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Co-sintering a cathode material and garnet electrolyte to develop a bulk-type solid-state Li metal battery with wide electrochemical windows†

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Co-sintering a cathode material and the $\text{Li}_7\text{La}_3\text{Zr}_2\text{O}_{12}$ (LLZ) electrolyte can assist in fabricating bulk-type all-solid-state batteries (ASSBs). However, owing to the use of low temperatures to prevent reactions between the different materials, this process can result in low relative densities and ionic conductivities. To overcome this trade-off between sintering temperature and electrochemical performance, a highly sinterable Ca–Sb-doped LLZ electrolyte integrated with the sintering aids Li_7SbO_6 and Li_3BO_3 was developed in this study. A composite powder of $\text{Li}_{6.5}(\text{La}_{2.925}\text{Ca}_{0.075})(\text{Zr}_{1.425}\text{Sb}_{0.575})\text{O}_{12}$ (denoted as LLZ–CaSb) and $x\text{Li}_7\text{SbO}_6$ ($x = 0–0.14$), whose composition differed from that of stoichiometric LLZ–CaSb, was prepared, with Li_3BO_3 added prior to sintering. The disk incorporated with the sintering aid $0.08\text{Li}_7\text{SbO}_6$ ($x = 0.08$) and then sintered at $750\text{ }^\circ\text{C}$ exhibited a relative density and ionic conductivity of 87% and $3.1 \times 10^{-4}\text{ S cm}^{-1}$, respectively, indicating its superior sintering properties. The formation of a low-melting-point liquid phase, generated by the mutual reaction between Li–Sb–O- and Li–B–O-type oxides, promoted sintering. The developed electrolyte exhibited a wide potential window (5 V vs. Li/Li^+). The ASSB assembled using the co-sintered LiCoO_2 /electrolyte system and Li metal anode maintained 98.6% of the initial discharge capacity (120.7 mA h g^{-1}) after 60 cycles at $25\text{ }^\circ\text{C}$ and 0.1C and performed adequately over a wide temperature range (-25 to $120\text{ }^\circ\text{C}$). The method reported herein has potential for achieving cost-effective large-scale ASSB production.

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Introduction

The use of renewable electricity and the electrification of automobiles have become crucial strategies for realising a carbon-neutral society. Consequently, the demand for high-capacity lithium-ion batteries (LIBs) is increasing. Traditional LIBs comprise a positive active material, organic liquid electrolyte, separator, negative active material, and current collector. However, the organic electrolyte component poses a fire hazard, especially in high-capacity batteries.^{1–3} Therefore, all-solid-state batteries (ASSBs) with non-flammable inorganic solid electrolytes are gaining attention for their ability to safely achieve high capacities.⁴

To realise high-capacity ASSBs, improving the electrode properties by, for example, increasing the cathode voltage and anode capacity, is crucial. Therefore, solid electrolytes with

a high lithium-ion conductivity and wide electrochemical window are required. Additionally, the use of the Li metal anode, which has a low operating voltage and high capacity (3860 mA h g^{-1}),⁵ has drawn research attention. Among oxide-based solid electrolytes, the garnet-type series shows considerable promise because of its high stability against Li metal and relatively high room-temperature ionic conductivity ($\sim 10^{-4}\text{ S cm}^{-1}$).^{6,7} This has led to a surge in research on the formation of the highly stable low-resistivity $\text{Li}/\text{Li}_7\text{La}_3\text{Zr}_2\text{O}_{12}$ (LLZ) interface for developing LLZ-based ASSBs.^{8–11}

However, the formation of a cathode/LLZ interface with attributes that can facilitate battery operation, such as low interfacial resistance and good contact, is a major challenge in realising LLZ-based ASSBs. Therefore, strategies such as sintering at temperatures above $1000\text{ }^\circ\text{C}$ have been adopted for the densification of LLZ to enhance its ionic conductivity and stability against Li metal. However, this high-temperature sintering inevitably leads to the formation of a highly resistive electrode/LLZ interphase, resulting in poor battery performance.^{12,13} A cathode/LLZ interface can be formed *via* two routes. In the first path, which involves coating the cathode layer onto sintered LLZ disks, LLZ is densified at an elevated temperature and then coated with the cathode layer using methods such as screen printing and immersion, and the resulting cathode/LLZ system is co-fired at a low temperature to

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suppress the reaction between the cathode and LLZ.^{14,15} In the second route, which involves simultaneously co-sintering the cathode and LLZ, densification of LLZ and formation of the cathode and LLZ with low resistivity are achieved. Although this co-sintering approach is technically difficult to implement, the number of sintering stages is reduced. In particular, co-sintering can be considered an environmentally friendly process from the viewpoint of energy savings and low CO₂ emissions if a sufficiently low co-sintering temperature is used. Therefore, low-temperature co-sintering is worth investigating.

Numerous approaches such as hot pressing and field-assisted sintering have been reported for reducing the LLZ sintering temperature.^{16–19} However, these processes have high equipment and production costs owing to the requirement of batch processing. In contrast, the use of sintering aids such as Li₃BO₃, Li₃PO₄, and Li₄SiO₄ is an attractive strategy for achieving low-temperature sintering because it does not require special equipment.^{20–26} Furthermore, low-temperature sintering with sintering aids can also be applied to tape casting for mass-producing ceramic products. Therefore, this strategy can potentially facilitate cost-effective ASSB production.

Previously, we conducted low-temperature liquid-phase sintering of LLZ by targeting sintering aids, doping elements, intentional compositional deviation, and wettability of the liquid phase for LLZ particles.^{27–29} Consequently, we obtained Ca–Bi-doped LLZ (LLZ–CaBi) with a relative density and conductivity of 94% and 1.2×10^{-3} S cm⁻¹, respectively, by sintering at 750 °C and using an Li–Ca–Bi–O system as the sintering aid. However, this electrolyte was not stable against the Li metal anode, given the ability of Li metal to promote the reduction of LLZ depending on the doping element.^{30,31} According to Zhu *et al.*, the element used for doping the Zr sites in LLZ modulates the stability of LLZ against Li metal.³⁰ Therefore, the reduction of Bi⁵⁺ at the Zr sites evidently causes the instability against Li metal. Density functional theory (DFT) calculations have also predicted the instability of Bi-based garnets against Li metal.³² Consequently, the development of a Bi-free LLZ with high ionic conductivity, superior sintering properties, and high stability against Li metal is essential for realising co-sintered LLZ-based batteries.

Elemental substitution is an effective material design strategy for reducing the LLZ sintering temperature, as it can promote the formation of a liquid phase at lower temperatures; this liquid phase should exhibit good wettability against LLZ particles and dissolve part of the LLZ particles, as established in our previous study.²⁹ Low-melting-point oxides exhibit a low dissociation energy with respect to oxygen. According to Sun, As⁵⁺ and Sb⁵⁺ are pentavalent elements with lower dissociation energies than Zr⁴⁺.³³ However, substituting the Zr sites in LLZ with As⁵⁺ is difficult.³⁴ In contrast, doping LLZ systems with Sb is a relatively simpler approach, as evidenced by several studies conducted in this regard; for instance, as a separator for the Li metal anode, the Sr–Sb-doped LLZ sintered at 1100 °C has performed stably over 100 charge/discharge cycles.³⁵ These findings strongly suggest that Sb-doped LLZ is stable against Li metal. Furthermore, Ca²⁺ doping into La sites can help reduce

the melting point of oxide glass as a modified metal as well as preserve the Li carrier concentration.²⁸

Therefore, the present study was aimed at lowering the sintering temperature of Ca- and Sb-doped LLZ (denoted as LLZ–CaSb) using a Li–Sb–O-type oxide and Li₃BO₃ as sintering aids. Incorporating a large amount of the sintering aid can lead to its accumulation in grain boundaries and hindered ion conduction; therefore, adding a small amount of the sintering aid is vital for increasing the contact area between Li–Sb–O and LLZ–CaSb particles to induce dissolution and precipitation.^{28,29} Consequently, the Li–Sb–O sintering additive was integrated with the LLZ–CaSb particles. Furthermore, the effects of introducing the Li–Sb–O additive on the sinterability, electrical properties, and stability against Li metal were explored. Additionally, a co-sintered bulk-type ASSB featuring the new LLZ powder was constructed.

