2020 Roadmap on Solid-State Batteries

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Abstract

Li-ion batteries have revolutionized the portable electronics industry and empowered the electric vehicle (EV) revolution. Unfortunately, the traditional Li-ion chemistry is approaching its physicochemical limit. The demand for higher density (longer range), high power (fast charging) and safer EVs has recently revamped the interest in solid state batteries (SSB). Historically, research has focused on improving the ionic conductivity of solid electrolytes, yet ceramic solids now deliver sufficient ionic conductivity. The barriers lie within the interfaces between the electrolyte and the two electrodes, in the mechanical properties throughout the device, and in processing scalability. In 2017 the Faraday Institution, the UK’s independent institute for electrochemical energy storage research, launched the SOLBAT (solid-state lithium metal anode battery) project aimed at understanding the fundamental science underpinning the problems of SSBs, recognising that the paucity of such understanding is the major barrier to progress. The purpose of this Roadmap is to present an overview of the fundamental challenges that are impeding the development of SSBs, the advances in science and technology necessary to understand the underlying science, and the multidisciplinary approach that the SOLBAT researchers are taking to face these challenges. It is our hope that this Roadmap will guide academia, industry and funding agencies towards the development of the batteries in the future.
Keywords: solid-state batteries, lithium metal, interfaces.
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The solid-state battery (SSB) is arguably the most important challenge in battery research and development today[1]. Advances in SSBs would enable step changes in the safety, driving range, charging time and longevity of electric vehicles (EVs)[2]. In contrast to work on Li-ion batteries, SSB research stands out as long-term and high-risk, but potentially high-reward. Historically, SSB research has focused on improving the ionic conductivity of solid-state electrolytes (SSE). Ceramic solids, including garnet oxides and several sulphides, are now sufficiently conductive and electrolytes are no longer the biggest hurdle facing SSB development[3].

The current barriers are at the electrode-electrolyte interfaces, in the mechanics throughout the cell and in processing at scale (Figure 1).

In response to this diverse set of challenges, the Faraday Institution, the UK’s independent institute for electrochemical energy storage research, launched the SOLBAT (solid-state metal anode battery) project back in the spring of 2017[1]. We have assembled a multidisciplinary team of experimentalists and modelers, with expertise in mechanics, metals, ceramics, polymers, and interfaces from inside and outside the battery field. Our priority is to unravel the fundamental science underpinning the problems of SSBs, recognising that the scarcity of understanding is the major barrier to progress.

Here we report an overview of the fundamental challenges that are impeding the development of SSBs, the advances in science and technology necessary to understand the underlying science, and the multidisciplinary approach that the SOLBAT researchers are taking to face these challenges. The resulting Roadmap can be broadly divided in four areas, as schematically depicted in Figure 1.

We first introduce the challenges at the Li-metal/solid electrolyte interface, starting from the concept of critical current density and its connection to interfacial voids and lithium dendrites, the ultimate cause of failure in SSBs. We then discuss how voids and dendrites can be modelled mechanically: an accurate measurement of the mechanical properties of Li-metal, its wetting behaviour, and its visualisation by electron microscopy are all important factors that contribute to understanding the root causes of their formation. The significance of a holistic electro-chemo-mechanical approach to both modelling and experiments in SSBs is then discussed. The characterization of the electrode-electrolyte interfaces will then be examined. We will discuss possible avenues to tackle the delamination and chemical degradation issues at the cathode/SSE interface, as well as the importance of solid-liquid, solid-polymer interfaces to their implementation. Strategies to synthesize and ameliorate the performance of the leading SSE materials to date (garnet oxides and sulphides) will then be introduced. A clear definition of the relevant key metrics in SSE is crucial as well as a novel approach to materials discovery. Finally, novel avenues for processing and manufacturing SSBs and the importance of x-ray imaging in characterizing their failure mechanisms will be considered.

It is our hope that this Roadmap will help guide academia, industry and funding agencies towards the development of the solid-state batteries of the future.

References

Figure 1: Schematic of the underlying scientific challenges hindering the development of solid-state batteries.

Critical Current Density in Solid-State Batteries

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Status

The critical current density of a battery is commonly defined as the current density above which the battery will short-circuit due to Li dendrite penetration through the ceramic electrolyte, but below which the battery can cycle with long-term stability. The importance of increasing the critical current density of a solid-state battery (SSB) can hardly be overstated, as the current densities achievable today are far below those required to overcome the challenges of modern battery applications, such as fast charging of electric vehicles. Our recent work reveals that there are two separate critical current densities; the critical current on stripping (CCS) and the critical current on plating (CCP) [1, 2]. Li-metal dendrites are observed to initiate and grow when plating the Li-metal anode during charge. The CCP is defined as the current density on plating above which the growth of dendrites initiates. Conversely, stripping Li-metal from the anode during discharge can lead to the formation of voids in the anode, resulting in a concentration of current at the remaining areas of contact. In such cases the local current density can exceed CCP even when the global current density is lower. This can therefore lead to dendrite formation on the subsequent charge (Figure 1). For solid electrolyte materials studied to date, CCS < CCP and so it is in fact the current density on stripping which is the practical limitation to the rate of cycling. Understanding the critical current densities on plating and stripping is vital in the approach to increasing battery power. The two modes of failure, void formation on stripping and dendrite formation on plating, occur by different mechanisms and so different parameters can be changed to mitigate each. Understanding and manipulating the factors which determine these critical current densities will enable SSBs which are able to achieve practically useful rates on both charge and discharge.

Current and Future Challenges

The critical current density on stripping is dependent primarily on mass transport toward and away from the interface with the solid electrolyte. For a morphologically stable interface to be maintained during cycling, the rate of Li diffusion/deformation to the interface must be greater than or equal to the rate of Li-ion transport away from the interface under the current load [1–4]. As such, the critical current density on stripping is dependent on two factors; the current density of discharge (i.e. the flux of Li-ions from the interface) and the rheological properties of the Li-metal anode (i.e. the rate of Li-metal transport to the interface). Therefore, if we desire to achieve a particular critical current density, we must choose conditions for the cell under which the Li-metal will diffuse/deform at a sufficient rate. The current challenge involves determining conditions that remain practically achievable for a commercial cell whilst enabling high current densities.

Plating Critical Current Density

Whilst the causes of failure on stripping are fairly well understood, the causes of dendrite penetration on plating are less so. There is no consensus in the literature on a mechanism by which low yield strength Li-metal could cause dendrite penetration through ceramics with high fracture toughnesses, although several theories based on stress-corrosion cracking [5] and pressure build-up [6] have been proposed. Therefore, the challenge to increasing plating critical current densities is to reach a fundamental understanding of how dendrite penetration occurs, so that the problem of dendrite penetration at high rates of charge can be overcome.
Advances in Science and Technology to Meet Challenges

Stripping Critical Current Density

As noted above, the stripping rather than the plating current density is the factor limiting the maximum rate of charge and discharge. To increase stripping critical current densities, the rate of mass transport of Li-metal to the interface must be increased. There are three possible modes of mass transport of Li-metal: self-diffusion, creep and plastic deformation. Recent work has targeted increasing the rates of self-diffusion and creep to enable higher current densities. In recent work by Janek and co-authors, the maximum current density achievable in a Li/Li$_7$La$_3$Zr$_2$O$_{12}$(LLZO)/Li symmetric cell under no external pressure was determined to be 0.1 mA cm$^{-2}$, meaning that self-diffusion alone was not able to transport Li to the interface at a sufficient rate to support higher power densities [7]. This result underlines the importance of pressure driven deformation in achieving targeted current densities of upwards of 5 mA cm$^{-2}$ [8, 9]. It is therefore clear that to achieve higher critical current densities, solid-state batteries will require the application of stack-pressure. Under pressures of a few megapascals, lithium deforms by creep, which is rate dependant. Therefore, to enable higher CCSs, higher pressures are required to deform Li-metal to the interface at a high enough rate to prevent the formation of voids [1, 4, 7]. One approach taken to increase CCSs has been alloying Li-metal with 10% Mg. The alloyed anode has higher rates of self-diffusion of Li and therefore CCSs are found to be higher in cells under no pressure. However, whilst alloying increases the rate of self-diffusion, it has no positive impact on the rate of creep of the metal. It was therefore found that Li-Mg anodes had no impact on cells under pressure [3]. An alternative approach is to switch to Na metal anodes rather than Li, as the rates of both self-diffusion and creep are higher, despite the sacrifice in anodic energy density. The higher rate of creep in Na metal enables higher CCSs. In a Na/Na$_x$β"'-alumina/Na cell, pressures of > 9 MPa enabled morphologically stable stripping at the high current density of 2.5 mA cm$^{-2}$[2]. Going forward, to push CCS towards the ultimate limit of CCP we may need to provide conditions such that the metal anode is not under rate-dependent creep, but under

Figure 1: Figure illustrating the failure of solid-state cells at current densities above the critical current on stripping. Voids form in the metal anode (grey) on stripping, leading to high local current densities and eventual dendrite penetration through the Li$_6$PS$_5$Cl ceramic (orange) on plating. Reprinted with permission from [1]. Copyright Nature Materials 2019.
rate-dependent plastic deformation. To achieve this may still require higher pressures or even operating SSBs under higher temperatures. Is this feasible in practice?

**Plating Critical Current Density**

As the relative mechanical properties of Li-metal and ceramic electrolytes lie at the heart of dendrite formation, an important avenue of research is to better understand the mechanical properties of solid electrolytes and Li-metal. A recent report that suggests micron-scale Li-metal has significantly higher yield strengths than the bulk metal, if confirmed, could have important implications for understanding interfacial mechanics under the operating conditions of a cell [10]. Many approaches to increasing plating critical current densities have focused on improving the fracture toughness of solid electrolytes to inhibit cracking and dendrite penetration. One approach is the use of a composite electrolyte in which a structural polymer is introduced to increase the fracture toughness and improve other mechanical properties of the electrolyte and so prevent electrolyte cracking. An example is shown in Figure 2 [11]. The alternative approach for increasing CCP by decreasing the yield strength of Li has also been explored, with reports that operating cells at elevated temperatures can significantly improve current densities [12]. Taken to the extreme, the use of liquid sodium anodes in Na-S and ZEBRA batteries can enable current densities > 1 A cm⁻².

**Concluding Remarks**

Increasing critical current densities is important if we are to achieve power densities of solid-state batteries that are competitive with Li-ion cells. While progress has been made on understanding some of the factors limiting critical current densities, notably the critical stripping current, the challenge of understanding dendrite penetration into solid electrolytes remains. Improved fundamental understanding of the ceramic/lithium interface is needed to enable solid-state batteries capable of sufficient current densities for commercialisation.

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**References**

Figure 2: Experimental results showing that increasing the fracture toughness of a solid electrolyte increases critical current density on plating. Schematic and SEM images detailing the preparation of structured ceramics are shown in (a). Hybrid ceramics show a greater fracture toughness (b) and so do not fail at a current density of 0.7 mA cm\(^{-2}\) whereas non-hybrid ceramics mechanically fracture at this current density. Figure reprinted with permission from [11]. Copyright Energy Environmental Science.


Mechanical Modelling of Dendrite and Void Formation

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Status

Ceramic electrolytes have potential in solid-state batteries (SSBs). When combined with Lithium (Li) anodes they can deliver enhanced safety and higher energy densities compared to liquid electrolyte Li-ion batteries. However, the charging of such cells at current densities greater than a critical value leads to Li-filled fissures, commonly termed as “dendrites”. The dendrites nucleate and grow from this Li-metal electrode across the electrolyte, and thereby short-circuit the cell. Dendrites can adopt a range of morphologies from a 3D ‘mossy’ form that is thought to originate from the filling of interconnected porosity, to planar fingers that involves fracture of the ceramic electrolyte. Characteristic features of this failure mechanism are now established through the recent work of Bruce and co-workers\cite{1}, and Sakamoto and co-workers\cite{2}. For example, (i) the critical current required to short the cell increases with decreasing resistance to the flux of Li$^+$ ions across the electrolyte/Li-metal electrode interface, and (ii) continued charging/discharging of the cell results in the formation of voids in the Li-metal at the interface with the electrolyte. Dendrites initiate and grow in the vicinity of the voids, see Figure 1. The application of an external pressure shrinks the voids (by diffusional flow and power law creep) and thereby elevates the critical current for dendrite formation within the electrolyte. A mechanistic understanding of these observations for both dendrite growth and void growth remain elusive.

Current and Future Challenges

Robust and rigorous models can provide insights into mitigation against failure modes in solid electrolyte cells such as dendrite growth from voided-interfaces. However, such models are not yet available. A number of approximate calculations have been performed to predict the growth of a pre-existing dendrite. The usual assumption is that the dendrite behaves as a Li-filled crack, with crack advance driven mechanically by pressurised Li within the crack. This approach is problematic, however, as the magnitude of the pressure required to attain the fracture toughness of the electrolyte is sufficiently high that the Li will instead leak into the soft Li electrode, thereby relieving this pressure. Moreover, there is insufficient electrical energy available to provide the elastic energy in the electrolyte that is associated with the pressurized crack. Recently, it has been suggested that dendrites grow as parallel sided dislocation-like features as this mode does not entail pressurisation of the dendrite at its mouth, and requires much less elastic energy storage within the stiff electrolyte. However, these calculations suggest that such dendrites are unlikely to grow without the electrical field concentration as generated by the presence of voids in the Li along the electrode/electrolyte interface \cite{3}. There thus exists a major challenge to understand the reasons why voids develop within a Li electrode. Preliminary models suggest that void growth occurs during stripping of Li from the electrode and is associated with a high concentration of Li flux from the electrode into the electrolyte at the periphery of the void. However, the fundamental mechanism for this flux concentration needs to be understood, along the following lines. The product of ionic interface resistance $Z$ and ionic conductivity $\kappa$ within the electrolyte defines a characteristic material length scale, and the degree to which Li flux is concentrated at the periphery of the void increases with the ratio of void dimension to this length scale $Z\kappa$. Typically, $Z\kappa\approx20\,\mu m$ and it is unclear why small voids (smaller than say 100 $\mu m$) are able to concentrate the Li flux and thereby induce void growth. While there exist useful data and
observations on macro-scale phenomena in such cells, model validation and mechanistic understanding also requires high resolution observations. For example, there is a challenge to observe the initiation and growth of a dendrite in order to give insights into the mechanism of dendrite formation. The thickness of dendrites is on the order of 20 nm, and so high-resolution methods are needed. Such observations will help resolve whether dendrites grow with a crack-like opening or in a dislocation mode with parallel-sided flanks. Again, while there is a well-documented link between the measured value of interface ionic resistance $Z$ (between Li electrode and the adjacent ceramic electrolyte) and the critical current density for dendrite formation, these $Z$ values are averages over the entire electrode/electrolyte interfaces. It is clear that large variations in the flux along the interface can trigger void growth but there is little information on the spatial distribution of $Z$ along the interface. Do variations in $Z$ along the interface explain the void growth observations?

**Advances in Science and Technology to Meet Challenges**

There is a clear need for increasing the resolution of x-ray computed tomography (XCT) methods and related microscopy in order to resolve dendrites, and to follow their growth. The availability of such data for liquid electrolytes spurred significant advances in understanding and model development. In parallel there is an urgent need for the development of theoretical frameworks for modelling the processes within solid-state cells. In such cells there is a strong coupling between mechanical loading (e.g. elastic straining of the electrolyte due to dendrite formation or power-law creep of the Li-metal electrode) and the associated electrochemical processes. These couplings often make inappropriate a number of the commonly used assumptions in the theory of electrochemical systems. For example, the Butler-Volmer equation is commonly expressed in terms of current density as a function of voltage jump across an interface. More accurately, the current density is a function of jump in electrochemical potential across an interface, and this jump in potential is related to jumps in stress state, strain state, vacancy content and so on, in addition to the jump in voltage. The development of appropriate new modelling approaches are expected to shed light into some of the puzzles alluded to above.

