Current status and future perspectives of Lithium metal batteries

Alberto Varzi, Katharina Thanner, Roberto Scipioni, Daniele Di Lecce, Jusef Hassoun, Susanne Dörfler, Holger Altheus, Stefan Kaskel, Christian Prehal, Stefan Alexander Freunberger

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Current Status and Future Perspectives of Lithium Metal Batteries

Alberto Varzi a,b,* Katharina Thanner a,b,c Roberto Scipioni d Daniele Di Lecce e Jusef Hassoun f Susanne Dörfler g Holger Altheus g Stefan Kaskel h Christian Prehal i,j Stefan A. Freunberger i,k

a Helmholtz Institute Ulm (HIU), Helmholtzstraße 11, 89081 Ulm, Germany
b Karlsruhe Institute of Technology (KIT), P.O.Box 3640, 76021 Karlsruhe, Germany
c BMW Group, Petuelring 130, 80788 München, Germany
d Department of Sustainable Energy Technology, SINTEF Industry, 7034 Trondheim, Norway
e Electrochemical Innovation Lab, Department of Chemical Engineering, University College London, Torrington Place, London, WC1E 7JE, United Kingdom
f Department of Chemical and Pharmaceutical Sciences, University of Ferrara, Via Fossato di Mortara, 17, 44121, Ferrara, Italy
g Fraunhofer Institute for Material and Beam Technology (IWS), Winterbergstraße 28, 01277, Dresden, Germany
h Technische Universität Dresden (TUD), Bergstrasse 66, 01069 Dresden, Germany
i Institute for Chemistry and Technology of Materials, Graz University of Technology, Stremayrgasse 9, 8010 Graz, Austria
j Department of Information Technology and Electrical Engineering, ETH Zürich, Gloriastrasse 35, 8092 Zürich, Switzerland
k IST Austria (Institute of Science and Technology Austria), Am Campus 1, 3400 Klosterneuburg, Austria

*corresponding author: alberto.varzi@kit.edu

Abstract

With the lithium-ion technology approaching its intrinsic limit, lithium metal is recently receiving renewed interest from the battery community as potential anode for next-generation rechargeable batteries. In this focus paper, we review the main advances in this field since the first (failed) attempts in the mid 1970s. Strategies for enabling reversible cycling and avoiding dendrite growth are thoroughly discussed, including specific applications in all-solid-state (polymeric and inorganic), Lithium-sulfur and Li-O2 (air) batteries. A particular attention is paid to review recent developments and current state-of-the-art of these battery technologies with respect to the 2030 targets of the EU Integrated Strategic Energy Technology Plan (SET-Plan) Action 7.

Keywords: Battery, Lithium metal, Lithium-sulphur, Lithium-air, All-solid-state
1 – The “holy grail” Li anode: brief history, early failures and future targets of rechargeable Li-metal batteries

Since the mid-20th century, lithium metal has been of high interest for high energy density batteries. In particular, its high theoretical gravimetric capacity of 3861 mAh g⁻¹, and the most negative standard reduction potential (−3.040 V vs. standard hydrogen electrode, SHE) make lithium an attractive anode material.[1][2] The historical development of lithium batteries has already been extensively covered by several recent reviews [3–5] and goes beyond the aim of this paper. Nevertheless, it is worth highlighting a few key events that determined the development of this field.

Following the pioneering work done in the late 60s and early 70s by Rüdorff, Rouxel, and co-workers on the intercalation of alkali metals in transition metal di-chalcogenides[4], it was Whittingham in 1976 (who was then working at Exxon) to patent the first rechargeable Li/TiS₂ rechargeable chemistry.[6] In the following years, several cathode materials have been proposed in combination with Li metal, including transition metal oxides (V₂O₅, V₆O₁₃) and metal selenides (NbSe₃)[7]. In the late 80s, the Canadian Moli Energy succeeded with commercializing the first rechargeable Li metal cells based on a molybdenum sulphide (MoS₂) cathode[8,9]. Unfortunately, millions of sold cells had to soon be recalled due to frequent fire accidents[10]. In fact, while potentially providing high gravimetric energy, the low standard reduction potential of Li lies well outside the stability window of most liquid organic electrolytes.[11] The electrolyte is therefore reduced by the lithium metal, leading to the formation of a Solid Electrolyte Interphase (SEI).[12][13][14] Due to newly forming the full volume of hostless lithium during charge (i.e., Li plating) the SEI can rupture and fresh lithium is continuously exposed. The fresh lithium consumes electrolyte, deteriorates coulombic efficiency, and increases cell impedance due to the increase in SEI thickness.[15] The ruptured SEI also provides an inhomogeneous surface during lithium plating, eventually resulting in dead lithium and dendrite formation. Sand’s equation states that the time for lithium dendrite formation is inverse proportional to the current density. Hence, a homogeneous distribution of the current is crucial to balance space-charge and to avoid local electric field build-up. Depending on the applied current density, dendrites either form as mossy dendrites (high current density) or needle-like dendrites (low current density).[2] The latter are more likely to penetrate the separator and contact the cathode, leading to short-circuit and thermal runaway, i.e., uncontrollable exothermal reactions between the cells components, raising the cell
temperature and forming highly flammable and toxic gases. The temperature increase in turn increases the reaction rate, speeding up the gas formation. Eventually the internal cell pressure leads to explosion and ignition.[16][17] This brought the safety issues of recharging Li metal cells to the public attention, driving the development of the much safer carbon anode, which finally resulted on what is nowadays known as the Li-ion battery (LIB).[7,18,19] Despite the incredible commercial success of LIBs having initially set aside the development of rechargeable batteries with Li metal anodes, the topic has recently experiencing a renewed interest motivated by Li-ion technology approaching its limit. Meanwhile, the academic interest in Li metal batteries has never waned and the understanding of beyond Li-ion systems, such as, for example, Lithium-Sulphur (Li-S) and Li-O_2 batteries, has substantially advanced in the past decade.[20,21] While for Li-O_2 systems many fundamental questions remain unanswered, the practical development of Lithium-Sulphur cells has already reached a relatively high TRL. In fact, OXIS Energy (UK) has been developing lithium-metal prototypes exploiting the sulphur cathode with a capacity ranging from 10 and 35 Ah, currently reaching a specific energy up to 400 Wh kg\(^{-1}\), which has been stated to increase shortly to 500 Wh kg\(^{-1}\) [22]. OXIS Energy and Codemge recently signed a lease agreement to build the world's first Li-S manufacturing plant[23]. In addition, plans to build lithium-sulphur gigafactories in Norway are underway[24].

Currently, substantial efforts are made to finally benefit from the advantages of Li metal anodes in commercial rechargeable cells, especially for electric vehicles (EV) applications. To accelerate this transition, several R&D programs have been launched in USA, China, Japan, etc.[25] Also in Europe, batteries are included among the key clean energy technologies of the Integrated Strategic Energy Technology Plan (SET-Plan) Action 7.[26,27] To become competitive in the battery sector, very ambitious targets have been set for performance (energy, power and lifetime), cost, and manufacturing volume.[26] In terms of battery chemistries, the transition to Li metal batteries (i.e., Generation 4: all-solid-state lithium metal; and Generation 5: lithium-air, lithium-sulphur)[27] is planned starting from 2025.[26] Certainly, large efforts are required to fill this technological gap and overcome the still existing challenge associated with the use of Li metal. This review covers all these aspects comprehensively.

2- The challenge of stabilizing Li metal anodes: general strategies
As recently discussed by Cui et al.[10], among all challenges identified in the past decades, two main issues need to be addressed to enable Li metal anodes: (i) the formation/disappearance of the full volume, and (ii) the high chemical reactivity.

Regarding volumetric changes, the morphology of the anode is key. Pristine Li metal foil is soft, ductile and both a good electronic and ionic conductor. Such features justify its traditional use in form of thin foil, without needing a current collector. However, a thickness change of tens of μm results from applying cathodes with practical capacities > 3 mAh cm$^{-2}$. To mitigate the Li interface movement during cycling, lithium metal powder has recently been considered as alternative. Lithium powder particles (~20 μm in diameter) compacted into a round disc (15 MPa, Ø 15 mm) contain roughly 4.5 times the surface area of a lithium metal foil disc of the same diameter.[28] According to the Sand’s equation, the increased surface area reduces the current density on the lithium surface, slowing down dendrite growth.[29] Additionally, the porous structure can accommodate part of the volume changes upon charge/discharge in the pore volume of the electrode.[30] However, lithium powder electrodes have significant disadvantages compared to foils as they are not freestanding and need a substrate, usually Cu-foil. The porosity of the powder electrode allows contact between the Cu and liquid electrolyte, resulting in galvanostatic corrosion (spontaneous lithium dissolution at the Cu/Li interface).[31] A similar effect has been seen at the Li/electrolyte interface, resulting in pits and voids. Both dissolution effects form “dead” lithium and deteriorate the lithium electrode, causing premature cell death.[31] A solid electrolyte instead may reduce the lithium dissolution at the Cu/Li interface, but causes issues at the lithium/electrolyte interface, discussed in detail later in section 3.1.2.[32]

The very low standard reduction potential of lithium is the root of its high reactivity. Even when stored under inert conditions, i.e., under argon, lithium readily reacts with trace residual atmospheric gases, resulting in a surface passivating layer.[33] This so-called “native SEI” consists mostly of Li$_2$O, LiOH and Li$_2$CO$_3$. While it enables handling of lithium metal in dry room conditions, its composition and morphology, can be influenced by production and storage conditions and is difficult to control. Meyerson et al. analysed the surface composition of a native SEI and determined a mostly inorganic surface (Li$_2$O and Li$_2$CO$_3$) with organic rich veins.[34] The inorganic sections were shown to be less reactive than the organic rich veins. Schmitz et al. additionally found Li$_3$N and Li$_2$C$_2$ when analysing the native SEI, yet their work does not mention distinct morphological differences.[35] Once the lithium electrode is exposed to the electrolyte, a “secondary SEI” forms on top of the electrode. The presence of the native
SEI, and its influence on the secondary one, is often neglected in literature. This complicates a thorough understanding of the Li surface and the development of suitable surface protection strategies.

To tackle the challenges associated with lithium metal, two main approaches have been considered, as shown in Figure 1. The first is to stabilize the lithium metal in the liquid electrolyte via a suitable SEI.[36] The SEI requires similar properties to that applied in state-of-the-art LIBs regarding high ionic conductivity, being electronically insulating and chemically stable.[37][38] Due to the much larger volumetric changes of lithium metal compared to the graphite anode, substantially higher mechanical stability is needed. Possible SEI formation routes include: (i) “in-situ” electrochemical SEI formation via a properly chosen electrolyte (solvent/salt/additive combination) and (ii) an “ex-situ” artificial SEI produced before cell assembly. The second approach is applying a solid instead of liquid electrolyte.[2] The high mechanical strength of solid electrolytes, either polymeric or inorganic, should suppress dendrite growth, therefore prolonging cycle life. Additionally, solid electrolytes improve the overall cell safety. Unlike liquid organic electrolytes, they are not flammable. Yet, solid electrolytes tend to have additional issues, discussed later in section 3. Here we will focus solely on general strategies to enable Li metal electrodes in liquid cells.
Figure 1. Schematic drawing showing the main stabilization routes for lithium metal in liquid and all-solid-state battery cells. For liquid cells, lithium metal can be stabilized with a host structure, “in-situ” SEI or “ex-situ” artificial SEI. All-solid-state cells can either use an inorganic or polymeric solid electrolyte to stabilize the lithium metal anode.

