





Doctoral Thesis

Interfacial Evolution and Engineering on LiMO₂ (M = Co, Ni, Mn) Positive Electrodes for All-Solid-State Li-Ion Batteries Using Sulfide Solid Electrolytes

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Approved by Advisor

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Abstract

Most inorganic solid electrolytes (SEs) suffer from narrow intrinsic electrochemical windows and incompatibility with electrode materials, which results in the below par electrochemical performances of all-solid-state Li-ion or Li batteries (ASLBs). Unfortunately, in-depth understanding on the interfacial evolution and interfacial engineering via scalable protocols for ASLBs to mitigate these issues are at an infancy stage.

In this dissertation, rationally designed Li₃BO₃–Li₂CO₃ (or Li_{3-x}B_{1-x}C_xO₃ (LBCO)) coatings for LiCoO₂ (LCO) in ASLBs employing sulfide SE of Li₆PS₅Cl (LPSCl) were reported. The new aqueous-solution-based Li₃BO₃ (LBO) coating protocol allows us to convert the surface impurity on LCO, Li₂CO₃ into highly Li⁺-conductive LBCO layers (6.0×10^{-7} S cm⁻¹ at 30 °C for LBCO vs. 1.4×10^{-9} S cm⁻¹ at 100 °C for Li₂CO₃ or 1.4×10^{-9} S cm⁻¹ at 30 °C for LBO), which also offer interfacial stability with sulfide SE. By applying these high-surface-coverage LBCO coatings, significantly enhanced electrochemical performances are obtained in terms of capacity, rate capability, and durability. It is elucidated that the LBCO coatings suppress the evolution of detrimental mixed conducting interphases containing Co₃S₄ and effectively passivate the interfaces by the formation of alternative interface phases.





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Nomenclature

_

SE	Solid electrolyte
ASLB	All-solid-state lithium-ion battery
LBCO	Li ₃ BO ₃ -Li ₂ CO ₃ or Li _{3-x} B _{1-x} C _x O ₂
LCO	LiCoO ₂
LPSCI	Li ₆ PS ₅ Cl
LBO	Li ₃ BO ₃
LIB	Lithium-ion battery
LE	Liquid electrolyte
SEI	Solid electrolyte interphase
bcc	Body-centered cubic
LIPON	Li _{3.3} PO _{3.9} N _{0.17}
RT	Room temperature
LGPS	$Li_{10}GeP_2S_{12}$
LLZO	$Li_7La_3Zr_2O_{12}$
LLTO	Li _{3x} La _{2/3-x/3} TiO ₃
LATP	$Li_{1.5}Al_{0.5}Ge_{1.5}(PO_4)_3$
LAGP	$Li_{1.4}Al_{0.4}Ti_{1.6}(PO_4)_3$
LISICON	Lithium super ionic conductor
a-LBCO	artificial-LBCO
c-bare	Cleaned bare
PEEK	Polyaryletheretherketone
GITT	Galvanostatic intermittent titration technique
XRD	X-ray diffraction
FESEM	Field emission scanning electron microscopy
BSE	Backscattered scanning electron
HRTEM	High-resolution transmission electron microscopy
STEM	Scanning transmission electron microscopy
EELS	Electron energy loss spectroscopy
XPS	X-ray photoelectron spectroscopy
TGA	Thermogravimetric analysis
LEIS	Low-energy ion scattering
SAED	Selected area diffraction
EIS	Electrochemical impedance spectroscopy
MCI	Mixed conducting interphase



1. Introduction

Over the past decades, rechargeable lithium-ion batteries (LIBs) have conquered the market of energy storage devices owing to their superior energy density to their competitors. However, harsh efforts to maximize the energy density of LIBs, such as the use of ultra-thin separators ($\leq 10 \mu$ m) and raising the upper limit of voltages, have brought unprecedented risks in safety, which originates from the use of flammable organic liquid electrolytes.¹⁻⁸ Moreover, the safety concerns of LIBs are imperative for emerging large-scale applications, such as battery-driven electric vehicles and grid-scale energy storage.^{9, 10} In this regard, solidifying electrolytes with nonflammable inorganic materials is one of the best solutions.¹⁰⁻¹⁶ Additionally, inorganic solid electrolytes (SEs) are considered enablers for next-generation electrode materials, such as Li metal and S (or Li₂S), which typically suffer from poor compatibility with conventional organic liquid electrolytes.^{10, 15-19}

Sulfide SE materials are some of the most promising candidates to realize high-performance all-solidstate batteries. Several state-of-the-art sulfide superionic conductors developed (e.g., $Li_{10}GeP_2S_{12}$,²⁰ $Li_{9.54}Si_{1.74}P_{1.44}S_{11.7}Cl_{0.3}$,¹¹ $Li_7P_3S_{11}$ ²¹) have shown impressively high ionic conductivities reaching the order of 10^{-2} S cm⁻¹ at room temperature with a single ionic transport nature, which implies the feasibility of all-solid-state batteries significantly outperforming conventional LIBs.^{22, 23} More importantly, sulfide materials are mechanically sinterable at room temperature and are thus beneficial for practical electrode fabrication.^{10, 24}

Recent theoretical studies showed that similar to organic liquid electrolytes for conventional LIBs, inorganic SE materials also have narrow intrinsic electrochemical windows, and that the passivation of SEs is necessary for the reversible operation of all-solid-state batteries.²⁵⁻²⁹ In particular, the adaptation of conventional $Li_{x}MO_{2}$ (M = Co, Ni, Mn) cathode materials to all-solid-state Li-ion or Li batteries (ASLBs) suffers from huge interfacial resistances, which could be attributed to multiple factors such as surface impurities on Li_xMO_2 ,³⁰ severe reactions between Li_xMO_2 and sulfide SEs,^{25, 29, 31} space charge layer effects,³² lattice mismatches,³³ and poor wetting of SEs.^{12, 13, 34} It is known that the formation of surface impurities, such as LiOH and Li₂CO₃ on Li_xMO₂ in ambient atmosphere conditions, causes the degradation of the electrochemical performances of conventional LIBs.³⁵⁻³⁷ When it comes to ASLBs, the poor ion-conducting properties of the surface impurities could be more problematic.^{30, 38} Since the first report in which it was demonstrated that interfacial engineering on LiCoO₂ using Li₄Ti₅O₁₂ could significantly lower the interfacial resistances in ASLBs,³² various protective coatings have been developed (Table 1) to date: $LiNbO_3$,^{11, 12, 39} Li_2SiO_3 ,³¹ Ta_2O_5 ,⁴⁰ Al_2O_3 ,⁴¹ and Li_3PO_4 .⁴² In most previous works regarding ASLBs using Li_xMO₂ and sulfide SEs, Li_xMO₂ coated with these materials were tested without placing a strong emphasis on the mechanistic details of the coatings.^{10-12,} $^{14,\,20,\,23,\,24,\,34} \quad \text{Moreover, to date, only a few in-depth and/or systematic studies on Li_xMO_2/SE interfacial}$ evolution/engineering have been reported.^{29, 31, 42-44} The general consensus from the previous reports



