Review Article

Porous Mixed Ionic Electronic Conductor Interlayers for Solid-State Batteries

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Received 11 December 2020; Accepted 4 February 2021; Published 29 March 2021

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Rechargeable solid-state batteries (SSBs) have emerged as the next-generation energy storage device based on lowered fire hazard and the potential of realizing advanced battery chemistries, such as alkali metal anodes. However, ceramic solid electrolytes (SEs) generally have limited capability in relieving mechanical stress and are not chemically stable against body-centered cubic alkali metals or their alloys with minor solute elements ($\beta$-phase). Swelling-then-retreating of $\beta$-phase often causes instabilities such as SE fracture and corrosion as well as the loss of electronic/ionic contact, which leads to high charge-transfer resistance, short-circuiting, etc. These challenges have called for the cooperation from other classes of materials and novel nanocomposite architectures in relieving stress and preserving essential contacts while minimizing detrimental disruptions. In this review, we summarize recent progress in addressing these issues by incorporating other classes of materials such as mixed ion-electron conductor (MIEC) porous interlayers and ion-electron insulator (IEI) binders, in addition to SE and metals (e.g., $\beta$-phase and current collectors) that are the traditional SSB components. In particular, we focus on providing theoretical interpretations on how open nanoporous MIEC interlayers manipulate $\beta$-phase deposition and stripping behavior and thereby suppress such instabilities, referring to the fundamental thermodynamics and kinetics governing the nucleation and growth of the $\beta$-phase. The review concludes by describing avenues for the future design of porous MIEC interlayers for SSBs.

1. Introduction

Remarkable breakthroughs in liquid electrolyte- (LE-) based rechargeable Li-ion batteries (LIB) have enabled modern portable electronics and electric vehicles (EVs). Such applications, however, now call for solid-state batteries (SSBs) due to safety concerns and the demand for higher energy density [1, 2]. In this respect, rechargeable alkali metal SSBs with ceramic solid electrolytes (SEs) are promising alternatives, but they have yet to achieve sufficient cycling stability [2]. This review thus is aimed at critically examining the need for SEs and other types of materials in developing safe rechargeable batteries with both high energy density and long-term stability.

With the mass marketing of portable electronics and EVs, the shortcomings of LE-based batteries became more apparent [1]. First, organic LE-based batteries have significant fire and explosion risks due to the high vapor pressure and combustibility of organic liquids when mixed with oxygen gas [3]. They usually have metal oxides with high oxygen content on the cathode and thermally unstable solid electrolyte interphase (SEI) on both electrodes, which is produced by organic LEs reacting with the electrodes. When these batteries are subjected to abuse by faulty operation or traffic accidents, severe thermal runaways can occur, triggered by the liquid’s high volatility (equilibrium vapor pressure) and fueled by the cathode’s oxygen inventory and SEI decomposition [4]. Second, aqueous LEs, while not as dangerous, traditionally have lower energy density because of a narrower electrochemical stability window $[U_{\text{lower}}, U_{\text{upper}}]$ where $U$ is the absolute electronic potential [5] that is related thermodynamically to a charge-neutral Li-atom’s chemical potential embedded in either electrode, beyond which an electrolyte is reduced or oxidized. Third, by dissolving a variety of unwanted stray ions from cathodes (e.g., Mn$^{2+}$ in Li-ion batteries (LIBs) [6] or polysulfides in Li-S batteries [7]) and
conducting them and their solution shells via vehicular diffusion, LEs often enable cathode-anode shuttling and parasitic reactions that reduce the cycle life and shelf life of batteries.

In contrast, SEs have clear advantages in terms of safety and charge-carrier-transport characteristics. First, most ceramic SEs are less prone to catching fire since they have low vapor pressure and are less flammable than organic LEs [2, 8]. Moreover, most of them can enable advanced battery chemistries that would offer higher energy density by forbidding unwanted ions to pass through them [1]. The total bulk electrical conductivity of any material can be partitioned into transference numbers for various charge carriers such as electrons, a primary cation (e.g., Li+ for LIBs and Na+ for sodium-ion batteries), and other ions: \( t = t_{\text{electron}} + t_{\text{Li+}} + t_{\text{other ions}} \). In electrolytes, which are electronic insulators, \( t_{\text{electron}} \) is essentially zero, and ions partition the bulk electrical conductivity. While LEs often have \( t_{\text{Li+}} \) less than 0.5 due to vehicular diffusions of all free ions with their solvation shells, \( t_{\text{Li+}} \) of most inorganic SEs is very close to 1 at service temperature [2], and hence, the flow of ions other than Li+ is strongly forbidden in SEs due to the exchange mechanism of diffusion in SEs. This \( t_{\text{Li+}} = 1 \) solves the cathode-anode shuttling and parasitic reaction problem generically, opening up more possible chemistries (like high-energy-density Mn-rich or sulfur cathodes) and a higher temperature range for stable batteries. Furthermore, there is a strong expectation that SEs may enable the use of high-energy-density alkali metal anodes, which can realize longer mileage of EVs [1, 9, 10]. This expectation arose from the SE’s relatively high elastic modulus, which was believed to hinder dendritic penetration of \( \beta \)-phase [11], as \( \beta \)-phase tends to be elastically quite soft. It later turned out that high elastic modulus alone cannot combat the dendrite problem when stress and corrosion attacks on SEs are combined, they can synergistically work to disrupt essential contacts or create unwanted contacts, sometimes even resulting in SE fracture. The design strategies for “3D” anodes have thus been of great interest, with an emphasis on manipulating the microscopic \( \beta \)-phase deposition and stripping behavior through controlling its nucleation and growth kinetics, that share many common features with classical physical metallurgy examples.

In this review, we begin by summarizing the mechanical-electrochemical origin of the aforementioned instabilities at the SE/\( \beta \)-phase interface. We then examine 3D open porous architectures constructed with different types of materials and discuss possible roles of phase transformation, wetting, diffusion, and \( \beta \)-phase creep/plasticity in mitigating driving forces of instabilities and thereby maintaining acceptable \( R_s \) as well as \( t_{\text{electron}} = 0 \) barrier function of SEs. Moreover, since SEs often need to transmit several MPa of pressure, we also assess the role of stack pressure in SSBs, and whether this pressure is absolutely necessary. The traditional components of anodes for rechargeable alkali metal SSBs require SE and metals (e.g., \( \beta \)-phase, and the Cu current collector, CC). The main thesis of this review is that more material classes are required in certain spatial arrangements to achieve long-term stability of rechargeable alkali metal SSBs. In particular, anodes would need to incorporate two additional classes of materials—a mixed ion-electron conductor (MIEC) and an ion-electron insulator (IEI)—of a certain architecture.