2.1 - In-situ SEI with additives/electrolyte

Understanding the SEI formation process has led to thorough research towards electrolyte optimization, to derive decomposition products desirable for the SEI. Therefore, electrolyte additives have gained great interest. They are usually divided into two main groups, reduction type and reaction type additives (Figure 2a-i).[39] Reduction type additives, have a relatively high redox potential and are reduced prior to the electrolyte depletion. Their decomposition products form an insoluble film, protecting the electrode/electrolyte interface. Reduction type additives are divided into two subgroups. The first subgroup consists of reactive compounds containing an unsaturated carbon bond. These reactive monomers form an electrochemically stable and organic rich polymer layer, upon electrochemical reduction at ~0.9 V vs Li/Li⁺. This group of additives contains, amongst others, vinylene carbonate (VC)[40][41], fluoroethylene carbonate (FEC)[42], vinylene ethylene carbonate[43,44], methyl cinnamate[45], vinyl-
containing silane-based compounds[46], and furan derivates[47]. The polymerization of vinylene carbonate (VC) occurs at the carbon-carbon double bond (C=C). The second subgroup are reductive agents aiding the SEI formation. The reductive agents too get reduced before the electrolyte and their decomposition products adsorb to the electrode surface. They additionally react with other species involved in the initial reduction process, reducing the overall amount of radicals present. Most common are sulphur containing additives such as sulfolane[48], ethylene sulfite (ES)[42], sulphur dioxide (SO2)[49] or 1,3-propane sultone (PS)[50]. Their reduction leads to the formation of Li2SO3 and (RSO3Li)2. The presence of (RSO3Li)2 additionally enhances the ionic conductivity of the SEI. The second group, reaction type additives tend to be so-called “scavenger” additives. They react with intermediate compounds or radicals, aiding the formation of a more stable SEI. Although most scavenger additives have been tested in lithium-ion batteries, their mode of operation should be identical in combination with lithium metal batteries. (Trimethylsilyl)isothiocyanate (TMSNCS) has a high electron donating ability and scavenges PF5 and HF in LiPF6 based electrolytes.[51] Phosphite containing compounds such as tris(2,2,2-trifluoethyl) phosphite (TTFP) and trimethyl phosphite are excellent PF5 scavengers, due to being highly nucleophilic, hence acting as a Lewis base.[52][53] P(III) acts as electron donor and forms a stable complex with PF5. Effective HF scavengers contain simple electron-donating sites and form a complex with HF.[54] Lithium hexamethyldisilylimide scavenges HF and produces NH3, LiF and trimethylsilyl fluoride.[55] Scavenger additives overall improve the stability of LiPF6 containing electrolytes and prolong cycle life. Lithium salts have also been used as additives (Figure 2a-i). Salts with an active multivalent cation (e.g., Mg2+, Ca2+, Zn2+ Fe2+, In3+ and Ga3+) form an intermetallic alloy phase with lithium on its surface.[56] The intermetallic alloy phase has a lower conductivity than lithium and hence lithium diffuses into the layer instead of plating on top, suppressing dendritic deposition of the lithium.[57]
Figure 2. Overview of the main stabilization methods for lithium metal anodes in liquid electrolyte. a) “in-situ” SEI, b) “ex-situ” artificial SEI and c) host structures. a) “in-situ” SEIs can be tailored via i) electrolyte additives or ii) ionic liquids. b) “ex-situ” artificial SEIs can be produced by i) atomic layer deposition, ii) gassing, iii) dip-coating or iv) cutting of lithium in a precursor solution. c) stabilizing host structures can consist of i) a carbon-sphere thin film, ii) a h-BN/graphene thin film, iii) hollow carbon nanospheres, iv) a ultrafine lithium seed layer or v) seeded carbon nanowires.
LiAsF$_6$ has also been investigated as lithium salt additive for organic carbonate based electrolytes.[58] LiAsF$_6$ is reduced in the electrolyte, forming an Li$_x$As alloy phase and LiF on the lithium anode, positively affecting lithium deposition and the surface morphology.[59] Overall, halogenated lithium salt additives are beneficial for improving long-term cyclability of lithium metal batteries. Lithium halides (LiF, LiBr and LiI) suppress dendrite formation. Even without good salt solubility, the anions (F$^-$, Br$^-$ and I$^-$) adsorb on the lithium surface and enhance the surface mobility of lithium ions.[60][61] Since halide salts cannot be reduced any further, they reduce or prevent reactions of lithium with other electrolyte components.

Ionic liquids (IL) have also been investigated as SEI precursors for lithium metal, yet many ionic liquids are not stable towards lithium metal (Figure 2a-ii).[62][63] Generally, ionic liquids are reduced at a more positive potential with respect to the potential of lithium plating. Adding a lithium salt, such as LiBF$_4$, LiPF$_6$ and LiTFSI to an IL is beneficial. By using either the FSI$^-$ or TFSI$^-$ anion, the stability window of the electrolyte is extended and it can be combined with lithium metal.[62][63] Since ionic liquids do not contain solvents, the anion plays the deciding role in the SEI formation and can be tailored accordingly. In the case of LiFSI-IL, the SEI consists of LiF, Li$_2$O, LiOH and FSI$^-$ decomposition products.[64] Once the cell is cycled, additional species associated with the cation are present.

Another example of safe electrolyte worth to be mentioned is the 1.2 M LiFSI in a mixture of triethyl phosphate (TEP) and bis(2,2,2-trifluoroethyl) ether (BTFE) reported by Chen at al. [65]. Besides being non-flammable, it produces a much thinner and dense SEI on Li metal compared to conventional carbonates, thus mitigating its continuous corrosion, which results on less surface available for SEI formation and other parasitic reactions. As shown by Niu et al.[66], when employed in a 1 Ah Li|NMC$_{622}$ pouch cell a gravimetric energy of 300 Wh kg$^{-1}$, this electrolyte substantially mitigates cell swelling under applied external pressure.

### 2.2 - Artificial SEI

As a measure to prevent dendrite formation and ensure long-term cycling stability artificial SEIs have been of particular interest. The artificial SEI is the passivate layer formed on top of the lithium metal anode before coming into contact with the electrolyte (Figure 2b). Depending on the processing method, the artificial SEI forms on top of pristine lithium or the native SEI. Stabilizing the anode surface before cycling allows the regulation of the SEI considering the
thickness, homogeneity and conformity. Artificial SEIs specifically for lithium metal electrodes are often formed by atomic-layer deposition, aeration or coating in a liquid.[67][68][69]

Atomic layer deposition (ALD) is an advanced thin-film fabrication technique, producing homogenous, conform, and ultra-thin films at temperatures below the melting point of lithium (Figure 2b-i).[70] The surface film needs to be as thin as possible to preserve high ionic conductivity, but be thick enough to protect the lithium metal surface. ALD films based on Al₂O₃ result in the lithiation of Al₂O₃ and the formation of a stable, ionically conductive LiₓAl₂O₃ alloy layer.[71] Kozen et. al. showed, that a 14 nm thick film, only contains the LiₓAl₂O₃ alloy phase in the 6 nm closest to the lithium metal surface. The top 8 nm consist of Al₂O₃ and undergo lithiation upon cycling, resulting in a pure LiₓAl₂O₃ alloy layer.[72] Combined with a sulfidic solid electrolyte the ALD Al₂O₃ protective layer prevents self-discharge during the rest period and reduces capacity loss by 40% after 100 cycles.[73] A subsequent study by Kazyak et. al. showed the beneficial effect of a significantly thinner ALD Al₂O₃ film of only 2-3 nm.[74] This film was beneficial for suppressing dendrite propagation and doubled the lifetime of lithium metal electrodes before short-circuiting. Despite the reduction of the Al concentration on the lithium metal surface, the more homogenous current distribution on the surface reduces dendrite growth significantly.

Another method of creating an artificial SEI is via reaction of lithium metal with gaseous species (Figure 2b-ii). The treatment with N₂ at room temperature results in a stable and dense Li₃N protective film.[75] Wu et. al. produced a highly conductive Li₃N layer with a thickness of 159 nm. The protective layer effectively prevents side reactions between lithium metal and the electrolyte whilst Li₃N, due to its high lithium ion conductivity, provides barely any resistance towards lithium ion mitigation.[76] After 100 cycles the passivating layer is still stable and without cracks. Importantly, the exposure time of lithium to N₂ is the deciding factor towards performance and stability of the passivating Li₃N film. Alternatively, CO₂ has been used to passivate the lithium metal surface. Lithium exposure to a CO₂ atmosphere at room temperature leads to the electrode being coated with a Li₂CO₃ layer.[77] The protective layer improved the ionic conductivity and resistance compared to the native SEI on lithium. For the Li₂CO₃ layer formation, the native SEI has to be removed from the lithium surface via mechanical brushing. Without this step, the surface film would be dominated by Li₂O, resulting in reduced ionic conductivity. The high lithium ion exchange rate for Li₂CO₃ is based on the charge centre in the carbonate shifting from one oxygen atom to another, due to orbital interaction and charge delocalization.[78] Due to low ionic resistance, the Li₂CO₃ layer itself
is relatively stable and withstands high current densities of 20 mA cm\(^{-2}\) without cracking. [77] Sulfur gas has been also used to produce a stable Li\(_2\)S layer on lithium metal electrodes.[79] The gas phase reaction at elevated temperature (170°C) forms a homogenous and conductive layer. Due to its certain ionic conductivity (10\(^{-5}\) S cm\(^{-1}\)), the Li\(_2\)S layer can mitigate inhomogeneous lithium ion flux. Upon cycling the artificial SEI preserves its protective function by converting into a layered SEI, containing RCO\(_2\)Li, Li\(_2\)CO\(_3\), sulfonates and a Li\(_2\)S/Li\(_2\)S\(_2\) mixture. The Li\(_2\)S protective triples the cycle life compared to unprotected lithium at 2 mA cm\(^{-2}\).

Additionally, an artificial SEI can be fabricated by exposing lithium metal to selected liquid chemicals. One method is dip-coating lithium metal in appropriate SEI precursors (Figure 2b-iii). For example, dip-coating with polyphosphoric acid solution (0.4 wt% in DMSO) leads to the formation of an artificial Li\(_3\)PO\(_4\) SEI layer.[80] This method replaces the native SEI on the lithium surface with a uniform Li\(_3\)PO\(_4\) SEI, showing excellent chemical stability, a high Young’s modulus (10-11 GPa) and high lithium ion conductivity. Dip-coating lithium metal in a metal chloride solution (MCl\(_x\) in THF, M = In, As, Bi, Zn) forms a Li\(_x\)M\(_y\) alloy phase on the lithium surface.[81] This method utilizes the high lithium ion conductivity of the alloy phase and lithium ion from the underlying lithium metal. The formation of electronically insulating LiCl compensated the bulk alloy layer being electronically conductive, by establishing an electric field across the surface film, driving lithium mitigating through the protective layer. The layer prevents lithium reduction on the surface and suppresses dendrite growth sufficiently, allowing stable cycling at high current densities (2 mA cm\(^{-2}\)) for up to 1000 h. Using a dip-coating procedure to fabricate the artificial SEI has one major drawback. It produces the artificial SEI on top of the native SEI, making it difficult to unambiguously assign electrochemical properties. Furthermore, the composition of the native SEI depends on the lithium provider and storage conditions and can vary between lithium batches. Cutting the lithium directly in the precursor solution ensures the artificial SEI being produced on top of pristine lithium and enables improved investigation of the artificial SEI (Figure 2b-iv). This method was developed and used by Ding et al., to form a protective layer based on 1-pentylamine in pentane.[82] Pentane itself does not react with lithium; hence the resulting protective layer mainly consists of Li\(_3\)N. Li\(_3\)N has an exceptionally high ionic conductivity, not hindering lithium transport and produces a stable SEI with little resistivity, but it can also be a brittle solid.[83] It is important that the 1-pentylamine concentration is sufficiently high (1 M) to produce a stable, homogenous surface prolonging cycling stability.
2.3 – Host engineering

A different approach is to alter the lithium metal surface via nanoscale interfacial engineering. Mechanically and chemically stable frameworks are introduced at the surface to stabilize the SEI forming naturally during charge and discharge (Figure 2c).[84]

Coating the lithium metal surface with a monolayer of interconnected amorphous hollow carbon nanospheres guides lithium deposition and its nucleation within the hollow carbon spheres and on the copper substrate underneath (Figure 2c-i). During further lithium deposition, the layer lifts whilst remaining intact resulting in a continuously stable solid electrolyte interphase.[84] Additionally, lithium deposits in a column like structure rather than long filaments or protruding dendrites. The nanospheres layer enabled cycling at a current density of 1 mA cm⁻², whilst maintaining a coulombic efficiency of 97.5% for more than 150 cycles. Two-dimensional structures such as graphene or hexagonal boron have been proposed alternatively, as stabilizing structures (Figure 2c-ii).[85] During lithium deposition, the ions travel through point and line defects of the 2D layer and deposit underneath on the copper substrate. Both layers are chemically inert and stable against lithium metal. Even a single atomic layer has sufficient mechanical strength to suppress dendrite formation, due to strong intra-layer bonds, resulting in a Young’s modulus of up to 1.0 TPa, more than twice of lithium metal.[86] The graphene layer being a semimetal differs from the insulator hexagonal boron layer. Upon cycling the protective hexagonal boron layer mixes with the electrolyte producing a complex SEI and electrolyte layer. This mixing causes the Coulombic efficiency to rise from 87% to 97% within the first two cycles and remains stable for over 50 cycles at 0.5 mA cm⁻². The hexagonal boron layer protects the lithium anode, but the coulombic efficiency is not yet sufficient for full cells. The graphene layer on the lithium metal anode also enables stable cycling, but the coulombic efficiency is lower (95% at 0.5 mA cm⁻² for over 50 cycles), probably due to its reduced average thickness.[87] Alternatively, guided lithium deposition via pre-infusion or seeded growth has been examined. Pre-infusion host structures based on carbon[88][89], polymeric[90], ceramic[91] or others such as stable nickel foam[92] reduce the volumetric changes experienced by naturally hostless lithium metal and ensure homogenous lithium deposition. The host structure is infused with molten lithium driven by capillary force and provides an electrochemically and mechanically stable artificial interface. Lin et al. showed that such a composite anode has a reduced volumetric expansion of only ~20%, a low overpotential of ~80 mV at 3 mA cm⁻² and is able to retain ~3390 mAh g⁻¹ capacity.[88] Seeded growth entails guided lithium nucleation at chosen “seeds”, aiding homogenous deposition.[93] A plethora of
seeds have been investigated, including homogenous ultrafine lithium seeds (Figure 2c-iv) [94] and heterogeneous seeds such as hollow, amorphous carbon spheres containing gold nanoparticles (Figure 2b-iii)[95] or silver nanoparticles anchored onto carbon nanofibers (Figure 2b-v).[96] Pre-plated lithium seeds provide highly lithiophilic active sites, which significantly reduce the nucleation barrier promoting specific nucleation sites. The consequent homogenous lithium plating results in a dendrite-free surface for 350 h and low overvoltage of 20 mV at 3 mA cm$^{-2}$. [94] Both the carbon spheres and carbon nanofibers provide a 3D matrix in which the lithium nucleation occurs. In case of the amorphous carbon spheres, lithium initially alloys with the gold seeds, forming Li$_{x}$Au, before completely filling the carbon sphere as lithium metal. The carbon spheres are able to alleviate the volumetric expansion as well as protect the lithium from unwanted side-reactions with the electrolyte.[97]

3 - Generation 4: All-Solid-State Batteries (ASSB)

Conventional organic liquid electrolytes in rechargeable Li-ion batteries (LiBs) still pose one of the major safety hazard because of their flammability [98] and, with the development of up-scaled batteries for automotive or stationary application, the risk of fire and explosion has become a serious issue [98,99]. Replacing the flammable liquid solution with an inorganic solid electrolyte (ISE) or a solid polymer electrolyte (SPE) is considered an attractive strategy to mitigate the safety risks which impede the full commercialization of large-scale batteries [100,101]. Furthermore, the use of a solid electrolyte with higher thermal and mechanical stability would enable the use of lithium metal as anode, expediting the development of higher energy-dense batteries [100,101].

