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A perspective on the building blocks of a solid-state battery: from solid electrolytes to quantum power harvesting and storage

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Abstract. We present a comprehensive perspective on the fundamental components of a solid-state battery, starting from all-solid-state electrolytes and extending to quantum power harvesting and storage. Firstly, we enquire into the key characteristics that define an effective electrolyte. It is confirmed that the most promising electrolytes are semiconductors with a low work function, displaying behavior akin to anodes. Moreover, electrolyte classes composed of multiple dominant elements S, O, and Cl, such as argyrodites and oxyhalide electrolytes, show more significant potential than those with only one dominant element, like sulfide, oxide, and halide electrolytes. We also explore ferroelectric-electrolytes-based devices capable of both energy harvesting and storage. Ferroelectric materials possess exceptional properties that hold the promise to revolutionize the next generation of electronic devices. Likewise, we explore the realm of ferroelectric topological insulators (FETIs), which open up a new world of possibilities. The pioneering studies in this area are reported herein. Continuing our assessment, we investigate the domain of the small and provide an overview of the principles behind quantum batteries. We explain the general quantum model and present three different types of quantum batteries: the Dicke model, the spin-chain model, and the quantum thermal machines, whose function mechanism is based on Maxwell's demons (isothermal baths) and quantum refrigerators. By presenting these insights, we hope to contribute to advancing solid-state batteries and quantum power harvesting and storage technologies that maintain coherence even at macro sizes, paving the way for more efficient and sustainable electronic devices.

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1. Introduction

As we enter a new era, fossil fuels are no longer considered the forthcoming primary means of powering society. It has become imperative to seek sustainable technologies capable of producing and storing green energy [1]. Over time, batteries have undergone tremendous advancements and have now become the leading technology for energy storage. The most significant breakthrough in this field came with the development of Lithium-ion batteries (LIB), which have revolutionized energy storage, from portable charging devices to electric vehicles [2].

Liquid or gel electrolytes form the fundamental building blocks of LIBs, enabling Li⁺-ions to be transported from one electrode toward the other while cycling a battery. However, in some cases, when the charging process is too rapid, for example, due to low temperatures, cations cannot be inserted into the negative electrode. As a result, lithium accumulates at the surface, leading to the formation of dendrites. These dendrites are fractals and may eventually penetrate the separator, connecting both electrodes and creating a short circuit. The formation of dendrites poses significant safety risks due to the flammable character of liquid and gel electrolytes, the likelihood of thermal runways, and cathodes' oxygen release [3]. To overcome the challenges associated with traditional liquid and gel electrolytes, researchers have developed non-flammable solid-state electrolytes that exhibit cationic conduction, such as Li⁺, Na⁺, and K⁺ [4], [5]. Solid-state electrolytes are generally not flammable, offer increased stability to dendrite formation, and are compatible with high-potential electrodes, allowing for higher energy-density batteries as they permit using electrodes with higher theoretical capacity, such as Lithium (3860 mA.g⁻¹).

This perspective will start by looking into the driving forces of a solid-state battery, which are unified with the other energy harvesting, storage, and switching devices such as photovoltaics, thermoelectrics, capacitors, diodes, and transistors, among others, some of them punctually called to comparison here. A solid-state electrolyte is not just a variety of its liquid sibling; in solid electrolytes, usually, there is a main mobile species that leaves behind opposite sign-charging regions. The latter has implications for the velocity of these charge carriers and their position about the mobile ion, which affects how an electrical impedance spectroscopy EIS plot is analyzed, resulting in very different determinations of the ionic conductivity for similar data.

On the other hand, the polarization that has traditionally been tried to be avoided is not worth the effort, as polarization arises spontaneously from the difference in Fermi levels electrode/electrolyte that may not be overcome from the tunneling of electrons from the electrode to the electrolyte, as the electrolyte is an insulator with a specific relative position of its lower energy of the conduction band in regards to the Fermi level or electrochemical potential of the conductor-electrode. Therefore, it is paramount to remember these principles, which are shared across solid-state devices. Lately, it has been shown [6] that electrolytes may demonstrate themselves as p-type semiconductors.