Experimental

Synthesis of LLZ–CaSb and Li–Sb–O composite particles

The Li_{6.5}(La_{2.925}Ca_{0.075})(Zr_{1.425}Sb_{0.575})O₁₂ + xLi₇SbO₆ (x = 0, 0.04, 0.06, 0.08, 0.12, and 0.14) composite was synthesised *via* a solid-state reaction. LiOH(H₂O) (>99.95%) sourced from Sigma-Aldrich (Germany) and La(OH)₃ (>99.99%), ZrO₂ (>98%), Ca(OH)₂ (>99.9%), and Sb₂O₅ (>99.99%) obtained from Kojundo Chemical Laboratory (Japan) were used as starting materials. To compensate for Li loss, LiOH(H₂O) was added in excess (6 mol%). The aforementioned materials were ball milled in 2-propanol using 2 mm-diameter zirconia balls. The mixture obtained after solvent evaporation was calcined in an alumina crucible at 950 °C for 12 h in dry air. The resulting powder was further pulverised in 2-propanol in a uniaxial ball mill with zirconia balls.

Electrolyte preparation

LLZ–CaSb was mixed with Li₃BO₃ in a 4.75 : 1 ratio (wt%), with Li₃BO₃ synthesised according to a previously reported method.²⁸ After adequate blending, the mixed powder was pressed into pellets under a pressure of 98 MPa. The pellets were sintered at 750 °C for 24 h in dry air and then placed on Au sheets to prevent Al contamination from the alumina setter.

Material characterisation

Phase identification of the synthesised powders was performed by powder X-ray diffractometry (XRD) using a SmartLab device (Rigaku, Japan) with Cu K α radiation.

Microstructural analysis was conducted by field-emission scanning electron microscopy (FE-SEM; JEM-6510, JEOL, Japan), and the elemental distribution was analysed by energy-dispersive X-ray spectroscopy (EDS; QUANTAX Flat QUAD System Xflash 5060FQ, Bruker, Germany). The samples were polished *via* ion milling using an IB-19520CCP instrument (JEOL, Japan). Ultrafine microstructural analysis was conducted by scanning transmission electron microscopy (STEM; JEM-ARM200F, JEOL, Japan). Thin samples were prepared by focused ion beam treatment using a nanoDUET NB5000 system



(Hitachi High-Tech, Japan). Elements were detected by EDS and electron energy loss spectroscopy (EELS; GIF Quantum-ER, Gatan, Japan).

Relative densities were calculated from the size and weight of the pellets and the theoretical densities of the powders; the latter were determined using a gas displacement pycnometer system (AccuPyc II 1340, SHIMADZU, Japan).

To investigate the sintering behaviour of the composite powder pellets, thermomechanical analysis (TMA) was performed using a TMA-60 instrument (SHIMADZU, Japan). To that end, samples were heated to the target temperature at a constant heating rate of $10\text{ }^{\circ}\text{C min}^{-1}$ in dry air. A constant load of 5 g was applied to ensure that the detector adapted to the pellet shrinkage.

Electrical characterisation

The Li-ion conductivity of the prepared pellets was determined using Li-ion-blocking Au-sputtered electrodes with a two-probe AC impedance measurement system (4990EDMS; Keysight Technologies, USA) in the frequency range of 20 Hz to 100 MHz at temperatures ranging from 240 to 320 K. Prior to Au sputtering, the surface of each sample was adequately polished with emery paper to remove impurities, and the sample thickness was set to 0.8–1.0 mm.

Cyclic voltammetry (CV) profiles of Au|LLZ-CaSb + $x\text{Li}_7\text{SbO}_6$ + Li_3BO_3 |Li cells were obtained at a scanning rate of 1 mV s^{-1} in the range of -0.5 to 5 V (vs. Li^+/Li) using a potentiostat (1287A; Solartron Analytical, U.K.). A $1\text{ }\mu\text{m}$ -thick Li electrode was fabricated by vacuum thermal evaporation after polishing.

Symmetric Li|Au|LLZ-CaSb + $x\text{Li}_7\text{SbO}_6$ + Li_3BO_3 |Au|Li cells were fabricated for Li plating/stripping tests to be performed for 150 cycles at a current density of 0.05 mA cm^{-2} , with each polarisation step conducted for 30 min. The current density was increased in a stepwise manner from 0.1 to 0.57 mA cm^{-2} to investigate the critical current density. In this case, Li electrodes were formed by heating $\varnothing 8\text{ mm}$ discs of Li foil obtained using a hole punch at $185\text{ }^{\circ}\text{C}$ in a glove box. The sputtered Au layer was added to improve the wettability with Li metal.

Bulk-type all-solid-state battery performance

To ascertain the viability of co-sintering the cathode material and garnet electrolyte, the electrochemical cycling attributes and rate characteristics of the bulk-type cell were evaluated using a half cell fabricated *via* co-sintering. LiCoO_2 powder (Sumitomo Metal Mining, Japan), which was used in our previous studies,^{28,29} LLZ-CaSb powder ($x = 0.08$), and Li_3BO_3 were mixed at a volume ratio of 45:45:10 to prepare the cathode. LLZ-CaSb ($x = 0.08$) and Li_3BO_3 were mixed at a volume ratio of 90:10 to fabricate the separator. A mixed powder comprising 2.5 mg of the cathode material (final loading amount after sintering = 6.50 mg cm^{-2}) and 300 mg of the separator was filled in a $\varnothing 8\text{ mm}$ mould and compacted at 98 MPa . Subsequently, co-sintering was performed at $750\text{ }^{\circ}\text{C}$ for 24 h in dry air. Thereafter, Au was sputtered onto the cathode surface to form the current collector. Li foil was used as the anode, and an Ag-sputtered layer was inserted to enhance

contact between LLZ and Li metal. The Li electrode was formed by heating at $185\text{ }^{\circ}\text{C}$. Battery tests were performed using a potentiostat (1287A; Solartron Analytical, U.K.). Cycling tests were performed at 0.1 C for 60 cycles at $25\text{ }^{\circ}\text{C}$. The high and low cutoff voltages were set to 4.2 and 3.0 V , respectively. Rate performance tests were conducted at 0.1 C , 0.2 C , and 0.3 C for five cycles at each rate. Additionally, the dependence of the battery performance on the operating temperature was assessed at -25 , -1 , 70 , and $120\text{ }^{\circ}\text{C}$ for five cycles each at 0.1 C . AC electrochemical impedance spectroscopy was performed at frequencies ranging from 10^6 Hz to 0.1 Hz using a frequency response analyser (1260A; Solartron Analytical, U.K.).

Results and discussion

Properties of synthesised LLZ-CaSb + $x\text{Li}_7\text{SbO}_6$ powders

Powder XRD patterns of the LLZ-CaSb + $x\text{Li}_7\text{SbO}_6$ ($x = 0$ – 0.14) system were acquired after calcination at $950\text{ }^{\circ}\text{C}$ (Fig. 1). The cubic garnet phase with the space group $Ia\bar{3}d$ appeared predominantly in all samples, in accordance with ICSD entry 182 312. Moreover, the intensity of the Li_7SbO_6 -associated diffraction peak – conforming to ICSD entry 413 370 – increased with increasing x , especially at x values greater than 0.04 (see Fig. S1†). STEM-EDS images of the LLZ-CaSb + $0.08\text{Li}_7\text{SbO}_6$ particles (Fig. 2) were analysed because they exhibited the highest sinterability among the samples, as described later. Two regions with different contrasts were observed in the LLZ-CaSb particles. Notably, the elemental maps (Fig. 2c–f and k–n) indicated the absence of La and Zr and the presence of Sb and Li in this region. The acquired electron diffraction patterns (Fig. 2g, h, and o; corresponding to Points A, B, and C in Fig. 2b and i) were assigned to monoclinic Li_3SbO_4 ($P2/c(13)$), monoclinic Li_5SbO_5 ($C2/m(12)$), and trigonal Li_7SbO_6 ($R\bar{3}(146)$), respectively. Although the XRD results indicated that Li_7SbO_6 was present largely as a secondary phase, STEM analysis revealed a small number of different Li–Sb–O-type oxides.