This new generation of all-solid-state batteries (ASSB), also known as generation 4 (or generation 4b when a lithium metal anode is used), would potentially meet the demand for safer and higher energy-dense batteries for large-scale applications. However, several bottlenecks still impede the full commercialization [100,102–105]. Achieving an ionic conductivity comparable to classical liquid electrolyte systems (higher than 10$^{-3}$ S cm$^{-1}$) [106], and reducing the large impedance at the electrode-electrolyte interfaces are the main challenges to the full development. Furthermore, electrochemical stability against lithium metal is another major bottleneck in generation 4b batteries.
3.1 – ASSBs with inorganic electrolytes

Inorganic solid electrolytes (ISEs) are considered the most attractive option for generation 4 batteries, mainly because of their high thermal stability, ionic conductivity and cyclability [101]. Compared to solid polymer electrolytes (SPEs), ISEs can achieve a higher ionic conductivity at room temperature ($10^{-3}$-$10^{-4}$ Scm$^{-1}$ vs $10^{-5}$-$10^{-7}$ Scm$^{-1}$) and high Li-ion transference number [107]. On the other hand, they are characterized by a higher interfacial impedance (caused by a poorer solid-solid contact at the electrode/electrolyte interface) and electrochemical instability toward lithium metal, which is dependent on the ISE chemistry [101,108].

3.1.1 Inorganic solid electrolyte chemistries

Sulphide-based electrolytes are among the most promising ISEs for all-solid-state batteries, because their lithium-ion conductivity is comparable to most organic liquid electrolyte [101,107,109,110]. A new class of superionic conductor, based on Li$_3$PS$_4$, is recently being developed as materials of choice for ASSB because of their extremely high ion conductivity and good mechanical properties that allow good solid-solid contact with the electrode interfaces [101,109]. Li$_{10}$GeP$_2$S$_{12}$ (LGPS), in particular, has one of the highest Li-ion conductivity ever achieved in solid electrolytes at room temperature ($12$ mS cm$^{-1}$), which also exceeds the ionic conductivity of most conventional organic liquid electrolytes [107,111]. Contrary to oxide-based systems, sulfide-based ISEs are softer and more deformable, and can be cold-pressed into pellets with tightly connected electrolyte particles. This densely packed configuration has very low grain boundary resistance, and does not require sintering as in many oxide-based electrolytes [109]. On the other hand, sulfide-based systems (as more thoroughly described in 3.1.2), are characterized by high reactivity toward both lithium metal and high voltage cathode materials and are extremely hygroscopic.

Oxide-based electrolytes constitute a wide family of ionic conductor for all-solid-state batteries. The most attractive crystalline Li-ion conductors are garnet-type [112], perovskite-type [113,114], Natrium Super Ionic Conductor (NASICON) and Lithium Super Ionic Conductor (LISICON)[106,115]. Even though their ionic conductivities are usually lower than sulfide-based electrolytes, oxide-based systems are among the most investigated because of their better electrochemical stability with lithium metal and lower degradation at high voltage[101]. Garnet-type conductors are promising candidates to be used in solid state batteries[116], and are finding wider application as inorganic fillers to improve the ionic conductivity and mechanical properties of many solid polymer electrolytes [101]. Although lithium-garnet
electrolytes like Li7La3Zr2O12 (LLZO) exhibit a relatively low ion conductivity (10^{-6} – 10^{-4} Scm^{-1}), this can be enhanced to 10^{-3} Scm^{-1} when the cubic phase is stabilized after Al-doping [101,117]. Contrary to many sulfide-systems, Garnet solid electrolytes are stable at high voltage and when in contact with lithium metal [101]. They are also relatively stable in air, but are very sensitive to water and CO2 which usually cause the deposition of low-conducting side products on the surface (e.g. carbonates)[106,107]. Unfortunately, they are characterized by high resistance at the grain boundaries, whose formation is hardly avoided when synthesized [101,112]. NASICON-type and perovskite-type conductors possess a relatively high ionic conductivity (in the order of 10^{-3} S cm^{-1}) [106,115]. Their fast lithium ion conductivity correlates strictly to their large lattice volume. However, modifications that cause an increase in the channel width for lithium transport are always needed. In a NASICON conductor like LiZr_{x}Ti_{2−x}(PO_{4})_{3}, lithium cannot diffuse fast in a framework mainly consisting of ZrO_{6} octahedra and PO_{4} tetrahedra, but, when Zr is replaced by Ti, the conductivity reaches 10^{-3} Scm^{-1} [118]. Increasing the lattice volume works well also for perovskite such as Li_{3x}La_{2/3−x}TiO_{3} (LLTO) where partial substitution of La with larger Sr ions can enhance the ionic conductivity to 1.5 x 10^{-3} S cm^{-1}[109,114]. LISICON-type lithium conductors possess very high ionic conductivity at high temperature, but relatively poor at room temperature. Li_{3.5}Zn_{0.25}GeO_{4} is reported to have the highest conductivity (0.125 Scm^{-1}) at 300 °C, but only 10^{-7} Scm^{-1} at room temperature[106]. Furthermore, LISICON electrolytes suffers from decrease of the ionic conductivity with time at low temperature because of the formation of Li_{4}GeO_{4}, a complex which traps the mobile lithium ions[106,109,119].

3.1.2 Electrochemical and mechanical stability at the interfaces

Several improvements were done in the enhancement of the ionic conductivity of many inorganic solid electrolytes and results comparable (or even higher) to conventional liquid electrolyte systems were reached for many ISEs such as thio-phosphates, NASICON and perovskite-conductor. However, many other challenges like poor electrode/electrolyte solid-solid contact and electrochemical instability of the solid electrolyte in contact with lithium metal or the high voltage cathode still hamper solid-state batteries full commercialization.

Reactivity toward lithium metal is one of the main drawbacks of sulphide-base solid electrolytes [101,109]. The solid electrolyte/lithium metal interface is very unstable and multiple solid phases (like Li_{2}S, Li_{3}P, Li_{17}Ge_{4}, and polyphosphide compounds) with limited ionic transport properties are usually formed [120,121]. Furthermore, sulphide-based electrolyte are extremely
hygroscopic and can react with moisture producing toxic H₂S[101,122]. Depositing surface coatings or artificially fabricated SEI layers are the most common solution to stabilize the interface [101]. Many sulfide-electrolytes are also electrochemically instable when in contact with high voltage cathode materials[101,109]. Some glass-ceramics thiophosphates (LPS) like Li₇P₃S₁₁ can react with the layered oxide cathode to form metal sulfides (of Co, Mn and Ni, e.g.) with consequent high interfacial impedance. The high electrode potential tends to deplete lithium ions, making the interface highly resistive. For this reason, the sulfide solid electrolyte needs to be protected by the high cathode potential with the deposition at the interface of a buffer layer which needs to be electronically insulating and ionically conductive [101,109]. A thin film of an oxide-based electrolyte is usually used, acting as a buffer against lithium depletion and lowering the interfacial resistance[101,109]. The layer is deposited on the cathode active material surface, before contacting the electrolyte surface. Several ternary metal oxide buffer layers like LiNbO₃, Li₂ZrO₃, Li₂SiO₃, and LiTaO₃ have been successfully used as protective layers on the surface of LiNi₀.₅Mn₁.₅O₄ (LNMO), LiNiₓMnₓCo₂O₂ (NMC), LiNi₀₈Co₀₁₅Al₀₀₅O₂ (NCA), and LiCoO₂ (LCO), significantly reducing the electrode/electrolyte interfacial impedance [123–130]. However, a recent study from Zhang et al.[131] on an NMC-LPS system shows that lithium borates like Li₃B₁₁O₁₈ (LBO) have better stability at high voltage than lithium zirconate Li₂ZrO₃, being promising coatings for thiophosphate systems.

Similarly to sulphide-electrolytes, super ionic conductors like NASICON and perovskite-type electrolyte are also characterized by a bad stability at the lithium metal interface [107,132]. Electrolytes like Li₁₊ₓAlₓTi₂₋ₓ(PO₄)₃ (LATP) and LLTO contain tetravalent Ti, which can easily be reduced when in contact with low-potential anodes. West et al. [133] found that a dark non-metallic insulating layer is usually formed on LATP when in contact with lithium metal. However, the deposition of 1µm-thick lithium phosphorus oxynitride (LiPON) protective layers increases the chemical stability and reduces the reactivity with Li metal [106,133]. Zhou et al. [134] protected LATP from both cathode and lithium metal interface, preparing a ceramic membrane sandwiched with a cross-linked poly-(ethylene glycol) methyl ether acrylate (CPMEA), on both sides. The polymer layer at the solid electrolyte/lithium was observed to suppress dendrite formation, provide a higher wetting ability and protect LATP from Ti reduction caused by contact with lithium metal.

Garnet-type systems are among the most stable inorganic solids against lithium metal [112]. LLZO has a very low interfacial resistance with lithium metal [101,112], but it has to be
protected against humidity and CO₂ during the synthesis, while the lithium surface has to be free of impurities (i.e. LiOH and carbonates) [101,135,136]. Another important aspect that needs to be considered is the lithium wettability on the solid electrolyte. LLZO, like many ceramics, has a poor lithium wettability but it can be improved by sputtering a “lithiophilic” coating as a buffer layer to maintain contact between the lithium anode and the oxide surface. Interesting results were obtained after coating a dense/porous LLZO electrolyte with an ALD-deposited ZnO layer and infiltrating the molten lithium in the LLZO pores [137]. The lithium anode could be cycled for 300h at 0.5 mAcm⁻² without significant dendrite induced polarization. Utilization of a lithium-metal alloys (with Mg[138] or Al[139] as metals) in contact with LLZO is also an interesting strategy that showed a reduction of the contact loss at the solid-solid interface during lithium stripping. The garnet-cathode interface is not exempt by high interfacial resistance. Kato et al. [140] investigated the use of a thin Nb layer (∼10 nm) to reduce the resistance at the interface between LLZO and a LiCoO₂ cathode. The Nb layer was observed to produce an amorphous Li–Nb–O structure, which is reported to be Li⁺ conductive, reducing the interfacial resistance and improving both the battery cyclability and rate capability.

### 3.1.3 Fabrication of All-solid-state Batteries with ISE

Contrary to conventional liquid electrolyte batteries with porous electrodes, all-solid-state batteries require dense electrolyte and electrode layers. While the energy density of the battery can benefit from this, assuring a good ionic and electronic transport becomes challenging.

Fabricating the composite electrode containing cathode active material and solid electrolyte particles is one of the main strategies to provide an ionic and electronic network, while having an intimate contact between cathode and solid electrolyte. The difficulty of preparation of these composite electrodes can vary significantly according to the inorganic electrolyte chemistry. Sulfide-based electrolyte are easier to process because of their softness and deformability [109,141–143], allowing cold-pressing of the composite electrode. The main drawback is the low mechanical strength to lithium dendrite growth [109]. Oxide-based electrolytes possess instead higher mechanical strength; however, they require high temperature sintering to reduce the grain boundary and electrode/electrolyte interface resistances [106,109].

