcogenides

Chemical and Biological Engineering, Colorado School of Mines, Golden, Colorado 80401, USA. E-mail: cwolden@mines.edu

are typically formed through cation/anion doping of parent ternary compounds. The first reports of experimental and theoretical investigation of quaternary compounds in the Na-Sn-P-S system were inspired by highly conductive Li quaternary with Li₁₀MP₂S₁₂ (M = Si, Ge, Sn) composition. ^{18,19} These efforts resulted in the synthesis of Na₁₁Sn₂PS₁₂ (NSnPS), a slightly different stoichiometry than the original assumption, as the phase pure product of the Na-Sn-P-S system by different groups. 14,15 NSnPS possessed a newly explored tetragonal phase with I4₁/acd space group and achieved high ionic conductivity (1.4-3.7 mS cm⁻²) owing to the presence of vacancies in the crystal structure and three-dimensional network of Na+ migration pathways.

Similar to ternary Na₃PS₄, the substitution of the phosphorous central atom with other pnictogens was attempted to address the air and moisture sensitivity of NSnPS according to the "hard and soft acid-base" (HSAB) theory. Using "softer" Sb resulted in an isostructural Na₁₁Sn₂SbS₁₂ (NSnSbS) with different sodium site occupations and vacancy, resulting in a slightly lower ionic conductivity $(0.56 \text{ mS cm}^{-2})$. The difference in conductivity is attributed to changes in local bonding (P-S vs. Sb-S) and the lower electronegativity of Sb which leads to stronger Na-S coulombic attractions in the Sb quaternary.²⁰ In related work, Heo et al. 21 also reported Na_{4-x}Sn_{1-x}Sb_xS₄ $(0.02 \le x \le 0.33)$ electrolytes derived through systematic doping of Na₄SnS₄ with Sb atoms. Moreover, the substitution of S atoms with Se was designed to improve the transport

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properties of quaternary chalcogenides through lattice softening resulting in $Na_{11.1}Sn_{2.1}PSe_{12}$ composition.²² However, structure-transport studies of $Na_{11}Sn_2SbS_{12-x}Se_x$ (x=1, 6, and 12), solid-solution showed that the ionic conductivity decreases as the Se content is increased due to the change in the prefactor of the Arrhenius equation.³

In the majority of studies to date the quaternary chalcogenides were synthesized through high-temperature reactions between solid precursors in quartz ampoules, which is energy and time-intensive, and not amenable for industrial scale-up. This has prompted the development of liquid-based approaches which mitigate mass transfer limitations and enhance contact between reacting phases. In addition, the size and morphology of SSEs can be tuned by controlling liquid phase parameters such as solvent type, reagents chemistry, and reaction temperature. Hence, the choice of solvent becomes pertinent since aspects such as solubility and stability of reactants in the solvent and possible solvent complexes formation need to be considered.^{23,24}

The liquid-phase reaction of chalcogenide SSEs can be divided into two categories: "suspension" and "solution". In the "suspension" approach, binary/elemental precursors are dispersed in organic solvents that facilitate the desired reaction by forming soluble reactive complexes. Following the reaction, the SSE precursor is isolated, and post-synthetic processing like annealing is typically required to induce crystallization or phase transformation. In contrast, the "solution" approach is built on the full dissolution of precursors and simultaneous formation of SSEs in polar protic solvents (*i.e.* water, EtOH) or amine–thiol co-solvents. The latter provides a homogenous solution that is useful in composite cathode processing. Also, crystallization obtained at low temperatures eliminates the need for further heat treatment. ^{25,26}

In this context, Yoon group attempted to extend their aqueous synthesis route of Na₃SbS₄ to Na_{3.75}Sn_{0.75}Sb_{0.25}S₄ (or Na_{11,25}Sn_{2,25}Sb_{0,75}S₁₂). As the stoichiometry is slightly different than the more stable 11-1-2-12 phase, a major phase of Na₃SbS₄ was yielded at 200 °C while a higher annealing temperature was required to induce the quaternary phase.²⁷ Hartmann and coworkers also explored the Sb doping in the Na₄SnS₄ compound through co-decomposition of the hydrated phase of Na₄SnS₄ and Na₃SbS₄ at 170 °C. Both Na₄SnS₄·14H₂O and Na₃SbS₄·9H₂O were crystallized from aqueous solutions of respective precursors.²⁸ In the case of Na₁₁Sn₂PS₁₂, Weng et al. reported the first liquid-phase synthesis by utilizing 1,2dimethoxyethane (DME) that achieved 0.173 mS cm⁻¹ conductivity. Though the samples sintered at high temperatures (>400 °C) showed the formation of tetragonal Na₁₁Sn₂PS₁₂, the as-recovered powder from the liquid phase reaction still contained a significant quantity of unreacted raw materials (i.e., Na₂S and SnS₂) after a long reaction time. This observance along with the formation of unknown phases in the sintered sample indicates that the quaternary formation mainly occurred through sintering rather than liquid phase reaction.²⁹ In another pioneering study, the Yoon group extended the use of alkahest solvents (amine-thiol mixture) to a series of Li and Na

SSEs including NSnPS using ethylenediamine (EDA) and 1,2-ethanedithiol (EDT) co-solvents. The as-synthesized NSnPS contained significant carbonaceous products upon high-temperature annealing due to the high boiling point of dithiols. Consequently, the obtained NSnPS had higher electronic conductivity than ionic conductivity.²⁵ Even though this approach was shown to be useful in producing composite cathodes, it is deemed unsuitable for the synthesis of pure NSnPS. The number of reports on the solution synthesis of quaternary chalcogenides is very limited and selenide quaternaries, in particular, have not yet been synthesized in the liquid phase. In addition, there is a lack of understanding about the underlying mechanism of these approaches. This inspired us to investigate the impacts of the various liquid-phase synthesis routes for a series of quaternary chalcogenides.

In this work, we first demonstrate the synthesis of phase pure SnCh₂ (Ch = S, Se) via a simple metathesis reaction between hydrated tin chloride and sodium chalcogenide solutions. The resulting SnCh₂ is then used as a precursor in the synthesis of $Na_{11}Sn_2PnCh_{12}$ (Pn = P, Sb, Ch = S, Se) electrolytes through liquid-phase reactions. We compare the efficacy of "suspension" and "solution" routes using aprotic, protic, and alkahest solvents in the context of these compounds. For each material, we outline the design principles and examine how key parameters such as reagent choice, solvent chemistry, and thermal post-treatment influence the formation mechanism, using a series of complementary characterization techniques such as XRD, FTIR, and Raman spectroscopy. Finally, the transport properties and electrochemical stability are probed through electrochemical impedance spectroscopy, DC polarization, and linear sweep voltammetry. Our findings identify Na₁₁Sn₂SbS₁₂ as the fastest ionic conductor and most stable quaternary composition for use in sodium all-solid-state batteries. This study demonstrates that Na-based quaternary chalcogenides can be designed and produced systematically through solvent-assisted approaches, further advancing the scalable production of solid-state electrolytes for Na ASSBs.