**Concluding Remarks**

SSBs offer significant benefits in terms of energy density and power density, but this can only be achieved when potential failure modes are eliminated. There is a need to develop quantitative models at the meso-scale that are consistent with the governing
field equations. Such models require experimental validation but have the potential to vector material developments by explaining the relationship between material properties and failure mechanism. The problems are challenging and require a grounding in both electrochemistry and solid mechanics.

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References

Mechanical Properties of Metallic Lithium

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Status

The leap forward in energy density allowed by solid-state batteries (SSBs) is thanks to the use of a metallic anode significantly increasing the number of possible charge carriers [1]. This brings new challenges as the historic lack of any structural application means the mechanical properties of alkali metals are under investigated. Recent work on metal anode systems for SSBs illustrated that applying an external pressure to cells has a beneficial effect [2, 3]. This is seen to reduce the formation of pernicious “dendrites”. The cause of “dendrites” is currently an unexplained phenomenon where the “soft” lithium penetrates the “hard” ceramic electrolyte [4]. These issues have sparked recent interest to fully mechanically characterise metallic lithium. This has led to a number of recent papers looking to measure the tensile, compressive and time-dependent mechanical properties of lithium with a spread of results [5–7]. However, this field is still in its infancy and requires more work as developing a full understanding of the mechanisms behind the mechanical properties of lithium metal are a missing keystone of the new battery revolution.

Current and Future Challenges

Sample Preparation

Li-metal is air sensitive, readily forming compounds with oxygen, water, carbon dioxide and nitrogen. This means the material must be kept in an atmosphere exempt from these common gases and a chemical or mechanical method is required to remove any residual surface layers to create the smooth surface required for most characterisation techniques. The low hardness of Li-metal means mechanical polishing via abrasion aiming to remove material causes significant plastic deformation and geometric change to the surface. Chemical cleaning for standard metals utilises acids to etch which would cause a violent reaction with lithium. The final alternative for surface preparation is through the use of an ion beam, however, this can lead to local heating/melting and suspected implantation of these ions into the material. There are concerns that legacy work on lithium (especially where handling of samples was not reported) maybe affected by the formation of passivation layers, leading to uncertainties in this data.

Experimental Testing

Standard mechanical testing requires high precision test rigs which combat compliance, thermal drift and noise in displacement readings by being large (~2m tall). For air sensitive materials smaller test set-ups hosted within gloveboxes are required, leading to compromise in the reliability of data. Cutting or forming lithium test specimens can introduce large numbers of dislocations which in turn lead to work hardening. The low yield stress relative to the Young’s modulus means that only strain between 0.0025% and 0.01% (dependent on orientation) is needed to lead to plasticity. These issues combine to make measuring a “yield stress” in pure lithium difficult. The orientation dependence comes from the anisotropy of elastic moduli in the bcc crystal structure of Li-metal. This is significant in lithium due to the low melting point leading to large grain microstructures in room temperature samples. The effect of texture is the suspected root cause of the variability in reported elastic properties [5, 6].

Understanding

The effect of low stress plasticity and the ease of plastic flow needs to be understood for localisation of plasticity relative to the application of force. It is
Figure 1: Challenges: Li-metal reacts to form a bi-product with atmospheric gases, understanding the localisation of plasticity the role of work hardening and the anisotropy of the elastic modulus.

possible that the weak material at the surface will accommodate the applied stress through plasticity leading the bulk of the material to experience little to no stress. Additionally, the low melting point of lithium means room temperature ($\sim0.6T_m$) creep is likely, causing time dependent plasticity. How the creep mechanism is affected by strain rate, temperature and crystallography is still unknown with little mechanistic understanding reported. All this work is geared toward investigating Li-metal in isolation, but the true impact will come when observing the role of mechanical properties in a working cell which will bring more challenges experimentally and cognitively.

Advances in Science and Technology to Meet Challenges

The ubiquity of gloveboxes of all shapes and sizes has allowed for an increased number of table top mechanical testing techniques to be utilised on Li-metal. These small scale tensile and compression tests combined with optical monitoring has given some enhanced insight into how lithium acts under stress [5],[6]. This increased the amount of mechanical data on Li-metal but lacks some information on microstructure to allow for comparison between literature due to the anisotropy discussed above. Going to smaller scale test using nanoindentation and pillar compression allows for characterisation of pseudo single crystals avoiding the
effect of any preferential texture on recorded data [7]. The downside of these tests is in controlling the volume of material tested as it is unconstrained leading to some questions around the plasticity being left unanswered. The use of gallium and argon plasma focused ion beams (FIBs) in future could allow for testing of known volumes of material through cantilever or tensile tests on the microscopic scale. However, there are questions about the effect of such ions beams on materials physical and electronic properties. This requires the joint technologies of cutting-edge microscopy/lithography and the ability to do such work at cryogenic temperatures to avoid the local melting referred to when discussing ion surface preparation. Further advancements could be made to conduct tests in situ with Scanning Electron Microscopy (SEM), Transmission Electron Microscopy (TEM) and x-ray diffraction XRD to characterise crystallographically the plastic changes in real time helping to understand the stress/strain evolution. Critical in this understanding is the interplay between dislocation nucleation and dislocation motion, the latter having huge consequences for the time-dependent plasticity referred to as creep. Comprehensive mechanical test data will help to unpick the fundamental character of Li-metal under stress allowing for great advances in modelling of the materials and complement the wealth of knowledge on the other aspects of Li-metal. The pinnacle of understanding could be reached by realising in-operando tests allowing direct observation of how the mechanical properties of lithium change as it acts in battery cells.

Concluding Remarks

Advancement in lithium mechanical property characterisation has not happened in an intellectual vacuum, it has come as a necessity. As such quick advancements were made in terms of working with air-sensitive materials from the current knowledge of the battery community but the next step change will be driven by the application of metallurgists and material scientists who have the language and tools to understand the complex elastic and plastic nature of Li-metal. The understanding and application of the deformation modes of Li-metal will help modellers, designers and builders of high energy density battery technology long into the future.

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References

Wetting Behaviour of Li-metal

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Status

In current commercial designs of Li-ion batteries, the graphite anodes are almost at the limit of their theoretical specific capacity (372 mA h g\(^{-1}\)) [1]. Li-metal anodes (LMAs) are considered the most promising alternative in future cells because of their high theoretical specific capacity (3860 mA g\(^{-1}\)), low electrochemical redox potential (−3.04 V vs the standard hydrogen electrode), and low density (0.53 g cm\(^{-3}\))[2]. A major hurdle to the successful commercial application of LMAs is the formation of needle-like Li dendrites during charging/discharging cycles. Once the Li dendrites detach from the bulk LMA, they become ‘dead Li’, i.e., they are no longer electrochemically active, reducing the Coulombic efficiency of the cell [3]. More crucially, the sharp dendrites can penetrate the separator, creating a short circuit with associated overheating and even an explosion risk [4]. Solid electrolytes (SEs) have been shown to mechanically suppress the dendrite growth [5] and are more inert toward metallic Li than their liquid counterparts. However, dendrites are still observed to grow from grain boundaries and other interfacial defects in SEs [6]. Another issue that restricts the application of SEs is their high interfacial resistance, and therefore a high pressure is required to maintain close contact with the electrodes. Therefore, an improved understanding of the binding or wetting between the LMA and SE is also essential to developing lower-resistance interfaces. Two possible scenarios are schematically shown in Figure 1, which also includes the copper (Cu) current collector at the anode [7]. Ideally, we would like complete wetting to occur between both SE/Li and Li/Cu (Figure 1a), to maximise the electrical contact. However, in practice partial wetting is likely, where Li grows into three-dimensional (3D) islands between the SE and Cu (Figure 1b). The aim of Li wetting studies is to find processing parameters that will allow the current collector, the LMA, the SE, and any interphases between these materials to have sufficiently low interfacial energies to enable the formation of flat interfaces to be thermodynamically favourable. This will hopefully act as a way of preventing Li dendrite formation and increasing the ionic conductivity between the anode and the SE. The strategy is to investigate all these interfaces individually by forming them under ultrahigh vacuum (UHV) conditions and to extract interfacial energies from these studies.

Current and Future Challenges

The degree of Li wetting is determined by the interfacial energies between the three materials, i.e., \(\gamma_{SE/Li}\), \(\gamma_{SE/Cu}\), and \(\gamma_{Li/Cu}\), as labelled in red in Figure 1c. At the triple junction, the three interfacial energies are related by geometry at equilibrium:

\[
\gamma_{SE/Cu} = \gamma_{SE/Li} \cdot \cos \theta_1 + \gamma_{Li/Cu} \cdot \cos \theta_2
\]

Therefore, to achieve complete wetting (Figure 1a), the \(\theta\) angles have to be zero and the following relationship needs to be satisfied:

\[
\gamma_{SE/Cu} \geq \gamma_{SE/Li} + \gamma_{Li/Cu}
\]

The value of \(\gamma_{SE/Li}\) can be worked out by depositing Li on the SE material and observing the behaviour of Li island growth. The other two interfacial energies, \(\gamma_{SE/Cu}\) and \(\gamma_{Li/Cu}\), can be found in a similar way, by depositing the relevant materials on top of each other. However, this type of physical vapour deposition (PVD) study of Li is rare [8]. Although Li wetting on ceramics has not been studied before, there are many reports on the interaction between other metals and ceramic substrates. For a single crystal ceramic substrate, the main mechanism by which it influences the morphology of the supported...
metal islands is via their interfacial energy ($\gamma_i$) and the substrate surface energy ($\gamma_s$). For example, Figure 2 shows five different degrees of wetting of a supported platinum (Pt) crystal, from no wetting (first crystal) to partial wetting (middle three) to complete wetting (last crystal) [9]. Higher values of $\gamma_s$, $\gamma_{Pt}$ (surface energy per unit area of Pt) and lower values of $\gamma_i$ encourage the crystal to wet. More interestingly, two-dimensional (2D) wetted islands of gold were also reported to coexist with gold crystals on various oxide substrates. For example, they are stabilised by the $(2 \times 1)$-reconstructed (001) surface of strontium titanate (SrTiO$_3$) [10]. Based on the above, for LMAs, the research question is to develop a fundamental understanding of the interface between Li and the electrolyte ceramic material. This will also include a study of the chemically distinct interphases that form at their interface.

**Advances in Science and Technology to Meet Challenges**

The study of the SE/Li interface requires a combination of characterisation techniques. As an example, Li can be deposited by PVD in UHV onto a ceramic substrate. As a starting point, a model ceramic oxide can be used, whose surface structure must be well known, e.g., Nb-doped single crystals of SrTiO$_3$ [10]. The wetting behaviour of Li can then be established at the atomic and microstructural length scales, by scanning tunnelling microscopy (STM) and scanning electron microscopy (SEM), respectively. The reactivity of Li with the substrate can be investigated using X-ray photoelectron spectroscopy (XPS), which provides information on the chemical environment of the buried Li-metal at the interface with the ceramic. STM results obtained from a similar system, gold on SrTiO$_3$, clearly illustrate the morphologies of both 2D and 3D metallic islands [10]. The shapes of
3D nanocrystals can be measured accurately and used to calculate the interfacial energy $\gamma_{\text{SrTiO}_3/\text{Li}}$, according to the Winterbottom construction (Figure 2). In Ref. [10], a square pattern of spots was also obtained on 2D gold islands under STM, which possibly results from the frustrated commensurate epitaxy between gold and the substrate. These all demonstrate the potential of STM in investigating the SE/Li system. Similarly, STM characterisation of Cu deposited on SrTiO$_3$ and Li deposited on Cu will provide values of $\gamma_{\text{SrTiO}_3/\text{Cu}}$ and $\gamma_{\text{Li/Cu}}$, which together will establish the Li wetting scenario in the SrTiO$_3$/Li/Cu system. Once the model experiments have been performed, the Li interface with technologically pertinent solid electrolyte and cathode materials can be studied. These include Li$_7$La$_3$Zr$_2$O$_{12}$ (LLZO), LiMn$_2$O$_4$ (LMO), LiCoO$_2$ (LCO), LiNi$_{0.8}$Co$_{0.15}$Al$_{0.05}$O$_2$ (NCA), etc. An additional challenge is that for these insulating oxides to be characterised by STM, they need to be prepared in the ultra-thin-films form as epitaxial overlayers on a conducting substrate, e.g., on Au(111). This can be achieved either through UHV evaporation and oxidation of the elemental materials or via pulsed laser deposition of the target oxide. Again, interfacial energies ($\gamma_{\text{SE/Li}}$, $\gamma_{\text{SE/Cu}}$, $\gamma_{\text{Li/Cu}}$) can be obtained to study Li wetting. Special attention should also be paid to the effects of the ceramic surface defects on the Li binding, which play a role in Li dendrite propagation.

Concluding Remarks

Ultimately, the techniques described above (STM, SEM, and XPS) will provide valuable insights into the fundamental processes taking place at the Li/electrolyte interface. In particular, the interfacial energies involved in the SE/Li/Cu setup will allow us to work out the degree of Li wetting and, in turn, identify SE materials with optimised electrode/electrolyte binding. In addition, study of the SE/Li interfacial processes can shed light on the origin and propagation of Li dendrites. We can then derive novel strategies to create more resilient ion-conductive ceramics for the best possible performance of a solid-state battery. The successful implementation of LMAs with SEs is a critical step for electric vehicle improvement, and will result in safer cars with batteries superior in performance to current Li-ion batteries.

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References


Electron Microscopy: Imaging Li-metal

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Status

Lithium (Li) is the key element in Li-ion batteries (LIBs) and is thus contained in the cathodes and (solid/liquid) electrolyte and there is significant potential for Li-metal to be used as high-capacity anode for next-generation batteries. Techniques to image Li in battery materials therefore form an important part of the portfolio of battery materials characterisation methods. In particular, imaging with high spatial resolution will allow a deeper understanding of Li function in battery materials. Here we discuss techniques of electron microscopy that are being developed and applied to study these materials. Early work applied transmission electron microscopy (TEM) to the study of the motion and interaction of dislocations in Li-metal using the dark-field mode [1]. More recently, imaging has been used to understand the mechanical interactions at the electrode/electrolyte interface in a battery [2]. Imaging of Li can give insight into the electrochemical reaction, leading to greater understanding of the degradation mechanisms. In charged cathode materials, transition metal movement into the Li locations results in cation disorder degradation of the cathode [3]. Imaging of Li and transition metals in cathode materials reveals the new structures formed and the defects that are responsible for the reduced reversibility. At the anode interface, Li dendrites grow during cycling and lead to safety issues. Scanning electron microscope (SEM) imaging of the dendrites has helped to understand the formation mechanism at different plating/stripping conditions [4] and validates the strategies to mitigate dendrite growth. Solid electrolytes (SEs) contain disordered structures and vacancies to ensure Li conduction. Scanning TEM (STEM) imaging of Li in SE revealed the atomic arrangement of the existing Li-rich and Li-poor phases and vacancy clusters [5]. Li aggregation close to the vacancies was suggested to enhance the crystal distortion and affect the Li-ion migration pathways [5]. The solid electrolyte interphase (SEI) is a chemically-formed passivating film between the electrolytes and electrode, critical to the reversible battery operation. Imaging of Li in SEI has atomically resolved the Li arrangement at the bottom of SEI layer and Li-based nanocrystals in the film. The atomic structures of both the nanocrystals and Li dendrite nanowires on the layer illustrate tuneable SEI configurations arising from the change of electrolytes [6].