In general, increasing the electrode surface is crucial to assure a high solid-solid contact between the cathode and the ISE (Figure 3a,b) [144], which is fundamental to achieve full active material utilization at high areal loadings (thick electrodes) [145–147], and limit the
amount of solid electrode required in the cathode composite. Obtaining a high surficial contact is possible thanks to mechanical ball milling of the active material, carbon black, and the inorganic solid electrolyte [105,144]. Infiltration of a solution of the soluble electrolyte, usually sulfide-based, in the porous electrode (with consequent removal of the solvent) is another interesting strategy for improved surface contact [105]. However, wetting agents can ensure intimate connection between particles when an insoluble compound like an oxide-based electrolyte is used. Materials like Li$_3$BO$_3$ (LBO) are used as wetting agents that, melting at 700 °C, improving the interfacial contact between garnet-type electrolytes like LLZO [105,148]. Active material coating, as described in 3.1.2, is usually necessary to protect the electrolyte from degradation when in contact with high voltage cathodes (e.g. LiNbO$_3$ at the LiCoO$_2$/sulfide interface [123]), or to create buffer layers with improved ionic transport (e.g. thin Nb layers on the garnet-conductors surface [140]) (as shown in Figure 3c,d). Pulsed layer deposition (PLD) coating of sulphide electrolyte on cathode active material is an interesting technique that allow the preparation of densely packed electrodes with increased energy density[149].
Figure 3. Optimization of the cathode- and anode-electrolyte interfaces in ASSBs. Schematics of lithium ion and electron transport in a sulphur-based composite cathode consisting of large particles with (a) non-homogeneous distribution and (b) small particles with homogeneous distribution. Reprinted with permission from [144]. © 2017 Elsevier. Schematic illustrations of (c) non-modified and (d) Nb-modified LLZO/LiCoO₂ interfaces. The mutual diffusion between LLZO and LiCoO₂ produces non-Li⁺-conductive phases such as La₂CoO₄. The Nb-modified surface suppresses the mutual diffusion and produce Li⁺-conductive amorphous phase. Reprinted with permission from [140]. © 2014 Elsevier. (e) Schematic illustration of the fabrication of an ultrathin electrolyte all-solid-state Li/LLZO/LiFePO₄ battery. Reprinted with permission from [150]. © 2017 American Chemical Society. (f) Schematic illustration of the lithium melt infusion into 3D porous garnet with or without...
The dense solid electrolyte layer is the core of the solid-state battery. It must be intimately in contact with the cathode materials, assure fast ionic conduction (while being an electron insulator), have a good wettability with the lithium anode and protect from dendrites growth and puncturing. At the same time, to achieve specific gravimetric energy (Wh kg\(^{-1}\)) and volumetric energy density (Wh L\(^{-1}\)) comparable to conventional liquid electrolyte LIBs, the solid electrolyte layer must be thinner than a critical value, called break-even thickness [105,151,152]. The break-even thickness is usually dependent on the ionic conductivity of the electrolyte material and on the cathode material loading. It is indeed higher in sulfide-based electrolyte like LGPS (~70-250\(\mu\)m) than in garnet-type conductors like LLZO (~25-80\(\mu\)m)[151]. For instance, if a 60\(\mu\)m-thick cathode with a 15 mg cm\(^{-2}\) loading is used, the break-even thicknesses for LLZO, LATP and LGPS would be ~41 \(\mu\)m, ~75 \(\mu\)m, and ~115 \(\mu\)m, respectively [152]. Fabricating electrolyte thinner than the break-even thickness by traditional fabrication methods is rather challenging; however, Yan et al. [150] reported the preparation of a ultrathin nanoscale LLZO electrolyte for application in an all-solid-state Li/LLZO/LiFePO\(_4\) battery (Figure 3e). They produced a solid electrolyte layer significantly thinner (3-5\(\mu\)m) than the break-even thickness for LLZO by conventional slurry ball milling and tape casting onto the composite LiFePO\(_4\)/LLZO cathode, obtaining high performance in term of cyclability (capacity loss during cycles 2–100 was only 0.06%). Production of even thinner dense electrolyte layers requires the use of advanced techniques such as pulse layer deposition (PLD) [149,153], atomic layer deposition (ALD) [154], sol-gel[155,156], aerosol deposition[157,158]. Unfortunately, the main obstacle of these advanced methods is the high cost for up-scaled production.

Finally, fabrication of a rational interface with the lithium metal anode is crucial for a high energy dense battery with long cycle life. As already mentioned in 3.1.2, protective nanocoating are necessary to protect lithium from reactive electrolytes or simply improve the wettability of the ceramic surface rendering the lithium-ion flux at the interface more homogeneous, facilitating a more homogeneous deposition of lithium, with the final goal of preventing the formation of dendrites [101,108,159]. Expensive advanced fabrication methods like ALD or PLD are usually employed to deposit inorganic nanolayers of ZnO [137], Al\(_2\)O\(_3\) [160] or Si
on the electrolyte surface. However, alternative solutions like using a polymer/ceramic/polymer sandwich can be interesting to improve the adherence to the lithium metal surface. Preparation of a composite lithium metal anode by melt infusion in a 3D scaffold is an attractive strategy for having a homogenous lithium-ion flux. Wang et al. successfully infiltrated melted lithium metal in a 3D garnet-based scaffold ALD coated with a lithiophilic ZnO layer. The tight contact between the lithium metal and the electrolyte significantly decreased the interfacial impedance (from ~2000 Ωcm$^2$ to 20 Ωcm$^2$).

Developing an anode-free cell, where the formation of a lithium metal anode occurs in-situ using the cathode as the only source of lithium while charging, is currently the most promising strategy for increasing the energy density and facilitate the cell manufacturing. Furthermore, removal of the lithium metal foils during the cell assembly process reduces the battery cost. Interesting studies were recently done on the in-situ growth of a lithium metal anode on different substrates (Cu, Au, or preexisting Li) using LLZO garnet-type or LIPON electrolytes. But the best results so far were obtained by Samsung, who developed a Li metal-free Ag-C/ Li$_3$PS$_4$Cl/ NMC ASSB, where lithium is grown in-situ on the silver-carbon composite electrode. The Ag-C layer regulates the Li deposition, leading to longer electrochemical cyclability. The Samsung battery, with an impressive energy density >900 Wh L$^{-1}$, showed a stable Coulombic efficiency > 99.8% and long battery lifetime >1000 cycles. Despite many challenges still need to be faced for large-scale fabrication, these results prove that high energy densities and long battery lifetime are achievable by ASSB. Nevertheless, as nicely outlined in the benchmark study published by Randau et al., optimized cell designs are required to reduce internal cell resistance and improve the power density. Developing electrochemical compatible high conductive solid electrolyte (>10 mS cm$^{-1}$, e.g. sulphide-based) with reduced thickness, accessing the full theoretical capacity of the cathode active material and implementing a new generation of anodes for in-situ lithium growth are the main pathways to follow.

### 3.2 – ASSB with polymer electrolytes

#### 3.2.1 – Science and Technology

Solid organic polymers dissolving lithium salts represent an alternative to ISE, ensuring adequate safety level and possibly, better scalability. Dry polymer membranes with suitable physicochemical characteristics may be indeed processed into thin separators acting as
host for lithium ions, which can move under an electric field [169]. These so-called solid polymer electrolytes (SPEs) allow dissociation of the lithium salt due to favourable coordination of the electrolyte species and a Li⁺ transport assisted by segmental motion of the organic chains [170]. Accordingly, the cation motion mostly occurs within the amorphous fraction of the polymer matrix above the glass transition temperature (Tg) [170], although a few studies described lithium-ion conductivity in crystalline polymer phases [171,172].

Among the SPEs, those based on poly(ethylene oxide) (PEO) [173] revealed the most promising features in terms of applicability and scalability. Indeed, PEO with solid amorphous state dissolves a large variety of LiX salts, where X is typically a voluminous anion such as perchlorate (ClO₄⁻)[174], trifluoromethanesulfonate (CF₃SO₃⁻) [175], bis(oxalato)borate (BOB⁻)[176], and bis(trifluoromethansulfonyl) imide (TFSI⁻) [177,178]. These salts form complexes with PEO in which the anion is almost trapped by the polymer backbone, while lithium cation can move either through the helicoidal ether chains between them by hopping, thus allowing the ion migration through the membrane under an electric field [178,179]. These solid solutions may exhibit relatively high ionic conductivity (above 10⁻⁴ S cm⁻¹) at temperatures higher than the polymer transition point from crystalline to amorphous state [180,181], which can range from 60 to 75 °C depending on the PEOs’ chain length [182,183]. Improved conductivity and mechanical stability can be actually achieved by employing ceramic fillers of various nature [184], e.g., Lewis acid or bases, such as Al₂O₃ [179], ZrO₂ [185], TiO₂ [186], and SiO₂ [187], as well as functionalized fillers [188] and nano-sized oxides to obtain nanocomposite polymer electrolytes (NCPEs, Figure 4a) [189]. Fillers can in fact enhance the membrane strength[190], facilitate self-standing configuration, and increase at the same time the amorphous fraction into the polymer, thereby definitively improving the ionic conductivity and the lithium transference number of the electrolyte [168]. Following this trend, inert inorganic fillers may be replaced with either crystalline or glassy lithium-ion conducting nanoparticles or nanowires to decrease the operating temperature, as demonstrated by promising results obtained in laboratory-scale cells [191]. However, solid PEO electrolytes normally achieve an amorphous condition with ionic conductivity suitable for use in a battery only at medium-high temperatures (typically exceeding 65 °C) [192], thus limiting the application range. The inclusion of plasticizers such as organic [193] and ionic liquids [194,195] to the polymer electrolytes (Figure 4b) [196] allows an increase of the room-temperature conductivity, in spite of a decrease in mechanical strength.
Figure 4 - (a) Temperature dependence of the ionic conductivity of various nano-sized oxide-PEO composite polymer electrolytes [189]; (b) free-standing PEO-LiTFSI membrane containing 150 wt.% PYR$_3$TFSI and solid-state LMP battery consisting of a Cu current collector, Li metal anode, SPE, composite cathode and Al current collectors [196]; (c) temperature dependence of the ionic conductivity for several single-ion block copolymer electrolytes comprising polystyrene segments \[i.e., \text{poly(styrene trifluoromethanesulphonylimide of lithium)}\], P(STFSILi)], with (inset) isothermal conductivity at 60 °C according to the wt% of the P(STFSILi) block [197]; (d) voltage profiles of a Li|LiTFSI-PEO|LFP cell using an amorphous polymeric interlayer at the anode side at 70 °C under a constant current of C/10 [198]; (e) voltage profiles of a Li|PEGDME|LFP cell [where PEGDME is poly(ethylene glycol) dimethyl ether] at 50 °C under constant current rates of C/5
and at 50°C [199](f) voltage profiles of a Li|poly(ethylene ether carbonate)-based electrolyte|LiNi_{0.6}Mn_{0.2}Co_{0.2}O_2 cell at 25 °C under a constant current rate of C/10 [200].

It is worth mentioning that gelled membranes with similar characteristics to typical liquid solutions, also named as gel polymer electrolytes (GPEs), are widely employed in commercial Li-ion configurations [201]. GPEs are commonly modified carbonate-based membranes, in which the liquid phase being responsible for the ion conduction (such as a EC:DMC – LiPF_6 solution) is embedded into a polymer matrix, e.g., based on polyvinylidene fluoride (PVDF), PVDF-hexafluoropropylene (PVDF-HFP) or polymethyl methacrylate (PMMA) [202]. Accordingly, these electrolytes suffer from similar safety issues to conventional polypropylene separators trapping liquid carbonate solutions, thereby hindering a possible application in Li-metal batteries [203].

Solvent-free polymers can increase the safety level, and allow scaling up to high-energy, laminated systems by partially exploiting the current roll-to-roll lithium-ion battery manufacturing line[204]. According to the abovementioned approach, polymer electrolyte and cathode slurries are deposited onto a polypropylene support and a current collector foil, respectively, and laminated after drying. The polymer cell is then assembled using ultra-thin lithium foils prepared by extrusion and rolling/calendering[204]. However, various issues beside the high operating temperature still hinder the large-scale diffusion of such an attracting battery system. These are: i) relatively low cation transference numbers, ii) possible dendrite growth at the lithium anode leading to decay in efficiency and poor cycle life [198,205], and iii) electrochemical instability above 4 – 4.1 V along with poor film forming properties, particularly beyond 65 °C [206], which currently prevent using high-voltage layered LiCoO_2 as well as its high-energy analogues (e.g., NMC materials with various compositions ranging from 1:1:1 to 8:1:1 and NCA) [207]. In this regard, layered cathodes may undergo phase change, release oxygen, and delaminate on charge in PEO-based electrolytes. On the other hand, LiFePO_4, working at 3.5 V vs Li^+/Li, is fully compatible with the lithium-metal polymer configuration, further benefiting from a high thermal stability due to the strong polyanionic framework, which fully enables application at elevated temperatures [208]. In particular, PEO-based solvent-free polymers have shown suitable practical features in Li|LiFePO_4 batteries with a maximum specific capacity of 170 mAh g^{-1} as referred to the cathode mass [209]. Notably, lithium-metal polymer batteries may ensure a gravimetric energy density as high as 300 Wh kg^{-1}, that is, a value approaching that of high-performance lithium-ion systems [210,211], despite the use of low-voltage LiFePO_4 and a relatively low volumetric energy density ranging
from 500 to 600 Wh l\(^{-1}\) [210]. Indeed, cell thickness and weight may be reduced by moving from the conventional lithium-ion configuration to a dry-polymer, laminated geometry employing thin electrode and electrolyte foils [204]; furthermore, the high-capacity metal anode does not require a heavy Cu current collector [210]. Therefore, the use of high-energy NCM and NMC 811 electrodes in a lithium-metal polymer cell might actually lead to gravimetric energy density values within 400 and 450 Wh kg\(^{-1}\), as well as volumetric energy density between 700 and 850 Wh l\(^{-1}\) [210]. Significant enhancement might be also achieved by solid polymer batteries using LiMn\(_2\)O\(_4\) and LiNi\(_{0.5}\)Mn\(_{1.5}\)O\(_4\), that is, within approximate ranges of 300 – 400 Wh kg\(^{-1}\) and 600 – 700 Wh l\(^{-1}\) [210]. However, the abovementioned limited anodic stability of the electrolyte represents a serious drawback to be addressed for boosting the cell performance up to that required in long-range electric cars.