The analysis of the electrolytes will be based on the type of elements they contain and the impact on the DOI: 10.1039/D31A04228F their properties. Their crystalline structures or other properties, such as ferroelectricity, will also distinguish them for the same reasons pointed out before, similar to what happens in the photovoltaics world.

The quantum batteries will be discussed in all their known variations. Their macro-size versions are already working prototypes and may harvest thermal energy besides electrical energy storage through coherent properties.

1.1 Features enabling energy storage

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The ionic conductivity in solid electrolytes depends on parameters such as the mobile ions' concentration, size and mobility, lattice structure and defects, and conduction mechanisms. Performing solid electrolytes, like liquid electrolytes, display mass and charge transport. The solid electrolytes herein typically exhibit cation conductivity as opposed to liquid electrolytes, which count with both ion mobilities, thus creating the need for intensive research on ionic transport in solid electrolytes to obtain fast ionic conductivity at room temperature (RT).

In solid electrolytes, ions move through disordered sublattices within an enclosed, cage-like framework. The equation expresses the ion conduction in solid-state electrolytes $\sigma_i = n_i q \vartheta_i$ where n_i , q, ϑ_i are the mobile ion concentration, the elementary charge, and the ionic mobility. These parameters are dependent on the temperature following an Arrhenius-type behavior. The ionic conductivity is related to the diffusivity of mobile ions - which is given by the Nernst-Einstein equation, $D_i = \frac{\sigma_i k_B T}{n_i z_i^2 q^2}$, where k_B and T are the Boltzmann's constant and the absolute temperature, z_i is the number of transferred charges per ion, and e is the unitary charge of the electron.

Ionic diffusion in inorganic materials consists of ions hopping between the adjacent available sites, overcoming a higher energy barrier, the activation energy, due to geometry constraints and electrostatic interactions. Generally, two different hopping mechanisms can be distinguished: the direct and knock-off. In the direct hopping mechanism, an individual ion hops from one lattice site to its adjacent vacant site. In contrast, the knock-off-hopping is due to an indirect interstitial mechanism where an interstitial atom pushes the matrix atom toward another interstitial site and realizes ionic transport.

In solid electrolytes, creating highly disordered defects for ions to move, designated Schottky and Frenkel point defects, is necessary to enhance the ionic conductivity. According to the Arrhenius model, the ionic conductivity in crystalline solids is given by the equation $\sigma_i = \frac{A}{T}e^{-\frac{E_A}{k_B T}}$, where *A* is a pre-exponential factor, and *E*_A is the activation energy of diffusion. The predominant carriers for ion conduction are point defects in solids; this characteristic excludes most inorganic materials due to the lack of functional defects for ion migration or high activation energy. Thanks to the advances in

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crystal structure engineering, it was discovered that only a limited number of materials sharing nume particular types of structures could diffuse Li⁺ and Na⁺ ions efficiently. However, minor concentration substitutions of the monovalent mobile cation by multivalent cations with a high ionic radius may tailor new solid electrolyte solutions.

However, the fundamentals of solid-state batteries surpass the properties associated with the internal impedance associated with ionic transport, which is a consequence, not the driving force for the ionic and electric mechanisms within the cell.