is that the interfacial resistance of ASLBs is inversely proportional to the Li⁺ conductivity of the coating materials.⁴² For example, using an amorphous Li_{3.5}Si_{0.5}P_{0.5}O₄ coating with a high Li⁺ conductivity of 1.6×10^{-6} S cm⁻¹ at room temperature resulted in a promising electrochemical performance of LiCoO₂/In ASLBs,⁴² though the high ionic conductivity of the coating material could be achieved only for its amorphous form, derived by a costly vacuum deposition process. LiNbO₃ is one of the most frequently practiced coating materials for sulfide ASLBs because of its high Li⁺ conductivity of ~10⁻⁶ S cm⁻¹ at room temperature and easy preparation protocol based on a wet method using alcohols (Table 1).^{10-12, 14, 20, 23, 24, 34, 39} However, Nb is not earth-abundant and the use of flammable alcohol in the coating process would be a concern when scaling up. While these findings on the correlation between the Li⁺ conductivity of the coating materials and the electrochemical performance aid in the design of alternative coating materials, it should be noted that the multiple aspects of not only Li⁺ conductivity, but also scalable preparation and cost-effectiveness should be carefully considered. Moreover, a detailed understanding on the evolution at electrode-SE interfaces affected by protective coatings is required. These aspects are imperative for the practical development of high-performance ASLBs.

From this background, Li₃BO₃ (LBO) has caught our attention. Despite its relatively low Li⁺ conductivity $(1.4 \times 10^{-9} \text{ S cm}^{-1} \text{ at } 30 \text{ °C}$, measured in this work), LBO has been investigated as a sintering aid for oxide SE materials, such as Li₇La₃Zr₂O₁₂, for oxide-based ASLBs, as it can help lower the sintering temperatures for the oxide SEs because of its low melting point (700 °C).45-50 However, until now, there has been no report on the application of LBO or LBO-derived materials for sulfide-based ASLBs. Herein, we report the development of rationally designed Li₃BO₃-Li₂CO₃ (LBO-LCO or $Li_{3,x}B_{1,x}C_xO_3$ (LBCO)) protective coatings prepared via a simple and scalable wet protocol using water, which drastically enhances the electrochemical performances of LiCoO₂ for ASLBs using sulfide SEs. The surface impurity on LiCoO₂, Li₂CO₃, generally impedes Li⁺ transport at the interfaces, but, after the aforementioned wet-coating process for LBO, it is converted into highly Li⁺ conductive LBCO coating layers. Complementary analyses reveal that the as-derived highly conductive, thick, and high-surface-coverage LBCO coatings for LiCoO₂ effectively suppress the formation of detrimental Co₃S₄ phase and form good passivating layers comprised of phosphates, thus minimizing interfacial resistances. This is also supported by our thermodynamic computational results based on first principles calculations regarding various states of mixed phases. Compared with other coating materials, LBCO and its precursor are cost-effective and environmentally benign (Table 1). Moreover, the use of water as a solvent is a significant advantage which avoids the use of flammable solvents employed in typical coating procedures.



			Preparation method				
Material	σ_{RT} [S cm ⁻¹]	Crystallinity	Reported method	Availability of wet-method	Solvent	Price [USD/ton] ^c	Ref
Li ₄ Ti ₅ O ₁₂	-	-	Wet method	0	EtOH	2.5–2.8 k (TiO ₂) 3.5–3.6 k (Ti(OiPr) ₄)	32
LiNbO ₃	~10 ⁻⁶	Low	Wet method	0	EtOH	120–180 k (Nb ₂ O ₅) 1000–1500 k (Nb(OEt) ₅)	39
Ta ₂ O ₅	10 ⁻⁶		Solid-state reaction followed by wet-method	Δ	-	169–390 k (Ta ₂ O ₅)	40
LiTaO ₃	~10 ⁻⁶	Low			-	169–390 k (Ta ₂ O ₅)	51
Li ₅₆ Nb ₂₂ Ta ₂₂ oxide	4.2×10^{-6}	Low	Physical vapor deposition	Δ	-	169–390 k (Ta ₂ O ₅)	51
Li ₄ SiO ₄	1×10^{-8}	Low	Pulsed laser deposition	Difficult	-	3.2–4.0 k (SiO ₂)	42
Li ₄ GeO ₄	7×10^{-8}	Low	Pulsed laser deposition	Difficult	-	900-1200 k (GeO2)	42
Li _{3.5} Ge _{0.5} P _{0.5} O ₄	2×10^{-7}	Low	Pulsed laser deposition	Difficult	-	900–1200 k (GeO2)	42
Li ₃ PO ₄	5×10^{-7}	Low	Pulsed laser deposition	Difficult	-	0.7–0.9 k (H ₃ PO ₄)	42
$Li_{3.5}Si_{0.5}P_{0.5}O_4$	10^{-6}	Low	Pulsed laser deposition	Difficult	-	3.2–4.0 k (SiO ₂)	42
Al ₂ O ₃	-	Low	Atomic layer deposition	0	EtOH	0.4–0.6 k (Al ₂ O ₃)	41
Li ₂ O-ZrO ₂	~10 ^{-9a}	-	Wet method	0	Propanol	15–30 k (ZrO ₂) 100 k (Zr(OPr) ₄)	52
Li ₃ BO ₃	1×10^{-9} $(2 \times 10^{-8})^b$	High	Wet method	0	Water	0.6–0.8 k (H ₃ BO ₃)	This work
$Li_{3-x}B_{1-x}C_xO_3$	6×10^{-7} $(1.2 \times 10^{-6})^b$	High	Wet method	0	Water	0.6–0.8 k (H ₃ BO ₃)	This work

Table 1. Characteristics of various coating materials for LiCoO₂ for ASLBs.