2. Evolution of Instabilities in Alkali Metal SSBs

\( \beta \)-phase often corrodes the surrounding material components in rechargeable alkali metal SSBs. In Figure 2, we show that materials can be generally classified into four quadrants according to whether they conduct free electrons and/or the
primary ion. A metal (M) conducts free electrons but not ions. A solid electrolyte (SE) conducts $\text{Li}^+$ or $\text{Na}^+$ but not free electrons. A mixed ion-electron conductor (MIEC) conducts both $\text{Li}^+$ or $\text{Na}^+$ and electrons (i.e., $t_{\text{ions}} > 0$ and $t_{\text{electron}} > 0$), while an ion-electron insulator (IEI) conducts neither. It is possible to find materials in each of the M, SE, MIEC, and IEI categories which are thermodynamically absolutely stable against LiBCC or NaBCC, but there are many more counterexamples in each category also, because the $\beta$-phase is chemically so reactive.

The corrosiveness of $\beta$-phase is reflected by the fact that only a small set of compounds among the 200,000+ known crystals in the Inorganic Crystal Structure Database (ICSD) have a direct tie-line to the $\beta$-phase on the multi-element phase diagram (e.g., ternary and quaternary phase diagrams shown at https://materialsproject.org). If a solid phase $\text{A}_m\text{B}_n\text{C}_q$ of interest does not have a direct tie-line to LiBCC in a Li-A-B-C phase diagram at the temperature of interest, the mixture of $x\text{A}_m\text{B}_n\text{C}_q$ and $y\text{LiBCC}$ by definition would prefer to react and turn into a combination of other intervening phases. That is to say, $\text{A}_m\text{B}_n\text{C}_q$ would have no thermodynamic immunity against $\beta$-phase corrosion. If we think of a ternary phase diagram as a “flight map,” then only the phases with “direct flight” to the city

Figure 1: Electrochemical stability window [$U_{\text{lower}}, U_{\text{upper}}$] of various solid electrolytes. Those of the corresponding binary inorganic solids are also provided for comparison. Reproduced from Ref. [18] with permission from American Chemical Society. Further permissions related to this figure should be directed to the American Chemical Society.
of \( \beta \)-phase (e.g., tie-line) are stable when spatially put side-by-side with the \( \beta \)-phase. This also means all other phases with minimally more than one transit to reach the \( \beta \)-phase would be thermodynamically unstable against it.

If the SE in a rechargeable SSB is \( A_2B_mC_y \) that does not have a direct tie-line to the \( \beta \)-phase, the corrosive \( xA_mB_nC_q + yLi_{BCC} \) reaction happens and gives rise to different instabilities depending on the transference numbers of the reaction products. First, suppose that the reaction products are either M or MIEC, which have nonzero \( t_{electron} \). If not kinetically stopped, the \( xA_mB_nC_q + yLi_{BCC} \) reaction will proceed in a self-perpetuating manner during the charging process. This is because \( xA_mB_nC_y \) will then have kinetic access to both \( Li^+ \) (SE or MIEC) and \( e^- \) (M or MIEC), which would combine to produce additional charge-neutral Li atoms under a cathodic current by the following charge-transfer half-cell reaction \( Li^+ (SE, MIEC) + e^- (M, MIEC) = Li \) atom in the \( Li_{BCC} \) phase. As the anode’s potential drops below 0 V versus Li upon charging of the full cell to give a driving force for \( Li_{BCC} \) deposition to happen, there will be a continuous supply of Li atoms into the \( xA_mB_nC_y + yLi_{BCC} \) system at a chemical potential \( \mu_{Li} \) equal to or lower than that of Li atoms residing in the \( Li_{BCC} \) phase. In other words, \( y \) will continue to increase, and the \( xA_mB_nC_q + yLi_{BCC} \) reaction will keep proceeding forward, further converting SE to more M or MIEC. Such reductive metallization may occur preferentially along the grain boundary (GB) network of SEs and can eventually cause SE failure and short-circuiting, which is often diagnosed as showing near-zero open-circuit voltage in battery full cells.

In contrast, suppose that the reaction products have \( t_{electron} = 0 \) and have direct tie-line to the \( \beta \)-phase themselves so they are stable side-by-side with \( \beta \). They may then form a kinetic passivation layer, cutting off the supply of free electrons, which would have otherwise combined with \( Li^+ \) to offer Li atoms to the \( xA_mB_nC_y + yLi_{BCC} \) system. If this layer is sufficiently compact and adherent, it will stop corrosive reactions kinetically, being the reductive analog of \( Al_2O_3 \) on Al in an oxidizing environment [19]. This layer can, however, have much poorer ionic conductivity than the original SE—\( Li_2O \) could form, for example—or even become an IEI. Such conversion would cause an increase of \( R_n \) in future cycles. Therefore, it is often an “either a drought or a flood” problem when corrosion of original SE occurs. Also, it is usually the case that this passivation layer does not last very long due to the mechanically stressful \( \beta \)-phase.

Next, we address the stressful aspect of the \( \beta \)-phase and the resultant electrochemomechanical instabilities. Stress can be easily generated in the course of \( \beta \)-phase deposition and stripping, as seen in the thermodynamics of the reaction:

\[
Li^+ (SE or MIEC) + e^- (U) = Li \text{ atom in } \beta \text{-phase} \quad (1)
\]

where \( U \) is the local electronic potential in M or MIEC that feeds the electron to the site of the reaction above. For the right-hand side, there is

\[
\mu_{Li} (\text{Li atom in } \beta \text{-phase}) = \mu_{Li}^{\ast} + k_B T \ln (y_{Li}X_{Li}) + \Omega_{Li}P(x) \quad (2)
\]

to leading order, where \( \mu_{Li}^{\ast} \) is the reference chemical potential of Li atom in pure \( Li_{BCC} \) in stress-free condition, \( y_{Li}X_{Li} \) is the composition-dependent chemical activity, and \( \Omega_{Li} \) is the partial molar volume of Li atom (~21.6 Å³) in \( \beta \)-phase. \( P(x) = -Tr(\sigma(x))/3 \), where \( \sigma(x) \) is the position-dependent mechanical stress field. Equating the left- and right-hand side of Equation (1) shows that an increase of the local overpotential by just 0.135 V can cause GPa-level stress inside the \( \beta \)-phase if a thermodynamic equilibrium is to be reached [20].