Current and Future Challenges

Direct imaging of the chemically reactive and beam sensitive Li-metal is challenging. Li is a light element so the high-energy electrons used in TEM imaging are scattered weakly and the primary effect is a small phase shift of the transmitted electron wave. The high-angle annular dark field (HAADF) images commonly used in STEM for atomic resolution imaging give contrast approximately as the square of $Z$, and so are insensitive to light elements, such as Li. Furthermore, the low contrast of Li tends to be unobservable when in proximity to the heavier transition metals because the strong signal of heavy elements may swamp the signal from light elements. Coherent bright field (BF) STEM images and high-resolution TEM (HRTEM) images provide phase contrast of both light and heavy elements but have restricted requirement on the specimen thickness. Annular bright field (ABF) STEM images make use of the annular detector located in the BF region and can simultaneously visualize both light and heavy elements over a wide range of specimen thickness. ABF images show a combination of weak $Z$-contrast and phase contrast imaging; however, the ABF setup requires well-aligned microscope optics and is challenging to use for quantitative measurements. Damage of Li is due to the high-energy electrons and radiation...
sensitivity of samples. The main damage mechanisms include knock-on, radiolysis and possible sample heating. Metallic bonded Li-metal is rich in free electrons and radiolysis can be quenched quickly [8]. The low mass of Li and low melting point of Li-metal make Li-metal particularly susceptible to the damage from electron sputtering (knock-on) and heating. Beam damage of Li-metal reduces the achievable resolution in imaging and reduces the detectability of Li using spectroscopy. Electron energy loss spectroscopy (EELS) is challenging because of the small inelastic cross section (low signal) and low energy of Li K-edge. To increase the signal-to-noise ratio in EELS, higher electron dose is needed resulting in higher damage. The Li K-edge is close to the plasmon region in the EELS spectrum, which can mask the Li K-edge. In thick specimens multiple scattering is prevalent and will obscure the edge further [8]. Li also has a low x-ray yield, resulting in poor detection efficiency for energy-dispersive x-ray spectroscopy (EDX). Characterizing x-rays of Li and mapping Li distribution can be challenging.

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Although imaging of Li has been proven challenging, recent advances in electron microscope design, specimen-transfer holders and imaging methods are now enabling reliable characterization of Li-metal. The radiation damage of the sample is highly dependent on the beam energy and optimal selection of the accelerating voltage in the microscope can mitigate beam effects. Modern (S)TEM instruments can be operated over a wide range of accelerating voltages and voltages below 100kV can reduce the knock-on damage on Li [6]. Li and most Li-containing battery materials are air-sensitive and to avoid air contamination air-free sample transfer to the microscope is needed. Vacuum-transfer holders are now available to mount the specimen and transfer it from an argon-flowing glove box to the microscope. Low electron-dose experimental conditions are necessary to lower the radiation damage. However, imaging at high-spatial resolution requires a relatively high electron energy and high dose rate, inevitably leading to increased sputtering and heating damage. Because of this, cooling the sample using a cryo-transfer holder has been proven helpful to reduce beam damage of Li and preserve atomic resolution (Figure 1) [7, 8]. It has also been suggested that the ice layer during cryogenic processing may protect the Li sample from air. Li K-edge EELS and mapping have also been used to analyse the composition of Li dendrites and electrochemically-deposited Li-metal in cryo-STEM [9]. Electron ptychography is a method to reconstruct the phase information from the complex specimen exit wave function and has been shown to be a dose-efficient method. This technique makes use of direct electron detectors that have a very high detection efficiency. The high-efficiency phase reconstruction by ptychography allows to reduce the
beam currents down to the sub-picoampere range, along with post-acquisition correction of residual aberrations. Ptychographic phase reconstruction of the charged Li-rich cathode has demonstrated sensitivity to heavy and light elements simultaneously with minimum beam damage [10] (Figure 2).

Concluding Remarks

High spatial-resolution imaging of Li-metal is of increasing importance battery function and degradation, for example through the potential of Li-metal as a high-energy battery anode despite the fact that the problems of dendrite growth and low coulombic efficiency are still unresolved. Imaging and spectroscopy of Li-metal are critical to the understanding of its physicochemical properties and the interface that forms with the electrolyte. Recent progress in direct electron detectors and data processing has highly improved the dose efficiency and lowered the beam damage in imaging. This is a rapidly developing area and forthcoming powerful detectors can create opportunities for further imaging and spectroscopy of Li-metal. A growing number of new capabilities (e.g. cryo-TEM) and low-dose imaging methods are being progressed for biological materials, the ideas of which can be translated into the study of Li-metal but need trials to validate their applicability to battery materials. Results of imaging and spectroscopy of Li-metal will provide new information about Li-metal in the batteries and future characterization will highly assist in the fundamental study of Li-metal chemistry.

Acknowledgements

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References

3. Mukherjee, P. et al. Surface Structural and Chemical Evolution of Layered LiNi_{0.8}Co_{0.15}Al_{0.05}O_2 (NCA) under High Voltage and Elevated Temperature Conditions. Chemistry of Materials 30, 8431–8445 (2018).
Figure 2: ABF-STEM micrographs (left) and ptychography reconstructed phase image (right) of a charged Li$_{1.2}$Mn$_{0.6}$Ni$_{0.2}$O$_2$ particle along the [010] direction taken with beam currents of (a) 6, (b) 2, and (c) 0.4 pA. Grayscale: (a) 0 to 0.98, (b) 0 to 1.48, and (c) 0 to 1.70 rad. Reproduced from ref[10]. with permission from American Chemical Society, Copyright (2018).
Status

The development of commercial electric vehicle requires safer power batteries that achieve a specific energy of 235 Wh kg\(^{-1}\) and energy density of 500 Wh l\(^{-1}\) at cell level, with a reduction of pack cost to $125/kWh [1]. Solid-state batteries using solid electrolytes are a next-generation system that has been proposed to meet these requirements. Early research on solid electrolytes originated more than 40 years ago, with studies focused on the application of beta-alumina as a sodium-ion conductor and on space-charge models to rationalize Donnan potentials in ionically conductive single-ion conductors [2]. Inorganic solid electrolytes with sufficiently high room-temperature conductivity for lithium-ion battery applications have only been widely available for the past decade [3, 4]. It is widely believed that these solid electrolytes could naturally mitigate many problems that place limitations on today’s liquid-electrolyte lithium-ion batteries [5]. Solid electrolytes are generally nontoxic and not flammable, properties that would improve battery safety significantly. Solid ion conductors have exhibited a much wider viable range of working temperatures, and do not freeze at low temperatures or vaporize at high temperatures. Generally, solid electrolytes are single-ion conductors, which eliminates the voltage losses that occur due to concentration polarization when liquid electrolytes are operated at high power. The most popular examples are ceramics or ceramic glasses, either oxides or sulphides. Because they do not suffer from concentration polarization in the electrolyte domain, solid-state batteries can in principle include much thicker composite electrodes, which might enhance their energy density. Moreover, the huge stiffness of solid oxide or sulphide separator materials can suppress nucleation of dendrites due to interfacial morphological instability at high currents. This stiffness advantage potentially enables lithium-metal anodes. Solid electrolytes can block degradation mechanisms which occur due to interdiffusion of species between the electrodes of a battery: solid oxides may enable lithium/oxygen technology by preventing oxygen crossover to the lithium anode, and solid sulphides may resolve the issue of polysulphide shuttling in lithium/sulphur systems. In liquid-based batteries, the electro-chemo-mechanics of electrode materials has received substantial attention. It is already known that volume change during lithium intercalation causes particle or SEI cracking, both regarded as crucial sources of degradation in today’s batteries. The growing interest in solid-state batteries has further fuelled the community’s interest in solid-state electro-chemo-mechanics.

Current and Future Challenges

Stress accumulates in solid electrolytes. Interactions among electrical, chemical, and mechanical phenomena, especially near solid/solid interfaces, account for most of the major barriers to viable solid-state batteries (Figure 1). Dendrite formation at the metal/electrolyte interface limits power density [6]. Mechanical failure – delamination, cracking, etc. – at the cathode-particle/electrolyte interface leads to active-material loss and consequent capacity fade [7].

Stiff solid electrolytes support stable cycling performance when currents are sufficiently low, while dendrites have been observed to form when cells are cycled above a ‘critical current’. The mechanism of dendrite formation in ceramic electrolytes is still unclear, since solid electrolytes suppress both carrier polarization and morphological instability, which are the key reasons for dendrite nucleation in liquids [8]. Early experiments on sodium beta-alumina
Figure 1: (a) Lithium filament formed in Li$_7$La$_3$Zr$_2$O$_{12}$ garnet and its microstructure [6]. Reprinted from Electrochimica Acta, 223, Cheng et al., 85–91, Copyright (2017), with permission from Elsevier. (b) Contact loss in NCM-LiPS composite electrodes and (c) details at the NCM-LiPS interface [7]. Reprinted with permission from Chemistry of Materials, 29 (13), 5574–5582, Koerver et al., Copyright (2017), American Chemical Society.

suggested that there are two modes of dendrite growth in solids: crack propagation from the edge and bulk plating related to electron conduction [2]. Both modes have been observed in solid lithium-ion conductors. Griffith’s cracking model implies that the propagation of pre-existing microcracks at the edge of an electrolyte will unavoidably lead to electrolyte failure [9]. Thus research should focus on designing systems that impede dendrite nucleation altogether.

Cavities can form at the metal/solid electrolyte interface when lithium is being stripped, a problem that highlights the importance of transport in the metal, as well as in the electrolyte. The loss of interfacial contact during stripping decreases the critical current in subsequent plating steps. Application of a uniaxial stress to the electrode stack can slow cavity formation, perhaps by speeding up lithium diffusion or flow within the metal. Mechanical properties of lithium – especially those describing creep and plastic deformation at the nanoscale – are sparsely measured, a factor that has impeded the theoretical analysis of stripping critical currents at the metal/solid electrolyte interface.

Volume expansion in intercalation materials is probably the most significant barrier to solid-state batteries. Solid electrolytes are generally stiff, and some are brittle. Such materials have a limited ability to accommodate the strain caused by the intercalation process. Deformation may not be purely elastic, so particles may not recover their initial shapes during delithiation, resulting in cracking or interfacial delamination. Furthermore, some solid electrolytes require coatings to remain stable in contact with intercalation materials, another factor that makes the chemo-mechanical analysis of composite electrodes more complex. Stress can affect open-circuit potentials, and therefore may impact interfacial reaction rates and lead to stress diffusion of lithium within particles, both factors that affect power capability. The complex interaction mechanics in composite, dual-solid cathodes are inevitably significant, and are still not well described by models.
Advances in Science and Technology to Meet Challenges

Understanding the impact of mechanical state on interfacial electrochemistry is crucial to improving solid-state battery performance. Interfaces exhibit a complex coupling among space-charge effects, electrochemical reactions, and multicarrier transport phenomena. State-of-the-art solid electrolytes exhibit critical currents around $0.1 \text{mA cm}^{-2}$ at room temperatures. However, as described in chapter two, a critical current density of $5 \text{mA cm}^{-2}$ is the desired target for practical application. Experiments have illustrated the central role of metal/electrolyte interfacial impedance and interfacial contact in the determination of critical currents. Many interfacial treatments have been exploited to reduce interfacial resistance, including heat treatment, alloy coating, the addition of liquid additives, etc. The microstructure of solid electrolytes, i.e., porosity and grain size, also shows great impacts on power performance. Observation of bulk plating requires more understanding of how solid electrolytes may contain and conduct free electrons. A recent model by the authors of a study on dendrite nucleation and bulk plating has shown that large mechanical forces can arise as a consequence of the dielectric properties of solid electrolytes and interfaces, which correlate with critical currents [10]. Dielectric properties have been largely ignored to date, and merit further study.

Solid-state batteries with thick cathodes have been built and cycled. It has been suggested that a proper volume ratio of electronic conductive active materials and ionic conductive solid electrolytes is required to provide good percolating conductive paths for both ions and electrons. Volume expansion induced by intercalation needs to be managed carefully to minimize contact loss or mechanical failure of particles. Smaller particles are generally favoured to mitigate cracking. Theoretical study suggests that delamination may be suppressed by properly matching the mechanical properties of solid electrolytes and intercalation compounds.

Concluding Remarks

The coupling of mechanics and electrochemistry in solid electrolytes and interfaces is critical to the performance of solid-state batteries. Phenomenological theories that elucidate solid electrolyte failure (especially the initiation stages of such failure) and degradation are desired to guide battery design. Despite the fact that working current densities have been significantly enhanced through the use of high-quality, dense solid electrolytes with various interfacial treatments, the origins of failure modes are still unclear. On the cathode side, although many electrode designs have achieved acceptable cycling performance, both theoretical and experimental efforts are still needed to reduce the susceptibility to mechanical degradation.

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References

6. Cheng, E. J., Sharafi, A. & Sakamoto, J. Intergranular Li metal propagation through polycrystalline Li$_{x}$Al$_{0.25}$La$_{3}$Zr$_{2}$O$_{12}$ ceramic electrolyte. Electrochimica Acta 223, 85–91 (2017).


Characterisation of Electrode-Electrolyte Interfaces in Solid-State Batteries

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Status

Within solid-state batteries (SSBs) there exists numerous interfaces between electrode active materials and the solid electrolyte. For practical application of an SSB, minimal impedances between interfacial layers are required. The buried nature of these interfaces presents certain challenges in order to characterise them with traditional surface characterisation techniques, whether ex-situ, in-situ or operando. To design solid-state batteries that can optimise specific energy and longer life, it is important to understand the processes happening at the interface between the solid electrolytes and cathodes and adopt rational approaches to solve the problems that are causing the cell degradation. Thus, the development and exploitation of new and existing methods of characterising the interface within solid-state batteries, at both anode and cathode, is critically important for guiding future development strategies.

The origin of cell failure is due to both chemical and electrochemical interfacial instability, as well as mechanical robustness, where fracture will result in loss of contact between the electrode and electrolyte [2]. As in the case of liquid electrolytes, solid-state electrolytes have an electrochemical stability window outside of the potential range of the majority of anodes and cathodes, thus, solid electrolyte interphase layers will form upon both electrodes, chemically and/or electrochemically, as shown in Figure 1 [3]. The solid-solid contact between the solid electrolytes and cathodes can be lost due to volume change of cathodes during cycling. Lithium (Li) metal creeping behaviour is also influenced by void formation at the interface between Li-metal and solid electrolytes [4]. Understanding the complex nature and interplay of these various buried interfacial regions as they evolve as a function of time, rate of charge/discharge and potential is a significant challenge that requires a host of advanced characterisation techniques. Much work has already been carried out to develop novel characterisation methods and tools for the study of the solid electrolyte-electrode interfaces, both ex-situ, in-situ or operando within SSBs, as summarised within Figure 2 and within recent review articles [2, 5]. Below, recent progress in interface characterisation will be highlighted and future challenges and strategies discussed.