^{*a*} Obtained using the sample prepared by solid-state synthesis. ^{*b*} Glass-ceramic prepared by solid-state synthesis. ^{*c*} From https://www.alibaba.com



2. Background

2.1. Principle of lithium-ion secondary batteries

Lithium-ion secondary batteries (LIBs) are energy storage devices that store electrical energy as chemical energy and convert chemical energy into electrical energy when needed.⁵³ LIBs are made up of four major components: cathode active material, anode cathode material, electrolyte and separator. The schematic illustration of the LIBs is shown in Figure 1.⁵⁴ The electrolyte and separator, which allows lithium to move and pass through, respectively, cannot move and pass through electrons. So, they can separate the lithium-ion pathway from the electron pathway between the two electrodes; Lithium-ion moves inside a cell and electrons move to an external wire to create electrical energy.⁵⁴ LIBs were first commercialized in 1991 using the cathode material, LiCoO₂ and anode material graphite. The working voltage difference of lithium-ion batteries between the cathode and the anode is superior to other batteries, resulting in much higher energy.^{53, 54} However, the electrolyte decomposition reaction is inevitable because the electrochemical window of the organic liquid electrolytes used is narrower than the high voltage difference between cathode and anode. Fortunately, electrolyte decomposed products can make solid electrolyte interphase (SEI), a layer that can pass through lithium ion but cannot pass through electrons, to prevent further electrolyte decomposition reactions when the battery is driven. This allows us to re-charge and reuse lithium-ion secondary batteries that produce high energy several times.

Recently, higher energy density and power density of LIBs are required as lithium secondary batteries become larger. Minimizing the thickness of the separation film, not the energy-producing material, can greatly help increase the energy densities, but this resulted in stability problems. Not only is LIBs storing high energy, but it also contains all three elements of combustion inside the battery, so stability problems cannot be eliminated fundamentally.¹⁻⁸





Figure 1. Schematic illustration of the first Li-ion battery (LiCoO₂/Li⁺ electrolyte/graphite). Copyright 2013, ACS publications



2.2. All-solid-state Lithium-Ion batteries

All-solid-state Lithium-ion Batteries (ASLBs) are the batteries that replace organic liquid electrolytes (LEs) in conventional lithium-ion batteries with solid electrolytes. LEs account for most of the aforementioned three combustion elements inside the battery. But, replacing them with solid electrolytes brings ultimate safety.¹⁰⁻¹⁶ In addition, ASLBs have lots of possibilities of showing up high energy density and power density. Because all components of battery including electrolytes are solid, cells can easily be stacked without any external pouch as there is no leakage concern.¹¹ In addition, the heat management system can be minimized. High energy materials such as Li and S have had difficulty in commercializing in the conventional LIBs can also solve safety concerns and dissolution problems respectively in ASLBs.^{10, 15-19} These differences have great potential for improving the energy density of ASLBs. The power densities are expected to be greatly improved. The SEs having higher Li⁺ ion conductivity than LEs have been developed even though the development of ASLBs is shorter than that of LIBs. The SEs also has a lithium transfer number of 1, so there is no concern about the movement of the bulky negative ion that exists in liquid electrolytes.¹⁰⁻¹⁶ Solid electrolytes have excellent thermal stability, enabling elevated temperature (>70 °C) that was not possible in conventional LIBs.¹¹ Thanks to these potentials, interest in ASLBs is now greatly increased and lots of research is underway.

2.2.1. Solid electrolytes

Typical solid electrolytes used in ASLBs are divided into oxide-based, sulfide-based and polymerbased SEs, and research regarding halide and hydride-based SEs also has been actively carried out recently. The requirements of good SEs include 1) high Li-ion conductivity, 2) wide electrochemical window (oxidation and reduction stability), 3) high deformability, 4) chemical stability, and 5) cheap processing cost. The performance characteristics of each SEs are shown in in Figure 2.⁵⁵

Li ion transport in solid electrolyte crystals is an important process of the overall charge transfer reaction of ASLBs.⁵⁶ Three factors, such as carrier type, diffusion path and diffusion type, control the Li ion transport mechanism within the solid electrolyte. Carrier types are closely related to defect chemistry. The diffusion path has a large correlation with the anion arrangement.⁵⁷ It has been proposed that an anion sublattice having a structure similar with body centered cubic (bcc) having a low energy barrier is more advantageous for Li-ion diffusion than another close-packed framework.⁵⁷ The Li ion diffusion type also affects the ionic conductivity and has three types; direct interstitial hopping, interstitial knock-off and direct vacancy hopping.⁵⁶

2.2.2. Bulk-type all-solid-state lithium-ion batteries



Thin-film ASLBs using Li_{3.3}PO_{3.9}N_{0.17} are well known as commercial batteries with excellent performance. However, the use of thin film ASLB is limited to small applications such as smart cards and microelectronics devices due to the expensive vacuum deposition process required for production. ⁵⁸⁻⁶⁰ Recently, ASLBs as not only portable electronic devices but also large-scale batteries are drawing attention. ^{61, 62} As shown in Figure 3, an important feature of bulk-type ASLB is the composite electrode structure including active materials, conductive materials and SEs Bulk-type ASLBs don't need to be produced by expensive vacuum deposition processes unsuitable for large applications. ¹⁴ In bulk-type ASLB, SE particles replace the conventional LE of the LIBs. Thick composite electrodes of ASLBs indicate that increased energy density can compete with conventional LIBs, but this requires high ion conductivity of 10⁻⁴-10⁻² S cm⁻¹ at room temperature (RT). The oxide-based and sulfide-based SEs have been extensively investigated as suitable SEs for bulk-type ASLBs. Especially, the most important advantage of sulfide SE over oxide SE is that the SE powders can be easily transformed to pellet form simply by cold pressing not contain any other heat treatment. ^{10, 14}





Figure 2. Performance of different solid electrolyte materials. Radar plots of the performance properties of oxide solid electrolytes (panel a), sulfide solid electrolytes (panel b), hydride solid electrolytes (panel c), halide solid electrolytes (panel d), thin-film electrolytes (panel e) and polymer solid electrolytes (panel f). ASR, area-specific resistance. Copyright 2017, Springer Nature





Figure 3. Schematic diagram of bulk-type all-solid-state batteries. Copyright 2018, Wiley-VCH



2.2.3. Interfacial issues for bulk-type all-solid-state lithium-ion batteries

A very wide electrochemical stability window (0.0–5.0 V) is required for the ideal SEs to combine lithium metal anodes with high voltage cathode material to activate the highest voltage output of ASLBs. Electrochemical window of SEs was generally obtained by applying cyclic voltammetry to Li/SE/metal semi-blocking cells. As a result of testing with this method, very wide electrochemical window from 0V to 5V was reported to most sulfide and oxide SEs.^{20, 63} However, the electrochemical performance of bulk-type ASLBs assembled with these SEs is much worse than conventional LIBs based on LEs, even though SEs have high ion conductivity that can be compared with liquid electrolytes.^{20, 31} Recent thermodynamic calculations indicate that SEs have very narrow electrochemical windows, unlike conventional experimental results (Table 2).^{25, 27}

High interfacial resistance is often insisted as a major limiting factor in the performance of ASLBs.⁶⁴ Although not yet fully understood, the origin of interfacial resistance is often due to physical interfacial contact,¹² formation of space charge layers,⁶⁵ and/or formation of mutual interphase layers due to chemical/electrochemical reactions between electrolytes and electrodes.^{27, 28, 32, 44, 50} Although various interface processing techniques such as dynamic pressing,¹² nanosizing,⁶⁶ co-sintering⁶⁷ and surface coating^{12, 32, 39} have attempted to engineer interfaces between electrode materials and SEs, the performance of ASLBs is still significantly lower than that of a conventional LIBs. Understanding and resolving the interfacial problems between electrode materials and SEs will be the key to exceeding the performance of the conventional LIBs.