Experimental methods

Materials

Anhydrous Na_2S and highly pure Sb_2Ch_3 (Ch = S, Se) were produced according to previously reported works. 11,30,31 Phosphorus pentasulfide (P_2S_5 , 99%, Sigma Aldrich), and elemental sulfur (S, Sigma Aldrich, 99.998% trace metal basis), sodium borohydride ($NaBH_4$, >98%, Sigma-Aldrich), selenium (Se, 99.99%, UMC), sodium (Na, Sigma-Aldrich), tin powder (Sn, Alfa Aesar, -325 mesh, 99.8%), tin(Iv) chloride pentahydrate ($SnCl_4 \cdot 5H_2O$, 98+%, extra pure, Thermo Scientific Chemicals), sodium carbonate (Na_2CO_3 , Alfa-Aesar, anhydrous ACS, 99.5% min), manganese(III) oxide (Mn_2O_3 , Strem, 99%), iron(III) oxide (Fe_2O_3 , Sigma Aldrich, 99%), acetonitrile (ACN, anhydrous, 99.8+%, Thermo Scientific Chemicals), ethanol (EtOH, Sigma-Aldrich, anhydrous, $\geq 99.5\%$), 1,2-ethylenediamine (EDA, Alfa Aesar, 99%), 1-propanethiol (PT, Thermo Scientific Chemicals),

98%), and UHP grade argon (Ar, 99.999%, General Air) were used as received without purification. Se pellets were further ground using a mortar and pestle to reduce particle size. All procedures were conducted in an Ar-filled glovebox (<5 ppm of H₂O) except otherwise stated.

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Sb₂Ch₃ and Na₃SbCh₄ (Ch = Se, Se) were prepared as described in the literature. 11,31

 SnS_2 . ~1.1 g batches of SnS_2 were produced as follows: SnCl₄·5H₂O (2105 mg, 6 mmol) were dissolved in 20 ml H₂O. Next, Na₂S (937.2 mg, 12 mmol) powder was slowly added to the tin solution. The metathesis reaction was instantaneous, and the solution turned orange. After stirring overnight, the solution was decanted and washed with excess H2O multiple times to remove the NaCl byproduct. Extra grinding and washing steps were necessary to remove the remaining NaCl. The precipitate was dried under vacuum overnight and further annealed at 300 °C. Extra grinding and washing steps were necessary to remove the remaining NaCl (yield = 78.3%).

SnSe₂. First, the NaBH₄ solution was prepared by adding 529.1 mg (13.97 mmol) of NaBH₄ in 20 ml ethanol and stirring for 1 h to fully dissolve. Next, NaBH4 solution was added dropwise to 920.3 mg (11.66 mmol) of Se in a three-neck flask that could facilitate the release of hydrogen gas. After stirring for 2 h at RT, a clear solution of NaHSe was formed. 20% excess NaBH₄ was used to ensure the full reduction of Se powder. In another container, 2043.2 mg (85.83 mmol) of SnCl₄·5H₂O was dissolved in 10 ml EtOH. Then, the tin solution was slowly added to the NaHSe solution and stirred overnight to ensure reaction completion. The solution was centrifuged and washed with excess water and EtOH consecutively and dried at RT for 12 h in vacuum (yield = 76.4%). The recovered powder was annealed under Ar flow at 300 °C in a horizontal tube furnace with a heating rate of 5 °C min⁻¹ to remove the residual solvents and form crystalline SnSe₂.

Na₁₁Sn₂PS₁₂. 237 mg (3.04 mmol) Na₂S, 152 mg (1.10 mmol) SnS₂, and 61 mg (0.28 mmol) P₂S₅ were ground with a mortar and pestle. The mixture was dispersed in 10 ml of ACN and stirred at 50 °C for 2 days. The precipitate was recovered by multiple centrifugations and washing with excess ACN followed by vacuum drying at RT. The powder was pelletized and annealed under dynamic vacuum at various temperatures for 1 h with a fast heating rate (40 °C min⁻¹). A similar reaction was performed in EDA-PT mixture for 1 day at RT. In this case, the quaternary was recovered by discarding the undissolved precipitate and drying the supernatant at 150 °C under Ar flow in a tube furnace (yield = 90-94%).

Na₁₁Sn₂SbS₁₂. This reaction was conducted in EtOH, H₂O, and EDA-PT solvents. In this process, Na2S (5% excess), S, assynthesized SnS2, and Sb2S3 were added to a 20 ml glass container, and respective solvents were added to the mixture. The H₂O and EDA-PT reactions were carried out at RT for 1 day while the ethanolic solution was heated at 50 °C for 2 days to ensure completion. In the case of EtOH, the solution was decanted and washed with excess EtOH multiple times, and

the precipitate was recovered at RT under vacuum. In the case of H₂O, the solution was dried as-is at 150 °C under Ar flow in a horizontal tube furnace. The collected powder was dispersed in EtOH and washed multiple times with excess EtOH to recover the quaternary sulfide. A similar recovery procedure to the NSnPS (EDA-PT) sample was carried out to obtain NSnSbS (EDA-PT). Na₄SnS₄ was obtained by reacting 525.9 mg (1.5 mmol) of SnCl₄·5H₂O and 468.3 mg (6 mmol) of Na₂S in 5 ml H₂O. The solution was stirred for one day at 70 °C. To precipitate the Na₄SnS₄ crystals, 20 ml of MeOH was added to the aqueous solution and stored in the fridge at 4 °C for 2 days. The formed crystals were washed with excess MeOH and dried at 180 °C for 1 h under Ar flow (vield = 96.3%).

Na₁₁Sn₂SbSe₁₂

Aqueous reaction. First, NaHSe aqueous solution was prepared as follows: 454.2 mg (12 mmol) of NaBH4 in 5 ml of water was slowly added to 395 mg (5 mmol) of Se at RT to prepare the aqueous NaHSe in a 25 ml flask. 20% excess NaBH4 was used to ensure the full reduction of Se powder. After 1 h of reaction, the solution was cooled down to 4 °C and centrifuged to collect and discard the precipitate. Next, 364 mg (0.81 mmol) of Sb₂Se₃, 181.9 mg (4.55 mmol) of NaOH, 457.5 mg (1.65 mmol) of SnSe₂, and 65.3 mg (0.83 mmol) of Se were added to the NaHS solution. The solution was stirred at RT for two days. To collect the selenide precipitate, 8 ml of acetone was added to the solution and cooled at 4 °C for 1 day. After centrifugation and excess washing with acetone, the precipitate was dried overnight under vacuum at RT (yield = 42.6%) to obtain NSnSbSe powder. 2× NaOH was used to ensure a complete reaction.

Ethanolic reaction. Ethanolic NaHSe was prepared by first dissolving 124.8 mg (3.30 mmol) of NaBH4 in 10 ml of EtOH and slowly adding to 217.1 mg (2.75 mmol) of Se powder in a 25 ml flask. Next, 276.6 mg (1 mmol) of SnSe₂, 120.1 mg (0.25 mmol) of Sb₂Se₃, 220 mg (5.5 mmol) of NaOH, and 39.5 mg (0.5 mmol) of Se were added to the ethanolic NaHSe. The solution was stirred at 50 °C for two days, decanted, and washed with EtOH twice. The precipitate was dried overnight under vacuum at RT (yield = 48.1%).

EDA-PT reaction. 109.2 mg (0.227 mmol) of Sb₂Se₃, 251.5 mg (0.91 mmol) of SnSe₂, 233.3 mg (2.99 mmol) of Se, and 115 mg (5 mmol) of Na were dissolved and reacted in 10 ml of EDA-PT (4-1) at RT inside the glovebox. After one day, the EDA-ET solution was centrifuged, and the collected supernatant was dried at 150 °C for 2 h under Ar flow in a tube furnace (yield = 73.3%). Recovered NSnSbSe was pelletized and further heattreated at various temperatures for 1 h under vacuum in a custom-built quartz tube. Na₄SnSe₄ was obtained by reacting 276.6 mg (1 mmol) of SnSe₂, 157.9 mg (2 mmol) of Se, and 92 mg (4 mmol) of Na in 5 ml of EDA-PT (4-1) at RT. A similar recovery procedure was employed to obtain powder Na₄SnSe₄ (yield = 77.3%).