The Li\(^+\) transport within dry SPEs may be enhanced by controlled copolymerization of selected monomers that can ensure anion immobilization along with suitable mechanical and electrochemical properties (Figure 4c) [197,206]. Moreover, appropriated amorphous polymeric interlayers may improve the lithium-metal plating/stripping process, thus enhancing cyclability and coulombic efficiency of the Li|LiFePO\(_4\) cell at 70 °C (Figure 4d) [198]. While the high operating temperature matches the typical requirements of the automotive and stationary storage markets, a widespread application in portable electronics is currently less realistic. Notably, high molecular weight end-capped glymes, that is, short-chain polymers and oligomers based on the ethylene oxide group, are characterized by a lower melting point compared to that of conventional PEO [199]. Despite suffering from limited mechanical stability, solid glyme-based solutions can be actually considered as a possible polymer electrolyte enabling a relatively low operating temperature to the lithium cell (Figure 4e) [199].

Several other chemistries, including poly(methyl methacrylate) (PMMA) [212] as well as cross-linked polymers and copolymers [213], have been proposed for allowing a lithium-metal ASSB; however, various issues, such as low conductivity, modest chemical stability, and scarce mechanical strength, hindered their diffusion and practical application. Among the alternatives to PEO, polyethylene carbonate (PEC) [214] is one of the most promising candidates since it is characterized by similar physicochemical features and higher ionic conductivity at lower temperatures. Furthermore, PEC-based electrolytes are more stable than PEO against high voltage layered cathode, as shown in Figure 4f [200]. However, issues in terms of mechanical and chemical stability, as well as relevant dendrite growth at the lithium side, still prevent a practical application of these electrolytes [215]. It is worth mentioning that a possible approach to mitigate the low chemical and/or electrochemical stability of SPEs, as well as uneven lithium
plating, mainly observed in solid poly-glymes, PEC, PAN and PMMA, is represented by the addition of a sacrificial film-forming agent, such as LiNO$_3$, for efficiently passivating and protecting the metal surface from side processes [199]. In contrast, increasing the battery voltage above 4 V [206] appear much more challenging, since it may involve a considerable change of the chemical nature of the SPE both to allow a wide electrochemical stability windows and to ensure a stable cathode/electrolyte interphase.

3.2.2 - The lithium metal polymer battery: a practical reality
Figure 5: (a) schematic of a Licerion® cell employing an ionically conductive ceramic/polymer barrier protecting the lithium electrode and a high-voltage metal-oxide intercalation cathode, (b) corresponding discharge voltage profile at various current rates ranging from C/3 to 3C and voltage vs. current at steady-state conditions [216]; (c) schematic of the lithium-metal battery prototype developed in the 1990s, comprising a Li anode, a PEO-LiClO4/LiTFSI electrolyte, and a VOx cathode [217]; (d) schematic of assembly of a LMP® battery pack (developed and commercialized by the Bolloré Group) from the cell level [204]; (e) Bluecar and (f) Bluebus (using the LMP® technology) commercialized by the Bolloré Group [218].

In spite of a great deal of efforts from both academia and industries for achieving commercially viable lithium-metal batteries, a few practical demonstrations have been reported to date [211]. Indeed, scaling up from laboratory-prototype coin and pouch cells to more realistic configurations, as well as module and battery packs, often faces substantial barriers [219].
Suitable mathematical models may assist an evaluation of the actual applicability of innovative cell chemistries so far investigated in proof-of-concept and fundamental studies, although various assumptions based on the current lithium-ion battery market may affect their long-term reliability [210]. Moreover, pre-commercial cells and emerging technologies lately on the market represent useful examples to identify the main obstacles that have to be overcome in the short-to-medium term for matching the economic and environmental targets of the European SET Plan. In 2015 Sion Power Corp. (US) announced a transition of their activity from lithium-sulfur batteries to a rechargeable lithium-metal oxide technology [216]. Interestingly, their Licerion® cell employs an ioni
cally conductive ceramic/polymer barrier for protecting the lithium electrode (see Figure 5a) and enabling reversible metal plating within a wide current range due to low interphase resistance, as well as a high-voltage metal-oxide intercalation cathode (see Figure 5b) [216]. Sion Power Corp. demonstrated an energy density exceeding 500 Wh kg$^{-1}$ and 1000 Wh l$^{-1}$ in 0.4 Ah cells in 2018 [216], as well as 800 full depth-of-discharge cycles to 70% of the nominal capacity for 1.8 Ah cells in 2020, estimating an energy density for EV applications of 420 Wh kg$^{-1}$ and 700 Wh l$^{-1}$ when scaled to commercial designs [220].

ASSBs based on the above discussed Li/PEO/LiFePO$_4$ technology have been successfully launched onto the market as Lithium Metal Polymer (LMP®) batteries by the Bolloré Group (France) over the past decade [204]. This battery is an evolution of a prototype developed in the 1990s by Hydro-Québec (Canada) and 3M (US) [204], which involved a lithium-metal anode, a PEO membrane dissolving LiClO$_4$/LiTFSI, and a VOx cathode (see Figure 5c) [217], thus ensuring an energy density of the order of 100 Wh kg$^{-1}$ with a life of 600 cycles at 80% depth of discharge [204]. A Bolloré’s subsidiary, Blue Solutions, commercialized a Li/PEO/LiFePO$_4$ battery with an energy density of 180 Wh kg$^{-1}$ which delivers over 1300 cycles within 60 and 80 °C, formed by ultrathin cathode, polymer electrolyte, and anode films. Cells are connected in series in a module; several modules are then connected in series in a full battery pack as shown in Figure 5d[204]. This technology is suitable for the automotive market, as demonstrated by the successful launch in 2011 of a car sharing program called Bluecar[204], employing small EVs (Figure 5e) [218] with a maximum speed of 120 km h$^{-1}$ and drive range from 150 to 250 km [221]. Recently, the Bolloré Group commercialized through its Brittany division a bus in two formats (6 and 12 m long, see Figure 5f) [218], named Bluebus, using an LMP battery [221,222]. The automotive applications of the LMP battery suggest the ASSBs with polymeric electrolytes for diverse applications that do not require low-temperature. Although, the targets established in the European SET Plan could be hardly achieved with the
current Li|LiFePO$_4$ chemistry (>400 Wh kg$^{-1}$ and >750 Wh l$^{-1}$ at the cell level as well as >250 Wh kg$^{-1}$ and >500 Wh l$^{-1}$ at the pack level by 2030). For instance, the LMP 63 pack proposed by the Bolloré Group is formed by 9 ASSB modules and has an operating voltage range from 450 to 648 V, an energy of 65 kWh, and an overall weight of 450 kg (42 kg per module) [223]. The same company developed through its Bluestorage division battery packs for stationary energy storage, i.e., the Blue LMP 250 and Blue LMP 400 252 and 392 kWh with overall weight of about 2250 and 3270 kg, respectively, (corresponding to 740 kg and 920 kg per rack and 42 kg per module) [224]. Therefore, the relevant safety and high-thermal stability of lithium-metal polymer configurations appear particularly adequate for developing load-balancing battery packs integrated in smart grids as well as power storage systems coupled with intermittent renewable energy sources and off-grid generators. The replacement of LiFePO$_4$ with high-voltage insertion/intercalation compounds, to date only demonstrated in proof-of-concept studies (see Figure 4f), might further extend the applicability of ASSBs using polymeric electrolytes, possibly matching the present requirements for long range electric cars in terms of both energy density and cost, as well as the European SET Plan targets. In this regard, a remarkable breakthrough in the upcoming years might be achieved by developing lithium-sulfur cells using high-viscosity glyme oligomers or solid low-molecular-weight glyme polymers. However, as discussed in the following section, the current Li-S technology suffers from various shortcomings needing substantial efforts to obtain commercially relevant results.

4- Generation 5: lithium batteries based on conversion cathodes

4.1 - Lithium-Sulphur Batteries

Lithium-Sulphur-Batteries can achieve high specific energies (> 450 Wh/kg [225]), are based on low cost raw materials and thus, are a highly attractive generation 5 cell technology [226]. Li-S-cells use lithium metal anodes, liquid electrolytes and conversion cathodes based on elemental sulfur mixed with carbon. The overall reaction is $S_8 + 16 e^- + 16 Li^+ \leftrightarrow 8 Li_2S$ with an equilibrium potential of 2.1 V vs. Li/Li$^+$. Typically cells are assembled in the charged state, and during discharge, lithium is stripped while sulfur is converted to lithium sulfide involving several electrochemical steps and various intermediate sulfur species (polysulfides). During charging, the cathode conversion is reversed back to elemental sulfur and lithium is plated on the anode. Hence, the anode chemistry in Li-S-batteries is per se very comparable to other lithium metal batteries. However, the sulfur conversion chemistry causes several specific characteristics, which need to be considered for lithium anode design. Electrolytes are typically
based on LiTFSI in ether based solvents (DME, DOL), while carbonate solvents are mostly
avoided due to decomposition reactions with polysulfides when not confined in the cathode
porosity. Furthermore, polysulfides are soluble in the electrolyte and can diffuse and participate
in side reactions on the anode surface. This involves continuous passivation of the lithium
surface and the reduction of dissolved long-chain polysulfides to short-chain species causing
self-discharge and low charge efficiency of the cells, also known as „polysulfide shuttle“ [227].
Lithium nitrate was found to be an effective additive in participating in anode surface passivation, thereby reducing the polysulfide shuttle current and enabling high coulombic efficiencies [228]. Thus, the combination of DME/DOL, LiTFSI, and LiNO₃ is a well-
established electrolyte system for exploring lithium-sulfur batteries. Under lean electrolyte
conditions (< 3 µl/mg sulfur), Li-S cells suffer from a fast capacity fade, and prototype cells
typically do not achieve more than 100 charge/discharge cycles. The low cycle life is still the
major obstacle for the technology breakthrough.

Consumption of electrolyte components in general and polysulfide in particular through
reduction at the anode surface and the structural anode degradation are known to be the major
failure mechanisms [229]. For these reasons, innovations in anode protection or structural
design are of high relevance for improving Li-S cell performance. This section provides a
review on the most promising lithium anode concepts from a holistic point of view and assesses
critical parameters to be considered for application-relevant cells.

Several review articles on metallic lithium anodes for lithium-sulfur cells have been published
[230–237] describing the major obstacles and first approaches how to tackle the complex issue
of a highly reactive anode and sulfidic intermediate species intrinsically derived from to the
conversion mechanism. It is vital, however, to evaluate the investigated approaches in regard
of a multilayered cell as a system comprising both active and inactive components [238–241].
In addition, comparability of results is frequently hampered as the electrochemical evaluation
is often generated in coin cells with varying electrolyte amounts (sometimes in excess),
separator types & thicknesses, and cathode porosities [240].

4.1.1 Concepts for lithium metal anodes in Li-S batteries

4.1.1.1 - Electrolyte adaption

In contrast to present lithium ion batteries, the electrolyte mass portion of presently developed
Lithium sulfur batteries is as high as ~ 50 wt-% [240]. In addition, the electrolyte according to
the state of the art dissolved a high amount of highly reactive polysulfides that indirectly stress
the anode. It is known that LiNO$_3$ in combination with the lithium polysulfides play an important role to passivate the lithium anode [242–245]. This effect is also dependent on the sulfur loading in the cathode [246]: below a certain threshold concentration of sulfur species, polysulfides do have a beneficial effect, similar as already mentioned in chapter 2.2 describing Li$_2$S as a stable layer. Above a certain sulfur amount, the current density is increased, so the probability of local spots rises where the electric field is increased and cannot be fully compensated by the anions and cations of the electrolyte [247]. Thus, dendrite formation or mossy lithium growth is accelerated [240]. Electrolyte additives such as lanthanum nitrate have a certain beneficial effect on the lithium stability as well given that it decreases the reducibility of metallic lithium and slows down the electrochemical dissolution/deposition reaction [237]. Hence, adapting the electrolyte is a key parameter to indirectly reduce anode corrosion by employing new solvents and new additives apart from lithium nitrate. So-called “sparingly (polysulfide) solvating electrolytes” are discussed as a sensible and effective approach in order to reduce the corrosive species on the anode side [248–253]. Significant improvement of the cycle stability due to fluorinated ether [254], even in multi-layered pouch cells has been successfully shown [255]. However, fluorinated solvents usually have a high mass density being detrimental for the overall specific energy. Nevertheless, electrolyte with high mass density can potentially tackle the issue of a low volumetric energy density of Li-S cells. Moreover, in order to implement a more stable carbonate-based electrolyte and to utilize the sulfur almost to a theoretical extent, confining of sulfur into polymers like polyacrylnitrile (PAN) [256,257] or microporous carbons [258] is possible, but mostly lacks of low sulfur weight portion in the cathode being detrimental for the overall energy density of prototype cells. A prototype cell using carbonates comprising 280 Wh/kg was introduced [259]. Another LiNO$_3$-free electrolyte in lean electrolyte regime and limited polysulfide solubility based on DOL was implemented in a multi-layered pouch cell reaching 300 Wh/kg [260]. The combination of symmetric and “non-symmetric” ethers with varying alky chains also adjust the polysulfide solubility [261]. A special case of sparingly polysulfide electrolytes are solid state concepts, the most promising ones are sulfide-based inorganic glasses (P$_2$S$_5$–Li$_2$S) leading to almost theoretical sulfur utilization [262] and reasonable power capability [263].