The insertion volume and generated stress mechanically load up SSBs’ solid components, thereby driving fatigue and failure. Some of the in situ formed passivation layers may not be able to withstand such stresses and end up being spalled off as electron-insulating debris. This spallation would lead to capacity loss as the active Li would be

| Ion conductor | Ion insulator |
|---------------|---------------|
| Ion conductor and electron conductor | Ion insulator and electron conductor |
| a.k.a. Mixed ion-electron conductors (MIEC) | a.k.a. normal metal (M) |
| e.g., LiC\(_n\), LiAl, Li\(_{2+y}\)Si\(_x\) for Li\(^+\) | e.g., Ni, Stainless steel |

![Figure 2: Four different types of materials classified according to whether they conduct electrons and/or primary ions. Examples of mixed ion-electron conductors (MIEC) and ion-electron insulators (IEI) are provided, taking Li\(^+\) as primary ions.](image-url)
consumed to form a new passivation layer, and the debris can also disrupt or block the electron-conducting paths. Moreover, since the stresses with an opposite sign arise during discharging, the solid components of SSBs are subjected to cyclic stresses throughout the operation, which can deteriorate interfacial contact and drive fatigue crack growth. Particularly, \(\beta\)-phase deposition at preexisting flaws on the surface of SEs generates crack-tip stress, which drives crack propagation [12]. As the surface flaws and cracks are favorable nucleation sites [21], this attack generally occurs in a self-reinforcing manner, forming dendrites, and results in SE fracture and short-circuiting.

The pressure in the \(\beta\)-phase can be relaxed by plastic deformation driven by a pressure gradient \(\nabla P(x)\) and shear stress—it can be shown that there has to be deviatoric shear stress whenever there is a \(\nabla P(x)\)—that allow the atoms in \(\beta\)-phase to either convect or diffuse into adjacent spaces. Such stress relaxation requires either prearranged empty space (reserved porosity) or working against the stack pressure to create the extra space for insertion. If the former route is taken, the convection or diffusion of Li atoms in the \(\beta\)-phase, or at the interface with the \(\beta\)-phase, must be facile enough to cover the distance between where reaction (1) happens and where the reserved porosity is. This process then involves either displacive deformation or diffusion of the metal atoms. In addition, nucleation of the \(\beta\)-phase is also required, and standard metallurgical treatment of the interfacial wetting, heterogeneous nucleation, and Gibbs-Thomson effect induced coarsening/ripening needs to be discussed. Lastly, we note that the reserved pore spaces with vacuum or inert gas can be considered a type of IEI, although typically we take IEI to mean the binder solid phase between SE and MIEC [20].

3. Materials and Architecture

One promising approach adopted to relieve the corrosion and stress attacks from the \(\beta\)-phase on the move is to accommodate it in a 3D MIEC architecture with preserialized porous regions. For instance, Xu et al. [22] introduced a MIEC framework, where a 3D porous garnet structure is covered by a conformal carbon nanotube coating (Figure 3(a)), and achieved stable cycling over a few tens of cycles at a high current density of \(\geq 1\) mA cm\(^{-2}\). The garnet conducts \(\text{Li}^+\), whereas the carbon nanotube conducts both electrons and \(\text{Li}^+\) upon lithiation. Alloy-type MIECs have also been employed. For example, Yang et al. [23] cycled Li-Mg random-solid-solution alloy, which has an appreciable solubility of \(\text{Li}\) atoms and thus can be considered as a MIEC, and reported stable cycling over a few hundred cycles at a high current density of \(\geq 1\) mA cm\(^{-2}\). Furthermore, Zhu et al. [24] utilized a Sn-Ni alloy-coated Cu nanowire network as an anode (Figure 3(b)), where Li-Sn intermetallic compound forms upon lithiation and functions as a MIEC. They demonstrated a notable improvement in rate capability—a doubling of capacity at 5C and a more than five times increase in cycle life at 1C as compared to the Li metal anode—without detrimental corrosion or stress-induced collapse.

MIECs have been chosen as the base materials for the porous interlayer or 3D electrode by virtue of their conduction properties. First, having kinetic access to both \(\text{Li}^+\) and electrons, they can provide sites for reaction (1) to occur, and hence curtail \(\text{Li}\) atom insertion at the SE/\(\beta\)-phase interface; the large surface area of the 3D porous MIEC gives it an advantage. It has thus been expected that 3D MIEC architectures would alleviate spallation at the SE/\(\beta\)-phase interface and the associated capacity loss [26]. Meanwhile, the spallation at the MIEC/\(\beta\)-phase interface has generally been thought to be negligible since the formation of interphase (e.g., passivation layer) itself can be prevented by employing the MIECs that are thermodynamically absolutely stable (not just kinetically passivated) in the range of operating \(\text{Li}\) potentials [26]. Second, porous MIECs can secure electron-conducting paths even when there exists electron-insulating debris spalled off from the SE/\(\beta\)-phase interface as they conduct electrons in a redundantly percolating fashion. As the rate at which \(R_{\text{ct}}\) increases upon cycling can be slowed down in 3D porous MIEC-assisted SSBs, better rate capability can be anticipated [22–24].