Current and Future Challenges

Compared to traditional lithium-ion cells containing liquid electrolytes, SSBs present new challenges for the interfacial characterisation methods and tools. The surface/interface areas between the electrodes and electrolytes of cells made with liquid electrolytes can be easily exposed when the cells are disassembled and the separators are removed. In-situ or operando characterisation can also be performed by positioning an optical window at the appropriate point on the cell body. Due to the buried nature of the interfaces within the SSBs, it is challenging to separate and characterise a clean interface between the solid electrolytes and electrodes, in particular after cycling. All the sample handling should be carried out in an air-proof environment due to the hygroscopic nature of most solid electrolytes and electrode materials, which provides another challenge for the development of sample transport tools and holders to be mounted onto various surface characterisation techniques. Ex-situ studies on cathode/SE interface
that highlight the complexity of the interfacial layers in SSB will now be discussed. Recently Yildiz et al. [6] reported detrimental interphase formation caused by Co and La inter-diffusion and \( \text{Li}_2\text{CO}_3 \), \( \text{La}_2\text{Zr}_2\text{O}_7 \), and \( \text{LaCoO}_3 \) formation at the interface between \( \text{LiCoO}_2 \) and \( \text{Li}_7\text{La}_3\text{Zr}_2\text{O}_{12} \) during the annealing process, which is a crucial step during the preparation of the oxide-based solid-state cell. To understand these phenomena a variety of ex-situ techniques were used, including x-ray diffraction, x-ray photoelectron spectroscopy (XPS), secondary ion mass spectroscopy, x-ray absorption spectroscopy and hard x-ray photoelectron spectroscopy (HAXPES). Wang et al. [7] observed the interface between the deposited \( \text{LiCoO}_2 \) cathode and lithium phosphorus oxynitride (LiPON) solid electrolyte with in-situ scanning transmission electron microscopy (STEM) coupled with electron energy loss spectroscopy (EELS). A chemically-formed disordered interfacial layer was identified between \( \text{LiCoO}_2 \) and LiPON, even within the pristine cell. This layer was found to evolve and form \( \text{Li}_2\text{O} \) and \( \text{Li}_2\text{O}_2 \) and cause high impedance at the interface and subsequent capacity decay.

**Advances in Science and Technology to Meet Challenges**

In-situ and operando characterisation tools are being developed to examine SSB’s under real world working conditions to reflect actual processes ensuring that the experimental conditions eliminate the interference and artefacts generated on the interface due to cell breakdown, sample handling and transport. Reducing the high-energy x-ray and electron beam damage effects in XPS and transmission and scanning tunnelling electron microscopy (TEM/STEM) and improving the acquisition sensitivity and accuracy of these surface characterisation techniques is also expected. Furthermore, techniques that provide spatial characterisation of interfaces either in 2D or 3D are being developed and exploited. Yamamoto et al. [8] mapped the electric potential distribution across the interface between the pulsed laser deposited \( \text{LiCoO}_2 \) and \( \text{Li}_{1+y}\text{Ti}_{2-y}\text{Si}_x\text{P}_{3-x}\text{O}_{12} \) solid electrolyte when the cell was cycled within a transmission electron microscope. \( \text{Co}^{3+} \) was found to be oxidised to \( \text{Co}^{4+} \) at the cathode side during charging. 3D \(^7\text{Li}\) magnetic resonance imaging (MRI) was employed by Chien et al. [9] to examine the \( \text{Li}^+ \) concentration at the \( \text{Li}_{10}\text{GeP}_2\text{S}_{12} / \text{Li} \) interface in a \( \text{Li} / \text{Li}_{10}\text{GeP}_2\text{S}_{12} / \text{Li} \) symmetrical cell. \( \text{Li} \) depletion at the interface caused potential barrier and electric double-layer effect and was found to be mitigated with poly(ethylene oxide) coating at the interface. The interphase formation also matched with the impedance growth of the cell monitored by in-situ time-resolved electrochemical impedance measurements.

Concentrating on the Li-metal /SSE interface, an in-situ XPS cell was designed by Janek et al. [10] to study the \( \text{Li}_{10}\text{GeP}_2\text{S}_{12} \) surface during \( \text{Li} \) deposition. Here \( \text{Li}_3\text{P}, \text{Li}_2\text{S} \) and \( \text{LiGe} \) alloy were observed to form at the \( \text{Li}_{10}\text{GeP}_2\text{S}_{12} / \text{Li} \)
interface. Optical techniques such as Raman or infrared have not yet been fully exploited in SSB research. *In-situ* Raman spectroscopy was used to evaluate potential-dependent changes of sulphide based solid electrolyte/Au interface during Li deposition and stripping [11]. Raman technique could be further improved via use of shell-isolated nanoparticles for enhanced Raman spectroscopy (SHINERS) to enhance the Raman signal to detect weakly scattering interfacial species. The technique has been used to study the interfacial reactions at the electrode interfaces in metal-air batteries [12]. Raman can also be exploited as an imaging technique particularly *in-situ* Raman imaging for the direct chemical visualisation of the evolution of electrode-solid electrolyte interface under potential control.

### Concluding Remarks

Much progress has already been made in the characterisation and understanding of the complex electrode-solid electrolyte interfaces within SSB, although significant challenges remain, particularly in understanding longer term interfacial changes within cells. Numerous interfacial degradation products and Li depletion at the interface phenomena have been identified, and correlated with observed severe interfacial impedance growth, cell decay and eventual failure. These have been achieved with traditional *ex-situ* and *in-situ* materials and surface characterisation techniques, such as STEM, XPS, EELS, and MRI, while alternative advanced spectroscopic techniques are being explored and developed. Detection and identification of gas release during cycling is an area of interest to fully categorises all side reaction products and initial studies in this direction have been reported [13]. The knowledge achieved so far have been valuable with design strategies to mitigate and removal or prevention of unwanted interfacial reactions, such as coating the cathode with particle or polymers coatings or atomic layer deposition and Li-metal surface protective layer to aid progress towards practical future SSBs.

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References


Hybrid Solid-Liquid Electrolytes: The Importance of the Solid-Liquid Electrolyte Interphase

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Status

A relatively recent concept is a battery based on a hybrid solid-liquid electrolyte. In this format, a solid electrolyte-based layer is used to enable the reversible cycling of Li-metal anodes thanks to its superior mechanical properties, close to unity transference number and stable solid electrolyte interface. A liquid electrolyte is employed to prevent the solid electrolyte shortcomings on the cathode side: it guarantees intimate contact and wetting with the thick porous cathode electrode upon cycling, high oxidative stability, high lithium-ion and low electronic conductivity. Unfortunately, a large impedance at the solid-liquid interface is generated, limiting practical application of this concept.

Current and Future Challenges

A schematic of a protected lithium metal anode in a hybrid solid-liquid electrolyte cell is illustrated in Figure 1. The solid electrolyte layer is employed as a protective barrier between lithium metal anode and reactive liquid electrolyte. Abe et al. investigated the resistances present in these hybrid solid-liquid electrolytes.[1] The well-characterized resistances due to charge-transfer ($R_{CT}$), bulk solid electrolyte ($R_{SE,bulk}$), grain boundaries in the solid electrolyte ($R_{SE,gb}$), and bulk liquid electrolyte ($R_{LE}$) are present. However, a new resistance was attributed to the solid electrolyte–liquid electrolyte interface, $R_{SE/LE}$, with corresponding activation energies ($E_a$) ranging from 30-100 kJ mol$^{-1}$ derived for several hybrid electrolytes. In follow up papers, Abe et al. demonstrated that $R_{SE/LE}$ is influenced by the concentration of the liquid electrolyte and solvent composition of the liquid electrolyte.[2, 3] From this work, a large $E_a$ was attributed to ion-ion interactions in lithium salts and desolvation of lithium cations in the liquid electrolyte.

Janek et al. provided new insight into the origin of $R_{SE/LE}$ by observing the formation of a solid-liquid electrolyte interphase (SLEI) at this solid electrolyte-liquid electrolyte interface.[4] That is, the liquid electrolyte can react with the solid electrolyte to form an interphase layer between the solid electrolyte and liquid electrolyte. This SLEI can have a profound influence on the performance of the battery, as the transport of lithium ions through the SLEI is not well understood. For example, $E_a$ derived from $R_{SE/LE}$ were highest compared to $E_a$’s derived from $R_{SE,bulk}$, $R_{SE,gb}$, and $R_{LE}$ in this work. Recent work has provided more insight into the complex nature of the SLEI and its impact on performance of hybrid solid-liquid electrolyte based batteries.[5–8] These studies observed that the SLEI originated from chemical and electrochemical reactions of the liquid electrolyte on the surface of the solid electrolyte and, depending on the solid electrolyte and liquid electrolyte, high $R_{SE/LE}$ values of 100-1000 Ω cm$^2$ were measured. Resistances of this magnitude cause large potential drops across the SE/LE interface, significantly decreasing the rate capability of hybrid solid-liquid electrolyte batteries. Similar to the heavily investigated Solid Electrolyte Interphase (SEI) [9], these SLEIs predominately contain Li$_2$CO$_3$, with Li$_2$O, LiF, and other decomposition products of the liquid electrolyte. Therefore, engineering the composition and morphology of the SLEI should also be applicable to the engineering of the SLEI. The major challenge for the future will be designing and synthesizing an SLEI that has high lithium-ion conductivity, low electronic conductivity, good wettability of both solid and liquid electrolytes, and is stable over a large
Advances in Science and Technology to Meet Challenges

Electrochemical and structural characterization of lithium ion battery materials has advanced significantly since the inception of commercial lithium ion batteries in the 1990’s. High precision potentiostats are readily available to probe electrochemistry of these materials and can be coupled with a variety of characterization techniques. For example, microscopic and spectroscopic techniques with high spatial resolution are capable of resolving features at the atomic scale. Coupling these techniques will allow for reliable characterization of the SLEI. The application of a four-probe (4P) electrochemical impedance spectroscopy (EIS) measurement is vital for accurately determining the magnitude of $R_{SE/LE}$, as shown by Abe et al. and Janek et al. The key advantage with 4P EIS is its ability to exclude resistance contributions from the counter electrodes during measurement, providing accurate quantification of $R_{SE/LE}$. Using electrode materials that have minimal reactivity with the liquid electrolyte, such as LiFePO$_4$, will further improve quantification. Furthermore, two-probe and three-probe EIS, potentiostatic, and galvanostatic methods will continue to play an important role in evaluating the performance of hybrid solid-liquid electrolyte batteries. The composition of the SLEI can be determined by the many spectroscopic techniques that are used to characterize the SEI. For example, x-ray photoelectron spectroscopy (XPS) can be used to determine elemental composition and oxidation states of surface species, with an analysis depth on the order of tens of Å. Coupling XPS with techniques such as Infrared Spectroscopy (IR) can help identify specific molecules, especially useful for organic/polymeric species, that may play a key role in the function of the SLEI. Other techniques, for example, Nuclear Magnetic Resonance (NMR) Spectroscopy, Raman Spectroscopy, Gas Chromatography-Mass Spectrometry (GC-MS), etc., can be used to identify liquid and gaseous products formed from the decomposition of electrolyte species, allowing for
the complete SLEI reaction to be characterized. The in situ analogues of these techniques are under current development, which will investigate these reactions in real time.[10] There is also opportunity to understand morphology of the SLEI with microscopic methods. In particular, but not limited to, Transmission Electron Microscopy (TEM), Atomic Force Microscopy (AFM), and Scanning Electron Microscopy (SEM) are used to investigate important interfaces in battery technology.[10] Perhaps TEM is the most versatile, as it can be used to provide structural information with Selected Area Electron Diffraction (SAED) and compositional information with spectroscopic techniques such as Electron Energy Loss Spectroscopy (EELS) and Energy Dispersive Spectroscopy (EDS), with atomic resolution. Interestingly, AFM can not only image the surface with high precision, but also can be used to extract mechanical properties of the SLEI, which has been recently demonstrated for the SEI.[12] In situ analogues of these techniques are also being developed [10], which can reveal nucleation and growth mechanisms during electrochemical reactions, applicable to growth/nucleation of the SLEI.

Concluding Remarks

Hybrid battery concepts based on solid-liquid electrolytes are an exciting technology that can enable next generation high energy density batteries. A promising route in particular would be through the development of protected lithium metal anodes. With this new technology there are novel opportunities for exploring fundamental science, such as exploring the design and construction of the SLEI in order to minimize \( R_{SE/LE} \). A suite of analytical techniques will be required for complete electrochemical and structural characterization of the SLEI, providing the possibility for interdisciplinary work. In general, research regarding these solid-liquid interfaces will be valuable for improving the performance of energy storage technology and provide insight into fundamental surface science.

Acknowledgements

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References

1. Sagane, F., Abe, T., Iriyama, Y. & Ogumi, Z. Li\textsuperscript{+} and Na\textsuperscript{+} transfer through interfaces between inorganic solid electrolytes and polymer or liquid electrolytes. *Journal of Power Sources* **146**, 749–752 (2005).

2. Sagane, F., Abe, T. & Ogumi, Z. Li\textsuperscript{+}-ion transfer through the interface between Li\textsuperscript{+}-ion conductive ceramic electrolyte and Li\textsuperscript{+}-ion-concentrated propylene carbonate solution. *Journal of Physical Chemistry C* **113**, 20135–20138 (2009).

3. Yamada, Y., Sagane, F., Iriyama, Y., Abe, T. & Ogumi, Z. Kinetics of lithium-ion transfer at the interface between \( \text{Li}_0.35\text{La}_{0.55}\text{TiO}_3 \) and binary electrolytes. *Journal of Physical Chemistry C* **113**, 14528–14532 (2009).


7. Liu, J. *et al.* The Interface between \( \text{Li}_{6.5}\text{La}_{3}\text{Zr}_{1.5}\text{Ta}_{0.5}\text{O}_{12} \) and Liquid Electrolyte. *Joule*, 1–8 (2020).

8. Naguib, M. *et al.* Interfacial Reactions and Performance of Li\textsubscript{7}La\textsubscript{5}Zr\textsubscript{2}O\textsubscript{12}-Stabilized Li-Sulfur Hybrid Cell. *ACS Applied Materials and Interfaces* **11**, 42042–42048 (2019).


Polymer and Composite Solid Electrolytes: Role of Polymers in Solid-State Batteries

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Status

Polymers have many attractive properties for application in solid-state batteries. For example, their flexibility improves interfacial contact with electrode materials as well as enhancing stability to electrode volume changes upon cycling. They show lower flammability compared to liquid electrolytes, which ensures safer performance. Potential for low cost and light-weight polymers coupled to facile processability are promising for realisation of scale-up and fabrication of thin, flexible batteries with increased energy density. Polymers could play a focal role as bulk solid polymer electrolytes, connecting anode and cathode, or as crucial interface modifiers between an electrode and inorganic solid-state electrolyte as well as functional (conductive or elastomeric) polymeric binders or coatings in composite materials. Dating back nearly 50 years to the initial discovery of lithium-ion conductivity of poly(ethylene oxide) (PEO), PEO and its derivatives still remain the dominant polymer class used in battery applications. This is attributed to the high Li-ion conductivity at temperatures >60°C (10^{-3} – 10^{-4} S cm^{-1}) owing to its low glass transition temperature coupled to the high oxygen density. Extensive research has focused on addressing the limiting room temperature ionic conductivity of PEO (10^{-7} – 10^{-8} S cm^{-1}), a result of its high crystallinity as well as imparting desirable mechanical properties, overcoming the low lithium-ion transference number (0.2–0.3) and widening the electrochemical stability window which is limited to 4 V. Various strategies have been used to improve upon the properties of PEO/salt electrolytes, including: copolymerisation, crosslinking, blending with other polymers, composites with inorganic materials and modification to yield polymer single-ion conductors.[1, 2] Polymers other than polyethers have been investigated including (but in no way exhaustive) polyalcohols, polyamines, polynitriles, polysiloxanes and polycarbonates.[3, 4] As yet, these materials show insufficient room temperature ionic conductivity and/or poor mechanical stability highlighting the importance of continued research efforts within the field. Polycarbonates are of particular focus in this work for application in next-generation batteries owing to their enhanced oxidative stability (4.5-5 V) which could enable the use of high-voltage cathode materials. Generally, higher lithium ion transference numbers (≥0.5) are also reported and are important for improving cell performance.