Na-Sn alloy. Na-Sn anodes are prepared by milling the stoichiometric amounts of finely cut Na and Sn powder (total mass of ~ 2 g) in a planetary mill (Across International, PQN2)

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at 510 rpm for 20 h. The milling cycles consisted of 15 min of milling and 15 min of rest (yield = 65%).

Materials characterization

Thermogravimetric analysis and differential scanning calorimetry (TGA/DSC) were performed on a TA Instruments SDT-Q600 model. For a typical run, 10 mg of sample was loaded into a pre-cleaned alumina pan and heated under flowing Ar from RT to 600 °C at 10 °C min⁻¹ rate and then cooled down naturally. X-ray diffraction (XRD) was performed with a Philips X'Pert X-ray diffractometer with Cu K α radiation (λ = 0.15405 nm) between 10 and 60° at a scan rate of 5° min⁻¹. Samples were prepared on a glass slide with a protective tape covering the material to prevent undesired reactions with ambient air. XRD background subtraction was performed using HighScore software. Raman spectroscopy was conducted with a WiTec alpha 300 M confocal microscope/Raman spectrometer employing a 100 mW 532 nm laser. Samples were mounted on a glass slide and sealed under a 0.1 mm quartz cover slip. The laser was focused through the coverslip onto the sample using a 20× objective, and spectra were collected using a CCD detector (Andor Technologies) at -60 °C. HighScore software was used to subtract the Raman background.

Field emission scanning electron microscopy (FESEM) images were collected on a JEOL JSM-7000F FESEM instrument equipped with energy-dispersive X-ray spectroscopy (EDX) for compositional analysis. To prepare the samples for SEM and EDX measurements, powder samples were immobilized onto an aluminum stub using double-sided carbon tape. An accelerating voltage of 5 kV was used for the SEM image, while a higher voltage of 15-20 V was employed for EDX spectra collection. All sample preparation was done in an Ar glovebox.

Electrochemical characterization

Conductivity measurements. Pellets of quaternary SSE were prepared for conductivity measurement via conventional uniaxial pressing. For this purpose, 150-250 mg of the electrolyte was loaded into a 12 mm PEEK split cell with stainless steel plungers under a uniaxial fabrication pressure of 270 MPa and held for 5 min. Pellets were typically 0.9-1.1 mm thick (\sim 75–90% densification). The SSE pellets were contacted using stainless steel plungers as ion-blocking electrodes for electrochemical characterization. The pressure was released for 1 min, then increased again to 75 MPa. A Gamry Interface 1000E potentiostat was used to perform electrochemical impedance spectroscopy (EIS) measurements across a frequency range of 1 Hz to 1 MHz with a 10 mV perturbation. Temperaturedependent EIS testing was performed by heating the split cell apparatus with an electrical heating element and allowing it to stabilize at the target temperature for 1.5 h. DC polarization measurements were performed by applying multiple-step potentials to the sample (0.1, 0.2, 0.3, 0.4, and 0.5) and recording the transient current. The steady-state current was recorded after 2 h at each step potential, and the electrical conductivity was calculated using Ohm's law.

Symmetric cell cycling. Sodium plating/stripping cycling was performed in symmetrical Na-Sn|SSE|Na-Sn cells with a current density of 0.02 mA cm⁻² at 60 °C. To assemble the symmetric cell, 150 mg of SSE was cold pressed in a PEEK split cell at 270 MPa. Then, 50 mg of Na-Sn anode powders were dispersed on both sides of the pellet and pressed together at 340 MPa for 5 min. The finished cell was clamped in an inhouse built pressure jig fixated at 50 MPa. Galvanostatic charging and discharging (30 min per step) were performed after aging the cell at 60 °C for 2 h by a battery tester (MTI Corporation, BST8-MA).

Linear sweep voltammetry (LSV). LSV was performed on two Na-Sn|SSE|SS-SSE cells at RT using Gamry potentiostat. To fabricate the cells, stainless steel (SS) powder and SSE were hand-ground with a 21:9 ratio in a mortar to form the working electrode. The separator layer was formed by pressing 100 mg of the electrolyte at 150 MPa (5 min) in a 12 mm PEEK cell. Next, 20 mg of the working electrode was dispersed on the separator carefully and pressed at 275 MPa for 5 min. Lastly, the reference/counter electrode was formed by adding 60 mg Na-Sn on the other side of the separator layer and pressing the system at 275 MPa for 5 min. The stacking pressure of 50 MPa was applied during measurements. The cathodic and anodic scans were measured on the ranges open circuit voltage (OCV) -5 V and OCV -0 V at a scanning rate of 0.1 mV s⁻¹, respectively.

Results and discussion

SnCh₂ metathesis

One key challenge in developing Na-based solid-state electrolytes (SSEs) is the limited accessibility of their binary precursors. Binary chalcogenides, such as Na₂Ch, Sb₂Ch₃, and SnCh₂, are essential components and significant cost drivers for various components of Na all-solid-state batteries (ASSBs). However, their synthesis is typically expensive and primarily available for research purposes. 11,31,32 SnCh2 compounds, in particular, feature layered structures held together by van der Waals forces, which impart excellent optical and electronic properties. Wet chemical methods, including solvothermal^{33–36} and colloidal approaches, 37-40 have been reported for synthesizing SnCh₂. Solvothermal synthesis typically requires autoclaves capable of withstanding high pressure and temperature, while the colloidal method uses high boiling point solvents at elevated temperatures. Although these techniques yield size-controlled and highly pure tin chalcogenides, their complex reaction mechanisms, specialized equipment, and toxic precursors hinder their viability for large-scale production.

In this context, we employed simple metathesis reactions in polar solvents (H2O and EtOH) followed by mild heat treatment to produce the SnCh2 required for quaternary synthesis. Metathesis involves the exchange of counterions between two chemical species, often resulting in the precipitation of one product, which offers favorable energetics and facile separation. The salts used in this process are generally low-cost and readily available. 41,42 The reactions used for the synthesis of

SnS₂ and SnSe₂ in this work are outlined in eqn (1) and (2), respectively.

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$$2Na_2S_{(aq)} + SnCl_4 \cdot 5H_2O_{(aq)} \rightarrow SnS_{2(s)} + 4NaCl_{(aq)} + 5H_2O_{(l)}$$
(1)

$$2\text{NaHSe}_{(\text{sol})} + \text{SnCl}_4 \cdot 5\text{H}_2\text{O}_{(\text{sol})} \rightarrow \text{SnSe}_{2(\text{s})} + 2\text{NaCl}_{(\text{sol})}$$
$$+ 2\text{HCl}_{(\text{sol})} + 5\text{H}_2\text{O}_{(\text{I})} \tag{2}$$

Fig. 1a shows the XRD patterns of SnCh₂ compounds obtained from their respective reactions. For SnS2, the reaction was performed in H₂O, and the resulting precipitate was thoroughly washed with excess water to remove the NaCl byproduct. The sulfide was then recovered by vacuum drying at room temperature. Since the reduction of Se in water tends to produce persistent borax impurities, the SnSe₂ reaction was instead conducted in ethanol, following a similar recovery process as for SnS₂.31,43 The amorphous SnCh₂ powders obtained at room temperature were further heat-treated to enhance their crystallinity (Fig. S1 and S2, ESI†). Thermogravimetric analysis (TGA) revealed that a temperature of 300 °C was necessary to completely remove the solvate complexes in SnS₂ (Fig. S1a, ESI†).