Based on MIECs’ conduction properties and 3D architectures’ open porosity, 3D MIEC architectures have been expected to realize pragmatic rechargeable SSBs by accomplishing high energy density, long-term stability, and rate capability simultaneously. That being said, most of the 3D MIEC architectures did not prevent direct contact of \(\beta\)-phase with an SE, leaving the progressive evolution of interfacial instabilities still possible, especially when high stresses are involved. The bulk ionic conductivities of inorganic SEs should be orders of magnitude smaller than the bulk electronic conductivities of many MIECs. Electrons should thus arrive “earlier” at the SE/MIEC interface and “wait” for \(\text{Li}^+\) (or \(\text{Na}^+\)) that travel through the SEs to arrive, thereby potentially creating \(\beta\)-phase at the SE/MIEC interface and engendering undesired stress. Mitigating this concern fundamentally demands a better understanding of the \(\beta\)-phase deposition/stripping kinetics based on the nucleation and growth of metals and alloys, and size dependence of these behaviors. Note that we use the word “mitigate” instead of “eliminate,” as \(\beta\)-phase may never be completely eliminated at/near the MIEC/SE interface. However, as long as those \(\beta\)-phases are small in quantity and do not have high mechanical stress, they will not be as aggressive and damaging to the SE.

A mechanistic understanding of how 3D MIEC architectures manage \(\beta\)-phase “pumping” has been provided using a simple structure consisting of aligned open-porous carbon tubules by Chen et al. [20]. Figure 3(c) shows the cross-sectional schematic of the adopted cell configuration on the anode side. The MIEC tubules are colored red. The ends of the tubules are rooted in the SE and the metallic current collector (CC), respectively. CC is typically copper or stainless steel foil. In this cell configuration, the neutralized metallic \(\text{Li}\) atoms are still produced mostly at the SE/MIEC interface (or SE/LpBCC interface in later stages). The continuous insertion of \(\text{Li}\) during plating, however, generates high compressive stress at this interface as discussed above. Since the other end of the LpBCC is the free surface with nearly zero
pressure, the pressure difference and the resultant gradient in the chemical potential of Li BCC develop throughout the tubules [20, 27, 28]. This gradient provides the driving force for the transport of metallic Li from the SE side to the CC side. Moreover, from the aspect of kinetics, it has been pointed out that the low melting point of Li BCC and large interfacial area that the 3D architecture provides would allow Li atoms to diffuse rapidly along the wall or surface of the MIEC [29] (namely, Coble creep), as illustrated in the boxed area of Figure 3(c).

These findings provide important boundary conditions regarding the β-phase deposition. First, they indicate that β-phase can nucleate at or diffuse into the interface between the roots of the MIEC and the SE, which is unfavorable from the viewpoint of structural stability. Provided that the MIEC has a good wettability by the β-phase, which is in fact a desirable property to offer appreciable interfacial solubility (i.e., segregation) and diffusion rate, the interface between the SE and the MIEC could also be an attractive site for heterogeneous nucleation of β-phase due to the low nucleation barrier therein. Subsequently, the soft β-phase infused between SE/MIEC can enable interfacial sliding like a lubricant, thereby making the MIEC roots mechanically unstable and facilitating interfacial decohesion upon stripping. Hence, 3D MIEC/SE architecture alone would be electrochemomechanically unstable unless large-scale β-phase invasion into MIEC/SE interface is averted [20, 26]. To circumvent this issue, it is proposed to cover the root with an IEI (colored yellow in Figure 3(c)), which is a class of materials that has not been used for anode construction [20]. The IEI layer is expected to prevent β-phase deposition at the root and function as a mechanical binder [20]. To function properly, an IEI...
should be not only insulating to both electrons and ions but also thermodynamically stable against β-phase [20]. It has thus been suggested that IEIs for this use should sit on a direct tie-line with the β-phase in the phase diagram with minimal Li solubility (lithiophobic [30]) and have a large electronic bandgap, like over 4.0 eV [20], to be sufficiently electronically insulating and stopping β-phase nucleation and/or aggression.

Second, if nucleation barriers at different interfaces are properly manipulated, it might be possible to induce β-phase to nucleate elsewhere other than the SE/MIEC interface or coarsen away the β-phase at SE/MIEC interface even if it is nucleated. In this respect, one 3D MIEC architecture worth noting is the Ag-C nanocomposite recently developed by Lee et al. [25], which has been verified to deposit β-phase at the MIEC/CC interface. It was constructed of carbon black powders and Ag nanoparticles with the diameter of ~60 nm. A schematic representation of the cross-section of the cell is provided in Figure 3(d). As the carbon black conducts both Li+ and electrons, this nanocomposite can be considered as an open-porous MIEC with the inner surfaces decorated with Ag. Interestingly, β-phase with dissolved Ag was formed between the MIEC and the CC as shown in the boxed area of Figure 3(d). The SE/MIEC interface remained largely free of direct contact with β-phase, unlike those in most of the MIEC-based 3D architectures, which establish direct contact with β-phase. Aided by the electrochemical and mechanical stability at the interface, this cell achieved Coulombic efficiency as high as 99.8% when averaged over 1,000 cycles, which is by far the best performance among the reported rechargeable alkali metal SSBs in terms of cycle life and long-term Coulombic efficiency [31, 32].

4. Mechanisms

To understand the mechanism underlying the β-phase deposition at the MIEC/CC interface, the evolution of microstructures upon charging/discharging was investigated with a focus on the room-temperature redistribution of both metallic Li and Ag [25]. Figure 4 shows the schematic of β-phase deposition/stripping through the open-porous MIEC interlayer with metal nanoparticles and microstructure evolution therein (Figure 4(a)) as well as the experimental observations (Figures 4(b) and 4(c)). First, the electron energy loss spectroscopy analyses and selected area diffraction pattern analyses with a transmission electron microscope (Figure 4(b)) revealed that Ag nanoparticles undergo morphological and structural changes during cycling, which suggests the formation of Li-Ag alloys inside the MIEC interlayer. Second, the energy dispersive spectroscopy analyses in a scanning electron microscope (Figure 4(c)) showed that most of the metallic Ag moves to the CC side (instead of the SE side) and forms β-phase there during charging. It goes back to the MIEC interlayer upon discharging; however, Ag that was dissolved in the β-phase remains mostly at the CC side of the MIEC interlayer, making the distribution of Ag inside the MIEC interlayer more and more asymmetric. Lastly, in terms of morphology, Ag nanoparticles in the MIEC layer were found to become fragmented upon cycling, but no instabilities such as pores were observed at the solid-solid interfaces. Based on these experimental findings, we describe next the thermodynamic driving forces that may be responsible for the microstructural evolution as well as the kinetic feasibility of such asymmetry development.