	Reduction Potential (V)	Phase equilibria at the reduction potential	Oxidation potential (V)	Phase equilibria at the oxidation potential
Li ₂ S	-	Li ₂ S (stable at 0 V)	2.01	S
$Li_{10}GeP_2S_{12}$	1.71	P, Li ₄ GeS ₄ , Li ₂ S	2.14	Li ₃ PS ₄ , GeS ₂ , S
$Li_{3.25}Ge_{0.25}O_{0.75}S_4$	1.71	P, Li ₄ GeS ₄ , Li ₂ S	2.14	Li ₃ PS ₄ , GeS ₂ , S
Li_3PS_4	1.71	P, Li_2S ,	2.31	S, P_2S_5
Li_4GeS_4	1.62	Li ₂ S, Ge	2.14	GeS ₂ , S
$Li_7P_3S_{11}$	2.28	Li_3PS_4 , P_4S_9	2.31	S, P_2S_5
Li ₆ PS ₅ Cl	1.71	P, Li ₂ S, LiCl	2.01	Li ₃ PS ₄ , LiCl, S
$Li_7P_2S_8I$	1.71	P, Li ₂ S, LiI	2.31	LiI, S, P_2S_5
LIPON	0.68	Li ₃ P, LiPN ₂ , Li ₂ O	2.63	P ₃ N ₅ , Li ₄ P ₂ O ₇ , N ₂
LLZO	0.05	Zr ₃ O, LaO ₃ , Li ₂ O	2.91	Li ₂ O ₂ , La ₂ O ₃ , Li ₆ Zr ₂ O ₇
LLTO	1.75	Li ₄ Ti ₅ O ₁₂ , Li _{7/6} Ti _{11/6} O ₄ , La ₂ Ti ₂ O ₇	3.71	O ₂ , TiO ₂ , La ₂ Ti ₂ O ₇
LATP	2.17	P, LiTiPO5, AlPO4, Li3PO4	4.21	O ₂ , LiTi ₂ (PO ₄) ₃ , Li ₄ P ₂ O ₇ , AlPO ₄
LAGP	2.70	Ge, GeO ₂ , Li ₄ P ₂ O ₇	4.27	O ₂ , Ge ₅ O(PO ₄) ₆ , Li ₄ P ₂ O ₇ , AlPO ₄
LISICON	1.44	Zn, Li ₄ GeO ₄	3.39	Li ₂ ZnGeO ₄ , Li ₂ GeO ₃ , O ₂

Table 2. Electrochemical window and phase equilibria at the reduction and oxidation potentials of the solid electrolyte materials. Copyright 2015, ACS publications



3. Experimental

3.1. Preparation of materials

The LBO and LBCO powders were prepared by dissolving a stoichiometric amount of LiOH (99.995%, Alfa Aesar), H₃BO₃ (>99.5%, Sigma-Aldrich), and Li₂CO₃ (99.997%, Sigma-Aldrich) in deionized water. The water was evaporated under a vacuum at 60 °C using a rotary evaporator, followed by a heat treatment at 600 °C for 5 h in air. The cleaned bare LiCoO₂ powders (c-bare) were prepared by heat treatment at 600 °C for 10 h in air. The LBO- and LBCO-coated LiCoO₂ powders were prepared using an aqueous LBO solution. After the bare LiCoO₂ powders were added into the coating solution prepared by dissolving a stoichiometric amount of LiOH (99.995%, Alfa Aesar) and H₃BO₃ (>99.5%, Sigma-Aldrich) in deionized water, the solvent was evaporated under a vacuum at 60 °C using a rotary evaporator, followed by a heat treatment at 600 °C for 10 h in air. To obtain the LBO- and LBCO-coated LiCoO₂ powders, c-bare and bare LiCoO₂ powders were used, respectively. For the LBCO-coated LiCoO₂ powders, the surface impurity, Li_2CO_3 , serves as the source for the coating materials. In contrast, the artificial-LBCO-coated (a-LBCO) LiCoO₂ powders were prepared using c-bare LiCoO₂ and a coating solution, prepared by dissolving LiOH, H₃BO₃, and Li₂CO₃ (99.997%, Sigma-Aldrich) in deionized water. The LPSCl SE powders were prepared by ball milling a stoichiometric mixture of Li₂S (99.9%, Alfa Aesar), P₂S₅ (99%, Sigma-Aldrich), and LiCl (99.99%, Sigma-Aldrich) at 600 rpm for 10 h with ZrO₂ balls.²³ Then, the ball-milled powders were heat-treated at 550 °C for 5 h in an Ar atmosphere.

3.2. Thermodynamic calculations

Potential reactions at the interfaces were considered as chemical reactions between two corresponding compositions at the interfaces.^{27, 28} Multi-dimensional compositional phase diagrams were constructed, and then pseudo-binary phase diagrams that have the two target compositions as end points were extracted from the multi-dimensional phase diagrams. The potential decomposition reactions were examined along the pseudo-binary phase diagrams with varying fractions of reactants. Most of the energy values used for constructing phase diagrams were obtained from the Materials Project database.⁶⁸ However, the energies of unstable target materials, such as layered Li_{0.5}CoO₂ and Li₆PS₅Cl, were corrected by making their decomposition energies become zero, as previously suggested.²⁵ Additionally, the energy of LBCO (Li_{3-x}B_{1-x}C_xO₃, x = 0.80) was evaluated as a linear combination of Li₃BO₃ and Li₂CO₃ because calculating the exact energy of the phase is computationally impossible. Despite these assumptions, we believe that the error of the calculated decomposition energy does not significantly affect the outcome of this study.