Works on other lithium conductive salts than LiTFSI, such as lithium trifluoromethyl-4,5-dicyanoimidazole were presented to restrict the solubility of polysulfides as well [264], and lithium bis(perfluoroethylsulfonyl)imide LiBETI is known to form more stable thin and compact SEI film containing mainly LiF on lithium [265]. The beneficial effect of lithium halides, especially LiF, was already discussed in chapter 2.1. Summarizing, for the holistic
development of enhanced Li-S cells the electrolyte takes a very complex role. Besides its interaction with the anode interface, possible limitations for maximum content and utilization of sulfur need to be considered. The power capability may be limited by the electrolyte conductivity depending on temperature and might vary over state of charge. Furthermore, the content and the specific mass density of electrolytes may have a drastic impact on the specific energy of Li-S cells.

Figure 6. Overview of the main stabilization methods for lithium metal anodes in liquid electrolyte being adapted for Li-S batteries. a) "in-situ" SEI with carbonates for confined sulfur species in the cathode pores, additives for in-vivo SEI formation and sparingly polysulfide solvating electrolytes as approach to intrinsically hinder anode corrosion by polysulfides. b) “ex-situ” artificial SEI by gassing, sputtering of inorganic layers, and application of polymers. c) host engineering by conductive and inconductive frameworks, spacer concepts, and alloys.
4.1.1.2 - Coatings

Inorganic ceramic coatings on lithium anodes - Intuitively, one approach is to generate a dense electric inductive but ion-conductive protection layer on metallic lithium. Li$_3$N is a potential candidate (compare chapter 2.2) and can be applied via simple reaction with nitrogen at room temperature.[266] However, Li$_3$N decomposes at ambient conditions and needs to be carefully handled. In addition, it is also a quite brittle solid leading to cracks during stripping and plating of lithium. Likewise, P$_4$S$_{10}$ was evaluated for having a positive effect on the lithium anode in lithium sulfur cell [267]. However, the employed electrolyte amount or rather excess being used for obtaining this finding should be considered as well. The in-vivo generation of Li$_3$PS$_4$ by using polysulfides and P$_2$S$_5$ had a beneficial effect on the cycling stability in symmetric cells [268]. A further candidate is Li$_{1.5}$Al$_{0.5}$Ge$_{1.5}$(PO$_4$)$_3$ being already implemented in a one-layered pouch cell [269]. LIPON (lithium phophorus oxynitrides) as system was implemented in a multilayered pouch cell reaching promising 300 Wh/kg [270]. In addition, mixed ion- and electron-conductive layers are discussed for Li-S cells, although the electrolyte amount was not stated and this approach might not be generally able to compensate the volume change during plating and stripping [271].

Polymeric or polymer-like coatings with and without fillers - In order to tackle the obstacle of the volume changes during lithium plating/stripping, the application of a polymeric protecting film is obvious. Nafion-coated separators are known to prevent polysulfide diffusion to the anode at least partially [272], and dual-functional polymer coating consisting of Nafion/polyvinylidene difluoride (PVDF) minimized leakage currents leading to decreasing diffusion of soluble polysulfides and thereby suppressed self-discharge [273].

In order to transfer these films into a Li$^+$-ion-conductive film, usually, lithium conductive salts are implemented. LiTFSI and PEO is a very common combination [274], however, those films per se can be only run at elevated temperatures [274]. At elevated temperature, the PEO chains show similar discharge slopes like an ether based solvent which is a strong hint that they dissolve lithium polysulfides. The beneficial role of ceramic fillers, both ion-conductive and non-conductive [275–278], has not been fully understood yet. It should be pointed out that fillers with high mass density increase the ionic conductivity, but might lower the overall energy density of the final prototype cell [279]. In combination with liquid electrolyte [280], polymers can also swell and might not provide decent protection then [281]. A further approach is to use lithium surfaces being treated with polysiloxane [282], with a crown ether [283], or with a
organosulfide-plasticized solid electrolyte interphase [284] leading to improved cycle stability at least compared to the respective reference materials.

**Lithium alloys with other elements** - Beneath constant electrolyte depletion and SEI formation, the volume change during plating and stripping causes breathing of the cells and dead lithium. Hence, metallic lithium can electrochemically alloy with the other elements in organic electrolyte at ambient temperature [285], and various alloys of lithium have been extensively investigated as anode materials for many years [286]. In order to balance the high electrochemical capacity of Li₂S, lithiated silicon was discussed as one promising material [287]. However, pre-lithiation has been a laborious process, and an all-over Cu foil current collector needs to be employed limiting the specific energy on Li-S battery pouch cell level. Therefore, employing alloys of lithium with metals such as magnesium or [288] can have a beneficial effect on the lithium stripping and plating as generic concept for lithium metal-based batteries. However, it should be noted that alloys increase the anode potential and lower the overall voltage window, so treating only the lithium metal anode’s surface with the alloy impacts less the overall energy density than using a completely alloyed anode. In addition, the electrolyte excess being employed during these studies might mask effects as well. Especially for the Li-S battery system, the alloyed anode (coating) needs to be compatible with the sulfur-species being dissolved in the electrolyte. The alloying element should not leak into solution and the resulting passivation layer should provide certain lithium ion conductivity.

**Scaffold, spacer, and filler concepts** - As already mentioned in chapter 2, plain metallic lithium has the intrinsic above-mentioned properties, such as continuous electrolyte depletion and SEI formation due to the highly reactive surface and volume changes during cycling. It is known from the Sand’s equation (vide supra) that the time for lithium dendrite formation is inverse proportional to the current density. Hence, a homogeneous distribution of the current is crucial to balance space-charge and to avoid local electric field build-up. Consequently, functional frameworks have been discussed as stabilizing scaffolds to facilitate lithium plating. These scaffolds can be divided into two main different types: (i) electrically non-conductive frameworks and (ii) electrically conductive frameworks.

Non-conductive frameworks have the beneficial effect that no SEI is formed due to the framework material itself and that the lithium can be plated on the bottom. For example, a fibrous Li₇B₆ matrix was presented in order to entrap lithium [289]. However, no decreasing of the areal current takes place and the lithium can push away those frameworks like a separator. Consequently, electronically conductive frameworks have some benefits despite the SEI.
formation. Conductive frameworks potentially enable a decrease of the areal current, and they are also potentially able to activate dead lithium [229]. However, plating on top of the framework is very likely and should be inhibited. In order to keep the mass fraction of inactive materials as low as possible, carbonaceous materials are ideal candidates in particular carbon fibers [290–295], graphene [293], graphene oxide [294] or hard carbon/stabilized lithium particle composites [296,297]. Hybrid lithophilic and lithiophobic gradients have been already presented in single-layered pouch cells [298]. Very often, the deposition of lithophilic sides, such as zinc oxides or silver are needed though. In contrast to sulfur-free battery systems, these frameworks should be compatible with the sulfur species in the electrolyte when implemented in Li-S cells. Moreover, the framework porosity should not take up additional excess of electrolyte resulting in a lower overall energy density on cell level.

Li₂S-based “anode free” concept - Lithium anode foils are commercially only available in thicknesses exceeding 50 µm. In addition, the handling and processing of the foil causes issues due to high ductility of this metal. For lithiated nickel-manganese-cobalt-oxides, it is known that lithium can be plated from this cathode material onto a lithophilic anode current collector (vide supra). A similar approach was published using Li₂S as lithium source [299]. Especially the volumetric energy density could be increased by plating only the required amount of lithium, however, the handling and/or encapsulation of Li₂S is challenging and the lithium ion loss during first charging for the SEI formation on anode side needs to be compensated.

4.1.2 Critical parameters for Li anode design in Li-S-Batteries

As explained above, the lithium anode design has a crucial impact on Li-S cell performance and often determines cycle life, power capability and energy efficiency. On the other hand, lithium excess as well as protective coatings, frameworks, new electrolytes etc. may impact the energy density of the cells. Therefore, the anode needs to be tailored considering the application-specific requirements.

Limiting factors for specific energy are inactive materials that have a high mass fraction in Li-S-cells, e.g. any copper/nickel current collector, especially when employed all over. In addition, electrolytes with high salt concentrations, high density solvents or rather fillers, and high overall electrolyte content drastically decrease the specific energy.

Limiting factors for volumetric energy density are inactive materials having a high volume fraction. E. g., thin current collectors (Ni, Cu), even when used all-over the area, can enhance
the volumetric energy density, as they allow a minimization of Li excess. The reaction between heavy metals such as Cu with the polysulfide should be considered though. In addition, Li and electrolyte excess limit the volumetric energy density of today’s Li-S-cells. First reports on “Li-free” anodes demonstrate the feasibility and potential of that concept, especially for established (lithiated) nickel-manganese-cobalt-oxide cathodes. The applicability of using Li₂S as single lithium source still needs to be further evaluated.

Limiting factors for power density is mossy lithium growth that depends on a critical areal charging current and eventually limits the charge rate of Li-S-cells. The above discussed 3D-framework-concepts are promising, however, the impact on energy density needs to be estimated as porous frameworks take significant weight and volume fractions of the cell, should to be wetted with electrolyte inactive mass, and also practically implemented in the prototype cell. The limiting current density and depletion of electrolyte especially negatively affects the cycle stability of lithium anodes. The limiting factors for production are techniques for thin or 3D Li film application as Li-S requires a specific range of film thickness (15-30 µm). Importantly, handling of sensitive films requires further development. Lithium passivation coatings for a better handling might differ from those suggested for other battery types to enhance the anode performance and in particular require compatibility with polysulfides.

4.2 - Lithium-air (Li-O₂) batteries

Lithium-air (Li-O₂) batteries operate by reversibly forming/dissolving Lithium peroxide (Li₂O₂) in the pores of a carbon cathode, while drawing/releasing oxygen (O₂) from air. The overall reaction is O₂ + 2 e⁻ + 2 Li⁺ ↔ Li₂O₂ with an equilibrium potential of 2.96 V vs. Li/Li⁺ [21,300–302]. Besides poor rechargeability due to parasitic reactions [303,304], practical realization relies on fully utilizing the high theoretical capacity of Li₂O₂ [305]. Only if the electrode porosity is filled by a large fraction with active material, significant capacity improvements can be achieved [305,306]. Yet, large Li₂O₂ pore occupation impedes mass and electron transfer. Achieving high reversible capacity requires maximizing Li₂O₂ packing densities through a detailed understanding of the oxygen reduction and evolution mechanism. Equally, parasitic chemistry is now understood to be curbed only through understanding its mechanism.

4.2.1 - Li-O₂ performance
Realistic capacities of Li-O₂ cathodes cause lots of confusion. This is because formal capacity (1168 mAh·g⁻¹, 2500 mAh·cm⁻³ Li₂O₂) is confused with theoretical capacity (Li₂O₂ including the minimum electron and ion conductor for Li₂O₂ to take place) and achieved true capacity (Li₂O₂ including the used electron and ion conductor)[305]. Given the positive active material O₂ is absent in the as made charged cathode, relating the capacity to the mass of carbon has become habitual, resulting in more than 10,000 mAh g⁻¹. As full capacity cycling is difficult, cyclability is often shown at, e.g., a fixed 1000 mAh g⁻¹, i.e. often < 10% depth-of-discharge. However, highly porous cathodes are back-filled with electrolyte. Figure 7a shows the true capacity per total electrode mass for three initial porosities, which are filled up to 80% with Li₂O₂. To achieve truly higher capacity than intercalation cathodes, filling the available pore space to the widest possible extent is crucial. Overly restricted depth-of-cycling results in no advantage. Fairly assessing true energy and cyclability requires values reported with respect to full electrode mass and volume. Li-O₂ cathodes could achieve higher true capacity than intercalation also in practice; key is high active material packing density and a small inactive/active material ratio.
Figure 7: a, True capacity of a Li-O\textsubscript{2} cathode as a function of capacity per mass of carbon for three cases of initial porosity (given by percentages above the curves), which is filled to 80\%. The black squares and red circles at 1,000 mAh g\textsubscript{carbon}^{-1} illustrate that respective true capacities vary strongly with electrode architecture. Value for the intercalation material LiFePO\textsubscript{4} is given by the dashed line for comparison. The insert shows the space filling of spherical Li\textsubscript{2}O\textsubscript{2} particles inside the porous electrode and the displaced electrolyte volume at 25,000 mAh g\textsubscript{carbon}^{-1} for 92\% initial porosity (indicated by the green circle). Adapted with permission from Ref.\[305\], NPG. b, Typical moles of O\textsubscript{2} and Li\textsubscript{2}O\textsubscript{2} involved upon discharge and charge. Adapted with permission from Ref.\[307\], NPG. c, Thermodynamics of alkali peroxides and superoxides. Standard potentials of the O\textsubscript{2}/MO\textsubscript{2} and O\textsubscript{2}/M\textsubscript{2}O\textsubscript{2} redox couples on the M/M\textsuperscript{+} scales with M = Li, Na, K. The scales are brought to a common scale based on their M/M\textsuperscript{+} standard potentials. The dashed horizontal line indicates the O\textsubscript{2}/KO\textsubscript{2} couple. The O\textsubscript{2}/LiO\textsubscript{2} potential is adopted from Ref.\[308\]. With permission from Ref.\[309\], Royal Society of Chemistry. d, Overview of the discharge and charge process with dominant disproportionation steps for the 2\textsuperscript{nd} e\textsuperscript{-} transfer and concurrent \textsuperscript{3}O\textsubscript{2}/\textsuperscript{1}O\textsubscript{2} release (centre). From Ref.\[309\], Royal Society of Chemistry. Steps are more detailed for discharge/charge on top/bottom. e, Parameters determining product morphology and degree of pore filling. f, Processes upon mediated discharge and charge and open questions regarding \textsuperscript{1}O\textsubscript{2} formation.
4.2.2 - Li-O₂ discharge

O₂ reduction during discharge in Li-O₂ batteries proceeds in two consecutive steps (Figure 7d) [310,311]. First, O₂ is reduced to superoxide (O₂⁻) at the carbon-electrolyte interface to O₂⁻*, which associates with Li⁺ to form the surface species LiO₂*. The desorption/adsorption equilibrium \( \text{LiO}_2^* \rightleftharpoons \text{Li}^+_{(\text{sol})} + \text{O}_2^{(\text{sol})} \) defines the extent to which associated LiO₂* is adsorbed at the surface or dissolved as \([\text{Li}^+ \cdots \text{O}_2]_{(\text{sol})}\), which can be anything between solvated free ions and solvated ion pairs or clusters [310]. Second, solid Li₂O₂ is believed to either form via electroreduction of LiO₂* or via disproportionation of the dissolved species. The former leads to a conformal Li₂O₂ coating up to a few nanometers in thickness [312], the latter to disc-like Li₂O₂ crystallites[313] that may assemble to toroidal particles of several hundred nanometers in size [314,315]. Disproportionation takes place via associated LiO₂ or clusters in solution or adsorbed on existing Li₂O₂ crystals [316,317]. While chemical experiments suggest that disc-like crystallites are a unique signature for disproportionation [318], the exact mechanism of toroidal particle formation remains to be clarified.