4.1. Li Deposition at MIEC/CC Interface. One of the most important yet puzzling features is that the β-phase layer is formed at the MIEC/CC interface as shown in Figure 4(c) [25]. It is generally thought that the rate-limiting factor in rechargeable SSBs is long-range ion transport in an SE [33, 34]. However, if the long-range transport was the dominant factor, β-phase should have nucleated at the SE/MIEC interface instead of the MIEC/CC interface, since the SE’s ionic conductivity should be several orders of magnitude smaller than the MIEC’s electronic conductivity. Such an issue was also revealed from a careful intrinsic rate capability study of rechargeable Na-ion SSBs [35]. Their rate performance was greatly improved when the \( R_{ct} \) was lowered, while the change of the SE’s bulk ionic conductivity in the range of 0.48–0.12 mS cm⁻¹ caused no difference. These observations indicate that the charge transfer reactions on the anode side can be governed by short-range, reaction-limited kinetics (phase nucleation barriers, interfacial wetting, Butler-Volmer electron transfer rate, etc.), given that SE’s bulk ionic conductivity is already in an acceptably high range.

Indeed, the open porous MIEC interlayer architecture with Ag creates an environment in which short-range \( R_{ct} \), not long-range transport, can manipulate LiBCC nucleation sites. First, the argyrodite-based SE has a bulk ionic conductivity greater than 1 mS cm⁻¹ [25], which satisfies the minimum requirement on the conductivity to allow other factors to be a determinant. Second, the MIEC interlayer’s porosity can decrease \( R_{ct} \); \( R_{en} \) is the only term that physically should scale with the true contact area in most equivalent-circuit models of the battery. As the porosity provides a large surface or interfacial contact area, it would help lower \( R_{ct} \). Finally, Ag nanoparticles can reduce the β-phase nucleation barrier. In rechargeable alkali metal SSBs, charge transfer reactions involve β-phase nucleation, in which the energy barrier depends on the interfacial wetting [21]; the greater the wettability (i.e., the smaller the wetting angle), the lower the barrier. The wettability of Ag nanoparticles by LiBCC is better than that of the MIEC by virtue of their metallic character and Ag’s solubility in LiBCC (~1 at.% at room temperature [36, 37]). They can thus lower the nucleation barrier and \( R_{ct} \). The recent finding that the overpotential required for β-phase nucleation on substrates with definite solubility in LiBCC (e.g., Au, Ag, Mg, Al, and Pt) is nearly zero provides a proof of principle for this effect [38] of lowering the β-phase nucleation barrier. Hence, \( R_{ct} \) might have motivated LiBCC to nucleate first at the inner surfaces of the MIEC interlayer.

That being said, \( R_{ct} \) alone cannot explain the β-phase location after first charging. Following the LiBCC nucleation inside the MIEC, coarsening would occur due to the Gibbs-Thomson effect [21]; different surface curvatures of the deposits raise the chemical potential of the components in each deposit to different extents, thereby making them favor
some deposits over the others. As the curvature term is inversely proportional to the radius of the deposits [21], this term is essentially zero for a flat surface. Hence, the deposits would eventually reside at either the SE/MIEC interface or the MIEC/CC interface that are flatter rather than on the inner surface of the MIEC, which is wavier. The MIEC/CC interface, however, does not have an advantage over the SE/MIEC interface in terms of the transport distance. It is thus likely that other factors such as interface chemistry play some role. One possible scenario is that metallic Li or Ag atoms near the SE/MIEC interface react with the SEs and form IEs (e.g., Li$_2$S or Ag$_2$S) during a warm-isostatic-pressing process, which is conducted to improve the contact at the SE/MIEC interface. The presence of IEs may provide a stronger adhesive force while lowering electronic conductivity, thereby making the interface more Li-repellant.

On the other hand, β-phase deposition at the MIEC/CC interface would be achieved more easily in the subsequent cycles owing to the extensive transport of metallic Ag to the CC side during the first cycle. The current collector (made of stainless steel or copper) may have a better wetting angle with Ag and is more “argentophilic” (silver is argentum in Latin and argunas in Sanskrit) than the SE. As shown in Figure 4(c), metallic Ag in the MIEC interlayer moves to the CC side during charging, but it does not restore its uniform distribution upon discharging. Instead, it shows a higher concentration near the MIEC/CC interface. Considering that metallic Ag lowers $R_{ct}$, this movement would result in a further reduction of $R_{ct}$ at this interface, and thus, metallic Li would favor the MIEC/CC interface in the later cycles. This conjecture is also in good accordance with the relatively low $R_{ct}$ value (~5 Ω cm$^2$ at 60°C) provided in the paper [25].
which was measured via the electrochemical impedance spectroscopy after charging.

4.2. Ag Transport at Room Temperature. The next step is to ascertain why and how metallic Ag moves at room temperature (RT). To determine the thermodynamic driving force for the diffusion of metallic Ag, the phase where Ag atoms reside should first be identified. The maximum Ag content is limited by the amount of Ag employed in cell preparation, which is 8–16 mg Ah\(^{-1}\). Since 1 Ah corresponds to 259 mg of Li, the maximum content of Ag in Li\(\beta\)-BCC at the MIEC/CC interface is approximately 0.4 at.%. The Li\(\beta\)-BCC with Ag atoms would thus be in the solid-solution \(\beta\)-phase regime (\(\beta\)-Li\(\beta\)-Ag) shown in the Li-Ag phase diagram [36, 37] (Figure 5(a)), where \(y\) is greater than 249.

The driving force for Ag diffusion is expected to arise from the decrease in total chemical potential change upon alloying/dealloying of Ag into \(\beta\)-Li\(\beta\)-Ag. The chemical potentials of metallic Li and Ag in the \(\beta\)-Li\(\beta\)-Ag phase regime have not been measured experimentally. They can, however, be estimated via Raoult’s law and Henry’s law in solution thermodynamics. Raoult’s law states that the solvent of a nonideal solution approaches ideal behavior when its concentration becomes much more stable as more and more metallic Ag enters the \(\beta\)-Li\(\beta\)-Ag, while Li atoms in the \(\beta\)-Li\(\beta\)-Ag become only slightly less stable. When the amount of each element is considered, the change in total chemical potential of Ag (\(\Delta\mu_{\text{Ag}}\)) is likely to be at least two times larger than that of Li (\(\sim X_{\text{Li}}\Delta\mu_{\text{Li}}\)). This change would establish a driving force for rapid inward diffusion of metallic Ag into the \(\beta\)-Li\(\beta\)-Ag phase along the porous MIEC and the grain boundaries.