3.3. Materials characterization

The XRD measurements were conducted using a D8-Bruker Advance diffractometer under Cu K_a radiation (1.54056 Å). To avoid exposure to air, the samples were sealed with a Be window. The FESEM and BSE measurements were carried out using Quanta 200FEG (FEI). The accelerating voltage and emission current were fixed at 1 kV and 10.5 μ A, respectively. The HRTEM images and their corresponding selected-area electron diffraction (SAED) patterns and EELS spectra were obtained using JEM-2100 (JEOL) and JEM-2100F (JEOL). The XPS data were collected with a monochromatic Al K_a source (1486.6 eV) at 72 W, 12 kV, and 6 mA using an X-ray photoelectron spectrometer (ThermoFisher). For the ex-situ XPS measurements, the collected samples were loaded in an Ar-filled dry glove box and loaded into the XPS equipment shortly while minimizing exposure to air. The TGA measurements were conducted using 0500 (TA Instrument Corp.). The weight fraction of the coating materials was determined using inductively coupled plasma optical emission spectroscopy (ICP-OES, 720-ES, Varian). The LEIS measurements were carried out using Qtac100 (IONTOF GmbH).

3.4. Electrochemical characterization

For the measurement of Li⁺ conductivity, LBO and LBCO pellets were prepared by cold-pressing of the powders at 370 MPa and subsequent sintering at 600 °C for 10 h in air. The as-prepared pellets were subjected to measurements of Li⁺ conductivity by the AC impedance method (Iviumstat, IVIUM Technologies Corp.) using symmetric Li-ion blocking carbon-coated Al (c-Al)/pellet/c-Al cells. The LiCoO₂/Li-In all-solid-state cells were prepared as follows.^{12, 23} Partially lithiated indium (Li_{0.5}In, nominal composition) powders were prepared by mechanically milling a mixture of In (Sigma Aldrich, 99.99%) and Li (FMC Lithium corp.). After the SE layer was formed by pressing 150 mg of LPSCl powders, the electrode mixtures of $LiCoO_2$ and LPSCI (70:30 weight ratio) were spread on one side of the SE layer, followed by pressing. Then, the as-prepared $Li_{0.5}In$ powders were put on the other side of the SE layer. Finally, the whole assemblies were pressed at 370 MPa. The mass loading of LiCoO₂ was 8.3 mg cm⁻². All the pressing was carried out in a polyaryletheretherketone (PEEK) mold (diameter = 13 mm) with Ti rods as current collectors. All the electrochemical tests were conducted at 30 °C. The C-rate of 1C corresponds with 161 mA g^{-1} . The GITT measurements were carried out at a pulse current of 0.5C for 90 s and a rest for 2 h. The EIS measurements were performed from 1.5 MHz to 5 mHz with 10 mV of amplitude after discharging the cells to 3.9 V (vs. Li/Li⁺) at 0.2C at the second cycle.



4. Results and Discussion

4.1. Thermodynamic calculations of coating materials

In our screening process for potential coating materials, we first carried out computational investigations to examine the intrinsic interfacial stability between the cathode and SE material, along with the effects of applying coating materials on it. First, the electrochemical window and possible oxidized products of SE and coating materials were calculated in Table 3. Various possible reactions at the interfaces before and after introducing coating materials were probed by calculating the thermodynamic reaction energies, as illustrated in Figure 4. The blue dashed line in the Figure presents the interfacial reaction energy as a function of the atomic fraction of the SE materials (Li₆PS₅Cl (LPSCl)) surrounding the cathode (LiCoO₂), which models the various local compositional inhomogeneities in the composite electrode. These analyses reveal that the interface between the cathode and SE material is not intrinsically stable, but undergoes a spontaneous decomposition with negative reaction energy, which becomes maximum (-320 meV atom⁻¹) when LPSCl and LiCoO₂ react at a ratio of approximately 1:1. Moreover, the decomposition reaction is further promoted when the SE materials are in contact with the delithiated cathode ($Li_{0.5}CoO_2$), with a maximum energy of -450 meV atom⁻¹, as displayed by the red dashed line, indicating more serious side reactions during the charging of ASLBs. This decomposition reaction deteriorates the interface properties and often leads to an increase of cell impedance and the loss of active materials in the electrochemical reaction. However, we observed that the stability of the SE can be significantly enhanced when it is alternatively in contact with coating materials, such as LBCO or LBO. The solid lines show that the decomposition of electrolytes can be mitigated by coating layers with a substantially reduced reaction energy. Even though the decomposition reaction is still thermodynamically favorable, the driving force is reduced by less than one fifth. Moreover, the interfaces between LiCoO₂ and LBCO (or LBO) were found to be stable without decomposition or with negligible decomposition energies (Table 4), which indicates that the surface degradation of $LiCoO_2$ can be suppressed by incorporating LBCO (or LBO) as coating layers. As a result, the incorporation of LBCO (or LBO) as a coating layer is expected to suppress the decomposition reactions of both the cathode and the SE materials at the interface of them.



Table 3. Intrinsic electrochemical window and oxidation reaction of SE and coating materials

Sample	Electrochemical window [V vs. Li/Li ⁺]	Reaction at oxidation potential
LPSCl	1.72-2.14	$Li_6PS_5Cl \rightarrow Li_3PS_4 + 0.25LiS_4 + LiCl + 1.75Li$
LBO	0.28-3.47	$Li3BO3 \rightarrow 0.25Li6B4O9 + 0.345O2 + 1.5Li$
LBCO	1.27-3.47	$\begin{array}{c} Li_{2.2}B_{0.2}C_{0.8}O_3 \rightarrow 0.05Li_6B_4O_9 + 0.8Li_2CO_3 + 0.075O_2 \\ + 0.3Li \end{array}$





Figure 4. Calculated mutual decomposition energy of Li_6PS_5Cl with pristine and delithiated $LiCoO_2$, LBO (Li_3BO_3), and LBCO ($Li_{3-x}B_{1-x}C_xO_3$, x = 0.80) at various phase fractions of Li_6PS_5Cl in the mixed compounds.



Table 4. Calculated maximum mutual decomposition energy of the coating materials with pristine and delithiated LiCoO₂.