The annealed SnS₂ exhibited a cubic structure (space group Fd3m), with the presence of pure cubic NaCl confirmed in the supernatant (Fig. S1b, ESI†). Additionally, the Raman spectra of the annealed SnS2 showed the characteristic SnS2 peak at 313 cm⁻¹, verifying its phase purity (Fig. 1b).⁴⁴ The annealed SnSe₂ sample displayed a CdI2-type hexagonal layered structure with P3m1 symmetry. Raman analysis also confirmed the presence of SnSe₂, with peaks at 187 cm⁻¹ and 117 cm⁻¹ corresponding to the A_{1g} and E_g modes of SnSe₂, respectively. 35,40 In addition to SnCh₂ compounds developed here, other binaries precursors, such as Sb₂Ch₃, were synthesized using the metathesis method as described in detail in our previous works. 11,31

Synthesis of Na₁₁Sn₂PS₁₂

The quaternary reaction to synthesize NSnPS was performed with Na₂S, SnS₂, and P₂S₅ as precursors according to the following stoichiometry (eqn (3)):

$$11Na_2S + 4SnS_2 + P_2S_5 \rightarrow 2Na_{11}Sn_2PS_{12}$$
 (3)

The choice of solvents for chalcogenide synthesis depends on factors like dielectric constant, polarity, and molecular composition, which influence the solubility and stability of reagents in the solvent. The dielectric constant (ε_r) reflects the solvent's ability to dissolve precursors. Polar solvents, such as water (H₂O) and ethanol (EtOH), have high dielectric constants due to hydrogen bonding. However, they are unsuitable for P-containing chalcogenides like Na₃PS₄ and Na₁₁Sn₂PS₁₂, as they have been shown to decompose P2S5 and PS43- moieties (Table S1, ESI†). 45 In contrast, aprotic solvents such as DME, diethyl ether (DEE), ethyl propionate (EP), ethyl acetate (EA), and ACN have been used in the Na₂S-P₂S₅ system. ⁴⁶⁻⁴⁹ These solvents are capable of forming reactive complexes without hydrolyzing the P₂S₅ precursor. ACN, in particular, offers both a high dielectric constant and easy solvent removal due to its low boiling point. Additionally, reagents like P₂S₅ and SnS₂ are shown to be stable and well-dispersed in non-polar ACN, promoting uniform mixing.

Building on these findings, we selected ACN as the reaction medium for the "suspension" synthesis of Na₁₁Sn₂PS₁₂. In this experiment, stoichiometric amounts of Na₂S, P₂S₅, and freshly prepared SnS2 were dispersed in ACN and reacted at 50 °C for 2 days. The resulting precipitate was recovered and washed with excess ACN before drying at room temperature under vacuum. The obtained powder was then heat-treated at various temperatures under vacuum to promote the solid-state reaction and enhance crystallinity. XRD and Raman analyses were conducted to assess the phase purity of the synthesized NSnPS materials and to investigate the formation mechanisms of the ternary and quaternary phases (Fig. 2a and b). The XRD pattern of the sample recovered at room temperature shows the

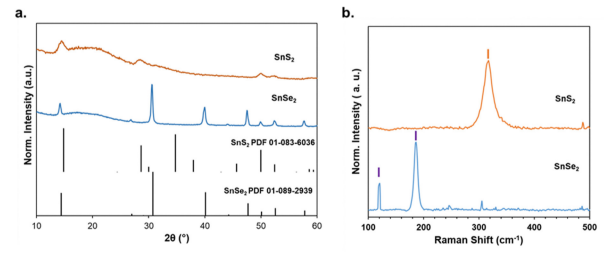


Fig. 1 (a) XRD and (b) Raman spectra of SnCh₂ (Ch = S, Se) recovered from metathesis reactions after annealing at 300 °C.

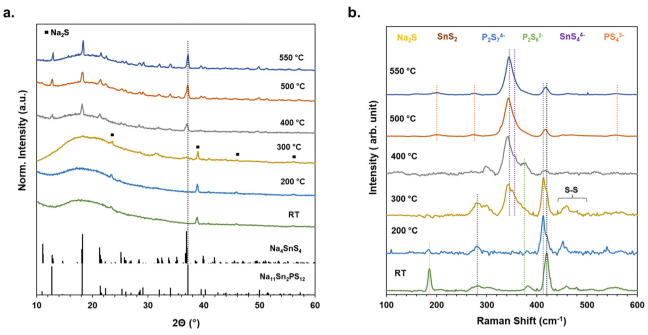


Fig. 2 (a) XRD, and (b) Raman spectra of NSnPS recovered from ACN and heat-treated at different temperatures.

presence of unreacted Na₂S; however, the Raman and FTIR spectra (Fig. S3, ESI†) indicate the formation of PS₄³⁻, P₂S₇⁴⁻, and P₂S₆⁴⁻ polyanions along with small traces of Na₂S and SnS₂, S-S bonds. The deconvoluted Raman spectra are presented in Fig. S4 (ESI†). Annealing at 200 °C did not cause any significant change in the XRD pattern of the NSnPS sample, but the Raman peaks attributed to P-S bonds indicated a potential shift in the intensity of [PS₄⁴⁻] and [P₂S₇⁴⁻] polyhedra, suggesting a possible transition from amorphous Na₃PS₄ to Na₄P₂S₇. ⁵⁰⁻⁵⁴ This reaction, which involves the release of Na₂S, can be represented as follows:

$$2Na_3PS_4 \rightarrow Na_4P_2S_7 + Na_2S \tag{4}$$

At 300 °C, a major peak at 342 cm⁻¹, accompanied by a strong shoulder at approximately 356 cm⁻¹, indicates the formation of $Sn_2S_6^{4-}/SnS_4^{4-}$ anions. However, there is no crystallographic evidence of Na₄SnS₆ formation in the XRD pattern.⁵⁵ Since Na₄SnS₄ and Na₄Sn₂S₆ exhibit very similar Raman vibration modes, it is challenging to definitively rule out the formation of either or a potential mixed polyanion compound without additional characterization techniques such as NMR.55,56 In contrast, Duchardt et al.57 demonstrated that the formation of NSnPS is accompanied by a blue shift to higher wavenumbers in the symmetric stretching vibration of Sn-S (345 cm⁻¹) and P-S (417 cm⁻¹) bonds compared to their original position in the corresponding Na₄SnS₄ (340 cm⁻¹) and Na₃PS₄ (412 cm⁻¹) compounds. The presence of a doublet peak in the 342-356 cm⁻¹ range may indicate the onset of NSnPS formation. Additionally, the peaks at 280.7 and 299.7 cm⁻¹ could be assigned SnS2 and Sn-S-Sn bonds, respectively.55 Although Na₂S remains the major crystalline phase, broad peaks associated with Na₄SnS₄/NSnPS appear in the XRD pattern, further confirming the Raman observations. Therefore, we can speculate that a minimum temperature of 300 °C is necessary for the onset of quaternary phase formation.