Current and Future Challenges

A challenge in the design of polymers for battery applications is the simultaneous optimisation of both ionic conductivity and mechanical properties. Synthesis of well-defined block copolymers is an effective approach to decouple ionic conductivity and mechanical properties. Microphase-separation of block copolymers into soft ion-solvating segments (e.g. PEO) and hard, mechanically rigid domains (e.g. polystyrene, PS) leads to enhanced mechanical stability whilst retaining the soft phase for ion transport. Typically, the reinforcement block for mechanical stability is non-conductive but recently Cao et al. investigated the introduction of conducting poly(propylene monothiocarbonate) (PPMTC) as an additional solvating block in block copolymers with PEO.[5] The double conductive phases in PPMTC-b-PEO/LiTFSI gave higher lithium-ion conductivities (2 × 10^{-4} S cm^{-1} at 25°C compared to PS-b-PEO/LiTFSI electrolytes with
a single conductive phase ($\sim 10^{-5} - 10^{-7}$ S cm$^{-1}$ at 25°C) and storage moduli ($G'$) up to 4 times greater than neat PEO/LiTFSI systems. A future direction may be to introduce a more rigid second conductive phase as the $G'$ was lower than PS-b-PEO/salt electrolytes due to the softer nature of PPMTC compared to PS. Another recent example of the utility of block copolymers in forming mechanically robust polymer electrolytes is the incorporation of random copolymers of trimethylene carbonate and \v{z}-caprolactone as soft amorphous blocks in hard-soft diblock copolymers employing poly(benzyl methacrylate) hard blocks. With 17 wt% LiTFSI an ionic conductivity of $9.1 \times 10^{-6}$ S cm$^{-1}$ at 30°C was reached with a storage modulus ($E'$) of 0.2 GPa, comparable to polystyrene. The solid polymer electrolytes showed oxidative stability up to \~{}5 V and an apparent transference number of 0.64.\[6\] Single-ion conductors consisting of a weakly coordinating anion anchored to the polymer backbone is a common strategy to achieve cation transport numbers close to unity and prevent concentration gradients during cell operation. The current issue limiting their practical application is the low ionic conductivity, particularly compared to the corresponding polymer/salt systems. Tethering of the weakly coordinating anion trifluoromethanesulfonylimide (TFSI) to the polymer chain has been widely studied. Recently, single-ion poly(ethylene oxide carbonates) were investigated by Meccerreyes and coworkers who were aiming to combine into one material three successful components of polymer electrolytes: ethylene oxide units, carbonate groups and lithium-sulfonimide. The copolymers were synthesised by polycondensation between polyethylene glycol, dimethyl carbonate and a functional diol bearing a pendant sulphonimide anionic group and lithium cation. The optimised copolymer had an ionic conductivity of $1.2 \times 10^{-4}$ S cm$^{-1}$ at 70°C with 0.89 transference number.\[7\] This is comparable to the previously reported best electrochemical performance for these types of systems which were based on PEO and methacrylic sulphonamide blocks.\[8\] In contrast, earlier work with similar UV-cross-linked poly(ethylene oxide carbonates) and LiTFSI salt (dual-ion conductors) gave higher ionic conductivities of $1.3 \times 10^{-3}$ S cm$^{-1}$ at 70°C. The optimised copolymer, comprised of PEO linked by polycarbonate segments with 10 wt% UV cross-linkable methacrylic pendant groups to form free-standing polymer films, also showed a high lithium transference number of 0.59.\[9\] In contrast to polycondensation routes for the synthesis of polymer electrolytes, living ring-opening polymerisation strategies require less energy intensive reaction conditions and offer better control over polymer properties such as molecular weight, molecular weight distribution and end-group fidelity. Particularly attractive from a raw materials availability consideration of next generation batteries, are those prepared by the controlled ring-opening copolymerisation of CO$_2$ and epoxides. For example, Meng and coworkers prepared functional CO$_2$-based polymers by terpolymerisation of propylene oxide, allyl glycidyl ether and CO$_2$ catalysed by zinc glutarate. Efficient, facile post-functionalisation with a lithium carboxylate yielded stand-alone polymer films exhibiting electrochemical stability up to 4.3 V vs. Li$^+/\text{Li}$ and a high lithium transference number of 0.86 (though moderate ionic conductivity).\[10\] Poly(propylene carbonate) synthesised by the alternating copolymerisation of CO$_2$ and commercially relevant propylene oxide has attracted a lot of interest. Deng et al. developed composite electrolytes with poly(propylene carbonate) and LLZTO showing oxidative stability up to 4.6 V, ambient ionic conductivity of $5.2 \times 10^{-4}$ S cm$^{-1}$, high ionic transference number (0.75) and reasonable mechanical strength (6.8 MPa).\[11\]

**Advances in Science and Technology to Meet Challenges**

Advances in our understanding of the behaviour of polymers within the battery environment is critical to designing polymers for solid-state batteries that meet the combined requirements of high ionic conductivity, high cation transference number, excellent mechanical properties and electrochemical stability. This includes a deeper consideration of the structure-property relationships of polymer-based materials and in particular polymer degradation mechanisms. Towards this, Oyaizu and co-workers recently compiled a large database of lithium-ion conducting polymers (104 entries) containing information about chemical structure and conductivity. Machine learning models suggested unexpectedly that more glassy polymers rather
than the more traditionally focused upon rubbery polymer electrolytes could offer improvements in ionic conductivity and decoupled from polymer chain segmental motion.[12] DFT studies were carried out to investigate the interface between various solid polymer electrolytes and lithium. Calculated absorption energies indicated stronger adhesion of ester, carbonate and nitrile polymer functional groups to lithium metal compared to PEO and poly(vinyl alcohol). However, a higher reactivity was predicted for polycarbonates and polyesters to form CO and alkoxide products.[13] Experimentally, end-capping of poly(ethylene carbonate) based electrolytes with acetate groups led to improved oxidative tolerance (5.4 V vs. Li⁺/Li) with 120 mol% LiTFSI compared to the non-end-capped (hydroxyl-terminated) polymer (~5.0 V). This implies alkoxide backbiting could accelerate polymer degradation under oxidative conditions and can be inhibited by a simple modification of the polymer chain end-group.[14]

Acknowledgements

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References


11. Zhang, J. et al. High-voltage and free-standing poly(propylene carbonate)/Li$_{6.75}$La$_3$Zr$_{1.75}$Ta$_{0.25}$O$_{12}$ composite solid electrolyte for wide temperature range and flexible solid lithium ion battery. *Journal of Materials Chemistry A* **5**, 4940–4948 (2017).


Garnet Solid Electrolytes

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Status

The garnet stoichiometry of $A_3B_2C_3O_{12}$ presents an exciting playground for the crystal chemist investigating dopant or vacancy effects on conductivity properties in these promising candidate solid ceramic electrolytes \cite{1-4}. The capacity for incorporating a wide range of cations into various sites in the garnet structure, shown in Figure 1 (a,b), has afforded a wealth of compositional variability, recently captured in a review by Thangadurai and co-workers \cite{5}. The promise of garnets lies in their high ionic conductivities ($\sim 10^{-3} \text{ to } 10^{-4} \text{ S cm}^{-1}$ at room temperature) as well as their apparent lack of reactivity with lithium metal, though recent reports challenge us to reconsider the basis for this stability. Opportunities for further development remain, namely, in the scaled-up synthesis of high-performance materials that afford compositional and microstructural control, in realising the chemical stability of these materials under ambient conditions, and in understanding and manipulating local interfacial structure to improve transport properties and avoid degradation.

Visualising and fine-tuning the ion mobility in lithium-stuffed garnet frameworks $Li_xLn_3Zr_2O_{12}$ (where $Ln$ is a lanthanide and $x \geq 3$) demands an intimate assessment of the cation distribution. It is well-established that the zirconium-based $Li_7La_3Zr_2O_{12}$ (LLZO) garnet can adopt two polymorphs: the high ionic-conducting cubic phase, stabilised at room temperature through aliovalent substitution (e.g. with $Al^{3+}$ or $Ga^{3+}$), and the tetragonal phase whose ionic conductivity is lower owing to the absence of disorder which facilitates lithium hopping. Lithium occupational disorder may also be induced by substitution of $Ta^{5+}$ or $Nb^{5+}$ for $Zr^{4+}$ in LLZO, giving rise to high bulk conductivities on the order of $\sim 1 \text{ mS cm}^{-1}$ (Table 1).

The thermodynamic interface stability is highly dependent on the nature of the dopant introduced \cite{7,8}. The stability of doped variants against lithium metal is a property which continues to necessitate careful assessment, as the dopants themselves may be susceptible to oxidation state changes which has implications on the interphase formed. With the advent of machine learning techniques and high throughput screening, it is possible to investigate this cation dopant effect on interfacial stability and, on the basis of this, predict new potential structures which warrant further investigation \cite{9}.

Current and Future Challenges

To assess interphase formation and to determine critical parameters such as ionic conductivities, highly dense pellets of the parent powder are needed for which the primary particle size is a crucial determining factor. Typically, high sintering temperatures and prolonged reaction times are applied to densify garnet powders, yet these run the risk of lithium loss or formation of impurity phases both of which result in lower ionic conductivities. Spark plasma sintering (SPS) presents an approach which allows densification at lower temperatures and shorter times, but specialist equipment and expertise are needed. There are considerable opportunities, therefore, for developing synthetic strategies which permit microstructural control, as well as surface chemistry manipulation to avoid surface contaminants. Chemical stability and handling of garnet electrolytes in ambient conditions is another pertinent challenge. Reaction with
Garnet Family  
Argyrodite Family

<table>
<thead>
<tr>
<th>Property</th>
<th>Garnet Family</th>
<th>Argyrodite Family</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ionic conductivity (mS cm(^{-1}))</td>
<td>(\approx 10^{-3} - 1)</td>
<td>(\approx 10^{-7} - 10)</td>
</tr>
<tr>
<td>Activation energy (eV)</td>
<td>(\approx 0.19 - 0.58) eV</td>
<td>(\approx 0.16 - 0.68) eV</td>
</tr>
<tr>
<td>Electronic conductivity (S cm(^{-1}))</td>
<td>LLZO, (\approx 10^{-8} - 10^{-7})</td>
<td>(Li_6PS_5Cl, \approx 10^{-9} - 10^{-8})</td>
</tr>
<tr>
<td>Practical upper voltage window(^a)</td>
<td>LLZO, (\approx 3.6) V vs. Li(^+)/Li</td>
<td>(Li_6PS_5Cl, \approx 2.2) V vs. Li(^+)/Li</td>
</tr>
<tr>
<td>Shear modulus</td>
<td>LLZO, (\approx 60) GPa</td>
<td>(Li_6PS_5Cl, \approx 8) GPa</td>
</tr>
<tr>
<td>Stability in ambient conditions</td>
<td>Store under dry conditions to avoid Li(^+)/H(^+) exchange which can promote Li(_2)CO(_3) formaion.</td>
<td>Require dry atmosphere due to propensity for hydrolysis reactions.</td>
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</table>

Table 1: Physicochemical properties of garnet and argyrodite candidate solid electrolytes. \(^a\) Practical upper voltage windows are based on redox activity of the solid electrolyte during (de)lithiation [6].

moisture and CO\(_2\) can lead to a surface layer of Li\(_2\)CO\(_3\) driven by Li\(^+\)/H\(^+\) exchange, which has a deleterious effect on interfacial contact and the resulting ionic conductivity since Li\(_2\)CO\(_3\) will preferentially grow along grain boundaries [10]. Understanding and mitigating this surface effect demands further study. For example, faster cooling rates after calcination could avoid Li\(_2\)CO\(_3\) formation and alternative heat treatments such as microwave methods may provide this [11]. The primary grain size may also play a role where smaller grains may be less prone to Li\(_2\)CO\(_3\) formation, although the nature of this effect remains unclear. Hand-in-hand with crystal structure optimisation must come particle processing and microstructural tuning, whereby greater control over the interfacial resistances may be possible through architectural manipulation of particle surfaces and subsequent control over interphases. Alternatively, additives could be sought which promote stability as exemplified by the recent work of Goodenough and co-workers [12], who have demonstrated stabilisation at the interface in Ta-doped LLZO by the addition of 2wt% LiF. While the majority focus has been on elevating the garnet ionic conductivity, recent work [13] has taken a closer inspection of the role that electronic conductivity plays in the evolution of lithium dendrites (Table 1). The observation, using operando neutron depth profiling, of lithium dendrite formation in LLZO-based electrolytes highlights the need for established boundary conditions for electronic conductivities in candidate solid electrolytes.

**Advances in Science and Technology to Meet Challenges**

Synthetic advances that afford microstructural and primary particle size control could afford rational insights into how densification and interfacial properties could be governed. For example, reducing the Al-doped LLZO particle size has been shown to decrease the densification temperature for attaining free-standing thin films with fewer grain boundaries [14]. Our own efforts [15, 16] at developing sol-gel based chemistries have afforded shorter calcination times to access Al-doped LLZO, where densification is aided by the in situ formation of LiAlO\(_2\). Such routes have also seen dense sol-gel derived electrolytes which display total conductivities comparable to SPS-treated materials. The application of advanced microscopy methods has enabled finer inquiry of LLZO-interface formation. In an elegant examination by aberration-corrected scanning transmission microscopy, Chi and co-workers [17] have shown that the immediate surface (up to five unit cells) of Al-doped LLZO is reduced on contact with lithium metal to form the lithium-rich tetragonal LLZO phase. While the tetragonal phase is the low ionic-conducting polymorph, its presence here is favourable as it limits further interfacial reactivity. The reported ionic conductivities of ceramic solid electrolytes can vary considerably. The synthetic approach adopted may give rise to compositional or microstructural variability, inducing changes in lithium concentration or grain boundary surface area which would inevitably be reflected in the measured ionic conductivity. The measurement technique itself may introduce user-based differences.
since comprehensive universally-agreed procedures do not exist, for example for standard impedance analysis. Interestingly, this idea of interlaboratory reproducibility has been investigated by a team led by Zeier who demonstrate the considerable deviation in total ionic conductivities and activation energies for identical samples measured across multiple laboratories [18]. Such a call to the community for rigorous measurement methodologies and established testing criteria is well-timed. Advances in applying local dynamic tools such as muon spectroscopy to interrogate self-diffusion properties as well as *operando* measurements will overcome the sensitivities of grain boundaries or surface effects [19]. Developing large-scale processing techniques remains a crucial bottleneck to be overcome in the realisation of solid-state batteries. The inherent challenge here is to scale-up without compromising ionic conductivity. Recently, Rupp and co-workers [20] have developed a multilayer-based technique using PLD which employs Li$_3$N as a lithium reservoir that reduces the stabilisation temperature of Al-doped LLZO by several hundred degrees. What is particularly exciting about this development is its general applicability, which allows precise control over the lithium concentration, as well as lower processing temperatures.