The subsequent question to address is whether the long-range transport of metallic Ag at RT is kinetically

\[\mu_{\text{Li}} = k_{\text{B}}T \ln (1 - X_{\text{Ag}}) = -X_{\text{Ag}}k_{\text{B}}T,\]
\[\mu_{\text{Ag}} = k_{\text{B}}T \ln \gamma_{\text{Ag}} X_{\text{Ag}}.\]
feasible. Chen et al. [20] have shown that Li BCC, which has a low melting point (~453 K), can advance and retract within the 3D open-porous MIEC at RT via interfacial diffusion, at a timescale acceptable for a battery to function. The pure Ag in the FCC phase, however, has a relatively high melting point (~1235 K). Figure 6 shows the deformation mechanism map of Ag FCC [29]. Metals can deform via either a displacive mechanism (e.g., dislocation glide) or a diffusional mechanism. As the former is viable only when high stress exists (a thresholding, nonlinear, behavior), the latter governs the deformation in the low-stress regime, which is what we want for pumping the β-phase. Since the stress greater than 5 MPa is expected to severely degrade the SE life [40], we can assume that the stress in this rechargeable SSB was less than 5 MPa. When the shear modulus of Ag (30 GPa) and its homologous temperature \((T/T_M)\) are considered, \(\frac{\mu}{d}\) would be in the elastic regime of the map.

However, Ag transport may yet be carried out by the interfacial diffusion owing to the nanolevel microstructural length scales of the MIEC interlayer. The \(\dot{\varepsilon}\) that can be achieved via interfacial diffusion is inversely proportional to grain size \((d)\) to the power of 3 as seen from the equation [29]:

\[
\dot{\varepsilon} \equiv \left(\frac{Ca}{d}\right) e^{-Q/k_B T},
\]

where \(C\) is a constant, \(\sigma\) is the applied stress, \(Q\) is the activation energy, \(k_B\) is the Boltzmann’s constant, and \(T\) is the temperature. When the diameter of Ag nanoparticles (60 nm) is considered, the \(\dot{\varepsilon}\) would be on the order of \(10^{-8} \text{ s}^{-1}\). This is an appreciable rate, which pure metals with sub-mm grains in a low-pressure regime can achieve at temperatures near their melting points. Indeed, it has been demonstrated both experimentally and computationally that surface diffusion on sub 10 nm crystalline Ag particles is fast enough to mediate liquid-like pseudoelasticity [41] at room temperature. Its transport at RT can thus be regarded as feasible if nanostructuring of the MIEC interlayer is carried out.

Moreover, the low liquidus/solidus temperatures in the β-Li,Ag phase regime and grain boundary premelting phenomena may also facilitate the interfacial diffusion of both...
Ag and Li atoms. The atomic structure of surfaces, grain boundaries, and other interfaces tend to become increasingly disordered when the temperature approaches the bulk melting point, \( T_m \) [42]. In the case of alloys, the solidus temperature is considered instead of \( T_m \) [42], and that of the $\beta$-Li$_x$Ag phase is lower than the \( T_m \) of Li$_{BCC}$. Given that Ag atoms segregate to the grain boundaries, the effective composition along diffusion paths would be near the eutectic point, which has the lowest melting temperature. This means that it might be even easier to diffuse Ag atoms at this composition along MIEC surfaces/interfaces than Li atoms in pure Li$_{BCC}$. In addition, the atomic size mismatch and the electronegativity difference between Ag and Li may intensify the disordering [36, 42–44] and result in the early premelting. In fact, Ag is known for anomalous diffusion ability in various research fields such as electronic materials [45] and nuclear materials [46]; this ability may have to do with the fact that Ag forms eutectic points with many other metallic elements, making the resultant alloys prone to premelting and interfacial segregation on MIEC surface. This, combined with the possible “argenphilicity” of the current collector versus the “argenphobicity” of the SE side, may explain the gradual asymmetry of the Ag distribution as the MIEC interlayer pumps Li up and down many times with battery cycling [25].

4.3. Suppression of Morphological Instability at MIEC/ $\beta$-Phase Interface. Another promising feature to note in Ref. [25] is that the MIEC/$\beta$-phase interface is free of micro-scale pores even after 100 cycles at 0.5C, which corresponds to a high current density of 3.4 mA cm$^{-2}$. Rechargeable Li metal SSBs have often suffered from a pore formation problem due to the slow diffusion kinetics [23, 47, 48]. Upon applying anodic current, a neutral Li atom in the $\beta$-phase loses one electron and becomes Li$^+$, which then diffuses via the MIEC + SE to the cathode, leaving a vacancy. The difference in the vacancy concentration then drives the diffusion of Li atoms to the interface where the charge-transfer reaction takes place [47]. If the local current density (i) does not exceed the vacancy flux multiplied by Faraday’s constant (\( J_v \cdot F \)), the interface will remain in contact [47]. On the other hand, if \( i \) surpasses \( J_v \cdot F \), excess vacancies will form and coalesce into pores, thereby raising interfacial resistance and making high rate capability impractical [47]. Operating SSBs under stack pressure and elevated temperatures (e.g., 5 MPa and 60°C) has been effective in improving rate capability by inducing both displacive and diffusional motion of Li atoms [48]; nevertheless, it is considered an auxiliary solution due to the limitations in further increasing stack pressure and temperature [40, 48].