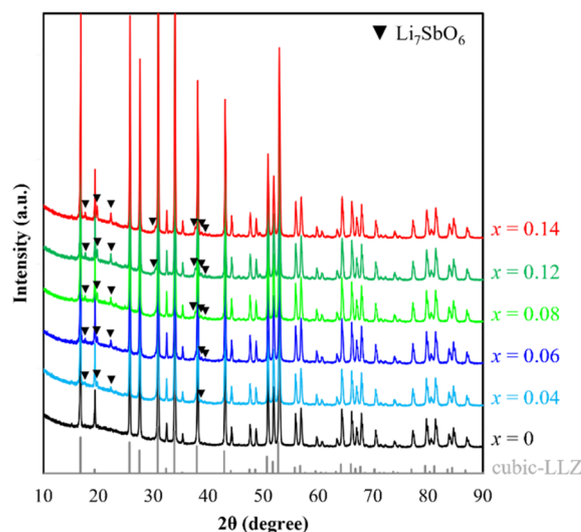


Fig. 1 XRD patterns of the synthesised LLZ-CaSb + $x\text{Li}_7\text{SbO}_6$ powders.



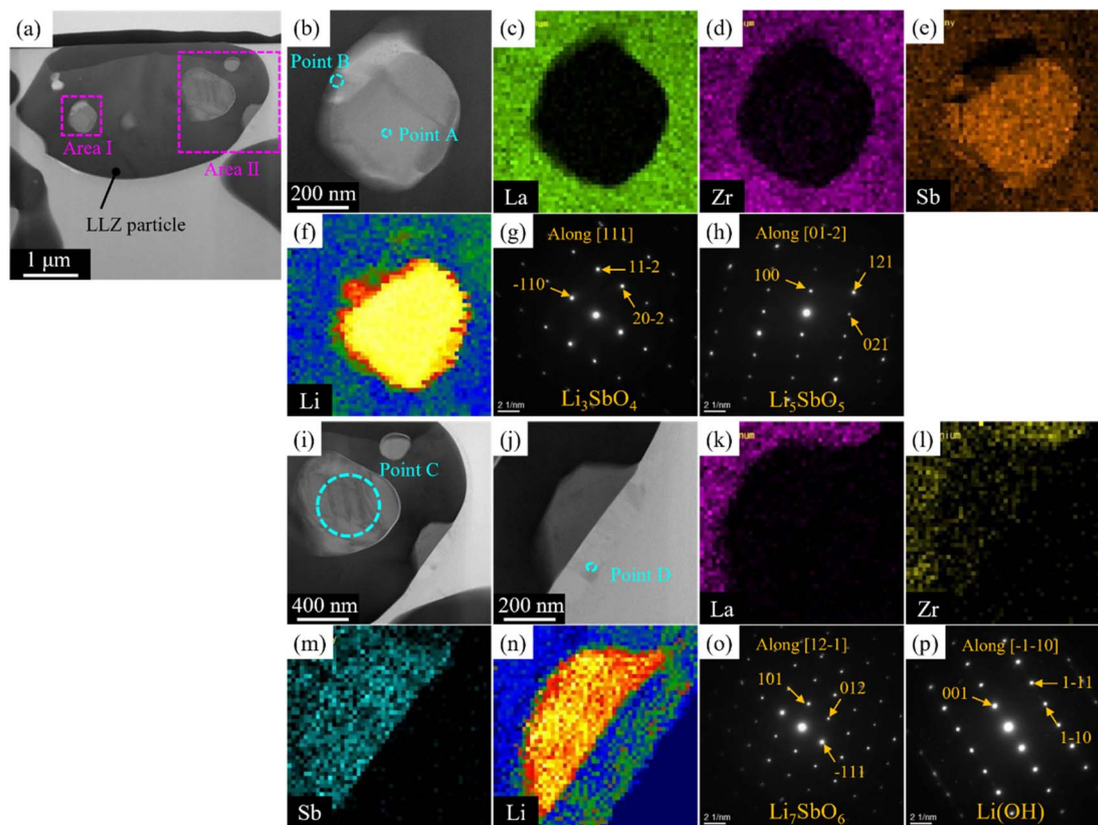


Fig. 2 (a) Bright-field (BF)-STEM image of an LLZ-CaSb + 0.08Li₇SbO₆ particle. (b) BF image and (c)–(f) elemental maps of Area I in (a) acquired through EDS and EELS. Electron diffraction patterns of (g) Point A and (h) Point B, marked in (b). (i and j) BF images and (k)–(n) elemental maps of area II in (a) obtained *via* EDS and EELS. Electron diffraction patterns of (o) Point C and (p) Point D, indicated in (i) and (j).

Moreover, the surface layer of the particles contained Li (Fig. 2j and n). The acquired electron diffraction pattern of the surface layer (Fig. 2p) was assigned to Li(OH) (tetragonal, $P4/nmm(129)$). In our previous study, this surface layer could not be confirmed through similar STEM observations of Ca–Bi-doped LLZ with Li–Bi–O sintering aids,²⁹ implying that the surface Li(OH) layer was probably created *via* excessive Li addition. Notably, Li₇SbO₆ as well as Li₃SbO₄ and Li₅SbO₅, which were produced through the decomposition of Li₇SbO₆, was detected in the present study through STEM analysis of the Ca–Sb-doped LLZ and Li–Sb–O sintering aids. The decomposition of Li₇SbO₆ was initiated by the generation of Li₂O, which easily reacted with humid air to form Li(OH); this functioned as a low-melting-point material that promoted the sintering of LLZ.³⁶ Thus, sintering of the Ca–Sb-doped LLZ with Li–Sb–O sintering aids can be influenced by Li(OH).

Sintering properties and characterisation of the LLZ-CaSb + xLi₇SbO₆ + Li₃BO₃ composite electrolyte

The manner in which the relative density of the sintered electrolytes depended on the sintering temperature was analysed, and SEM images of the fracture surfaces of samples with $x = 0$ and 0.08 were acquired (Fig. 3). At $x = 0$, the relative density of the specimen was low and numerous micropores were present. The relative density increased with increasing x , reaching

a maximum value of 87% at $x = 0.08$. Notably, no large pores were present in the sample corresponding to $x = 0.08$, and an intergranular fracture was partially observed, suggesting the progress of sintering (Fig. 3c).

Cross-sectional SEM-EDS images acquired after ion milling (Fig. 4) indicated that the LLZ-CaSb particles contained two differently coloured liquid-phase materials. Oxygen and boron were partially present in certain areas (black zones in Fig. 4b),

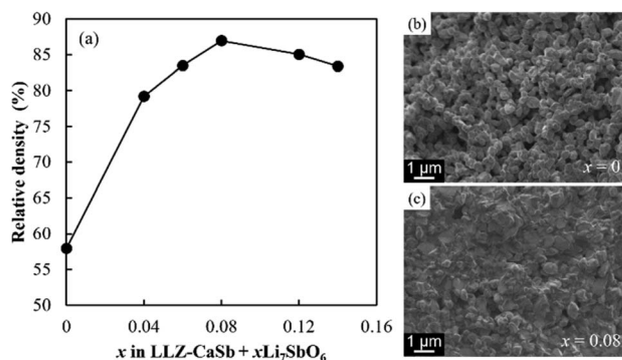


Fig. 3 (a) Dependence of relative density on the x value of the LLZ-CaSb + xLi₇SbO₆ + Li₃BO₃ composite electrolyte sintered at 750 °C. Cross-sectional SEM images of the electrolytes with x values of (b) 0 and (c) 0.08.