Sample	LiCoO ₂	Li _{0.5} CoO ₂
LBO	No reaction	$-4.4 \text{ meV} \text{ atom}^{-1}$
LBCO	No reaction	$-1.5 \text{ meV atom}^{-1}$



4.2. Characterization of coating materials and coated active materials

Inspired by the computational results, a series of LBCO-coated LiCoO₂ samples were prepared, along with the reference samples of $Li_{3-x}B_{1-x}C_xO_3$. The reference $Li_{3-x}B_{1-x}C_xO_3$ samples were obtained from a homogeneous aqueous solution containing LiOH, H₃BO₃, and Li₂CO₃. The phase-pure LBO samples (JCPDS no. 18-0718, Fig. S1a) exhibited a Li⁺ conductivity of 1.4×10^{-9} S cm⁻¹ at 30 °C (Figure 5, Table 5).⁴⁹ As Li₂CO₃ is added into LBO, the characteristic peaks for the isostructural phase with Li₂CO₃ (JCPDS no. 22-1141) evolved, as seen in the XRD patterns (Figure 6).^{49, 69} Correspondingly, Li⁺ conductivity was drastically increased to 6.0×10^{-7} S cm⁻¹ at x = 0.80 (Figure 5, Table 5), which is comparable to that of the state-of-the-art coating material for sulfide ASLBs: amorphous LiNbO₃ (Table 1).^{11, 12, 39} LBO-coated LiCoO₂ was fabricated using surface-cleaned LiCoO₂, referred to as "c-bare", which was obtained by a heat treatment at 600 °C in air, while the LBCO-coated LiCoO₂ was obtained using impurity-containing bare LiCoO₂, referred to as "bare". The characteristics of the LBO and LBCO coatings (weight fraction, thickness, and surface coverage) are provided in Table 6. Field emission scanning electron microscopy (FESEM) images of c-bare, LBO-coated (0.5 wt.%), and LBCO-coated (0.5 wt.% of LBO or 1.72 wt.% of LBCO) LiCoO₂ particles (Figure 7a-c, 8) showed no noticeable differences. However, the corresponding backscattered scanning electron (BSE) images reveal the inhomogeneous distribution of contrast in atomic numbers (Figur. 7a-c, 8), confirming the presence of the coating layers for LBO- and LBCO-coated LiCoO₂. Although a direct observation of the coating layers by high-resolution transmission electron microscopy (HRTEM) was hindered by the vulnerability of the low atomic-number constituents to electron beams, HRTEM images for LBO- and LBCO-coated LiCoO₂ particles showed lattice fringes corresponding with LBO ((020) plane) and LBCO ((-202) plane), as shown in Figure 9a, b respectively. Moreover, the presence of boron in the form of Li_3BO_3 on the surface of LBO- and LBCO-coated LiCoO₂ was corroborated by scanning TEM (STEM) images (Figure 10) and their corresponding electron energy loss spectroscopy (EELS) peaks at ~193 eV (Figure 9c).⁷⁰ In addition, compared with LBO-coated LiCoO₂, LBCO-coated LiCoO₂ exhibited a stronger carbon signature centered at ~292 eV.⁷¹

The presence of boron in coated LiCoO₂ was also confirmed by X-ray photoelectron spectroscopy (XPS) data for B 1s signals (Figure 11a). Both LBO- and LBCO-coated LiCoO₂ samples showed peaks at 191.5 eV corresponding to B^{3+} for Li₃BO₃.⁷⁰ The surface impurity on LiCoO₂, Li₂CO₃, was quantified by thermogravimetric analysis (TGA) in N₂. Whereas the c-bare sample showed no weight loss up to 850 °C, the bare sample started to lose weight at 700 °C, which is indicative of the thermal decomposition of Li₂CO₃ (Figure 11b).⁷² From the weight loss value, the amount of Li₂CO₃ on the surface of the bare sample was determined to be 1.1 wt.%. The thicknesses of the coating layers were estimated considering the surface area of LiCoO₂ powders obtained by N₂ adsorption-desorption isotherm measurements and are given in Table 6. Low-energy ion scattering (LEIS) measurements



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were carried out to analyze the conformality of the coating layers on LiCoO₂.⁷³ In LEIS, low-energy backscattered ions are analyzed, allowing the identification and quantification of the elements in the outermost atomic layer of a substrate.⁷⁴ Figure 11c shows the LEIS spectra for bare, c-bare, LBO-coated (0.5 wt.%), and LBCO-coated (0.5 wt.% of LBO) LiCoO₂ particles when using 5 keV Ne⁺ as incident ions. The strong peaks found at 1230 eV for the bare and c-bare samples correspond with the ions backscattered by Co in LiCoO₂. The lower intensity of the Co peak obtained for the bare sample compared with that obtained for the c-bare sample is due to surface impurities containing Li₂CO₃. Furthermore, the LBO- and LBCO-coated samples showed a much more attenuated Co-peak, indicating that Co atoms are well covered by the coating layers. Assuming that the surfaces of the c-bare sample are perfectly uncovered, the surface coverages of the other samples were determined by comparing the intensities of the Co peaks, and are shown in Table 6. For the bare sample, 21% of the surface is covered by impurities, such as Li₂CO₃. The surface coverages for LBO- and LBCO-coated samples turned out to be 79% and 87%, respectively. The higher surface coverage found for the LBCO-coated samples than for the LBO-coated one is attributed to the overall larger amount of coating materials.





Figure 5. Arrhenius plots of Li⁺ ion conductivities for Li_{3-x}B_{1-x}C_xO₃ (LBCO)



Table 5. Li⁺ ion conductivity at 30 °C and activation energy of LBCO.

x in Li _{3-x} B _{1-x} C _x O ₃	σ_{30} [S cm ⁻¹]	E _a [eV]
0.00	1.49×10^{-9}	0.60
0.20	1.18×10^{-7}	0.44
0.65	2.44×10^{-7}	0.41
0.80	6.05×10^{-7}	0.30





Figure 6. XRD patterns of Li_{3-x}B_{1-x}C_xO₃ (LBCO) and reference, Li₂CO₃ and Li₃BO₃



Comm1.	Sample $\frac{\text{wt.\% of the coatings}}{\text{Li}_3\text{BO}_3 \qquad \text{Li}_{3-x}\text{B}_{1-x}\text{C}_x\text{O}_3^a}$			Thickness of the	Relative surface coverage (%) ^c
Sample			- X III $L_{13-x}B_{1-x}C_xO_3$	coating (nm) ^b	
bare	0	-	-	-	21
c-bare	0	-	-	-	0
	0.05	0.06	-	1.0	-
LBO	0.1	0.15	-	2.5	-
	0.5	0.63	-	10.4	79
	0.1	1.24	0.10	21.5	-
LBCO	0.5	1.72	0.35	29.4	87
	1.0	2.18	0.50	37.0	-
a-LBCO	0.5	1.72	0.35	29.4	88

Table 6. Characteristics of LBO(-LCO) coatings for LiCoO₂

^a Obtained by ICP-OES, elemental analyzer, and TGA measurements. ^b Calculated based on the surface area of LiCoO₂, obtained by N₂ adsorption-desorption isotherm measurements (0.29 m² g⁻¹). ^c Obtained by LEIS measurements. Surface coverage for c-bare LiCoO₂ is assumed to be 0%.