On the other hand, $\beta$-phase can inherently introduce an additional driving force for diffusion. Upon stripping, the Li concentration near the interface decreases. If the stripping is limited by the mass transport, it will develop a chemical composition gradient [48]. This gradient can then drive further flux of Li atoms (\( J_{Li} \)), which is expressed as follows in the laboratory frame [49]:

\[
J_{Li} = -D \nabla c_{Li}(x),
\]

where \( \nabla c_{Li}(x) \) is the Li concentration gradient. Here, interdiffusivity \( D = X_{Li} D_{Ag} + X_{Ag} D_{Li} \), where \( X_{Li} \) and \( D_{Li} \) are the mole fraction and the intrinsic diffusivity of the chemical species in the subscript, respectively. It is thus possible that this flux has contributed to resisting the evolution of morphological instabilities at the MIEC/$\beta$-phase along with the flux driven by the vacancy concentration gradient and applied stack pressure. In fact, Krauskopf et al. [48] demonstrated that adding 10 at.% Mg into the Li metal anode can mitigate contact loss at the SE/$\beta$-phase interface during discharging, with the consideration of chemical diffusion coefficients. Moreover, it was shown that Li$_{1.75}$Ca$_{0.25}$Zr$_{1.75}$Nb$_{0.25}$O$_{12}$ garnet exhibits seamless interfacial contact with the Li metal anode when Ag is sputtered on its surface and thus forms Li-Ag alloy upon cycling; the reported interfacial resistance after 150 cycles is also ~7 times lower than the pristine garnet [50]. These findings suggest that additional driving force could indeed help alleviate morphological instabilities to some extent even when solubility in $\beta$-phase is limited. Similarly, the presence of alloying elements and the resultant establishment of concentration gradient may have contributed to suppressing morphological instabilities in this SSB.

Upon the removal of Li atoms, the remaining Ag would either stay dissolved in the $\beta$-phase or form intermetallic compounds depending on the local composition. Assuming mass-transport-limited stripping, the $\beta$-phase will have a concentration profile, where Ag concentration gradually increases and finally reaches the solubility limit (~1 at.% at RT [36, 37]). Interdiffusivity and mechanical properties will vary accordingly. In particular, the interdiffusivity and the mechanical properties of intermetallic compounds can differ greatly from those of the $\beta$-phase [51–53]. Nonetheless, provided that the particles of the intermetallic compounds formed are on the nanometer scale in size without agglomeration into microparticles, stripping of Li from these particles would be achieved at an appreciable rate without serious Li depletion near the interface and the resultant increase in interfacial resistance.

The change in alloy composition during stripping affects the alloy’s interdiffusivity and mechanical properties (e.g., elastic constants and strength), thereby changing Li transport kinetics [54]. However, its impact is expected not to be significant as long as Ag-rich alloy particles that form upon stripping are distributed uniformly in-plane near the CC side of the MIEC and remain at the nanometer scale in size after stripping. If the CC is perfectly argenphile, then the equilibrium wetting angle would be zero and the above can be easily achievable. But if it is argenphobic, dewetting and subsequent coarsening could leave big Ag particles at the CC side after Li stripping, which would be very bad for subsequent cycling from both the kinetics and mechanical robustness viewpoints.

5. Design Rules

With the aid of the mechanistic insights above, the rechargeable SSB with the Ag-C nanocomposite anode [25] has
shown outstanding Coulombic efficiency (over 99.8%) and long cycle life (1,000 times). This full-cell performance sheds light on rechargeable SSB development for use as EV batteries, where a cycle life of ≥5,000 is desirable (a minimum requirement is 800 deep charge-discharge cycles). Here, we discuss various features of an open-porous MIEC interlayer that would help to enhance the cycle life when optimized.

There has not been a detailed analysis of the failure mechanism in this SSB. It is nonetheless expected that the cyclic stress inevitable in the current cell configuration contributes to cell degradation. The maximum thickness of the β-Li,Ag layer is ~25 μm [25], whereas that expected from the areal capacity loading of 6.8 mAh cm⁻² is ~34 μm. This discrepancy implies that the β-Li,Ag phase is accommodated partially in the porous MIEC interlayer. The plasticity of the solidus temperature that is comparable to or lower than the ffi shown outstanding Coulombic efficiencies and their possible effects on stripping behavior should be investigated. With alloying elements that have limited solubility in Li, the interdiffusivity and mechanical properties of the remaining β-phase could change significantly upon Li stripping. For example, if Mg concentration in Li increases beyond 10 at.%, both displacive deformation and diffusional motion would become less effective as the homologous temperature decreases and the elastic constants/strength increase. It has indeed been shown that submicrometer pores form when cycling Li-Mg alloy and that they cannot be filled in by the external pressure of 15 MPa [48]. Thus, such a system is likely to form another MIEC interlayer constructed of solid-solution alloys, which can function as a 3D framework where β-phase resides [23]. This architecture could be beneficial for garnet-SE-based SSBs, where minimal volume change is desired.

Stack pressure may not be required at all if the structure of an open-porous MIEC interlayer with IEs is optimized. As discussed briefly in the previous section, it is generally thought that SSBs need stack pressure to avoid interfacial pores and contact losses caused by a large volume change since their solid components lack fluidity. It has been shown that a low stack pressure of ~5 MPa on SSBs can effectively lower the cell impedance and improve cell performance by increasing the interfacial contact area [40]. Higher pressure is, however, known to cause creeping of Li inside the SE and thereby short-circuiting [40]. Meanwhile, if the β-phase is guided to reside in the preserved porosity of a 3D MIEC interlayer, the degree of a volume change can be greatly reduced. Moreover, if IEs are decorated on the SE/MIEC interface, they can function as a strong mechanical binder that secures physical contact between SE and MIEC. These components could thus contribute to reducing the magnitude of the required stack pressure and hence lowering the chance of mechanically induced short-circuiting.

6. Outlook

Rechargeable alkali metal SSBs with SEs are an attractive candidate for the next-generation energy storage system. Fueled by the intense interests in them, there has been rapid advancement in SE development, but the overall rechargeable alkali metal SSB technology remains immature. The first task was to improve the bulk ionic conductivity of inorganic SEs, which was impractically low in the early years. This task has been accomplished, taking advantage of computational material genomics [15]. The current computational methods have, however, been a help only to a limited extent regarding interfacial problems, such as instabilities caused by swelling-then-retreating of highly corrosive and mechanically stressful β-phase [2]. In these interfacial issues, kinetic factors of which timescale is not achievable using first-principle calculations are involved, in contrast to the ionic conductivity problem [2]. Examples of such factors are the nucleation and growth of new phases at heterogeneous interfaces, the long-range diffusion of different elements [2], and stress relaxation by diffusional creep (or power-law creep, which
is a hybrid displacive-diffusional process that involves an intermediate level of shear stresses) mechanisms. Furthermore, the evolution of instabilities at the interface itself is considered hard to model due to the nature of the interface that can be summarized as high degrees of structural and chemical freedom.