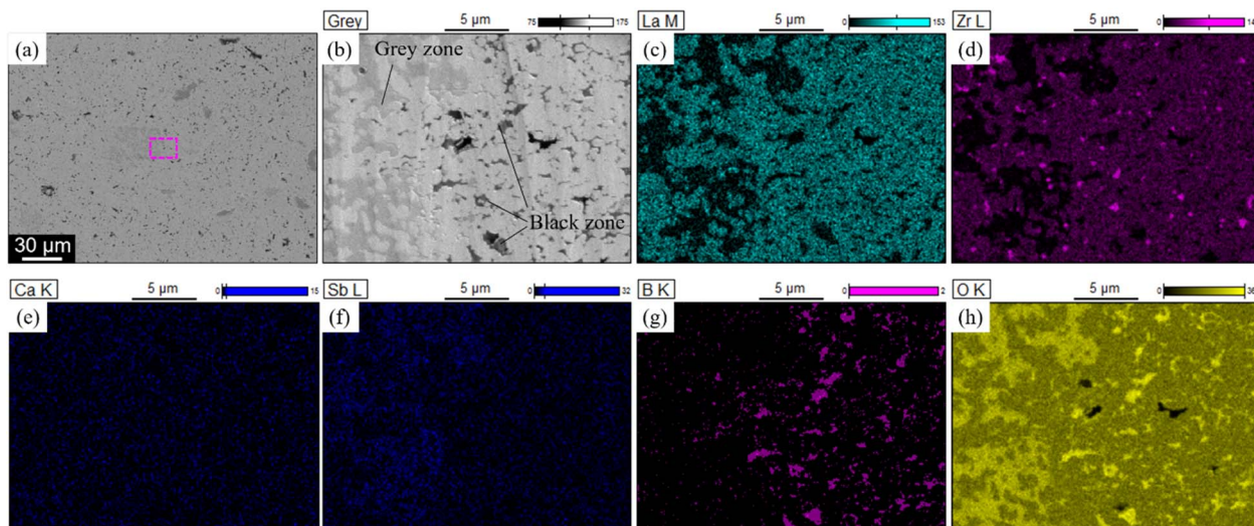


Fig. 4 (a) Low- and (b) high-magnification SEM images of the LLZ-CaSb + 0.08Li₇SbO₆ + Li₃BO₃ composite electrolyte sintered at 750 °C. (c)–(h) EDS maps of the region marked by the pink square in (a).

whereas oxygen and antimony were present in other regions (grey zones in Fig. 4b). Additionally, the Sb compound was spread around the LLZ-CaSb particles, whereas Zr-rich areas were present between the LLZ-CaSb particles. A similar imaging analysis of the electrolytes with $x = 0.04$ and $x = 0.14$ (Fig. S2[†]) suggested that the Sb-containing liquid-phase region broadened as x increased, aggregating over a wide area at $x = 0.14$. In all the investigated compositions, boron was absent in the areas where antimony was present, suggesting that the two liquid phases were immiscible. Therefore, Sb likely aggregated owing to its wettability with respect to the B-containing liquid phase. A more comprehensive analysis was performed by STEM-EDS imaging of the locations where the Sb-containing liquid phase was present between the LLZ-CaSb particles (Fig. 5). The acquired diffraction patterns indicated that the Sb-containing liquid-phase region and the Zr-rich zone comprised Li₇SbO₆ (triclinic, $P1(2)$) and Li₆Zr₂O₇ (monoclinic, $C2/c(15)$),

respectively (Fig. 5d and e). STEM images were also obtained for the other parts where Sb was not detected between the LLZ-CaSb particles (Fig. 6). The images were expected to showcase the Li₃BO₃ added as a sintering aid; however, the diffraction patterns suggested that Li₄B₂O₅ (orthorhombic, $Pca21(29)$) and Li₃B₁₁O₁₈ (monoclinic, $P21/a(21)$) were present in the electrolyte (Fig. 6c and f). According to the Li₂O–B₂O₃ phase diagram reported by Rousse *et al.* (Fig. S3[†]),³⁷ Li₄B₂O₅ has a eutectic composition and melts at a temperature (650 °C) lower than that of Li₃BO₃.³⁸ Moreover, Li₃B₁₁O₁₈ partially melts at 635 °C to form LiB₃O₅ and B₂O₃ liquid phases. The self-generation of this low-melting-point compound can potentially permit low-temperature sintering. XRD patterns of the LLZ-CaSb + x Li₇SbO₆ + Li₃BO₃ composite electrolytes (Fig. S4[†]) indicated that LiB₃O₅ was present in all the investigated samples ($x = 0–0.14$), with Li₇SbO₆ and La₂Zr₂O₇ also detected at $x \geq 0.12$. LiB₃O₅ was formed when Li₃B₁₁O₁₈ decomposed and melted, as described above. (The detailed reaction scheme is presented later.) In essence, Li₃BO₃ transformed into a Li-lean composition.

TMA was subsequently performed to clarify the sintering behaviour of the electrolytes (Fig. 7). A Li₃BO₃-free sample was also evaluated for comparison. The results indicated that all samples started expanding at ~ 350 °C. This trend was not observed in our previous study on Ca–Bi-doped LLZ,^{28,29} therefore, this feature was evidently associated with the composition adopted in this study. Through DFT calculations, Gaultois *et al.* determined that Li₇SbO₆ and Li₅SbO₅ absorb CO₂ at specific temperatures at $P_{\text{CO}_2} = 1 \times 10^4$ Pa *via* the following routes:³⁹



Based on this information, the aforementioned volume expansion between 400 and 530 °C was attributed to the volume

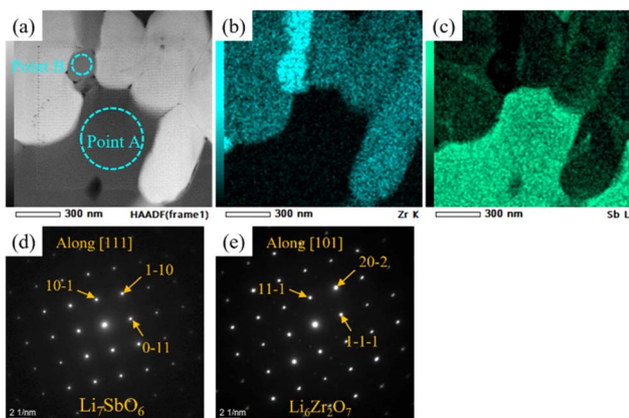


Fig. 5 (a) HAADF-STEM image and (b) and (c) elemental maps of the LLZ-CaSb + 0.08Li₇SbO₆ + Li₃BO₃ electrolyte sintered at 750 °C. Electron diffraction patterns of (d) Point A and (e) Point B, indicated in (a).