Increasing discharge capacities relies on facilitating solution growth since the Li₂O₂ volume formed via electrochemical reduction is limited by Li₂O₂’s poor electronic conductivity [300,303,311,319]. Using planar or low surface area electrodes, the capacity correlates directly with the formal Li₂O₂ layer thickness. In carbon cathodes with smaller pores, the Li₂O₂ particle size is limited to the pore size (Figure 7e). Whether the second electron transfer (surface mechanism) or solution mediated disproportionation (solution mechanism) prevails, is currently understood to be primarily controlled by the electrolyte’s solvation energy [310,311,320]. A high Gutman donor number (DN) of the solvent will drive Li⁺ solvation and consequently the solution mechanism [310]. Similarly, high DN anions [320] or trace H₂O [311] in the electrolyte shift the adsorption/desorption equilibrium towards dissolved species. In all cases, microscopy shows larger and less numerous Li₂O₂ particles in electrolytes with stronger solvation [311,320]. With decreasing current density, Li₂O₂ particles become larger and less numerous in line with nucleation and growth theory. At low currents or low overpotentials, solution mediated disproportionation generally dominates[321,322].

Concerning the surface mechanism, much speculation has been going on whether the higher conductivity in defect-rich or amorphous Li₂O₂ [323–325] could explain particle sizes of few tens of nanometers formed by electrochemical reduction. Actual electrochemical discharge, however, at realistic current densities and in prototype electrolytes for surface discharge
have established a maximum formal film thickness of about 5–10 nm, depending on the applied current. Theoretical studies explain the electron transfer by polaron-hole conduction and electron tunneling through crystalline Li$_2$O$_2$ [312,319,326]. Given the exponential tunneling resistance increase with Li$_2$O$_2$ thickness [312,327], particle growth by electrochemical reduction is self-limited, indicating that Li$_2$O$_2$ formed via the surface mechanism would always result in film-like morphologies.

Capacity of Li-O$_2$ batteries is primarily electron transport limited, i.e., electrode passivation with Li$_2$O$_2$ formed via the surface mechanism. At the end of galvanostatic discharge, widely Li$_2$O$_2$ covered carbon surface increases the local current density and the electrode potential drops[321,328]. The contribution of the surface mechanism would rise until all carbon surface is Li$_2$O$_2$ covered up to the maximum tunneling thickness. In electrolytes promoting solution discharge, also mass transport limitation through the increasingly tortuous network of Li$_2$O$_2$ and carbon is considered [329]. This suggests next to electrolyte solvation and current density the species mobility (O$_2^-$, Li$^+$, O$_2$) as a third parameter to control discharge capacities [330].

An interesting aspect arises from how the size and number density of Li$_2$O$_2$ particles varies with increasing solvation (solvent or anion DN, H$_2$O content), which is usually explained by the shifting partition from surface to solution mechanism. Less numerous, but larger particles are many times associated with an increased fraction of solution mechanism and accelerated disproportionation. However, this explanation contradicts nucleation and growth theory, if only homogenous nucleation in solution is considered: the concentration of dissolved Li$^+$ and O$_2^-$ and homogenous nucleation rates are highest in strongly solvating electrolytes. Consequently, the Li$_2$O$_2$ particle number density should be highest and the Li$_2$O$_2$ particle size smallest in strongly solvating electrolytes. Yet, just the opposite is observed [310,311,320]. In a multiscale modelling study Franco et al.[328] give a reasonable explanation for that: Li$_2$O$_2$ nucleation takes place via heterogeneous nucleation at the carbon surface, where nuclei form via the surface mechanism. Li$_2$O$_2$ particle growth above the tunneling limit takes place via solution mediated disproportionation. Hence, an increasing fraction of the surface mechanism would lead to more numerous and smaller Li$_2$O$_2$ particles, as observed experimentally.

Interestingly, some recent SEM [331–334] and TEM [314,315] studies raise doubts about the prevalence of the surface mechanism in low donor number electrolytes, although they were not interpreted this way. SEM and TEM micrographs of electrodes after discharge in supposedly prototype electrolytes for surface mechanism (dry DME, MeCN) show particles from tens to hundreds of nm, contradicting that they could have formed via the surface mechanism.
4.2.3 - Li-O₂ charge

Only recently, knowledge about the recharge mechanism has seen progress to a similar level as discharge. Generally, O₂ evolution starts just above 2.96 V with steadily rising voltage, sometimes with plateaus. It is agreed that oxidation has low kinetic barrier, and that the voltage rise stems mostly from accumulating side products and to a lesser extent from increasingly difficult electron transfer. Most recent understanding settled at a two-step process: First, formation of a superoxide intermediate, which may either be a Li-deficient Li₂₋ₓO₂ phase or LiO₂. Second, O₂ evolution via superoxide disproportionation (Figure 7d). Superoxide formation on charge can proceed at low voltages and has been proposed theoretically [335] and shown experimentally via XRD [336], PITT [337,338], XANES [339], and RRDE [338,339]. Whether O₂ evolution from the superoxide intermediate involves a second electron transfer or only disproportionation is still controversial, although evidence accumulates that the latter can fully explain things [309,338,339]. O₂ forms from the onset of charge and its fraction, being sensitive to the cations present (see next section), can only be explained with disproportionation [309]. While also solid superoxide may disproportionate[317], kinetically relevant appears soluble LiO₂. Hence, similar to discharge, the solvent becomes the governing factor as reported by Lu et al [338,339]. RRDE has shown that even in low DN dissolved LiO₂(sol) forms. XANES showed surface LiO₂ in high DN solvents but its absence in low DN solvents. Disproportionation in high DN solvents was shown via SEM, where after charging large Li₂O₂ particles, nanocrystalline, lamellar Li₂O₂ was seen. Similar recrystallization to nanocrystalline Li₂O₂ was also seen for low DN solvents by XANES [339]. Disproportionation as the O₂ release step is paramount for understanding parasitic chemistry, as discussed in the following.

4.2.4 - Li-O₂ parasitic chemistry

Parasitic chemistry keeps buzzing the community. The equation 2 Li⁺ + O₂ + 2e⁻ ↔ Li₂O₂ directly describes the ratios of charge passed and species converted, which have to match during discharge and charge. However, as illustrated in Figure 7c, typically the e⁻/O₂ ratio on discharge is ~2 while only ~50-95% Li₂O₂ form [309,340]. On charge, substantially less O₂ evolves as expected from the charge passed and the Li₂O₂ consumed. These discrepancies have for long been ascribed to the potential reactivity of superoxide and peroxide. However, theoretical calculations revealed prohibitively high barriers for the potential onset reactions: nucleophilic substitution, H-atom abstraction and H⁺ abstraction. Strategies to mitigate the irreversibilities using materials with higher stability against superoxide and peroxide proved only partially successful [341–344].
Singlet oxygen ($^1\text{O}_2$) could be the missing reactive species as first suggested by Hassoun et al. to possibly form upon Li$_2$O$_2$ oxidation at high voltages [345]. $^1\text{O}_2$ is the first excited state of ground state triplet oxygen being ~1 eV higher in energy. This idea was occasionally picked up, but experimental prove was hindered by the difficulty to detect $^1\text{O}_2$. Small amounts could first be shown to form upon charging between 3.55 and 3.75 V using operando EPR [346]. The used spin trap was, however, unable to measure during discharge and higher charge voltages. The finding could partly explain parasitic chemistry beyond 3.55 V. On discharge and from the onset of charge (where always less than 1 mol O$_2$ evolved per 1 mole of Li$_2$O$_2$ oxidized) parasitic chemistry could not be clarified [340,343,347].

To comprehensively investigate involvement of $^1\text{O}_2$, Freunberger et al. developed methods to sensitively and quantitatively detect $^1\text{O}_2$ over the entire relevant voltage range during discharge and charge of metal-O$_2$ cells [304]. The 1270 nm emission during the $^1\text{O}_2$ to $^3\text{O}_2$ decay gave direct unambiguous proof for $^1\text{O}_2$. To be more sensitive and quantifiable, 9,10-dimethylanthracene (DMA) was identified as a suitable $^1\text{O}_2$ trap, fulfilling all requirements in the cell environment. DMA forms with $^1\text{O}_2$ selectively its endoperoxide form (DMA-$^1\text{O}_2$); both species are stable in the relevant voltage range between 2 and ~4 V vs Li/Li$^+$; and the conversion can be measured using ex-situ HPLC of extracted electrolyte or by in-situ fluorescence.

$^1\text{O}_2$ has been shown to form both during discharge and from the onset of charge and with growing rate as the charge voltage rises, which resembles the rates at which parasitic reactions occur in Li-O$_2$ cells, Figure 7b. Given that $^1\text{O}_2$ accounts for the majority of parasitic reaction products on discharge and charge, $^1\text{O}_2$ arises as the biggest hurdle to cycle Li-O$_2$ cells by reversibly forming/decomposing Li$_2$O$_2$. $^1\text{O}_2$ not only decomposes the electrolyte [304,348], but also carbon [343,349] and redox mediators[350]. As a means to counter $^1\text{O}_2$-related side reactions, the DMA was shown to reduce parasitic products on discharge and charge by trapping $^1\text{O}_2$. Further, the $^1\text{O}_2$ quencher 1,4-diazabicyclo[2.2.2]octane (DABCO) was shown to even more strongly reduced parasitic chemistry by physically deactivating $^1\text{O}_2$. However, DABCO is unstable above 3.6 V. The mono-alkylated DABCONium proven as an effective, more oxidation stable quencher [351].

Formation of $^1\text{O}_2$ is now understood to predominantly stem from superoxide disproportionation

$$2\text{O}_2^- \rightarrow \text{O}_2^{2^-} + x\text{O}_2 + (1-x)^1\text{O}_2 \quad (1)$$

rather than direct 2 e$^-$ oxidation of Li$_2$O$_2$ [309]. Another source is superoxide oxidation above $E^0_{\text{O}_2/\text{LiO}_2} + E\left( ^1\text{O}_2 \leftarrow ^3\text{O}_2 \right) \sim 3.26 \ldots 3.56$ V [304] as well as Li$_2$CO$_3$ oxidation [352].
Superoxide disproportionation is involved on discharge and charge as discussed above. **Figure 7c,d.** With this recognition, a unified mechanism of $^1\text{O}_2$ generation has been established with the Lewis acidity of the cations involved in the disproportionation reaction governing the $^1\text{O}_2$ yield [309]. The cation controls the relative thermodynamic stability of (su)peroxide and hence the fate of the initial one-electron reduction product superoxide (**Figure 7c**). Li$^+$ or Na$^+$ as strong Lewis acids favour peroxide, albeit only slightly in the case of sodium [353]. K$^+$ and even weaker Lewis acids (e.g., quaternary ammoniums like tetrabutylammonium (TBA$^+$) and imidazolium) favour the superoxide [354]. The latter constitute often-used ionic liquid electrolytes.