**Concluding Remarks**

Extensive efforts have been made in enhancing ionic conductivities in garnet materials through compositional control and research continues in optimising particle morphology. Considerable gains can be made in engineering tailored garnet surfaces which stabilise the electrode-electrolyte interface without detriment to the ionic conductivity. Whether...
the greatest benefit comes from designer interfaces or coatings introduced post-synthesis or via a coating formed in situ during cycling remains to be seen. The recent interest in probing garnet electronic conductivity opens this up as a potential diagnostic tool for predicting lithium dendrite formation. Finally, engineering the scaled-up production of ceramic electrolytes for solid-state batteries will require some novel thinking such as smart multilayers, targeted morphologies for optimal packing or symbiotic electrolyte composites.

Acknowledgements

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References

8. Kim, Y. et al. Electrochemical stability of Li$_{6.5}$La$_3$Zr$_{1.5}$M$_{0.5}$O$_{12}$ (M = Nb or Ta) against metallic lithium. *Frontiers in Energy Research* **4**, 1–7 (2016).


Sulphide Solid Electrolytes

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Status

The search for new solid electrolytes is being driven by the desirability of metallic lithium anodes, and the associated increase in volumetric energy storage from obviating the need for graphitic anodes. Sulphides have some general advantages over oxide materials: the more polarisable sulphide anion should provide a lower electrostatic barrier to ion migration, weaker bonding leads to materials that densify at lower temperatures, and softer materials may be more forgiving of the changes in electrode volume during battery cycling. Key parameters are summarised in Table 1. Fast Li+ conduction is known in a number of glassy sulphides based on tetrahedral PS43– and di-tetrahedral P2S74– units that show variable thermal stability [1]. The glass-forming nature of these systems is a challenge and sulphide electrolytes have undergone a surge in interest following reports of fast Li+ conduction in related crystalline phases such as Li10GeP2S12 and Li6PS5Cl [2, 3]. The dependence of charge transport on chemical composition can be highly non-linear and extensive characterisation is necessary to avoid situations such as have arisen in Li7La3Zr2O12 (LLZO) and ‘beta-alumina’ where the initial chemical composition omitted the key enabling elements (Al and Na, respectively) that deliver the electrolyte properties. In the case of potentially glass-forming systems such as thiophosphates, it is crucial that crystalline structure derived from Bragg scattering is a meaningful approximation of all the sample and advances in pair distribution analysis has delivered great insight here [4]. A common component across fast ion conducting systems is a disordered crystal structure and in argyrodites the Li distribution and anion ordering can be manipulated by compositional adjustments of the halide or by replacement of P5+ by Si4+, Ge4+, Sb5+ to tune electrolyte performance [4, 5].

<table>
<thead>
<tr>
<th></th>
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<tr>
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<td>LLZO, ≈10−8−10−7</td>
<td>Li6PS5Cl, ≈10−9−10−8</td>
</tr>
<tr>
<td>Practical upper voltage window a</td>
<td>LLZO, ≈3.6 V vs. Li/Li+</td>
<td>Li6PS5Cl, ≈2.2 V vs. Li/Li+</td>
</tr>
<tr>
<td>Shear modulus (GPa)</td>
<td>LLZO, ≈60 GPa</td>
<td>Li6PS5Cl, ≈8 GPa</td>
</tr>
<tr>
<td>Stability in ambient conditions</td>
<td>Store under dry conditions to avoid Li+/H+ exchange which can promote Li2CO3 formaion.</td>
<td>Require dry atmosphere due to propensity for hydrolysis reactions.</td>
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Table 1: Physicochemical properties of garnet and argyrodite candidate solid electrolytes. a Practical upper voltage windows are based on redox activity of the solid electrolyte during (de)lithiation [6].

Current and Future Challenges

The major challenges facing sulphide electrolyte materials arise in chemistry, preparation, stability and interface management. Understanding chemical stability and solid electrolyte behaviour requires a detailed appreciation of the chemistry of the host lattice and its interactions with the mobile Li+. The argyrodite structure contains a face-centred...
cubic (fcc) anion lattice with PS\(^{4-}\) occupying the octahedral interstices. In Li\(_6\)PS\(_5\)I the relatively large size difference between S\(^{2-}\) (1.84 Å) and I\(^-\) (2.20 Å) drives complete anion-ordering, with iodide on the fcc sites and sulphur filling half of the tetrahedral interstices. Decreasing the halide size leads to partial anion mixing for bromide and sulphide. For Li\(_6\)PS\(_5\)Cl the smaller radius of Cl\(^-\) (1.81 Å) leads to an enrichment of the fcc sites with sulphur, with the chloride being the majority occupant of the tetrahedral interstitial sites. This change in anion distribution has a profound effect on the lithium arrangement, with Cl\(^-\) having a 50% occupied 48h site and the Br\(^-\) and I\(^-\) having more disordered arrangement of Li\(^+\) over two sites in the structure. Compared to Li\(_6\)PS\(_5\)Cl, the increasing lithium disorder and larger lattice parameter associated with bromide materials results in a slight decrease in activation barrier for Li\(^+\) mobility [4]. The conductivity falls away rapidly with the introduction of larger iodide which drives the anions into a fully-ordered arrangement. The lithium electrolyte performance of argyrodites is also manipulated via replacement of P\(^{5+}\) with isovalent Sb\(^{5+}\) or tetravalent Si\(^{4+}\) or Ge\(^{4+}\) affording a solid solution such as Li\(_{6+x}\)Si\(_2\)Sb\(_{1-x}\)S\(_5\)I [5]. The lower conductivity that is associated with the complete ordering of the I\(^-\)/S\(^{2-}\) ions is more than offset by the increased disorder in the lithium, with the additional Li\(^+\) cations being incorporated into the 24g site. This increases the conductivity to >0.01 S cm\(^{-1}\) but there is some indication that secondary impurity phases may segregate to grain boundaries, and limit the overall performance of the material. Impurities, grain boundaries and segregation are part of the considerable challenges of synthetic scale-up and electrolyte manufacture for the argyrodites. These materials are often made at gram-scale using extensive mechanical milling of chemical reagents for up to 10 hours. New simplified chemical pathways to these materials are still reliant on inert protective atmospheres and reactions performed inside sealed quartz tubes [7]. Long-term cycling stability with metallic lithium remains the great prize, but our understanding of failure modes of these solid electrolytes in a lithium battery is incomplete. A recent comparison of sulphide with other electrolytes concluded that electrical conductivity of the electrolyte material is a primary factor in determining whether metallic lithium is formed under cycling conditions [8]. This interesting proposition requires further testing.

**Advances in Science and Technology to Meet Challenges**

The demand for a solid electrolyte technology is driving scientific understanding of the key processes of lithium transport within and between particles. Recent examples have shown creative application of multiple properties measurements to determine where the lithium electrolyte performance is being limited. The incorporation of softer anions, e.g. by partial replacement of Cl\(^-\) with the larger Br\(^-\), is envisaged to reduce the barrier to Li\(^+\) migration between sites within the crystal. It has been less anticipated that softening the lattice reduces the phonon frequency and as a consequence the vibrational timescale is extended. This means that although the softer lattice increases the probability of a successful Li\(^+\) ion jump, the jump is attempted at a lower frequency and the ionic conductivity may be reduced [4]. Hence softening the lattice is associated with effects that both potentially enhance and diminish the rapidity of ionic migration through the crystal structure. As the conductivity of sulphides has increased, it has focussed attention on the requirements for operation in a solid-state battery, particularly the ability of Li\(^+\) cations to move through the macroscopic electrolyte assembly and into the electrode materials.

Although the lithium mobility in the crystal structure can match that of liquid electrolytes, there remain considerable challenges at managing grain boundaries and interfaces. Local probes, such as NMR can resolve the intra- and inter-cage Li\(^+\) transitions, with the latter being the rate-determining step for Li\(^+\) transport within a single crystal of Li\(_6\)PS\(_5\)Cl. Most importantly, measurements on a Li\(_6\)PS\(_5\)Cl/Li\(_2\)S electrolyte/electrode composite show that the key determinant of high rate battery performance is the exchange of Li\(^+\) cations between the argyrodite and the Li\(_2\)S electrode [9]. The power applications of a solid-state battery based on this chemical system may be limited by electrolyte/electrode interface and composite (nano)structure. However, the full extent of ion conductivity in argyrodite cannot be exploited without further engineering of particle/particle electrolyte/electrode interfaces. A great advantage of solid electrolytes is the wide thermal stability window. Operation both at low and
high temperatures indicate exceptional performance, including high charge rates (up to 18 C) can be realised [2, 5]. The dependence of battery degradation on charging rate and thermal history is not completely understood for conventional Li-ion batteries, and the potential for using thermal control to facilitate fast, safe charging of solid electrolyte systems may be even greater due to the extended thermal stability range of these electrolytes [10].

**Concluding Remarks**

Key challenges remain in the development of sulphide electrolytes suitable for solid-state battery, relating to stability, scale-up and electrolyte/electrode architecture. Recent advances in the experimental interrogation of interfaces and proposed models of electrolyte stability with metallic lithium suggest avenues for tailoring known phases towards materials more suited to meet these challenges.

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**References**


Crystal Structure Prediction and Experimental Realisation of Inorganic Solid Electrolytes

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Status

Starting from the periodic table of the elements and arriving at an entirely new commercial material satisfying specific performance requirements is one of the greatest challenges of modern materials science. Undoubtedly, the path between starting point and destination can be extremely complex, often demanding the development of new and novel cutting edge tools to address various challenges along the way. With its objective of finding novel inorganic solid electrolytes to enable step advances in solid-state Li-ion batteries, the materials discovery part of the SOLBAT project lies firmly in this realm. The ideal inorganic solid electrolyte should exhibit high room-temperature Li ionic conductivity (>1 mS cm\(^{-1}\)), zero electronic conductivity, wide electrochemical window for compatibility with both high-voltage cathodes and Li metal anodes, and mechanical/permittivity characteristics that suppress dendrite formation.[1] These (aspirational) requirements shape the main workflow for the materials discovery part of the SOLBAT project, Figure 1. The main tools employed lie in the disciplines of crystal structure prediction, synthesis, and characterisation. We have excluded the parallel effort of searching solid electrolytes from databases of known materials, either directly or by chemical analogy.

Crystal structure prediction (CSP) has developed to the point where it is now virtually an applied technology.[2] There are several approaches for implementing CSP including evolutionary (e.g., XTALOPT[3]), particle swarm (e.g., AIRSS[4]), and basin hopping (e.g., ChemDASH[5], an in-house code in our group) algorithms. ChemDASH has been the CSP workhorse for the SOLBAT project, employed within the philosophy of the probe structure approach.[6] This approach only aims to find a probe structure (i.e., a crystal structure with coordination environments that are representative of the ground state) rather than the ground state itself, thus reducing the cost of the CSP. The probe structure approach is intimately coupled with experimental synthesis. The search for the ground state is completed through synthesis trials at or near the chemical compositions where the most promising probe structures (based on energy above the convex hull) were found. If the experimental ground state is found, both computational and experimental tools are employed in tandem for the characterisation of the new compound. Computational tools for assessing ionic conductivity include bond valence sum mapping for preliminary screening, nudged-elastic band for energy barriers along predefined plausible migration pathways, and molecular dynamics (\textit{ab initio}, empirical or machine learning potentials) for a more detailed mapping of conduction pathways, energetics, and transport coefficients at finite temperature. On the other hand, electrochemical impedance and NMR spectroscopy techniques are commonly used to characterise ionic conductivity experimentally. Electronic conductivity is undesirable and standard electronic structure calculations are performed to determine the electronic band gap. Other key properties include the electrochemical stability and mechanical properties; both computational and experimental tools exist for their evaluation.

Current and Future Challenges

We mainly consider challenges around the CSP step because of its high cost. At the front end of the solid electrolyte discovery process is
Figure 1: Materials discovery workflow for the SOLBAT project. The main elements are material system (phase field) selection, composition space sampling within a phase field, crystal structure prediction for the different compositions based on the thermodynamic convex hull (stability test), synthesis trials for promising compositions, and characterisation of any compounds successfully synthesised. Information feedback is shown by the thin arrows: yellow – compositions failing the stability test guide further composition space sampling, dotted red – compositions failing the synthesis test lead to refinement of both the stability test and composition space sampling, red – successful compositions guide further composition space sampling, crystal structure prediction (they are added to the convex hull), and the stability criterion.

A vast two-tier chemical space representing all possible material systems (crystallographic phase fields) and all the different chemical compositions within a given phase field. Each of these exhibits a combinatorial explosion that precludes any possibility of exhaustive exploration, even with the most advanced high-throughput methods. Consequently, some scheme for sampling the vast chemical space is required to make progress and the question of how this sampling should be performed is increasingly looming large at the front end of CSP. Mathematically, phase field selection is a problem of sampling from the vast combinatorial manifold of all possible combinations of the chemical elements of the periodic table. Admittedly, this manifold is reduced by restricting the search to only chemically meaningful combinations but it remains extremely large. At this stage, the experienced chemist, based on domain knowledge, typically selects elements commonly found in known electrolytes and combines them in familiar ways. The notion of chemical similarity can be employed in a limited way to derive compounds analogous to known electrolytes. The validity of this approach as a reliable guide to materials discovery is increasingly coming under scrutiny. It unduly restricts the search space and could even perform worse than random selection,[7] making it ill-suited to novel materials.
discovery. Composition space sampling, i.e., choosing a tractable set of compositions to explore the phase field energy landscape is the second tier of the CSP front end. Not much is known yet about the structures of multinary phase field energy landscapes, especially at higher dimensions than ternary. The CSP step constrains the number of atoms, N, in a chemical composition (N∼100 atoms). However, even when N is constrained to be compatible with the CSP step, the space remains too large to search exhaustively. For example, in our study of a quinary system, we found \( \sim 10^5 \) compositions in the computable space \((N \leq 120)\) but only \( \sim 10^2 \) could be calculated. The key question is how to obtain these \( \sim 10^2 \) compositions so as to maximise the yield of synthetisable compounds from the phase field. Fundamentally, the search for novel solid electrolytes is a multiobjective problem requiring a search on the Pareto front of all the key properties. This is a major challenge with current tools and our approach is to prioritise only ionic conductivity at the front end of the search process. The hope is that if a good Li-ion conductor is discovered, it may be possible to engineer the other key properties into the material. Even with ionic conductivity as the only prioritised property, its computational evaluation can be significantly expensive if reliable finite temperature characterisation is required as this entails the use of ab initio molecular dynamics (AIMD). One way to reduce the cost of AIMD is to perform simulations at high temperatures (say \( > 500 \) K) but this is often still not enough to permit processing of a large number of systems. Instead one resorts to classical molecular dynamics but the force fields required may be difficult to construct and, in general, are not transferable.