Figure 7. Characterization of c-bare (cleaned bare), LBO-coated (0.5 wt %), and LBCO-coated (0.5 wt % of LBO) LiCoO2 by electron microscopy analysis. FESEM (upper) and the corresponding BSE (lower) images for (a) c-bare, (b) LBO-coated, and (c) LBCO-coated LiCoO₂ particles.





Figure 8. FESEM and the corresponding BSE images for a) c-bare, b) LBO-coated, and c) LBCO-coated LiCoO₂ powders.





Figure 9. HRTEM images for (a)LBO- and (b)LBCO-coated LiCoO₂ particles. (c) EELS for LBOand LBCO-coated LiCoO₂ particles. The corresponding RTEM images are provided in Figure 10.





Figure 10. STEM images for a) LBO-coated and b) LBCO-coated $LiCoO_2$. The EELS data in Figure 9c correspond to the red spots in (a, b). c) TEM image for LBO-coated $LiCoO_2$ and d) its corresponding SAED pattern, which corresponds with (-1 1 1) and (1 0 1) for Li_3BO_3 (JCDPS no. 18-0718).





Figure 11. Characterization of bare, c-bare, LBO-coated, and LBCO-coated LiCoO₂. (a) XPS spectra for B 1s signals. (b) TGA profiles for bare and c-bare LiCoO₂ in N_2 . (c) LEIS spectra for 5 keV Ne⁺ incident ions.



4.3. Electrochemical characterizations

The electrochemical performances of LiCoO₂/Li-In all-solid-state cells at 30 °C for LBO- and LBCO-coated $LiCoO_2$, depending on the weight fraction of the coatings, are shown in Figure 12 in comparison with those for the c-bare and bare samples. Compared with the c-bare LiCoO₂ samples, all the LBO-coated LiCoO₂ samples showed a lowered polarization in their charge-discharge voltage profiles (Figure 12a) and correspondingly higher capacities, especially at higher C-rates (Figure 12b), confirming the positive effect of the LBO coatings. The optimal performance obtained with 0.1 wt.% of LBO may reflect that an interplay between the lowered direct contact of LiCoO2-LPSCl and the nonimpeded Li⁺ transport through the LBO coating determines the overall kinetics.^{75, 76} The electrochemical performance was further improved by the LBCO-coating (Figure 13a, b). LiCoO₂ coated with LBCO with 0.5 wt.% of LBO exhibited the highest discharge capacities of 142 and 94 mA h g^{-1} at 0.2 and 2C, respectively, which are comparable to those of state-of-the-art LiCoO₂ electrodes in ASLBs.^{11, 12, 34} It should be noted that the LBCO coatings allow for a larger weight fraction (0.5 wt.% of LBO) than the LBO coatings (0.1 wt.%) to achieve an optimal rate capability, which can be attributed to the much higher Li⁺ conductivity of LBCO compared with of LBO. LBCO coating was also applied on c-bare LiCoO₂ using an aqueous solution containing LiOH, H₃BO₃, and Li₂CO₃; this sample is referred to as artificial LBCO-coated LiCoO₂ (a-LBCO). Consistent with the results of LBCO-coated LiCoO₂, a-LBCO-coated LiCoO₂ also showed an excellent rate capability. The trend of improvement, which goes in the order of bare (or c-bare), LBO-coated, and LBCO-coated samples, agrees well with the lowered polarization in the transient discharge voltage profiles obtained by galvanostatic intermittent titration technique (GITT) (Figure 14a) and the smaller interfacial resistances obtained from Nyquist plots (Figure 14b, 15, Table 7).

The cycling performances of LiCoO₂/Li-In all-solid-state cells at 0.2C and 30 °C using c-bare, LBOcoated, and LBCO-coated LiCoO₂ are shown in Figure 16. With an upper cutoff voltage of 4.3 V (*vs.* Li/Li⁺), the capacity retention for c-bare samples after 50 cycles, compared with that at the second cycle, was 88.8%. The coatings of LBO (0.1 wt.%) and LBCO (0.5 wt.% LBO) resulted in enhancements in capacity retention: 92.2% and 93.8%, respectively. When the upper cutoff voltage was raised to 4.5 V (*vs.* Li/Li⁺), more dramatic improvements in cycling performance caused by the coating were confirmed; the capacity retentions after 25 cycles, compared with that at the fourth cycle, were 81.6%, 88.7%, and 93.8% for c-bare, LBO-coated, and LBCO-coated LiCoO₂, respectively. Notably, the electrochemical performance of LBCO-coated LiCoO₂ for ASLBs appears to be superior to even that of the LiNbO₃-coated sample (Fig. S5). From the electrochemical results, the following features are summarized: i) the rate capability and cycling performances are enhanced, from worst to best, in the order of bare (or c-bare), LBO-coated, and LBCO-coated LiCoO₂, ii) compared with LBO coatings, thicker coatings are possible using LBCO thanks to its higher Li⁺ conductivity.





Figure 12. Electrochemical characterization of LiCoO₂/Li–In all-solid-state cells at 30 °C. Charge–discharge voltage profiles for (a) LBO- and (b) LBCO-coated LiCoO₂ varied by C-rate. The results for c-bare, bare, and a-LBCO-coated (artificial-LBCO-coated) LiCoO₂ are compared in (a, b).





Figure 13. Electrochemical characterization of $LiCoO_2/Li$ –In all-solid-state cells at 30 °C. Rate performances for (a) LBO- and (b) LBCO-coated $LiCoO_2$. The results for c-bare, bare, and a-LBCO-coated (artificial-LBCO-coated) $LiCoO_2$ are compared in (a–b).





Figure 14. Electrochemical characterization of $LiCoO_2/Li$ –In all-solid-state cells at 30 °C. (a) Transient discharge voltage profiles obtained by GITT. (b) Nyquist plots of $LiCoO_2/Li$ –In cells. The corresponding equivalent circuit model and interfacial resistances are shown in Figure 15 and Table 7, respectively.





Figure 15. Voigt-type equivalent circuit used for fitting the EIS data shown in Figure 14b.





Figure 16. Cycling performances for LiCoO₂/Li–In all-solid-state cells using c-bare, LBO-coated, and LBCO-coated LiCoO₂ at 0.2C and 30 °C. Discharge capacities as a function of the number of cycles in the voltage ranges of (a) 3.0–4.3 V (vs. Li/Li⁺) and (b) 3.0–4.5 V (vs. Li/Li⁺).