While stronger Lewis acids drive disproportionation, the $^1\text{O}_2$ fraction grows with decreasing Lewis acidity of the cation, causing insignificant $^1\text{O}_2$ with H$^+$ and strongly growing fractions with Li$^+$ and Na$^+$. Importantly, weakly Lewis acidic cations that alone do not drive disproportionation boost $^1\text{O}_2$ fractions when combined with strong Lewis acids. DFT calculations revealed that weak Lewis acids open pathways that bypass the otherwise most unfavourable reaction steps towards $^1\text{O}_2$. This allows TBA$^+$ to be used as a probe for disproportionation steps. Disproportionation must be involved if presence of TBA$^+$ increases the $^1\text{O}_2$ yield. This way, larger $^1\text{O}_2$ yields in mixed Li$^+/\text{TBA}^+$ electrolytes as compared to pure Li$^+$ electrolytes verified that disproportionation is the O$_2$ evolving step on both discharge and charge (**Figure 7d**) [309].

So far, parasitic chemistry remains the major concern in Li-O$_2$ batteries. Concluding about the impact of any measures (electrolytes, electrodes, catalysts, mediators, …) requires multiple quantitative analyses of the O$_2$ and Li$_2$O$_2$ inventory and of side products [303]. Qualitative measures cannot replace quantitative integral methods and cannot support claims of reversibility.

### 4.2.5 - Mediated Li-O$_2$ chemistry

The difficulties to reversibly fill the pore space with insulating Li$_2$O$_2$ at high rates and associated side reactions require countermeasures. Redox mediators could potentially mitigate all these problems by shuttling electrons between carbon surface and O$_2$ or Li$_2$O$_2$, thereby forming/decomposing Li$_2$O$_2$ distant from the surface at high rate and low overpotentials (**Figure 7f**) [355–358]. Upon discharge, mediators may act via outer or inner sphere pathways that differ in whether or not free superoxide is involved [358,359]. The relative absence of superoxide was suggested to mitigate side reactions on discharge. On charge mediators were
suggested to reduce side reactions by reducing the recharge potential [360]. Classes and some features of reduction and oxidation mediators have been reviewed comprehensively in, e.g., Refs [357,361]. However, the major open questions remain in the role of $^1$O$_2$ during mediated O$_2$ reduction and evolution. It is unknown whether inner sphere reduction that forms Li$_2$O$_2$ via disproportionation of LiMO$_2$ intermediates ($2$ LiMO$_2$ $\rightarrow$ Li$_2$O$_2$ $+$ 2 M $+$ $^1$O$_2$) [358] forms $^1$O$_2$ and, if yes, what governs its fraction. Equally, the mediated peroxide oxidation mechanism is unknown and whether the nature of the mediator may allow to suppress $^1$O$_2$ generation therefrom. More or less severe decay of the mediation effect suggests that $^1$O$_2$ is to some extent involved with both oxidation and reduction mediators [350]. Only detailed knowledge of the underpinning mechanisms will allow progress towards fully reversible Li-O$_2$ cells.

5 - General considerations towards SET plan targets for 2030

To enable the widespread commercialization of Li metal batteries, substantial efforts are required, in particular to stabilize the Li anode. Despite the multitude of protection strategies proposed so far, using highly reactive metallic Li in liquid cells still appears very challenging. Particularly because the safety issues associated with the presence of the flammable organic electrolytes remain. To guarantee safe operation of Li metal anodes, using non-flammable solid electrolytes is planned starting from 2025 with Generation 4 all-solid-state cells.

Lithium metal ASSBs with ISEs are considered one of the most promising energy storage technologies for automotive and stationary applications. Implementing an ISE with higher mechanical and electrochemical stability than organic liquid electrolytes would enable to use lithium metal as an anode (coupled with a high voltage cathode material), expediting the development of high voltage battery systems with enhanced energy density. The Interuniversity Microelectronics Centre (IMEC) of Leuven Belgium, after recently presenting an ASSB with a volumetric energy density of 400 Wh L$^{-1}$ at a charging speed of 0.5 C, aims to produce by 2024 a solid-state battery with an energy density of 1000 Wh L$^{-1}$ at 2-3 C (charging time of 20-30 min) [362]. These results bode well for the achievement of the performance targets in terms of charging time (12 min for 70-80% ΔSOC), volumetric (>750 Wh L$^{-1}$) and gravimetric (>400 Wh kg$^{-1}$) energy densities defined for a battery cell by the European SET-Plan Action 7 for 2030. Despite the significant improvements, the main challenge remains to stabilize the lithium metal and high-voltage cathode/electrolyte interfaces, considered crucial for long battery lifetimes. As highlighted by Randau et al. [163], further research is required to develop
protective coatings for high voltage cathode. Furthermore, achieving electrolyte thickness below 50µm, in-situ generation of the anode, and cathode areal capacities higher than 5 mAh cm\(^{-2}\) would be necessary to further improve the battery performance. An exceptional result was recently achieved by Samsung [141], where a 0.6 Ah pouch lithium metal cell (using a Ag-C nanocomposite anode for in-situ uniform deposition of Li metal) was recently developed. A record energy density of 900 Wh L\(^{-1}\), areal capacity >6.8 mAh cm\(^{-2}\), and lifetime of 1000 cycles was achieved. As previously described, several techniques (coatings, nanocomposite electrodes and alloys) are already available to obtain uniform lithium metal deposition. However, their high costs still hamper the scalability of the process, making challenging to simultaneous achieve the cost targets (75 €/kWh for an automotive battery pack, or 0.05 €/kWh/cycle for stationary) and battery lifetime (2000 cycles for BEV, or 10000 cycles for stationary). ASSB using lithium metal and a polymer electrolyte represents a very attracting energy storage system since it holds the potentialities for achieving high gravimetric and volumetric energy, long cycle life and remarkable safety. The favourable characteristics of this technology have been suggested by a large number of studies on laboratory-scale cells, which might achieve about 300 Wh kg\(^{-1}\) and 500 – 600 Wh L\(^{-1}\) in optimized conditions. Scaling up to practical solid-state LMP cells by various companies (i.e., Hydro-Quebéc, 3 M, and Bolloré Group) has led to actual commercialization for automotive applications. Yet, the European SET Plan targets for 2020 (i.e., 350 Wh kg\(^{-1}\) and 750 Wh L\(^{-1}\) at cell level as well as 235 Wh kg\(^{-1}\) and 500 Wh L\(^{-1}\) at the pack level) have not been achieved. Despite promising results suggesting large room for improvement by optimizing cathode, electrolyte, and anode interphase, the targets expected for 2030 (i.e., >400 Wh kg\(^{-1}\) and >750 Wh L\(^{-1}\) at the cell level as well as >250 Wh kg\(^{-1}\) and >500 Wh L\(^{-1}\) at the pack level) appear to be even more challenging.

Generation 5 batteries relying on conversion cathodes may be the key to achieve and, in theory, well exceed the performance target of the SET plan. Lithium-Sulfur batteries have been successfully demonstrated for an UAV application in 2014 with a specific energy of 350 Wh/kg [363]. Since then, further improvement led to specific energies up to 470 Wh/kg in prototype cells [22]. Thus, Li-S-technology clearly surpasses the SET plan targets in terms of specific energy. Considering the high content of excess electrolyte and lithium, further energy density enhancement is expected by improving the cell chemistry. While the volumetric energy of today’s Li-S-cells is limited to below 500 Wh L\(^{-1}\), reducing this excess and the amount of passive materials may lead towards 700 Wh L\(^{-1}\) in the future. Another major challenge is the limited cycle life (<100 for high energy cells) which is mainly caused by electrolyte and/or lithium depletion. Consequently, stabilizing the anode/electrolyte interphase is key for
progressing the Li-S-technology towards the SET targets in 2030. Novel electrolytes, protective coatings, and/or innovative electrode design are expected to be enablers for enhanced future Li-S-cells. On the other hand, Lithium-air (Li-O\textsubscript{2}) batteries, which operate by reversibly forming/dissolving Li\textsubscript{2}O\textsubscript{2} at the cathode are in a much lower stage of development. They have the highest formal energy amongst all battery systems. Sometime quoted figures of 3500 Wh kg\textsuperscript{-1} are based pure Li\textsubscript{2}O\textsubscript{2} and hence unrealistic. As outlined in Section 4.2.1, key for truly higher capacities compared to Li-ion is to achieve at the end of discharge a maximum of Li\textsubscript{2}O\textsubscript{2} volume occupation and hence high active-to-inactive volume and mass ratio. When doing so (e.g., 80% volume occupation of the initial pore space) and when accounting for the total mass and volume of cathode (active, binder, carbon, electrolyte), separator and anode, theoretical limits of 1700 Wh kg\textsuperscript{-1} and 1850 Wh L\textsuperscript{-1} excluding housing are obtained [305]. The main challenges to realize at least part of this promise are: First, reversibly electrodepositing insulating Li\textsubscript{2}O\textsubscript{2} and filling the porous electrode to the largest possible extent at high rates. Second, avoiding parasitic chemistry, which decomposes cell components and causes poor energy efficiency and cycle life. These problems are interrelated and can only be solved in conjunction. Controlling superoxide disproportionation is key for large discharge capacities and efficient recharge. At the same time, disproportionation is the major step forming singlet oxygen, which is now recognized to cause the vast majority of side reactions. Mediated Li-O\textsubscript{2} chemistry may mitigate all these problems in conjunction, but only if the mechanisms are clarified in detail. Overall, only detailed knowledge of the underpinning mechanisms will allow progress towards fully reversible Li-O\textsubscript{2} cells.

For sake of clarity, the current status of Li-metal batteries compared to the performance targets of the EU Integrated Strategic Energy Technology Plan (SET-Plan) Action 7 for 2030 is summarized in Table 1.
Table 1 – Current status of Li-metal batteries compared to the performance targets of the EU Integrated Strategic Energy Technology Plan (SET-Plan) Action 7 for 2030. Considering the relatively low TRL of these cell chemistries, cost and manufacturing targets are omitted. For the same reason calendar life is omitted from the performance targets and only values at the cell level are compared.

<table>
<thead>
<tr>
<th>SET Plan Targets 2030 (at cell level)</th>
<th>Current Status</th>
<th>Li Metal Batteries</th>
<th>Generation 4: ASSB</th>
<th>Generation 5: conversion cathodes</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>inorganic</td>
<td>polymeric</td>
</tr>
<tr>
<td>TRL</td>
<td>4-6</td>
<td>commercial</td>
<td></td>
<td>5-7</td>
</tr>
<tr>
<td><strong>ENERGY</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gr. (Wh/kg): &gt;400</td>
<td>Performance</td>
<td>450 Wh/kg / 900 Wh/l</td>
<td>Estimated (laboratory scale): 300 Wh/kg / 500 – 600 Wh/l</td>
<td>&gt;450 Wh/kg / &lt; 700 Wh/L</td>
</tr>
<tr>
<td>Vol. (Wh/l): &gt;750</td>
<td>Most limiting factor(s)</td>
<td>ISE stability towards high voltage cathodes</td>
<td>Operating temperature &gt; 60 °C</td>
<td>SPE stability towards high voltage cathodes</td>
</tr>
<tr>
<td></td>
<td>Suggested measures</td>
<td>Develop more effective coatings</td>
<td>Electrolyte additives</td>
<td>New cell chemistries</td>
</tr>
<tr>
<td><strong>POWER</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gr. (W/kg): &gt;700</td>
<td>Performance</td>
<td>&lt; 500 W/kg - &lt; 1000 W/l</td>
<td>170 W/kg / 170 W/l</td>
<td>&lt; 500 W/kg - &lt; 1000 W/l</td>
</tr>
<tr>
<td>Vol. (W/l) &gt;1500</td>
<td>Most limiting factor(s)</td>
<td>High cell impedance</td>
<td>Low Li$^+$ transference number</td>
<td>Cathode conversion kinetics and electrolyte resistance</td>
</tr>
<tr>
<td>Charge time (min): 12</td>
<td>Suggested measures</td>
<td>Reduction of SE thickness</td>
<td>New electrolyte formulations</td>
<td>Improved electrolytes / electrode design</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>CYCLE LIFE</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(to 80% DOD)</td>
<td>Performance</td>
<td>&lt; 1000</td>
<td>ca. 1300</td>
<td>&lt; 1000 (&lt; 100 for high energy cells)</td>
</tr>
<tr>
<td>BEV: 2000 Stationary: 10000</td>
<td>Most limiting factor(s)</td>
<td>Contact issue at interfaces and dendrite growth</td>
<td>Stability of electrode/electrolyte interphase</td>
<td>Electrolyte / Anode depletion</td>
</tr>
<tr>
<td></td>
<td>Suggested measures</td>
<td>Stable interlayers and highly dense SE / cheaper and more effective coating techniques for lithium metal interphase</td>
<td>Electrolyte additives / New electrolyte formulations</td>
<td>New electrolytes / improved Electrolyte-Anode interphase</td>
</tr>
</tbody>
</table>
Authors’ contribution

A.V. conceived the manuscript structure, wrote section 1, conceived Figure 1 and Table 1, coordinated and revised the entire manuscript. K.T. wrote section 2, conceived and realized Figure 2 and realized Figure 1. R.S. wrote sections 3 and 3.1, realized Figure 3, and contributed to section 5. D.D. and J.H. wrote section 3.2 and conceived and realized Figure 4 and 5. S. D. wrote the section 4.1, revised section 1 – 4.1, and conceived and realized Figure 6. H. A. revised the entire manuscript, and implemented values for Table 1 and wrote section 4.1. S. K. coordinated and revised the entire manuscript. C.P. and S.A.F. wrote section 4.2, realized Figure 7, contributed to section 5 and revised the entire manuscript.

All authors have revised and approved the final version of the manuscript before submission.

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