**Advances in Science and Technology to Meet Challenges**

While the challenges are clearly enormous, there are growing efforts to tackle them from a range of angles. One of the most promising avenues is machine learning (ML), driven by growing materials science databases and the emergence of automated ML workflows for materials discovery (e.g., ChemML) along with libraries of descriptors (e.g., DScribe) as well as the shift towards explainability of ML surrogate models. Some early promising results from ML in solid electrolyte research have been reported ranging from standard regression models through black-box optimisation algorithms to the prediction of conductors based on chemical composition alone. Efficient methods for sampling the front end of the CSP step are an indispensable part of the solid electrolytes road map. As ML approaches based on chemical composition alone continue to be developed, they could offer a promising route towards addressing this challenge. These approaches will need to be coupled with active learning ML tools to achieve an optimal trade-off between exploration and exploitation of these large chemical spaces. However, ML approaches in materials discovery still need to overcome the limitations of training data insufficiency (both in volume and diversity) and predictive uncertainty quantification. Various strategies are currently being investigated to alleviate the problem of small training datasets including data augmentation (e.g., using generative models) and transfer learning, in which models trained on physically related properties where data is sufficient are repurposed for the target property where data is insufficient. Nonetheless, it is apparent that the prospect of integrated platforms where the multiobjective problem is substantially automated centred around active learning ML protocols is becoming increasingly more likely. This is expected to accelerate breakthroughs in the search for novel solid electrolytes for Li-ion batteries. Lastly, as the balance gradually tips away from a primary dependence on domain expertise in materials discovery towards more objective and high-throughput ML models, the transition needs to be matched with new research funding models. Such funding models should deliberately create a space for novel research in the scenario of calculated risk offered by ML exploration.

**Concluding Remarks**

In conclusion, the immediate horizon should see a greater integration of CSP with more sophisticated ML tools, unlocking new and more efficient workflows for the accelerated discovery of inorganic solid electrolytes. However, CSP is expected to remain as a bottleneck to those workflows due to its high cost. Since CSP is merely a surrogate for predicting material synthesisability, workflows that partially or even completely eliminate it are expected to emerge in the long-term.
Acknowledgements
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References
Status

The manufacture of conventional Li-ion batteries involves separate lines of manufacture for anodes and cathodes coated onto foil current collectors, which are then integrated with a polymer separator, followed by various packaging operations including injection of the liquid electrolyte. The process is characterised by high productivity but a large number of process steps. So far the manufacturing approach for solid-state batteries (SSBs) has followed a similar approach of discrete manufacturing processes for anode, cathode and electrolyte: however, the electrolyte tends to be formed first and the positive electrode (a powder-based composite of the active cathode material, carbon and the solid electrolyte) and the Li-metal negative electrode (anode) are then added in separate operations. The electrolyte, usually either an oxide (e.g. Li$_7$La$_3$Zr$_2$O$_{12}$ (LLZO)) or a sulphide (e.g. Li$_6$PS$_5$Cl) is generally required to be largely pore-free to maximise ionic conductivity. Sulphides offer a manufacturing advantage because they can be pressed to a high density at room temperature. In contrast, oxides tend to require relatively high process temperatures (up to 1000 °C or even higher) and pressures (up to 500 MPa) for up to several hours sintering for useful density and ionic conductivity [1]. When the oxide electrolyte is mixed with carbon and an active material and consolidated to form a positive electrode, these high pressing temperatures tend to lead to excessive reactions and burn out of the carbon. Both oxides and sulphides have sensitivity to water vapour with LLZO forming unhelpful but essentially benign Li$_2$CO$_3$ whereas sulphides generate highly problematic toxic H$_2$S gas. Thus, for sulphide-based SSBs in particular, manufacturing must be performed in a dry room or under inert atmosphere.

Current and Future Challenges

Electrolytes

The most widely explored SSB electrolytes are fabricated from hot pressed powders. While hot pressing has been popular for research [2], it constrains options for scale-up beyond the laboratory. Dense oxide-based electrolytes produced by uniaxial hot pressing are usually disc-shaped pellets that are difficult to produce in diameters of 5cm or more without cracking. As-manufactured thickness is also typically and unhelpfully approximately 1 mm, which may be reduced by polishing to 100 µm or less, but avoiding cracking is extremely difficult. Approaches to reduce these problems include use of sintering aids to reduce hot press time and pressure, and tape casting. Another approach is composite electrolytes where, for example, a part-sintered porous LLZO layer is “back filled” with a polymeric Li-ion conductor such as those based on PEO [3]. These composite electrolytes may be more mechanically compliant and tougher, and thus realisable over larger areas with sub-millimetre thickness without excessive cracking.

Anode

The relatively low reactivity of solid electrolytes allows a high specific capacity Li-metal anode to be used, which is essential to compensate for the higher intrinsic density of solid compared with liquid electrolytes. A thin Li layer anode is usually applied by thermal evaporation and deposition under vacuum, or by pressing of a Li foil, onto the pre-formed electrolyte. Typically, an overall Li thickness of 10–30 µm is required to enable competitive cell-level volumetric and gravimetric energy densities. Thermal deposition requires the electrolyte surface to have good wetting...
properties, which may not always be applicable, and deposition costs become significant for practical anode thicknesses and areas. In many cases, interlayers are added to the anode/electrolyte interface (by evaporation, sputtering, pulsed laser deposition, atomic layer deposition, etc) in an attempt to improve wetting, stability and mechanical compatibility, and to reduce impedance [4]. While foil anodes are more straightforward, poor wetting (often due to Li surface contamination/reactivity) and resulting high resistance is typical. The price of Li foil also tends to increase with decreasing thickness and, so far, few manufacturers are able to produce sheets below 50µm. An alternative approach are so-called anode-free SSBs where the Li-metal anode is formed in situ by electrochemical plating during the first battery charge [5], which in theory at least is highly scalable and simple from a manufacturing standpoint.

Cathode
Key challenges for the manufacture of the composite cathode are to ensure simultaneous interconnectivity of the solid electrolyte for Li-ion movement throughout the cathode, interconnectivity of carbon for ubiquitous electron percolation, intimate contact of all active particles with the electrolyte (which is typically more difficult for “hard” oxides than “soft” sulphides), including during cycling when the active material typically swells/shrinks, and a minimised fraction of electrolyte and carbon overall to boost specific capacity. The target composite cathode thickness depends on the Li-metal anode thickness and the fraction of cathode active material, but typically lies in the range 30–150µm [6]. So far, most composite cathodes are manufactured by variants of slurry casting, with or without subsequent hot pressing.

Advances in Science and Technology to Meet Challenges
Given the restrictive nature of hot pressing methods for ceramic electrolyte or composite cathode layers (relatively small area, high thickness and low toughness/compliance), the use of polymer materials is a promising trend for enabling SSBs. Polymer engineering is mature and provides opportunities to simultaneously engineer mechanical and functional properties. For example, polymer ionic conductivities (10−8−10−4 S cm−1) may not yet match sulphides (10−3−10−2 S cm−1) but used as part of composite electrolytes (above), coatings on individual cathode particles (preferably using simple, scalable solvent based processing) or at the interfaces between the electrolyte and the composite cathode, they can confer improved cell-level mechanical and electrochemical stability [7]. Composites can also facilitate the use of non-hot press routes to SSBs. For example, Figure 1 is a scanning electron microscopy (SEM) image of the cross-section of a composite cathode based on active LiCoO2 (LCO) mixed with Ta-doped LLZO (LLZTO), onto which a LLZTO electrolyte layer was then deposited. Both cathode and electrolyte layers contained 5% inorganic binder and 5% Al2O3 as a sintering aid. Each layer comprised three relatively thin sub-layers of ~5µm, rapidly pressure-less sintered for the temperatures and times indicated, before the next layer was added. Marked densification was realised without application of pressure, while carbon was retained successfully. A cell was then formed simply by adding a Li foil as an upper layer. A similar approach has been used to produce high performance SSBs using a composite solid electrolyte based on PEO(LITFSI) mixed with Li1.5Al0.5Ge1.5(PO4)3 [8], as well as all-polymer symmetric solid-state cells in which a polymeric anode, electrolyte and cathode were sequentially sprayed to form an all-polymer cell in a single operation [9]. For these systems, aqueous and/or alcohol based suspensions, spray coating under ambient conditions and avoiding any hot-pressing provides for a potentially scalable approach for cathodes and electrolytes, guaranteeing fast processing while maintaining acceptably low levels of contamination. Sulphides and oxides can be processed by classical dispersion approaches under an inert environment, however some sulphides e.g. Li6PS5Cl can also be used in dissolution/reprecipitation methods in which the sulphide coats directly onto cathode particles on reprecipitation [10]. Even though in situ reprecipitation may reduce the ionic conductivity, electrolyte/active material contact can be improved throughout the cathode, giving an overall improvement in cycling performance. Figure 2 shows the Nyquist plot obtained for a 60µm thick binder-free sulphide film comprising 30 individual sulphide layers that were sprayed and re-precipitated in situ from an isopropanol/Li6PS5Cl mixture. This shows that dissolution/reprecipitation approaches may be attractive for manufacturing thin (15–40µm)
Figure 1: Three spray layers of a composite cathode containing LiCoO$_2$ (LCO), LLZTO, carbon, solid-state binder and sintering aid Al$_2$O$_3$, followed by three layers of a LLZTO based electrolyte layer. Each layer was pressureless-sintered for the temperature/time indicated.

Figure 2: Nyquist plot taken from a 60 µm thick LPSCI film comprising 30 individually spray deposited and re-precipitated layers, measured between two stainless-steel spacers under a uniaxial clamping stress of 400 MPa.

Concluding Remarks

Manufacturing research for SSBs is at a relatively early stage. Highest performing SSBs make use of evaporated thin Li-metal anodes, hot pressed electrolytes and slurry cast composite cathodes, based on oxide or sulphide electrolytes. Hot pressing produces dense electrolytes with the highest ionic conductivities but has restricted options for scalability, especially for high temperature oxides. Rolling may be an alternative to hot pressing, but hard oxide particles are difficult to consolidate by rolling alone, unless very high temperature thin glass-making approaches are used (but where stochiometric control is difficult). Softer, lower flow stress sulphides – despite their higher reactivity – may be more suitable for rolling, with acceptable density achievable at modest temperature (<300 °C).
Pressure-less sintering is also being explored and has shown some early promise where pulsed heating of thin layers, for times as short as 10 s, is being investigated. Where porosity remains in composite cathodes or electrolyte layers, “back-filling” or co-depositing of polymeric electrolytes to fill pores is emerging as a potentially acceptable approach, with ionic mobility traded for improved interfacial contact and toughness. Laboratory manufacturing research is producing SSBs with encouraging performance, at least at relatively low current densities, but tends to use processes that have challenge (cost, format, thickness, area, contamination, etc) for scale up and industrialisation. These challenges provide incentive and opportunity for process innovations that balance more holistically the needs of scalability and absolute performance requirement. Considerable scope remains for manufacturing innovations that will enable the cost-effective realisation of the full safety and performance benefits of SSBs at scale.

Acknowledgements

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References

8. Bu, J., Leung, P., Huang, C., Lee, S. H. & Grant, P. S. Co-spray printing of LiFePO4 and PEO-Li1.5Al0.5Ge1.5(PO4)3 hybrid electrodes for all-solid-state Li-ion battery applications. Journal of Materials Chemistry A 7, 19094–19103 (2019).
Processing of Oxide Solid Electrolyte Thin Films

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Status

Solid-state batteries (SSBs) are actively being developed due to their prospective safety and performance characteristics. Unlike their flammable liquid counterparts, solid electrolytes with good (electro)chemical stability allow the use of high capacity electrode materials such as lithium metal, thus promising increased energy densities whilst offering improved thermal stability. This will significantly lower risk of explosion or fire experienced every now and then in current battery technology using liquid electrolytes, therefore thermal management and safety would be less problematic for SSB packs [1, 2]. Solid electrolytes must be integrated into the battery as a thin film in order to maintain satisfactory battery performance [3]. Whilst oxide materials do not necessarily offer the highest conductivities, their processability and stability offer scalable solutions. In addition to large systems, solid thin film batteries can be applied also in microelectronics [4]. During 1990s and 2000s LiPON (Li$_x$PO$_y$N$_z$) electrolyte thin films were studied intensively and were widely used in commercial thin-film batteries due to good stability and cyclability with Li-metal with current densities over 1 mA cm$^{-2}$ without any Li shorting issues. However, its low Li$^+$ ionic conductivity (10$^{-6}$ S cm$^{-1}$ at room temperature) required fabrication as a thin film with thicknesses around 1 µm. Typical fabrication methods for LiPON thin films have been sputtering, vapour and pulsed laser deposition [2–4]. In the last decade researchers have focused on cubic garnet Li$_7$La$_3$Zr$_2$O$_{12}$ (LLZO) which is a promising candidate material for solid electrolyte due to its good stability and reasonably high Li$^+$ ionic conductivity (up to 10$^{-3}$ S cm$^{-1}$ at room temperature). LLZO electrolyte layer with thickness up to 10 µm enables acceptable ionic area specific resistance values (1 Ω cm$^2$) for SSB cells [3][4].

High quality dense LLZO thin films have been successfully fabricated using deposition processes like magnetron sputtering, pulsed laser deposition, chemical vapour deposition and sol-gel based wet coating [5]. Tape-casting was reported to be a good candidate technique for scalable fabrication of 10 µm thick LLZO layers with cost of <$10 $ m$^{-2}$ for industrial production. Therefore this fabrication method has the potential to contribute to commercialisation efforts of SSB technology.

Current and Future Challenges

Due to requirements for solid electrolyte membrane and specific material properties, development of scalable processing for LLZO thin films poses some challenges. A reliable LLZO electrolyte membrane needs to have high density with low grain-boundary resistance and defect-free microstructure to avoid any Li dendrite propagation [6]. Some precautions need to be taken during processing of LLZO material. It is well-known that LLZO is not stable in air atmosphere due to its reaction with ambient H$_2$O and CO$_2$, resulting in (Li$^+$/H$^+$) exchange in LLZO crystal lattice and formation of carbonate impurities on the surface. These processes can affect final sintering and conducting properties of the material and increase interfacial resistances in the battery cell [3, 7].

LLZO thin films have been prepared by using well-established deposition processes like magnetron sputtering and pulsed laser deposition. However, those vacuum-based deposition techniques are unlikely to be used for scaled-up manufacturing due to their high cost and slow deposition rates [1, 5]. Although the films deposited have well-controlled thicknesses, they are usually amorphous resulting in Li$^+$ ionic conductivity values several magnitudes lower than the ones achieved for bulk LLZO samples. Crystallization of the deposited LLZO
thin films is problematic as the high temperatures required (>900°C) will cause Li volatilization from the material resulting in changed stoichiometry [4]. Heat-treatment is even more problematic when LLZO electrolyte layers are prepared from crystalline LLZO ceramic particles, thus requiring even higher processing temperatures for long periods (several hours) to achieve satisfactory densification. Li loss is especially severe at these conditions for thin films because of their higher surface-to-volume ratios, resulting in formation of secondary phases mainly in the surface layers due to extensive Li loss [7]. Due to the brittle nature of refractory materials, standalone LLZO thin film can be mechanically too weak for handling any post-sintering steps like polishing to remove Li deficient surface layers which is usually done in case of LLZO pellets. Thus, some special measures have to be considered because of the challenges described above for developing practical and economical processing of LLZO thin films.

**Advances in Science and Technology to Meet Challenges**

Experience obtained from past studies about LLZO processing can be adapted for thin films. The issues related to carbonate surface impurities occurring during processing of LLZO can be alleviated by using an inert environment and some additional heat-treatment, although it would probably result in increased manufacturing cost. Mechanical properties of self-supported LLZO thin films can be improved by using flexible composite electrolytes of ceramic particles embedded into ionically conductive polymer network [8]. The advantage of ceramic+polymer composite layers is that they do not need any heat-treatment, thus the issues related to Li loss are not relevant in that case. Another approach is to create a multilayered structure by co-sintering LLZO thin film onto thick porous LLZO support which would act also as ionically conductive 3D scaffold for metallic Li electrode for example. The described cell architecture provides high interfacial surface area, enabling high operating current densities (10 mA cm\(^{-2}\)) [9].