Further increasing the temperature stabilizes the quaternary phase, with minor precipitation of the Na_4SnS_4 phase. Raman analysis also confirms the consumption of $[PS_4^{\ 3}^{-}]$ and $[P_2S_7^{\ 4}^{-}]$ polyanions, likely through a solid-state reaction with available Na_4SnS_4 and Na_2S to generate the quaternary phase (eqn (5) and (6)). At this stage, all unreacted SnS_2 is converted into ternary and quaternary phases, indicating the consumption of all binary precursors. The peak at $375~cm^{-1}$ could be attributed to the characteristic P–P stretching mode of $[P_2S_6^{\ 4-}]$ polyanion, which is likely generated as a side product of high-temperature annealing through the release of S atoms from $Na_4P_2S_7^{\ 51}$

$$Na_2S + SnS_2 \rightarrow Na_4SnS_4$$
 (5)

$$Na_3PS_4 + \frac{1}{2}Na_4P_2S_7 + \frac{1}{2}Na_2S + 4Na_4SnS_4 \rightarrow 2Na_{11}Sn_2PS_{12}$$
 (6)

Higher temperatures of 500 and 550 °C further reduce the concentration of Na_4SnS_4 , indicating its incorporation into the quaternary phase. As the concentration of Na_4SnS_4 decreases, the Sn–S vibrations of NSnPS shift toward higher wavenumbers. Nevertheless, the ternary phase remains present, as Rietveld refinement for the 550 °C sample indicates a 4.7 wt% Na_4SnS_4 (Fig. S5, ESI†). The as-synthesized NSnPS exhibits a tetragonal phase (space group $I4_1/acd$, PDF 04-025-2554) with lattice parameters of $a = b = 13.6472\ 1736$ Å, c = 27.3634 Å, and V = 5096.52 ų, which aligns closely with previous reports of NSnPS synthesized via solid-state reactions. The corresponding FTIR data also reveals small traces of P–O bonds, likely due to oxidation during sample handling (Fig. S4, ESI†).

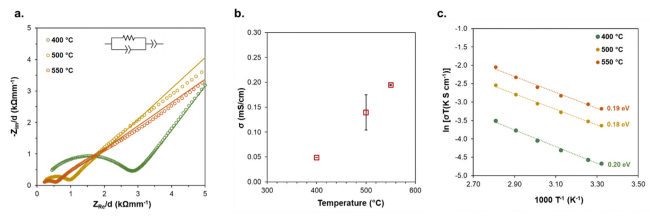


Fig. 3 (a) Nyquist plots, (b) corresponding ionic conductivity, and (c) Arrhenius plots of NSnPS synthesized in ACN as a function of annealing temperature.

The ionic transport properties of NSnPS recovered from ACN were investigated over a wide range of temperatures and frequencies using EIS. Fig. 3a shows the Nyquist plots of NSnPS heat-treated at different temperatures, while the extracted total conductivity values from equivalent circuit fitting are presented in Fig. 3b. The spectra were fitted to the equivalent circuit that consisted of a parallel combination of a resistor and constant phase element (CPE), which describes the suppressed semicircle, in series with another CPE that corresponds to electrode polarization (inset of Fig. 3a). Upon annealing from 300 °C to 550 °C, we observed an improvement in the ionic conductivity, correlating with the reduction in the concentration of Na₄SnS₄, highlighting the importance of phase purity on ionic transport in quaternary NSnPS. The highest ionic conductivity of 0.2 mS cm⁻¹ was achieved at 550 °C, which is in good agreement with the other reports of NSnPS synthesized through liquid phase routes (Table 1). 25,29 Even though the obtained conductivity is not explicitly superior to the other studies due to the presence of a low-conductive Na₄SnS₄ phase, the detailed investigation into the solvent influence and identification of potential impurities in this system is highly

valuable and could serve as a roadmap for the synthetic design of similar materials.

The kinetics of Na $^+$ transport were investigated through temperature-dependent EIS. Fig. 3c presents the Arrhenius plots for the corresponding annealed NSnPS, with activation energies extracted in the temperature range of 25–100 °C. The obtained activation energy values (0.18–0.20 eV) are slightly lower than those reported from other groups, potentially due to variations in experimental setups, off-stoichiometry, and the presence of impurity phases such as Na₄SnS₄ in the quaternary compounds (Table 1). The electronic conductivity of NSnPS compounds was measured using DC polarization in a symmetric cell configuration (Ti|SSE|Ti), similar to the EIS experiments. For the sample annealed at 550 °C, an electronic conductivity value of σ_e = 5.7 × 10⁻⁶ mS cm⁻² was obtained which was on the same order of magnitude as NSnPS derived from DME solvent in the report by Weng and coworkers.²⁹

We also explored the alkahest amine–thiol approach for the NSnPS reaction. For this, we selected a mixture of ethylenediamine (EDA) and propanethiol (PT) due to their combined lower boiling point compared to diamine–dithiol co-solvents. Lee

 Table 1
 Summary of quaternary synthesis parameters and performance reports in the literature

Quaternary	Synthesis method	Precursors	Heat- treatment (°C)	$\sigma_{\mathrm{Na^+}} \ \mathrm{(mS~cm^{-1})}$	$E_{\rm a}$ (eV)	Electrochemical window vs. Na ⁺ /Na (V)	Ref.
$Na_{11}Sn_2PS_{12}$	Solid-state (700 °C, 5 h)	Na ₂ S, P ₂ S ₅ , SnS ₂	_	1.4	0.25	_	14
$Na_{11}Sn_2PS_{12}$	Solid-state (600 °C)	Na ₃ PS ₄ , Na ₄ SnS ₄	_	3.7	0.39	_	15
$Na_{11}Sn_2PS_{12}$	Suspension (DME)	Na_2S , P_2S_5 , SnS_2	460	0.17	0.33	0.3 - 3.2	29
$Na_{11}Sn_2SbS_{12}$	Solid-state (700 °C, 5 h)	Na_2S , Sb_2S_3 , SnS_2 , S	_	0.56	0.34	_	20
$Na_{4-x}Sn_{1-x}Sb_xS_{12}$	Solid-state (450–550 °C, 12 h)	Na_2S , Sb_2S_3 , SnS_2 , S	550	0.51	0.39	$\sim 0.4 - 4.5$	21
$(0.02 \le x \le 0.33)$				(x = 0.25)	(x = 0.25)		
$Na_{11.1}Sn_{2.1}PSe_{12}$	Solid-state (600 °C, 12 h)	Na ₂ Se, Sn, P, Se	600	1.7	0.30	_	22
$Na_{11}Sn_2PSe_{12}$	Solid-state (900 °C, 48 h)	Na ₂ Se, Sn, P, Se	900	2.15	0.28	1.22 - 2.65	58
$Na_{11}Sn_2PSe_{12}$	Milling (400 rpm, 15 h), annealing (600 $^{\circ}$ C)	Na, Sn, P, Se	550	0.13	_	_	59
$Na_{11}Sn_2SbSe_{12}$	Milling (33 h, 400 rpm) solid-state (750 °C, 5 h)	Na ₂ Se, Sb ₂ Se ₃ , SnSe ₂ , Se	750	0.15	0.39	_	3
$Na_{11}Sn_2PS_{12}$	Suspension (ACN)	Na_2S , P_2S_5 , SnS_2	550	0.2	0.19	0.95-1.7 V	This
$Na_{11}Sn_2SbS_{12}$	Solution (H ₂ O)	Na ₂ S, Sb ₂ S ₃ , SnS ₂ , S	550	0.42	0.14	1.15-3.1 V	work
$Na_{11}Sn_2SbSe_{12}$	Solution (EDA-PT)	Na, Sb ₂ Se ₃ , SnSe ₂ , Se	550	0.07	0.22	0.85-2.0 V	

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et al.25 previously demonstrated that P2S5 fully dissolves in EDA-ethanethiol (ET) solvents without any noticeable hydrolysis. Our attempt with EDA-PT yielded a compound with an XRD pattern similar to what was reported in their study. A closer examination of the XRD pattern, coupled with Raman analysis (Fig. S6, ESI†), indicated the formation of ternary Na₄SnS₄ with a tetragonal phase (space group I4₁/acd), rather than the expected Na₁₁Sn₂PS₁₂ in the diamine-monothiol solvents. Interestingly, Hartmann et al.28 recently identified this new phase of Na₄SnS₄, which is isostructural to NSnPS and NSnSbS compounds, through controlled dehydration of Na₄SnS₄·14H₂O crystals.