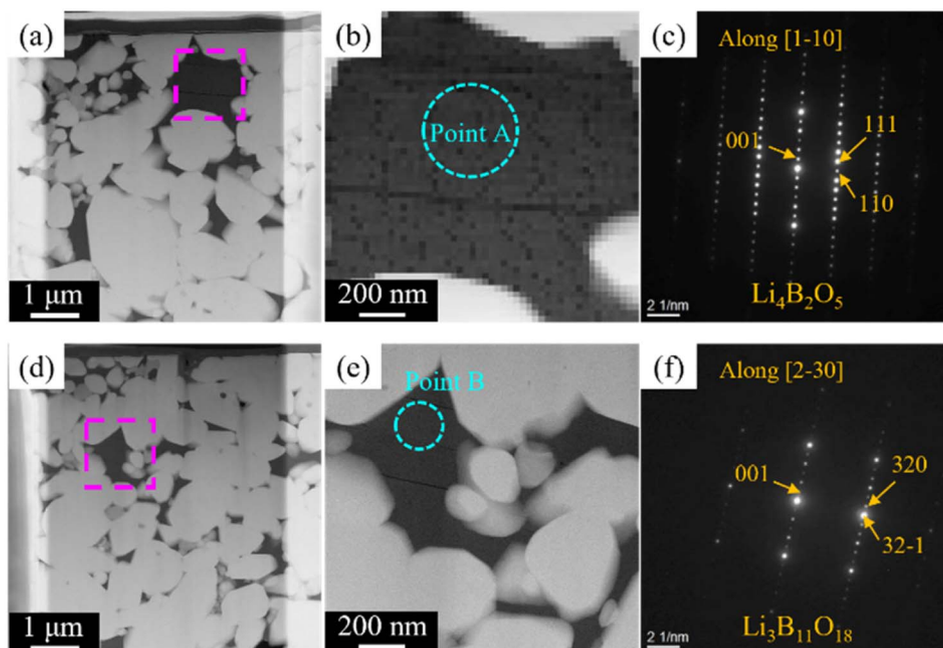


Fig. 6 (a, b and d, e) HAADF-STEM images of the LLZ-CaSb + 0.08Li₇SbO₆ + Li₃BO₃ electrolyte sintered at 750 °C. Electron diffraction patterns of (c) Point A and (f) Point B, marked in (b) and (e).

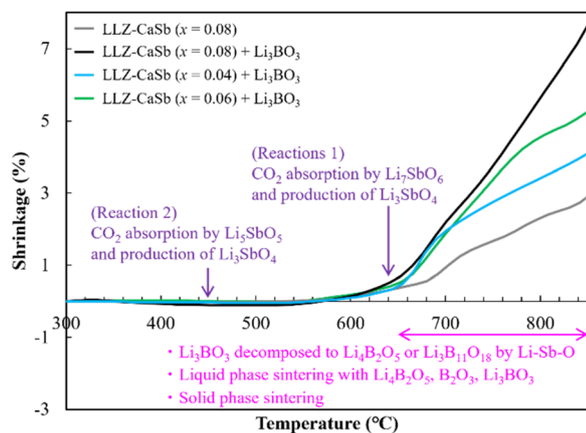
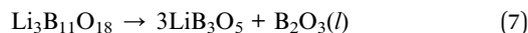
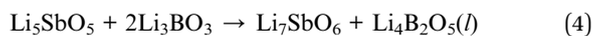
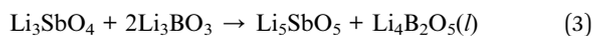


Fig. 7 Temperature-dependent linear shrinkage of LLZ-CaSb + x Li₇SbO₆ ($x = 0.04, 0.06, \text{ and } 0.08$) with and without the Li₃BO₃ composite pellets.

expansion induced by the CO₂ adsorption of Li₅SbO₅. However, all samples of LLZ-CaSb ($x = 0.04, 0.06, \text{ and } 0.08$) and Li₃BO₃ except LLZ-CaSb ($x = 0.08$) exhibited significant shrinkage at temperatures above ~650 °C. Therefore, based on the results shown in Fig. 5 and 6, reactions (3)–(7) transpired between Li₃BO₃ and Li₃SbO₄, and contraction occurred owing to the melting of Li₄B₂O₅ and Li₃B₁₁O₁₈. According to the Materials Project database, the formation energy of Li–B–O-type oxides (Li₃BO₃, Li₄B₂O₅, and Li₃B₁₁O₁₈) is lower than that of Li–Sb–O-type oxides (Li₇SbO₆, Li₅SbO₅, and Li₃SbO₄). Li₃B₁₁O₁₈ is the most stable compound among these oxides; therefore, Li₃BO₃ decomposes into Li₃B₁₁O₁₈.



Li₄B₂O₅, Li₃B₁₁O₁₈, LiB₃O₅, and B₂O₃ have B–O–B bonds in their structures, which can help extract Li₂O from LLZ.²² Consequently, Li₄B₂O₅, Li₃B₁₁O₁₈, LiB₃O₅, and B₂O₃ transform into Li₃BO₃, whereas LLZ forms an impurity phase of La₂Zr₂O₇ during sintering. The formation of La₂Zr₂O₇ was substantiated by the XRD patterns of the specimens with $x \geq 0.12$ (Fig. S4†). LiB₃O₅ exhibited its diffraction peak because it existed as a solid, given its melting point of 834 °C; therefore, solid-phase LiB₃O₅ showed poor reactivity and did not permit Li₂O extraction compared to the other compounds with B–O–B bonds. This was presumably due to the continuous occurrence of reactions (1)–(7) and the preferential reaction of Li₄B₂O₅, Li₃B₁₁O₁₈, and B₂O₃ with Li₂CO₃ produced continuously according to reactions (1) and (2). Additionally, one of the factors was the ability of the Li(OH) surface layer to prevent the direct reaction, as confirmed in Fig. 2. Reactions (3)–(6) suggest that for the reaction to proceed continuously, a sufficient amount of Li₃SbO₄ is required to extract Li₂O from Li₃BO₃. Notably, the slope of the shrinkage profiles at temperatures above 650 °C was relatively well maintained up to high temperatures as x increased (Fig. 7). For the specimens with $x = 0.04$ and 0.06 , the slope after the rapid contraction above 650 °C subsided was similar to that of



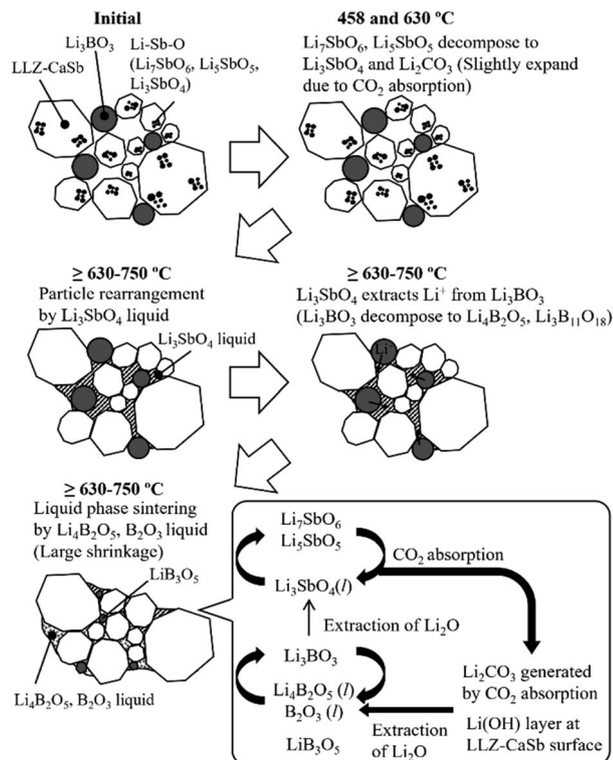


Fig. 8 Mechanism and scheme of reactions involving Li-Sb-O and Li_3BO_3 during sintering.

the Li_3BO_3 -free sample; therefore, the LLZ-CaSb particles shrank because of progress in the solid-phase sintering. However, for the sample with $x = 0.08$, the slope at temperatures above 650°C was constant. Therefore, at $x = 0.08$, the $\text{Li}_4\text{B}_2\text{O}_5$ and B_2O_3 liquid phases were continuously generated to promote reactions (3)–(7); however, at $x \leq 0.06$, the amount of Li_3SbO_4 was insufficient. These findings indicate that low-temperature sintering was achieved owing to the continuous occurrence of (i) the decomposition of the Li-Sb-O compounds by CO_2 absorption and (ii) the formation of the $\text{Li}_4\text{B}_2\text{O}_5$ and B_2O_3 liquid phases *via* the reaction between the decomposition product and Li_3BO_3 . As explained in Fig. S2† discussion, the Li-Sb-O and (Li)-B-O liquid phases were incompatible. Therefore, when $x \geq 0.12$, a large amount of the Li-Sb-O liquid phase inhibited the wetting of the (Li)-B-O liquid phase. Additionally, owing to the presence of a large amount of the $\text{Li}_4\text{B}_2\text{O}_5$ and B_2O_3 liquid phases at $x \geq 0.12$, the Li_2O derived from reactions (1) and (2) and the $\text{Li}(\text{OH})$ surface layer was insufficient, yielding $\text{La}_2\text{Zr}_2\text{O}_7$. Therefore, the sinterability diminished at $x \geq 0.12$. The sintering mechanism and scheme of the reactions occurring at $\sim 650^\circ\text{C}$ are presented in Fig. 8.