Heat-treatment of LLZO electrolyte can be
Figure 2: Fabrication of LLZO-based SSB by laminating and co-sintering the tape-cast layers, followed by integration of active electrode materials into the multi-layered LLZO architecture.

optimized to fine-tune its microstructure, e.g. minimize amount of grain-boundaries and pores, which would contribute to the resistance of electrolyte and serve as pathways for Li dendrite propagation. The most straightforward way to control Li content in the sintering environment would be the use of sacrificial cover powder [10]. Graphite substrates/envelopes can be used for sintering LLZO tape-cast layers in non-oxidizing gas atmospheres because of their non-wetting behaviour [7]. Thicknesses of LLZO electrolyte layer produced by tape-casting can be decreased down to 1µm. It requires careful control over starting powder (grain size and morphology), rheological properties of tape-casting slurry and casting parameters. Dense particle packing in the cast tape helps to achieve high final densities at moderate thermal treatment conditions with minimum time and cost. High density (99%) LLZO layers can be fabricated by using, for example, Li2O as sintering aid [10]. Short heat-treatment duration (<1h) was sufficient to achieve tape-cast LLZO layers with density of 95% and ionic conductivity of >1 S cm⁻¹ at room temperature. In this case the films were prepared from nanoparticles with adequate processing chemistry [7].

Concluding Remarks

Commercialisation of SSBs requires well-established scalable processing for producing solid electrolyte thin films. Fabrication of thin films made from LLZO, one of the main candidates for electrolyte
materials in future SSBs, offers some challenges for its processing. Several studies had previously focused on this matter, proposing different approaches to the issues related to handling and heat-treatment of LLZO layers. The solutions recommended include composite electrolytes, multi-layered structures, strict control over the characteristics of starting powder and processing chemistry, and adequate sintering setups. Scalable tape-casting technique is currently the main candidate method for feasible fabrication of dense oxide electrolyte layers with thicknesses down to 10 µm. Future work needs to demonstrate the functionality of these tape-cast LLZO membranes in the actual SSB cells with high performance.

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References


Thin-Film Solid-State Batteries

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Status

Thin-film solid-state batteries are fabricated by the successive deposition of electrode and electrolyte layers, each less than 10 µm thick, using techniques such as magnetron sputtering, pulsed laser deposition (PLD), vacuum evaporation, chemical vapour deposition (CVD) or atomic layer deposition (ALD). The first true thin-film cell, reported by Sator in 1952, featured a PbCl$_2$ electrolyte and Ag electrodes deposited by vacuum evaporation [1]. Interest in thin-film deposition grew with advances in solid-state batteries during the 1960s, driven by the need to minimise the contribution of low conductivity (~10$^{-7}$ S cm$^{-1}$) electrolytes to overall cell impedance. The first thin-film cell containing lithium (Li/LiI/AgI) was reported in 1969 [2], and was followed by improvements to both electrode and electrolyte materials over the next two decades, including the introduction of the TiS$_2$ intercalation cathode (~2.45 V vs Li$^+$/Li) and glassy electrolytes with ionic conductivities of ~10$^{-6}$ S cm$^{-1}$ [3]. The steadily increasing energy densities and stabilities of thin-film cells, along with the rapid development of integrated circuit technology, drove efforts to fabricate a thin-film “microbattery” directly on a microchip. Significant progress towards this goal was made in the 1990s at Oak Ridge National Laboratory with the development of a Li/lithium phosphorus oxynitride (LiPON)/lithium cobalt oxide (LCO) thin-film cell (Figure 1) [4].

This cell design was commercialised and remains the predominant thin-film system owing to its relatively high voltage and ability to survive several thousand charging cycles before failure [5, 6]. Next-generation thin-film cells with higher energy and power densities have the potential to enable a host of new technologies including implantable and wearable electronics and “Internet of Things” devices. As well as being key to microbattery production, thin-film cells offer several other benefits. These include a small volume fraction of inactive material due to the lack of binders and additives, low impedance interfaces and the attainment of high-density layers with negligible defect concentrations at much lower temperatures than required by the ceramic processing techniques used to fabricate bulk cells [4, 6]. These attributes result in a high theoretical energy density, long cycle life and simple, low-cost construction. Nevertheless, improvements in cell design, materials and processing are required before large-scale commercialisation will be attractive.

Current and Future Challenges

Many of the challenges facing the development of thin-film batteries apply to all types of solid-state battery. Important examples include obtaining cathodes with higher potentials vs. Li$^+$/Li, stabilising lithium metal anodes and finding solid electrolytes with greater ionic conductivities and wider electrochemical stability windows [6]. Challenges specific to thin-film cells mostly relate to defining the links between processing, structure and performance, the optimum cell architecture for maximising power density and volumetric capacity, and the viability of large-scale manufacturing. Electrolyte materials with low ionic conductivities but other favourable attributes (such as outstanding cycling stability and resistance to short circuiting in LiPON, which has an ionic conductivity on the order of 10$^{-6}$ S cm$^{-1}$ at room temperature [4]) may be viable in thin-film form. Nevertheless, it would be desirable to produce thin films of materials with high ionic conductivity in the bulk such as garnets based on Li$_7$La$_3$Zr$_2$O$_{12}$ (LLZO). Unfortunately, the room temperature ionic conductivities of garnet thin films are usually several orders of magnitude
lower than the bulk values ($10^{-7}$–$10^{-6}$ S cm$^{-1}$ rather than $10^{-4}$–$10^{-3}$ S cm$^{-1}$) [6]. Possible explanations include smaller grain sizes in thin films, lithium loss to the vapour phase during processing, and the creation of internal stresses during deposition. Most thin film cells suffer from low power densities and total capacities [7]. The absence of conductive additives and strain-absorbing fillers limits the cathode thickness that can be reached before the cell admittance and accessible capacity become unacceptably low. Furthermore, the flat cathode-electrolyte interface is a major lithium transport bottleneck; in conventional lithium-ion cells the liquid electrolyte penetrates the cathode, increasing the interfacial area and hence the energy available at high cycling rates. To be viable commercially, industrial-scale thin-film cell manufacturing must be both efficient and cost-effective. Techniques such as magnetron sputtering are used in the high-volume manufacture of functional thin films on touch screens and razor blades [8, 9], but little attention has been given to suitable mass production processes for thin-film batteries. Future thin-film cell development should focus on the optimisation of industrial deposition parameters and cell designs, since methods used in the laboratory may not be compatible with industrial processes.

Advances in Science and Technology to Meet Challenges

Thin-film deposition technology can create structures, and hence properties, that are not easy to achieve in bulk materials. For example, the amorphous structure of LiPON responsible for its outstanding cycling stability is formed at low temperatures by a simple magnetron sputtering process. The discovery of new amorphous electrolytes with the same mechanical stability as LiPON but higher ionic conductivities and wider electrochemical windows should be a focus of future research. Garbayo et al. [6] have already demonstrated the feasibility of producing amorphous garnet-type electrolytes with a degree of short-range ordering dependent on the deposition temperature. The highest ionic conductivity was measured in films deposited at 300$\degree$C – more than 700$\degree$C lower than the typical sintering temperatures required for the bulk garnet. With a similar ionic conductivity to that of LiPON, but a wider stability window, these films may perform better in cells containing high-voltage cathodes. In terms of cell design, thin-film deposition could allow the construction of many alternatives to the common planar structures. Architectures that maximise areal capacity and power density by increasing the cell layer area while keeping the cathode thickness and cell footprint area constant are of particular interest [7]. These designs belong to an emerging field of “3D thin-film batteries” which encompasses cells built on non-planar substrates. There are several promising processing routes for these cells; for example, a current collector can be deposited through a template to build up an array of nano/micro-rods onto which the subsequent cell layers are deposited (Figure 2) [10]. Alternatively, photolithography and etching or 3D printing can be used to create a three-dimensional substrate, or the cell layers can be deposited into a preformed structure with high internal surface area such as an aerogel or perforated membrane [7].
3D thin-film cells could present some manufacturing challenges since common deposition techniques such as magnetron sputtering may not be suitable for depositing uniform layers on a three-dimensional substrate [7]. ALD and electrochemical deposition have been reported to be more appropriate owing to their non-directional, self-limiting deposition characteristics. A significant amount of research will be required to determine the optimum designs and deposition parameters for high performance three-dimensional thin-film cells.

Concluding Remarks

Miniaturised electronic devices will underpin several technologies that promise to have a great societal impact. These devices will require safe, reliable and energy-dense power sources. Thin-film solid-state batteries should be ideally suited to these applications although several barriers to large-scale commercialisation currently exist. The overarching challenge is to improve our understanding of the processing-structure-properties relationships of thin-film cells, both at the materials and whole cell levels. In terms of materials challenges, studying the ionic conduction mechanisms in amorphous electrolyte materials will aid the selection of appropriate candidates to succeed LiPON. Whole-cell challenges centre on internal cell interfaces and the need to maximise areal capacity without sacrificing power density. Relatively few investigations have been performed on 3D cell architectures; future studies must seek to achieve the optimum balance between cell performance and processability, and overcoming these challenges is likely to be the focus of thin-film cell research for the foreseeable future.

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References


X-ray Imaging of Solid-State Batteries: Challenges and Opportunities

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Status

The development of X-ray imaging tools for applications in materials science and engineering is well documented [1]. Of the pantheon of microscopy tools, X-ray microscopy provides unique benefits, in particular relating to its non-destructive and multi-scale capabilities. X-ray computed tomography (CT) provides the opportunity to image materials in three-dimensions, and due to the non-destructive nature of X-ray imaging, investigations can be extended into the fourth dimension, to explore the changes in materials over time, and in response to a range of environmental stimuli.

X-ray imaging is compatible with laboratory and synchrotron sources; the latter providing substantial enhancements in X-ray flux; in their recent review Withers and Maire [1] catalogued a range of available X-ray tomography platforms. Whilst the benefits in temporal resolution for synchrotron sources are obvious, some laboratory tools provide competitive spatial resolution. It is possible to achieve sub-micron resolution by the implementation of optical architectures to focus the transmitted radiation, whilst scanning probe techniques and emerging techniques using ptychography and coherent diffraction imaging are also capable of providing further enhancements to spatial resolution. In spite of these advantages, the inherent resolution of X-ray imaging tools do not match those typically available from the suite of electron techniques, and consequently they are most powerful as part of a correlative microscopy toolbox [2].

The invention of X-ray computed tomography for medical applications earned its pioneers The Nobel Prize in 1979; by the early 1980s, these techniques were being actively applied to applications in materials engineering. However, it was not until much later that the first examples of X-ray tomography for applications in battery science and engineering emerged. Indeed, it was not until 2010 that images with sufficient resolution to characterise Li-ion battery electrode microstructure were published [3]. The pursuing decade has seen widespread adoption of tomographic tools for the study of battery materials and devices, which include multi-scale and in situ studies which span a range of chemistries and have been successfully incorporated into image based models. These achievements are summarised in a number of comprehensive, recent review articles [4–6].

Current and Future Challenges

The application of X-ray imaging to understand solid state batteries provides a significant opportunity, that will be summarised in the subsequent section. First, we will consider the hurdles that must be overcome for its effective implementation.

In X-ray absorption imaging the primary means of contrast generation is through attenuation differences, predominantly arising through electron density differences in the constituent materials. This is described by the Beer-Lambert law, from which we can derive a characteristic attenuation length for different materials. Solid state batteries commonly comprise a Li-metal anode with a dense ceramic electrolyte; for example, theoretical predictions from the Center for X-ray Optics database for the attenuation length of these materials at 20 keV incident beam energy are 117.307 µm for Li and 590 µm for LLZO.

This marked difference in the attenuation length of these materials gives rise to the first challenge:
the large density difference between Li and LLZO demand different requirements in the incident X-ray beam. The high mass density of the ceramic electrolyte requires high X-ray energy to provide sufficient transmission, whilst the limited X-ray interaction with metallic Li requires the use of lower energies, and alternative imaging modalities.

For some time, X-ray imaging of metallic Li was widely believed to be intractable. However, in recent years, the development of phase contrast imaging modalities has enabled their study. Phase contrast imaging relies not only on the absorption of incident X-rays, but also their phase shift; combined with phase retrieval algorithms, this imaging modality provides the flexibility to image samples with characteristically low densities. The first example of imaging metallic Li was presented by Harry et al. [10] in their study of metallic electrode growth through polymer electrolytes, and shortly thereafter by Eastwood et al. [11] who studied the geometric nature of dendrite growth in liquid electrolyte systems. Other examples include in situ imaging of metallic Li electrodes as a function of cycle life [11, 12].

Whilst initially, solid electrolytes were thought to block the passage of dendrites during stripping and plating, a growing body of evidence shows that this problem persists above critical current densities. The next major challenge for imaging concerns the multi-length scale nature of the ‘dendrite’ problem: the nucleation of Li plating is expected to occur at microscopic length scales, whilst the propagation of dendrites leading to final short circuit will have macroscopic effects. Moreover, these events may be ‘buried’ deep inside the cell. Consequently, there is a careful balance to strike between the sample volume analysed, and the resolution required; this tension is particularly acute for this inherently hierarchical problem.

**Advances in Science and Technology to Meet Challenges**

The growth in maturity of X-ray imaging technologies, and their flexibility as part of a portfolio of correlative characterisation techniques, provides a compelling opportunity to enhance the development and commercialisation of solid state batteries. Whilst historically the optimisation of battery technologies has been achieved through primarily empirical means, the availability of multi-scale, 3D imaging tools provides a toolbox for quantitative evaluation of materials and devices,
which can be readily fed back to the design process. It is increasingly possible to rapidly screen batteries and their constituent materials to evaluate new materials and device engineering approaches by providing rational design criteria and quantitative comparisons. In recent years, there has been significant improvement in the throughput of X-ray imaging systems, through improvements to hardware, imaging protocols, and reconstruction algorithms. Consequently, it may become tractable to routinely employ X-ray CT in the cell production process, as well as the materials supply chain, providing more robust quality control.

Owing to the non-destructive nature of X-ray imaging, its capacity to explore microstructure evolution in response to a range of environmental conditions is unparalleled. In the field of solid state batteries, there is an emerging opportunity to explore the changes in electrode and electrolyte morphology; for example to track the growth of metallic dendrites leading to short circuit, or the changes in electrode/electrolyte interface. In combination with the development of appropriate theory and modelling tools, this provides a toolbox for mechanistic studies with unprecedented insight.

Whilst the use of X-ray imaging in studies of solid state batteries is at a relatively nascent stage, finally we consider some examples from the literature: McDowell et al. have used X-ray CT to evaluate the mechanical stresses arising at the electrode/electrolyte interface which drive degradation of the battery [8], and similarly Bruce et al. have used in situ X-ray CT to explore the interface evolution, related to the critical current density for dendrite propagation [9]. Elsewhere, the evolution of connected porosity in garnet electrolytes has been explored [13], although critical challenges persist in the discrimination between electrolyte voids, and deposited lithium. These tools are increasingly being applied to emerging chemistries including solid-state Li-sulfur [14], and Na batteries [15] indicating the growing role of X-ray imaging to accelerate the development and commercialisation of next generation batteries.

Concluding Remarks

The emergence of X-ray tomography for applications in battery science and engineering has revolutionised our understanding of the performance/microstructure relationship for a range of materials and devices. With increasing demands on batteries across a range of applications, the drive towards post Li-ion chemistries is accelerating. In this context, X-ray imaging has a significant role to play to facilitate the commercialisation of these devices (including solid state batteries), by providing rational design guidelines to rapidly evaluate and optimise.

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References