Based on these observations, we can infer that the formation of Na₁₁Sn₂PS₁₂ in EDA-EDT as described by Yoon group, ²⁵ but not in EDA-PT, is closely related to the specific nature of P₂S₅ dissolution and the thiolate species produced in these systems. A recent study on chalcogen dissolution in amine-thiol solutions found that the type of amine and thiol can significantly impact the reactive intermediates formed, which, in turn, influences subsequent reactions.⁶⁰ Further investigations are needed to elucidate the specific reaction pathway that facilitates phosphorus incorporation to form NSnPS in diaminedithiol solvents. In conclusion, the suspension synthesis of NSnPS with ACN proves to be the optimal choice over the diamine-dithiol approach owing to its less complicated synthesis procedure, lower electronic conductivity, and use of benign and environment-friendly solvent. ACN offers a safer alternative compared to the relatively toxic amine-thiol cosolvents.

Synthesis of Na₁₁Sn₂SbS₁₂

Na₁₁Sn₂SbS₁₂ enhances the air stability relative to its isostructural analog, Na₁₁Sn₂PS₁₂. Sb-based and Sn-based chalcogenides such as Na₃SbCh₄ and Na₄SnCh₄ (Ch = S, Se) can be synthesized in polar solvents like H2O, MeOH, and EtOH. For example, the aqueous solution synthesis of Na₄SnS₄ has been reported with Na₂S·9H₂O, SnCl₄, or SnS₂ precursors. 61,62 We recently investigated different synthetic protocols for Na₃SbCh₄ compounds in EtOH and demonstrated that the formation of reactive Na chalcogenides is a critical step in producing highly pure Na_3SbCh_4 . 11,31,43 In these compounds, the nucleophilic attack of the chalcogenide ions on the binary chalcogenides, such as Sb₂Ch₃, promotes the formation of [SbCh₃]³⁻ anions, followed by the reduction of elemental chalcogen to produce [SbCh₃]⁴⁻ polyhedral anions. We hypothesized that a similar chemistry would be feasible in the synthesis of Na₁₁Sn₂SbS₁₂, utilizing the formation of [SbS₄]³⁻ and [SnS₄]⁴⁻ polyanions. Accordingly, the following reaction scheme was attempted in H₂O and EtOH:

$$11Na_2S + Sb_2S_3 + 4SnS_2 + 2S \rightarrow 2Na_{11}Sn_2SbS_{12}$$
 (7)

Sodium sulfide was prepared by dehydration and reduction of the low-cost technical grade hydrate (Na₂S·xH₂O)³⁰ and the other binary sulfide precursors were procured via metathesis as discussed above. Since NSnSbS is soluble in H2O, it is necessary to use an antisolvent, such as EtOH or acetone, to separate the quaternary compound. In this experiment, Na2S

(with a 10% excess), freshly prepared SnS2, Sb2S3, and S were dispersed in H2O and reacted at room temperature. The slight excess of Na2S was intended to ensure complete reaction. To recover NSnSbS, the aqueous solution was dried at 150 °C under an inert argon atmosphere and washed with excess EtOH to remove unreacted precursors and soluble intermediates.

The reaction in EtOH was conducted with a similar stoichiometry. Initial attempts indicated slow kinetics for NSnSbS formation in the less polar EtOH; therefore, the solution was heated to 50 °C for an extended period (2 days) to ensure a complete reaction. Since NSnSbS is insoluble in EtOH, this approach facilitated its direct precipitation at room temperature. The sample recovered from EtOH was further dried at 150 °C to eliminate any remaining solvent and to ensure consistent heat treatment with the aqueous experiment. Additionally, we investigated the applicability of amine-thiol cosolvents for this stoichiometry. Pioneering work by the Butchery group has demonstrated the dissolution and redeposition of binary chalcogenides, such as Sb₂Ch₃ and SnS₂, in aminethiol co-solvents. 63,64 Since all precursors are sufficiently soluble in the EDA-PT mixture, this reaction was performed at room temperature for 1 day, and the powder was recovered by discarding unreacted precipitate and drying the supernatant at 150 °C.

Fig. 4a shows the XRD patterns of compounds obtained from H₂O, EtOH, and EDA-PT solvents, with reference patterns for Na₄SnS₄ and Na₃SbS₄ included for comparison. Detailed synthesis protocols for Na₄SnS₄ and Na₃SbS₄ in aqueous solutions can be found in the Experimental section. The Na₁₁SnSbS₁₂ recovered from all solvents exhibits a tetragonal crystalline phase (I4₁/acd space group) consistent with the reported structure for this compound.20,21 The NSnSbS obtained from an aqueous solution is highly crystalline highlighting the effectiveness of highly polar solvents in promoting room-temperature crystallization of quaternary chalcogenides. In contrast, the powders recovered from EtOH and EDA-PT solvents exhibited lower crystallinity with small traces of Na₃SbS₄ detected. Since NSnSbS is isostructural with the Na₄SnS₄ recovered from the aqueous solution, XRD alone is inconclusive regarding the formation of NSnSbS.²⁸ Therefore, Raman spectroscopy was conducted to provide deeper insights into their molecular order and to assess the influence of Sb doping on the structure.

The corresponding Raman spectra are depicted in Fig. 4b. To accurately identify the existing vibrational modes, peak deconvolution was conducted for the Raman peaks of the samples recovered from H2O, EtOH, and EDA-PT (see Fig. S7, ESI†). The NSnSbS obtained from the aqueous solution displays peaks at approximately 341 cm⁻¹ and 363 cm⁻¹, which corresponds to the symmetric vibration (ν_s) of Sn-S and Sb-S, respectively. For the asymmetric stretching modes of the SbS₄ anions, expected peaks at $\nu_a = 385 \text{ cm}^{-1}$ and 403 cm⁻¹ were observed in the pure Na₃SbS₄ sample.⁶⁵ However, in the H₂Orecovered sample, these peaks merged into a broader peak at 399 cm⁻¹, indicating a change in the local structure due to the incorporation of Sn. Additionally, a strong peak at 372 cm⁻¹ can be attributed to the [SbS₄] polyhedra within the quaternary

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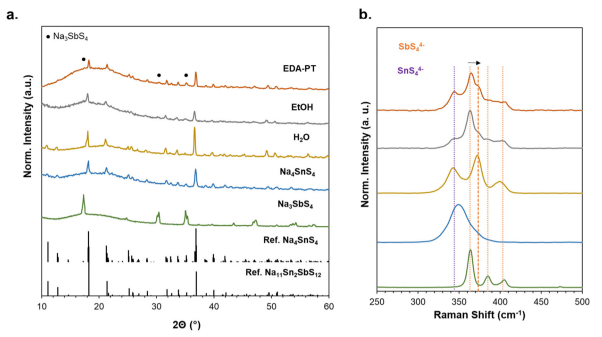


Fig. 4 (a) XRD, and (b) Raman of NSnSbS recovered from different solvent systems and dried at 150 °C

lattice framework. The NSnSbS structure features two types of isolated SnS₄ (or SbS₄) tetrahedra, positioned on distinct Snonly sites and an Sb mixed site.²⁰

The XRD pattern of NSnSbS obtained from EtOH and EDA-PT shows a strong presence of pure Na₃SbS₄, which is further emphasized by the high-intensity Sb-S stretching mode at 363 cm⁻¹ in the corresponding Raman spectra. The emergence of a peak at 372 cm⁻¹ as a smaller shoulder supports the direct correlation between the [SbS₄] peak signature and the extent of Sb incorporation into the quaternary lattice. Similar to the findings for NSnPS, a blue shift in the Raman peaks of the SbS₄ and SnS₄ polyanions was observed, likely due to alterations in the local bonding environment compared to the constituent ternary Na₃SbS₄ and Na₄SnS₄. ^{56,65} The Raman analysis provides clear evidence for the generation of quaternary Na₁₁Sn₂SbS₁₂ compounds; however, higher-resolution X-ray or neutron diffraction studies are necessary to fully elucidate the local crystal structure of the formed quaternary.