Electrical properties of the LLZ-CaSb + $x\text{Li}_7\text{SbO}_6$ + Li_3BO_3 composite electrolyte

Nyquist plots of the prepared electrolytes were acquired at 25°C (Fig. S5†). Two distinct semicircles were observed in the high- and intermediate-frequency regions, which represented the bulk resistance and grain boundary resistance, respectively. The

appearance of a tail in the low-frequency region suggested that the electrode blocked mobile Li ions. As x increased, both the bulk resistance and grain boundary resistance tended to decrease, and the magnitude of the grain boundary resistance was reduced. The lowered bulk resistance was due to the reduced tortuosity at a higher relative density (Fig. 3). Additionally, the progress in sintering contributed to the grain boundary resistance. Furthermore, sintering improved the interparticle contact and reduced the number of grain boundaries. A simple equivalent circuit expressed as $(R_e)(R_b//\text{CPE}_b)(R_{gb}//\text{CPE}_{gb})(\text{CPE}_{el})$ was used to fit the acquired data (solid profiles in Fig. S5†); here, R is the resistance; CPE is the constant phase element; and the subscripts b, gb, and el represent the contributions from the bulk, grain boundary, and electrode, respectively. The total Li-ion conductivity σ_{total} was calculated using R_e , R_b , R_{gb} , the electrode area S , and sample thickness t (Fig. 9a). Similar to the relative density, σ_{total} increased with increasing x and reached a maximum value ($3.1 \times 10^{-4} \text{ S cm}^{-1}$) at $x = 0.08$; this value is equivalent to that obtained previously by our group for the $\text{Li}_{6.5}\text{La}_{2.9}\text{Ca}_{0.1}\text{Zr}_{1.4}\text{Bi}_{0.6}\text{O}_{12}$ and Li_3BO_3 system at a sintering temperature of 750°C .²⁸ Arrhenius plots of the total Li-ion conductivity, bulk conductivity, and grain boundary conductivity were subsequently constructed (Fig. S6a–c,† respectively). The activation energies were calculated from the slopes of the lines, and the changes in bulk conductivity with x were analysed (Fig. 9b). The sample with $x = 0.08$ exhibited maximal bulk ionic conductivity and minimal activation energy E_{total} . This was caused more by the decrease in the grain boundary activation energy E_{gb} than by the decrease in the bulk activation energy E_{bulk} . At $x \leq 0.08$, the progress in sintering presumably enabled smooth ion transport at the grain boundaries. However, the E_{gb} values for the samples with $x \geq 0.12$ were large despite them

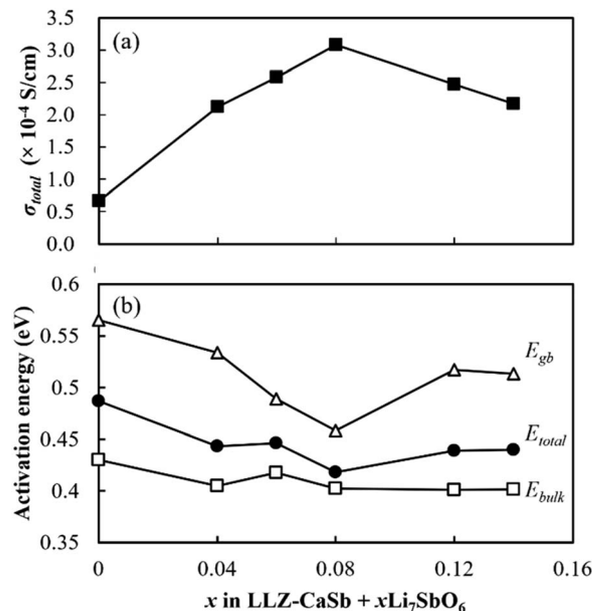


Fig. 9 Effects of x on the (a) total ionic conductivity at 25°C and (b) activation energy of the LLZ-CaSb + $x\text{Li}_7\text{SbO}_6$ + Li_3BO_3 composite electrolytes.



having the densities of the $x = 0.04$ and 0.06 specimens. Therefore, the grain boundary part of the electrolytes with $x = 0.08$ and 0.14 was subjected to STEM analysis to clarify this issue (Fig. 10). At $x = 0.08$, the grain boundary and bulk had the same contrast; however, at $x = 0.14$, a portion with a different contrast appeared at the grain boundary. This different-contrast area was poor in La and Zr but rich in Sb. The composition of the grain boundary differed significantly from that of LLZ at $x = 0.14$, which presumably increased the activation energy of the grain boundary in the region of the samples with $x \geq 0.12$. This compositional deviation was evidently caused by an increase in the Li-Sb-O liquid phase that entered between the LLZ particles, as illustrated in Fig. S7†

CV profiles were acquired to evaluate the electrochemical window (Fig. 11). The redox peaks appearing at ~ 0.2 V (vs. Li/Li⁺) were evidently due to the alloying of Li and Au and the stripping/plating of Li, as indicated by Yan *et al.*, who reported the alloying of Au and Li at 0.2 V.⁴⁰ Kundu *et al.* suggested that Li₇SbO₆ can undergo lithiation and de-lithiation at 0.8 V.⁴¹ However, the redox peak at ~ 0.8 V was not exhibited by the electrolyte reported herein. This was likely due to the presence of Li metal, which may have eliminated the need for using the Li from Li₇SbO₆, and the remarkably low volume of Li₇SbO₆ (~ 3.6 vol%), which prevented it from percolating, and consequently, functioning. Additionally, in the high-voltage range, no redox peaks were observed up to ~ 5.0 V. This tendency was also exhibited by other LLZ-CaSb + x Li₇SbO₆ ($x = 0.04, 0.06, 0.12,$ and 0.14) + Li₃BO₃ composites (Fig. S8†). Overall, these results suggest that the LLZ-CaSb + x Li₇SbO₆ + Li₃BO₃ electrolyte exhibited a wide electrochemical window.

Long-term Li plating/stripping tests conducted at 0.05 mA cm⁻² (Fig. 12) indicated that the plating/stripping could be performed for 150 cycles, and that the electrolyte was stable against Li metal. A comparison with previously reported Li plating/

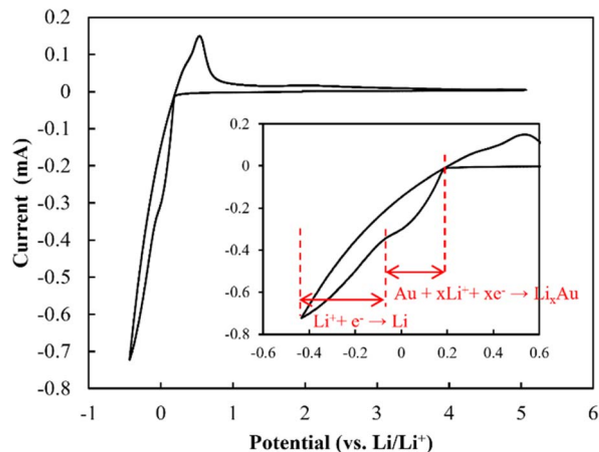


Fig. 11 Cyclic voltammetry profiles of the Au|LLZ-CaSb + 0.08Li₇SbO₆ + Li₃BO₃|Li cell at a scan rate of 1 mV s⁻¹.