Figure 17. Thermodynamic calculation results for LiNbO₃ and electrochemical characterization of LiNbO₃-coated LiCoO₂ (1.0 wt%). a) Calculated mutual decomposition energy of Li₆PS₅Cl with LiNbO₃. b) Charge-discharge voltage profiles at 0.2C and 2C and cycle performance for LiNbO₃-coated LiCoO₂ in LiCoO₂/Li-In all-solid-state cells at 30 °C. The results for bare, LBO-coated, and LBCO-coated LiCoO₂ are compared.



4.4. Ex-situ Surface analysis

As an attempt to gain mechanistic insights on the protective coatings on LiCoO₂ for ASLBs, ex-situ XPS analyses were carried out for c-bare, LBO-coated, and LBCO-coated LiCoO₂ electrodes before and after cycling to probe for changes at the electrode-SE interfaces. Because the mixture electrodes do not contain conducting carbon additives, any effects caused by carbon-SE interfaces could be ruled out. The signals for Co 2p, S 2p, and P 2p are shown in Figure 18-20. For the Co 2p spectra shown in Figure 18, the evolution of Co_3S_4 after cycling (shown in the deconvoluted peaks in violet) is noticeable.^{77, 78} Because the physical mixture sample of c-bare LiCoO₂/LPSCl does not show the signature of Co_3S_4 , the formation of Co_3S_4 is suspected to be electrochemically driven, which is consistent with the observation of interatomic diffusion of Co and S at the interfaces of LiCoO₂/Li₂S·P₂S₅ presented in a previous report.³¹ Because Co₃S₄ is electronically conducting (thus non-passivating), reactions at bare LiCoO₂/LPSCl interfaces occur progressively, which is detrimental to their electrochemical performance.^{25, 26, 29} In stark contrast, the Co 2p signal for LBO-coated $LiCoO_2$ after cycling shows a much lower intensity for Co_3S_4 . Moreover, LBCO-coated LiCoO₂ after cycling showed a negligible signature for Co₃S₄. This result reflects the excellent protection of $LiCoO_2$ provided by LBCO, which can be attributed to its high surface coverage (Figure 11c, Table 6) and its buffering effects, as our first principles computational results suggest (Table 4). In a consistent fashion, the suppressed evolution of Co₃S₄ after cycling from worst to best was confirmed to be in the order of c-bare, LBO-coated, and LBCO-coated LiCoO₂, as shown by the S 2p signals in Figure 19. As shown in the P 2p signals in Figure 20, the signature of phosphate (PO_4^{3-} , shown in the deconvoluted peaks in dark cyan) appeared for the LBO-coated sample and became more intense for the LBCOcoated one.^{43, 79} The phosphate species could be derived from the electrochemical reaction of LBO or LBCO with LPSC1. In contrast to Co₃S₄, the as-formed phosphates are good electronic insulators, thus effectively passivating to inhibit the continuous decomposition of the bulk SEs.^{25, 26, 29} The evolution of P₂S_{5+x} and S-S (bridging sulfur) after cycling observed in the S 2p and P 2p signals is consistent with previous reports.43,80,81

Based on the electrochemical characterization and the complementary analyses presented so far, the interface phases between cathode and SE material appear to be sensitively dependent on the coating materials used, as illustrated in Figure 21. The surfaces of bare $LiCoO_2$ are covered by the impurities, including Li_2CO_3 . More importantly, the electrochemically-driven reactions between $LiCoO_2$ and LPSCl form detrimental mixed conducting interphases (MCIs), as evidenced by the observation of Co_3S_4 , which shows a lack of passivating capability. The aqueous-solution coating process for LBO renders to form the LBCO layers. The high Li^+ conductivity of LBCO allows for the formation of thick and thus high-surface-coverage protective layers, which suppresses the significant decomposition



at the interface. Moreover, the electrochemical reaction of LBCO with LPSCl enables the formation of good passivating layers comprised of phosphates. As an overall consequence, LBCO coating on LiCoO₂ results in significant improvements in rate capability and durability.





Figure 18. XPS results of Co 2p signal for c-bare, LBO-coated (0.1 wt %), and LBCO-coated (0.5 wt % of LBO) LiCoO₂ for pristine powders and electrodes after cycling. The data for LiCoO₂/SE (Li₆PS₅Cl) mixtures is also shown for comparison.





Figure 19. XPS results of S 2p signal for c-bare, LBO-coated (0.1 wt %), and LBCO-coated (0.5 wt % of LBO) LiCoO₂ for electrodes after cycling. The data for LiCoO₂/SE (Li₆PS₅Cl) mixtures is also shown for comparison.





Figure 20. XPS results of P 2p signal for c-bare, LBO-coated (0.1 wt %), and LBCO-coated (0.5 wt % of LBO) LiCoO₂ for electrodes after cycling. The data for LiCoO₂/SE (Li₆PS₅Cl) mixtures is also shown for comparison.





Figure 21. Schematic diagram illustrating the different interface features of bare and LBCO-coated LiCoO₂ in all-solid-state-cell electrodes.



5. Conclusion

In summary, a new LBCO coating process on LiCoO₂ for sulfide-based ASLBs via a scalable aqueous-solution protocol was rationally designed, considering the formation of an interphase between the cathode and SE materials and was demonstrated to significantly improve electrochemical performances. Using the aforementioned aqueous LBO-solution process, the poorly Li⁺-conducting surface impurity on LiCoO₂, Li₂CO₃, could be converted into highly Li⁺-conductive LBCO (max. conductivity of 6.0×10^{-7} S cm⁻¹ at 30 °C), which could protect LiCoO₂ with thick and high-surface-coverage layers. More specifically, LiCoO₂/Li-In all-solid-state cells employing the proposed LBCO coating with 0.5 wt.% LBO showed discharge capacities of 142 and 94 mA h g⁻¹ at 30 °C at 0.2C and 2C, respectively, in contrast to the discharge capacities of 107 and 18 mA h g⁻¹ obtained for the ones using bare LiCoO₂. From the complementary analyses by electrochemical measurements, XRD, FESEM, BSE, HRTEM, EELS, TGA, LEIS, and ex-situ XPS, it was revealed that the LBCO coatings prevent the evolution of detrimental MCIs containing Co₃S₄ and can effectively passivate the interfaces by alternatively forming phosphate-based phases. We believe that our results provide not only an indepth mechanistic understanding on the interfacial evolutions for ASLBs, but also open up a new avenue to rationally engineer the interfaces for practical all-solid-state technologies.



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