To enhance the Na⁺ transport properties, the NSnSbS powders were annealed at 550 °C under vacuum. Fig. S8 (ESI†) shows the XRD patterns of annealed NSnSbS samples. The aqueous-recovered sample shows no apparent secondary phases, while the ethanolic NSnSbS exhibits traces of NaSbS2, likely formed from the decomposition of the Na₃SbS₄ phase at high temperatures. The sample recovered from EDA-PT indicates only traces of Na₃SbS₄. The ionic and electronic conductivities of the samples are summarized in Fig. 5a. As expected, the aqueousrecovered NSnSbS, having the highest crystallinity and phase purity, demonstrates a maximum conductivity of 0.42 mS cm⁻¹ at room temperature - approximately four times higher than the other samples containing NaSbS2 impurities. The electronic conductivity across all samples ranges from 3.8×10^{-7} to

 3.1×10^{-6} mS cm⁻¹, rendering them suitable superionic conductors.

Fig. 5b displays the Arrhenius plots for the NSnSbS samples, indicating a low activation barrier for the aqueous-recovered NSnSbS, attributed to its high crystallinity and phase purity. Additionally, Fig. S9 (ESI†) provides the XRD and Raman spectra from the heat treatment studies conducted on NSnSbS recovered from aqueous solutions. The trend of ionic conductivity as a function of temperature (Fig. S9c, ESI†) illustrates nearly a twofold increase in the ionic conductivity of the quaternary phase as the annealing temperature rises to 550 °C. This enhancement is likely due to the complete incorporation of Sb into the quaternary lattice and the removal of solvated complexes. Overall, these findings highlight the potential of aqueous synthesis methods for developing Sn and Sb-based chalcogenide solid-state electrolytes.

Synthesis of Na₁₁Sn₂SbSe₁₂

Lastly, we focused on quaternary selenides with the composition $Na_{11}Sn_2PnSe_{12}$ (where Pn = P, Sb). The common precursors for this class typically include elemental Na, P, Se, or binary selenides such as Na₂Se, SnSe₂ and Sb₂Se₃. As for the Sb analog, we explored various activated Se precursors in polar solvents, since aprotic solvents are not suitable due to the insoluble nature of elemental Se. Our attempts to synthesize NSnSbSe using aqueous and ethanolic solutions of NaHSe, following the reaction scheme outlined in eqn (8), led to the formation of Na₃SbSe₄. However, there was no apparent doping of Sn into the lattice structure of the resultant compound, as evidenced by the data presented in Fig. S10 (ESI†).

$$11\text{NaHSe}_{(\text{sol})} + \text{Sb}_2\text{Se}_{3(\text{s})} + 4\text{SnSe}_{2(\text{s})} + 3\text{Se}_{(\text{s})} + 11\text{NaOH}_{(\text{sol})}$$

$$\rightarrow 2\text{Na}_{11}\text{Sn}_2\text{SbSe}_{12(\text{s})} + 11\text{H}_2\text{O}_{(\text{l})}$$
(8)

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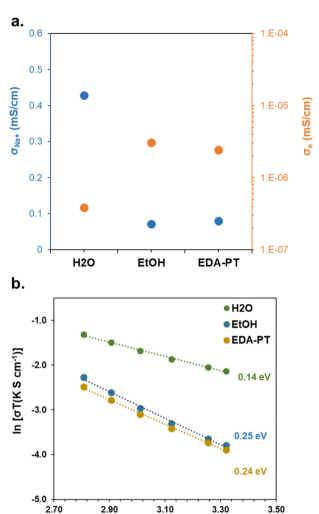


Fig. 5 (a) Ionic and electronic conductivity, and (b) Arrhenius plots of NSnSbS obtained from H_2O , EtOH, and EDA-PT solvents, and further annealed at 550 $^{\circ}$ C.

1000 T-1 (K-1)

Detailed descriptions of the reactions involved are provided in the Experimental section. Further annealing of the recovered powder from aqueous and ethanolic solutions resulted in the decomposition of the material into elemental and binary compounds.

To facilitate Sn incorporation, we turned to the use of amine-thiol solvents. Our previous work has demonstrated that amine-thiol solvents are particularly effective for synthesizing Na₃SbSe₄ by activating selenium through a redox process. The use of amine-thiol co-solvents allows for the dissolution of highly insoluble selenium at high concentrations, generating homogeneous inks suitable for the solution processing of inorganic semiconductors.^{66,67} In this process, an amine-thiol mixture produces highly reactive ammonium ions and thiolate species. The thiolate formation reduces Se to its anion form, while the complexation of selenium with ammonium ion yields soluble selenium in the form of [(Se⁻²)(R-NH₃⁺)₂]. Furthermore, this solvent combination has been shown to effectively dissolve a wide range of binary selenides, such as

 $Sb_2Se_3^{64,68,69}$ and $SnSe.^{70,71}$ Inspired by these studies, we attempted the NSnSbSe synthesis in EDA-PT (4:1 v/v) with the following stoichiometry:

$$22\text{Na}_{(\text{sol})} + \text{Sb}_2\text{Se}_{3(\text{sol})} + 4\text{SnSe}_{2(\text{sol})} + 11\text{Se}_{(\text{sol})}$$

$$\rightarrow 2\text{Na}_{11}\text{Sn}_2\text{SbSe}_{12(\text{sol})}$$
(9)

Fig. 6a displays the XRD patterns of NSnSbSe recovered from EDA-PT by drying at 150 °C. The predominant phase of NSnSbSe with considerable Na₃SbSe₄ secondary phase was observed. Upon annealing the sample at 550 °C, we found that the ternary phase incorporates into the quaternary phase through a solid-state reaction. However, attempts to anneal the sample at temperatures exceeding 550 °C led to significant evaporation of selenium species and melting of the material, complicating sample collection. To facilitate high-temperature reactions, melt-quenching techniques using enclosed quartz containers are required.

The Raman spectra of the synthesized selenide at 150 °C and 550 °C are compared with those of the parent ternary compounds, Na₄SnSe₄ and Na₃SbSe₄, in Fig. 6b. The Na₃SbSe₄ is produced through the reaction of Sb₂Se₃ with sodium and selenium dissolved in EDA-PT solvents, as detailed in our previous work. Similarly, the reaction involving SnSe₂, sodium, and selenium results in the formation of highly pure Na₄SnSe₄. The NSnSbSe compound exhibits strong Raman peaks at 197 and 230 cm⁻¹, which correspond to the vibrations of the Sn–Se bond, similar to those observed in other selenostannates such as Ba₂SnSe₄. The peaks at 206, 259, and 271 are attributed to the stretching modes of SbSe₄ tetrahedra, as seen in the Na₃SbSe₄ spectrum, which closely resembles other compounds like K₃SbSe₄. SlSe₄. Slate

The samples exhibit all the characteristic peaks associated with $[Sn/Sb]Se_4$ polyhedra, further confirming the successful formation of the quaternary phase. Consistent with the behavior observed in NSnSbS, the signature peak of SbSe₄ at 207 cm⁻¹ shifts to a higher wavenumber (\sim 212 cm⁻¹) upon annealing, indicating effective incorporation of Sb into the lattice structure. This trend is also noted for other vibrational modes of SbSe₄, reinforcing the notion of structural modifications upon thermal treatment.