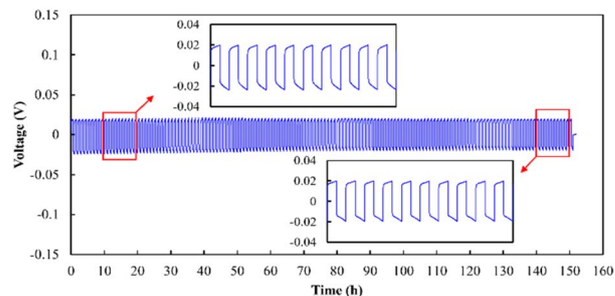


Fig. 12 Li plating/stripping performance of the Li|Au|LLZ-CaSb + 0.08Li₇SbO₆|Au|Li sample at a current density of 0.05 mA cm⁻².

stripping test results (Table S1†)^{25,42–48} suggested that the use of a low sintering temperature in the present study (750 °C) led to the same level of stability against Li plating/stripping as that of electrolytes sintered at higher temperatures. In addition to chemical stability, the wettability against Li metal is another crucial factor in achieving stable plating/stripping. Therefore, Li metal melting on pristine and Au-coated electrolyte specimens was analysed (Fig. S9†), and the results demonstrated the superior wettability of the Au-coated sample. This finding is consistent with previously reported alloying of Au with Li to improve wettability and reduce the interfacial resistance.^{44,46}

Characteristics of the co-sintered bulk-type all-solid-state battery

SEM-EDS images of the cathode layer were acquired, and cyclic charge-discharge tests were conducted at 25 °C (Fig. 13). The elemental maps showed a clear interface between LiCoO₂ and LLZ-CaSb, indicating that no impurity phases were formed. The O map suggested that the relative density was 93%, and the interfaces between LiCoO₂ and LLZ-CaSb particles were in close contact. Additionally, the initial discharge capacity was 120.7 mA h g⁻¹, and the cell could be charged and discharged for 60 cycles. Furthermore, the ASSB maintained a stable capacity over a long duration (Fig. 14a), with the discharge capacity retention rate

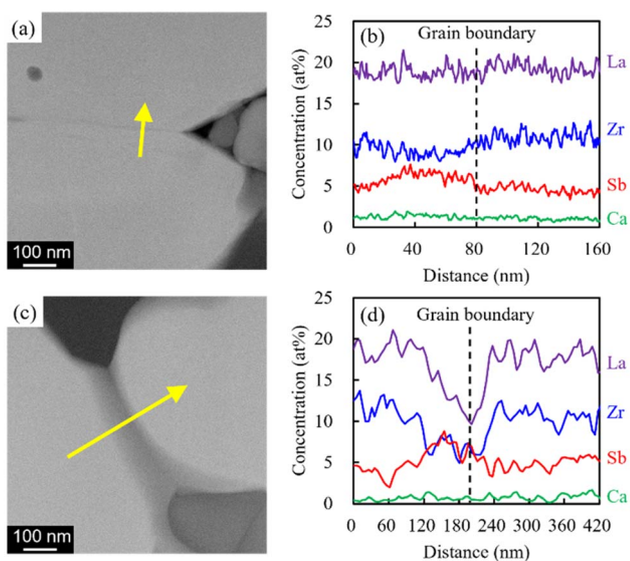


Fig. 10 HAADF images and changes in atomic concentration across the yellow line around the necking region for the electrolytes with (a) and (b) $x = 0.08$ and (c) and (d) $x = 0.14$.



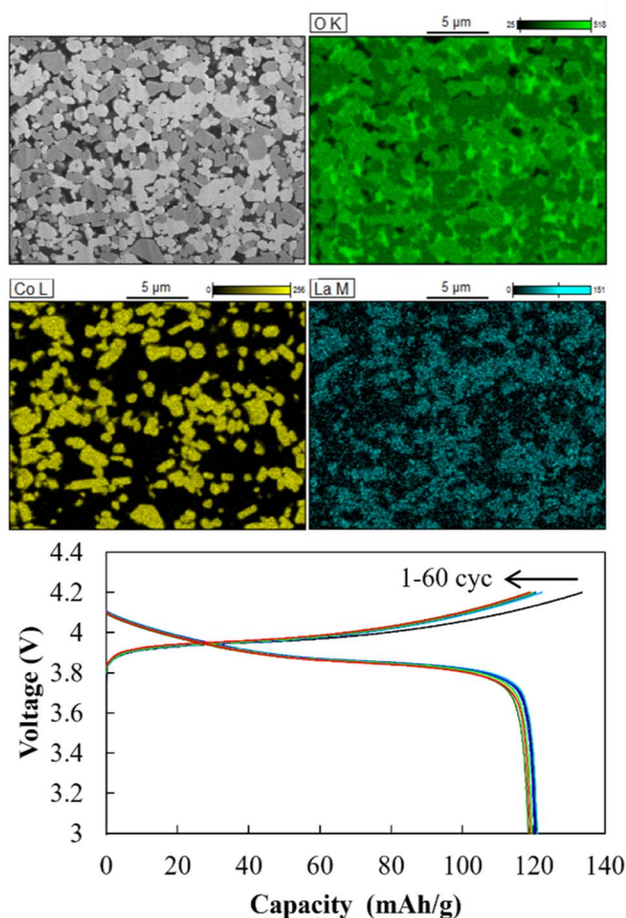


Fig. 13 Cross-sectional SEM-EDS images of the cathode layer in the $\text{Li}|\text{LLZ}-\text{CaSb} + 0.08\text{Li}_7\text{SbO}_6|\text{LiCoO}_2 + \text{LLZ}-\text{CaSb} + 0.08\text{Li}_7\text{SbO}_6 + \text{Li}_3\text{BO}_3$ ASSB, and charge–discharge curves acquired at 0.1C and 25 °C.

after 60 cycles being 98.6%. Table 1 lists the battery performance, manufacturing methods, and evaluation conditions employed in the present study and in previous investigations on the cycling characteristics of bulk-type ASSBs.^{14,15,28,49–51} Our previously reported system exhibited a noteworthy discharge capacity and cycling performance, with the battery comprising $\text{Li}_{6.5}\text{La}_{2.9}\text{Ca}_{0.1}\text{Zr}_{0.4}\text{Bi}_{0.6}\text{O}_{12}$ (LLZ-CaBi) and Li_3BO_3 exhibiting a retention rate of 92.8% after 40 cycles at 60 °C.²⁸ However, the electrolyte reported herein outperforms our previously reported system in terms of cycling behaviour; one of the reasons for this is the stability of $\text{LLZ}-\text{CaSb} + 0.08\text{Li}_7\text{SbO}_6$ against Li metal, which eliminates the need for a polyethylene-oxide (PEO)-based electrolyte in the buffer layer. Typically, PEO-based electrolytes are not stable against Li metal for prolonged durations, and their interfacial resistance increases.²⁸ To corroborate this result, the change in resistance after the first charge and after 60 charging cycles was monitored (Fig. 14b). The semicircular arc resistance that appeared at frequencies above 10^4 Hz increased after cycling. Nyquist plots were constructed after forming Ag electrodes on the electrolyte and then bonding Li metal to both sides of the same sample (Fig. S9†). In the profile of the sample with Li metal as the electrode, a semicircular arc was observed, which