Nonetheless, it has been recently demonstrated that the deployment of various types of solids altogether for the design of anodes can suppress the electrochemomechanical instabilities [20, 25]. Indeed, the resultant performance of some SSBs has been found to be sufficient for use in consumer electronics, and even vehicular applications (e.g., more than 800 deep charge-discharge cycles). First, a 3D open-porous MIEC architecture is used to host β-phase and enable fast stress relaxation [20, 25, 59, 60]. Second, the use of solid IEI (i.e., an ion-electron insulator) has been proposed as a binder between MIEC and SE to prevent the naked SE/MIEC interface from becoming mechanically unstable due to the deposition of soft β-phase [20] prone to be turning into effectively an adhesion crack. Third, solute elements with a certain degree of solubility in LiBCC were included, thereby enabling LiBCC nucleation instead of the MIEC interlayer rather than at the SE/MIEC interface [25], and the asymmetry that develops during battery cycling (attributed to argenphilicity/argenphobicity difference between SE and CC, and mechanical constructions) that reinforce lower $R_{ct}$ and β-phase nucleation barriers on the CC side compared to the SE side. Benefiting from these diverse factors, a long-cycling rechargeable alkali metal SSB has recently been developed [25]. In this cell, β-phase was deposited mostly at the MIEC/CC interface and some inside the porous MIEC, instead of mainly nucleating and growing at the SE/MIEC interface, thereby reducing the pressure on the SE and represing the evolution of electrochemomechanical instabilities inside the SE [25]. The porous MIEC interlayer can therefore be considered as a buffer and an asymmetric diverter that draws the aggression of the β-phase towards the more ductile and thermochemically stable metallic CC, rather than the fragile and thermochemically unstable ceramic SE. The use of electronically insulating, lithiophobic and argenphobic IEI as a binder between SE and MIEC and the use of lithiophilic and argenphobic CC materials provide driving force for this asymmetry.

Careful examinations of the open-porous MIEC interlayer-based anodes with outstanding full-cell performances provide insights for the further development of rechargeable alkali metal SSBs. First, for the minimization of stress that is exerted on the SEs, the 3D MIEC architectures should have open-porous channels, which can serve as a pipe where β-phase resides. Second, the β-phase deposition at the MIEC/CC interface when the inner surface of the MIEC is decorated with metal nanoparticles suggests that the modification of $R_{ct}$ can manipulate where large quantities of β-phase are first deposited that act as coarsening centers and prevent or delay nucleation/growth at the MIEC/SE side. Third, for the optimization of the rate capability, the composition of metal nanoparticles should be modified so that the resulting alloys have low eutectic melting points. This modification would enable constituting elements to diffuse along the solid interfaces at a rate relevant for industrial applications (e.g., EV), matching the expected areal current density. Finally, the use of IEIs at the SE/MIEC interface would improve mechanical stability, and tuning lithiophilicity/argenphilicity gradient would help establish the asymmetry. These design strategies are expected to further enhance SSB performance, especially in terms of cycle life and rate capability. They also provide opportunities for the development of Na or other alkali metal batteries.

The optimal design of such composite anodes is, however, challenging due to their multidimensional material space and complex temperature-pressure-chemistry window viable for composite processing. Even when only 10 candidates for each solid category are considered, the number of possible combinations already reaches 10³. Moreover, as metals with low melting points will be deployed inside an MIEC and the MIEC will have an open-porous 3D structure, there are not many methods available for the decoration of IEIs, which are mostly ceramics that require high temperature and pressure for sintering. The spray pyrolysis method [61] is one of the most promising methods that do not apply high temperature and pressure. However, the precursors adopted for this method often react with other solid components that constitute an interlayer. Therefore, thorough understandings of the mechanisms underlying the synergistic effect of combining different classes of materials (M, SE, MIEC, and IEI) and a broader material approach that includes processing and mechanics considerations should be established to narrow down the candidates for each category of materials. Furthermore, the simultaneous development of experimental processing techniques for such composites should be investigated.

Conflicts of Interest

The authors declare that there is no conflict of interest regarding the publication of this article.

Authors’ Contributions

S.Y.K and J.L conceived and wrote the manuscript.

Acknowledgments

This work is financially supported by the Samsung Advanced Institute of Technology. S.Y.K gratefully acknowledges partial financial support by the Kwanjeong Scholarship.

Supplementary Materials

The activity coefficient of Ag in β-Li$_x$Ag at infinite dilution ($y_{Ag}^{∞}$) was estimated using the CALculation of PHAse Diagrams (CALPHAD) technique in conjunction with Thermo-Calc software. TCAL6 databases for Al-based alloys were used as they include temperature and composition-dependent interaction parameters for the Li-Ag binary pair. $y_{Ag}^{∞}$ at 150 °C is used instead of that at room temperature as the databases were constructed based on the experimental data points [1] obtained at temperatures near or above
150°C [2]. At this temperature, the calculated chemical activity (Figure S1) is in good accordance with the experimental phase diagram (Figure S2) [3] in the Li-rich region ($X_L > 0.8$), which is the region of interest. We assume that $\gamma_{Ag}^{\infty}(T = 150°C) \approx \gamma_{Ag}^{\infty}(T = 25°C)$ as the temperature derivative of $\ln(\gamma_{Ag}^{\infty})$ is equivalent to $-H_{Ag}^{\infty}/RT^2$, [4] which is nearly zero at infinite dilution. Here, $H_{Ag}^{\infty}$ is the excess mixing enthalpy, $R$ is the gas constant, and $T$ is the temperature. $\gamma_{Ag}^{\infty}$ can then be estimated from the slope of activity versus composition relation at infinite dilution; thus, $\gamma_{Ag}^{\infty} = 0.00143$.

Figure S1: activity of Ag versus the bulk concentration of Li for the Li-Ag binary system. Figure S2: experimentally determined phase diagram of the Li-Ag binary system. Reproduced from Ref. [3]. (Supplementary Materials)

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