However, the uncorrected full spectrum reveals the presence of D and G bands, which are indicative of carbonaceous species formed during solvent removal, even at low-temperature drying (Fig. S11, ESI†). This presence of carbonaceous material is likely contributing to the elevated electronic conductivity observed for NSnSbSe, as shown in Fig. S12 (ESI†). EIS analysis indicates that the 550 °C sample achieves an ionic conductivity (σ_{Na^*}) of 0.07 mS cm⁻¹ and an activation energy (E_a) of 0.22 eV, which are considerably lower than those observed in other quaternary phases. This reduction can be attributed to the significant presence of the Na₃SbSe₄ secondary phase within the material. Previous attempts at the solid-state reaction of Na₁₁Sn₂SbSe₁₂ resulted in a substantial impurity phase of NaSbSe₂, which negatively impacted its ionic properties, yielding an ionic conductivity of 0.15 mS cm⁻¹. Despite the presence of

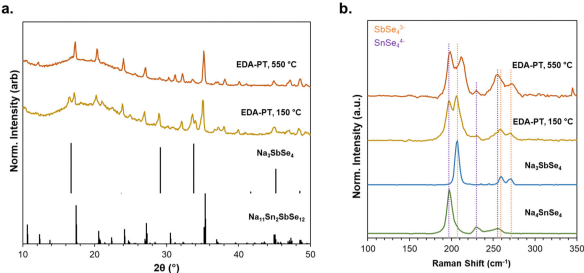


Fig. 6 (a) XRD, and (b) Raman of NSnSbSe recovered from amine-thiol solvents after annealing at 150 and 550 °C, respectively.

carbonaceous impurities, the current work represents the first liquid-based synthesis of quaternary selenide materials, to our knowledge. This pioneering approach could open new avenues for the development of high-performance solid-state electrolytes. Fig. S12 (ESI†) and Table 1 summarize the preparation conditions and transport properties of $Na_{11}PnCh_{12}$ (Pn = P, Sb; Ch = S, Se) produced in this work.

For the P-based selenide, the solution approach is highly challenging due to the extreme hygroscopic nature of white phosphorus (P₄). While red phosphorus serves as a more stable alternative, it is difficult to activate in organic solvents. Strumolo et al. 75 recently reported that red phosphorus can dissolve in an amine-thiol mixture, forming primarily cage-like polyphosphide anions (P₁₆²⁻). However, our attempts to utilize these polyphosphides in EDA-PT were unsuccessful, as significant portion of dissolved phosphorus precipitated out of the solvent mixture. The Raman analysis of the obtained supernatant (Fig. S13, ESI†) closely matched ternary Na₄SnSe₄, suggesting that phosphorus remained unreactive in these reactions, likely due to steric effects associated with the polyphosphide structure.

Electrochemical performance of quaternary chalcogenides

We assessed the electrochemical stability window assynthesized quaternary compounds using linear sweep voltammetry (LSV) in a Na-Sn|SSE|SS-SSE cell configuration at room temperature, with a scan rate of 0.1 mV s⁻¹. As shown in Fig. 7a, NSnSbS exhibits a wide electrochemical stability window, with reduction and oxidation processes occurring around 0.95 V and 2.9 V vs. Na-Sn alloy, respectively. A similar potential range was reported by Weng et al. through cyclic voltammetry, though the discrepancy in the low redox potential may stem from

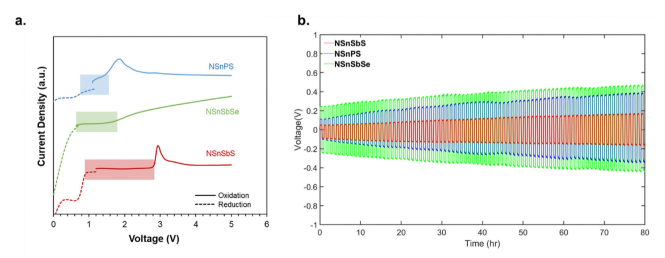


Fig. 7 (a) Linear sweep voltammetry of Na₁₁Sn₂PnCh₁₂ (Pn = P, Sb; Ch = S, Se) cells with a voltage range of 0-5 V at a scan rate of 0.1 mV s⁻¹, and (b) cyclability of Na-Sn|Na₁₁Sn₂PCh₁₂|Na-Sn symmetric cells at RT with a stacking pressure of 50 MPa.

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differences in establishing faradaic current onset in voltammetry data. To In contrast, NSnPS and NSnSbSe exhibit narrower stability windows of approximately 0.75–1.5 V and 0.65–1.8 V, respectively. Given the Na–Sn alloy potential of 0.2 V vs. Na+/Na, the updated stability windows are summarized in Table 1. To our knowledge, NSnSbSe's electrochemical stability has not been previously reported. Selenides appear more susceptible to oxidation than their sulfide counterparts at elevated potentials resulting a smaller electrochemical stability as observed in other selenide systems such as selenophosphate-based lithium argyrodites.

We also assembled symmetric Na–Sn|SSE|Na–Sn cells to evaluate the stability of the anode-solid electrolyte interface. As shown in Fig. 7b, NSnSbS exhibited the lowest initial overpotential of 50 mV, which gradually increased to ~ 169 mV over 90 hours of cycling. In contrast, NSnPS and NSnSbSe displayed significantly higher overpotentials of ~ 400 mV and ~ 470 mV, respectively. The transport property measurements (as shown in Fig. S13, ESI† and summarized in Table 1) confirm that aqueous-derived NSnSbS is the most suitable separator for Na all-solid-state batteries, owing to its high ionic conductivity, low electronic conductivity, and low activation barrier.

Conclusions

In conclusion, we presented the rational design of methods for the synthesis of quaternary chalcogenide solid-state electrolytes, $Na_{11}Sn_2PnCh_{12}$ (Pn = P, Sb; Ch = S, Se), using aprotic, protic, and alkahest solvents. The synthesis of P-containing quaternary is more complex due to the limited inventory of phosphorus reagents. Specifically, the high reactivity of P₂S₅ led us to use acetonitrile (ACN) with a high dielectric constant to ensure intimate mixing of the precursors, forming amorphous thiophosphate moieties, which were then converted to Na₁₁Sn₂S₁₂ through annealing. We found that heat-treatment at 300 °C is necessary to initiate the quaternary phase formation in this system, while higher temperatures (550 °C) are required to obtain phase pure Na₁₁Sn₂PS₁₂ with minimal Na₄SnS₄ impurity. For Sb-based analogs, polar solvents such as water, ethanol, and amine-thiol cosolvents facilitated the reaction and crystallization of Na₁₁Sn₂SbS₁₂ at a low temperature of 150 °C, with additional annealing needed to further enhance transport properties. The aqueous-recovered sample produced Na-based chalcogenides with performance comparable to that of energy-intensive, hightemperature solid-state reactions. Notably, we achieved the first liquid-phase synthesis of Na₁₁Sn₂SbSe₁₂ in this work, despite the presence of minor impurities. Overall, our findings demonstrate the potential of liquid-phase methods for the development of quaternary chalcogenides, offering advantages in achieving high ionic conductivity and phase purity, and paving the way for scalable production of Na-based solid-state electrolytes.

Author contributions

The manuscript was written through the contributions of all authors. All authors have approved the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

Data availability

The data supporting this article have been included as part of the ESI.†

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