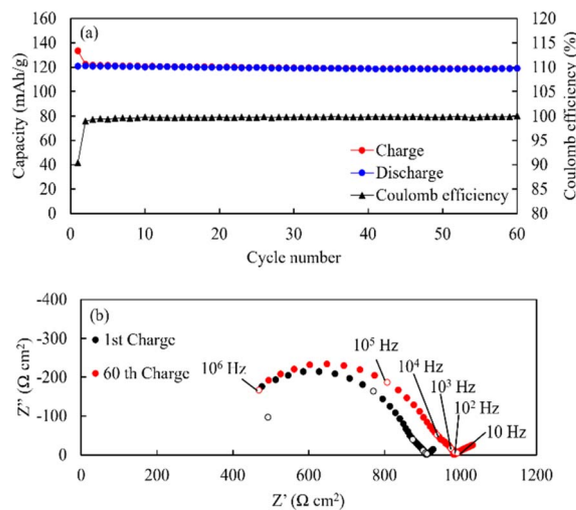


Fig. 14 (a) Long-term cycling behaviour, rate performance data, and (b) Nyquist plots of the $\text{Li}|\text{LLZ}-\text{CaSb} + 0.08\text{Li}_7\text{SbO}_6|\text{LiCoO}_2 + \text{LLZ}-\text{CaSb} + 0.08\text{Li}_7\text{SbO}_6 + \text{Li}_3\text{BO}_3$ ASSB obtained at 0.1C and 25 °C.

did not appear initially. Therefore, the resistance corresponding to frequencies $\geq 19\,952$ Hz was considered to be the grain boundary resistance of LLZ, and that corresponding to 6.3–19952 Hz was deemed the interfacial resistance between LLZ and Li metal. These results indicate that the increase in the grain boundary resistance of LLZ caused deterioration. This was presumably due to the change in the volume of LiCoO_2 during charging and discharging. Further research will help in shining more light on this matter.

Reducing the interfacial resistance between the cathode and LLZ is an important issue in oxide-based ASSB development, as reported by Kim *et al.*,⁵² who achieved a remarkably low resistance of $62\ \Omega\ \text{cm}^2$ by adapting a process to directly synthesise LiCoO_2 from infiltration metal salts in a porous LLZ scaffold. Zheng *et al.* reported that the interfacial resistance between the cathode and LLZ appeared at 1–150 Hz.⁵³ For the battery developed in the present study, the interfacial resistance after the first charge was $9.1\ \Omega\ \text{cm}^2$; this is the lowest value among similar previously reported systems. This implies that smooth ion transport can be realised at the interface, and a high-output battery can be developed. Therefore, rate performance was evaluated using batteries with the same configuration (Fig. S10†). The results indicated that charging and discharging could be repeated stably at 0.2C and 0.3C as well as at 0.1C. Furthermore, owing to its stability, the ASSB was expected to function at temperatures unconducive to traditional LIB operation. To substantiate this proposition, the charge–discharge properties were analysed at different operating temperatures (–25, –1, 70, and 120 °C; Fig. S11†). The results indicated that cyclic operation was possible at all the operating temperatures, although the capacity tended to decrease with decreasing operating temperature. To our knowledge, this is the first report on an ASSB with an LLZ electrolyte that can operate over a considerably wide temperature range, although several systems have been reported to operate at temperatures above 50 °C.^{15,18,54–58} This is because of the high battery resistance caused



Table 1 Manufacturing conditions, test environments, and performance data of previously reported bulk-type ASSBs

Reference	Cathode composition	Cathode sintering temperature (°C)	Sintering technique	Voltage range (V)	Rate (C)	Operating temperature (°C)	Initial discharge capacity (mAh g ⁻¹)	Number of cycles	Capacity retention rate
Han <i>et al.</i> (2018) ¹⁹	LiCoO ₂ + Li _{2.3} C _{0.7} B _{0.3} O ₃ + LLZ	700	Conventional sintering	3.0–4.05	0.05	25	94	100	88.3%
Wang <i>et al.</i> (2019) ¹⁴	LiNi _{0.6} Mn _{0.2} Co _{0.2} + Li ₃ BO ₃ + LLZ-Ta	700	Conventional sintering	3.0–4.2	0.05	25	106	30	70%
Feng <i>et al.</i> (2020) ⁵⁰	LiCoO ₂ + Li _{2.985} B _{0.05} OCl + LLZ	400	Hot pressing	3.0–4.1	0.05	25	71.3	20	70.1%
Guo <i>et al.</i> (2021) ¹⁵	LiCoO ₂ + γ-Al ₂ O ₃ + LLZ-Nb	850	Conventional sintering	2.8–4.15	0.2	100	93.6	16	92%
Akimoto <i>et al.</i> (2023) ⁵¹	LiNi _{0.6} Mn _{0.2} Co _{0.2} + LLZ-Ta	550	Hot pressing	2.5–4.2	0.01	25	125	26	85.6%
Hayashi <i>et al.</i> (2023) ²⁸	LiCoO ₂ + Li ₃ BO ₃ + LLZ-CaBi	750	Conventional sintering	3.0–4.2	0.1	60	100	40	92.8%
Present study	LiCoO ₂ + Li ₃ BO ₃ + Li ₇ SbO ₆ + LLZ-CaSb	750	Conventional sintering	3.0–4.2	0.1	25	120.7	60	98.6%

by the insufficient density of the cathode layer or the use of a large amount of a low-ionic-conductivity sintering aid. The electrolyte developed in this study enables densification of the composite cathode and exhibits high ionic conductivity even when sintered at low temperatures. Therefore, the LLZ reported herein is a stepping stone for the realisation of robust batteries that can operate in harsh environments.

Conclusions

The Li_{6.5}(La_{2.925}Ca_{0.075})(Zr_{1.425}Sb_{0.575})O₁₂ + xLi₇SbO₆ + Li₃BO₃ composite electrolyte sintered at a low temperature was found to be suitable for achieving a high relative density, high lithium-ion conductivity, and wide electrochemical potential. A relative density of 87% and conductivity of 3.1 × 10⁻⁴ S cm⁻¹ @ 25 °C were achieved as optimal values by the system sintered at 750 °C. Low-temperature sintering was realised by liquid-phase sintering with low-melting-point (Li)–B–O liquid phases (Li₄B₂O₅ and B₂O₃) produced by the reaction between Li₃SbO₄ and Li₃BO₃, with the former generated when Li₇SbO₆ absorbed CO₂. The (Li)–B–O liquid phase extracted Li₂O from LLZ-CaSb to form an impurity phase; this was subsequently impaired by the Li₂CO₃ generated when Li₇SbO₆ absorbed CO₂ and the Li(OH) layer produced on the LLZ-CaSb particles during the synthesis. When excess Li₇SbO₆ was added, the aggregated Li₇SbO₆ liquid phase entered between the LLZ-CaSb particles and obstructed the Li-ion motion, thereby increasing the grain boundary activation energy. Therefore, optimal performance was achieved at x = 0.08. Additionally, the electrolyte operated over a wide electrochemical window (>5 V vs. Li/Li⁺), allowing direct contact with Li metal. Therefore, the developed electrolyte can be used as a cathode layer, separator, and anode layer. The bulk-type ASSB fabricated by co-sintering exhibited outstanding charge-discharge performance. The new material reported herein shows considerable promise for use in ASSBs. Moreover, the new concept described herein can be a powerful tool for reducing the sintering temperature, because low-melting-point materials that react with LLZ and transform into high-melting-point materials can be used for liquid-phase sintering.

Author contributions

Naohiro Hayashi: conceptualisation; investigation; writing – original draft preparation. Ken Watanabe: formal analysis; supervision; writing – review & editing. Kengo Shimano: formal analysis; writing – review & editing.

Conflicts of interest

There are no conflicts to declare.

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