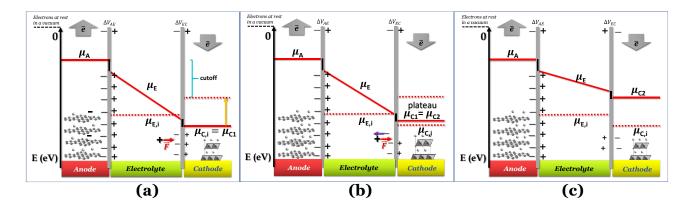


Figure 1. Schematic discharge of a battery as the function of the chemical potentials of the electrodes and electrolyte; **a** cell was charged and started to discharge; $\Delta V_{battery cell} = \Delta V_{AE} + \Delta V_{EC}$; **b** a new phase is formed at the cathode, and the two phases become at equilibrium ($\mu_{C1} = \mu_{C2}$), corresponding to the plateau voltage; $A^+ + \bar{e}$ (e.g. $A^+ = \text{Li}^+$) forms more μ_{C2} ; $\Delta V_{battery cell} = \Delta V_{AE} + \Delta V_{EC}$; **c** as the chemical potential overcomes the intrinsic chemical potential of the electrolyte, both surfaces of the electrolyte are charged positively; $\Delta V_{battery cell} = \Delta V_{AE} - \Delta V_{EC}$. The black lines at the interfaces are the walls of the potential quantum well/barrier related to interfacial impedances/resistances. **Note:** For simplification, no band gap is represented here for the electrolyte; however $\mu_{E,i}$.= $-(E_c + E_v)/2 = -E_g/2$, where E_c , E_v , E_g are the minimum energy of the valence band, and the band gap, respectively; $\mu_{E,i}$ is the chemical potential of the electrolyte, and cathode (phase 1). \vec{F} coulombic force. Schematics with the band gap of the electrolyte in Fig. 4.

The driving force in a battery is associated with the necessity of aligning the electrochemical potentials, according to the equation,

$$\bar{\mu}_j - \bar{\mu}_i = \mu_j - \mu_i + z_i e(\phi_j - \phi_i) \tag{1}$$

where *j* and *i* are different species in electrical contact, as in a heterojunction; $\bar{\mu}_j$, μ_j , and ϕ_j are the electrochemical potential or Fermi level for the solid species, chemical potential, and surface potential of species *j*; $z_i e \phi_j$ is an electrical work performed by *i* on the surface of *j*; z_i is the valence of the mobile charge (e.g., $z_{Li} = +1$). When the electrochemical potentials are equalized, $\bar{\mu}_j - \bar{\mu}_i = 0 = \mu_j - \mu_i + z_i e(\phi_j - \phi_i) \Rightarrow \mu_j - \mu_i = z_i e(\phi_i - \phi_j) = z_i e \Delta V_{ij}$. If both *i* and *j* are conductors, such as the electrodes, they may equalize their chemical potentials by exchanging electrons upon electrical contact. Nonetheless, if one of the species is an insulator or semiconductor, when possible, electrical double-layer capacitors EDLC of potential difference ΔV_{ij} will form at the interfaces to equalize the difference in chemical potentials. In a solid-state device such as a battery, the EDLCs store the

energy. Therefore, for energy storage to happen, an interface involving at least one semiconductor or insulator must be present. A battery cell's asymmetry relies on one or two double-layer capacitors formed at the interface negative electrode/electrolyte or electrolyte/positive electrode.

If the negative electrode is lithiated graphite, the battery cell may discharge without any change in the electrochemical potential of the negative electrode, as to the equilibrium $LiC_6 \Leftrightarrow (1 - 2x)LiC_6 + xLiC_{12} + x(Li^+ + \bar{e}), \ 0 \le x \le 0.5$ corresponds to a capacity of 177 mAh.g⁻¹, which is usually enough to discharge a traditional cathode with an experimental capacity inferior to 170 mAh.g⁻¹. At equilibrium, $xLiC_{12}$ and $x(Li^+ + \bar{e})$ form from LiC_6 , while the chemical potentials of the species are kept constant. Therefore, if the anode is lithiated graphite (LiC_6) , while discharging, the voltage versus time/capacity curve reflects the equilibria at the cathode side. The cathode's chemical potential increases while the electrons are transported to the cathode through the external circuit from the anode. Then, a new phase rich in mobile ions Li⁺ will start to form, and while the initial/charged cathode phase does not run out, the discharge curve is at a plateau because the chemical potential of both phases is equal (Fig. 1b), as pointed out earlier.

Here, we have considered $\mu_C \leq \mu_E \leq \mu_A$ for $E_0 = 0$ for electrons at rest in a vacuum at the surface of the correspondent material (Fig. 1a); in fact, the chemical potential of the electrolyte may also assume different relative values, $\mu_E \leq \mu_C \leq \mu_A$ and $\mu_C \leq \mu_A \leq \mu_E$, depending on the electrolyte [6], while the cathode and anode must have chemical potentials $\mu_C \leq \mu_A$ per definition of cathode and anode. If the electrolyte's potential $\mu_C \leq \mu_A \leq \mu_E$, the quantum well in Fig. 1 becomes a quantum barrier [6].

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When the Li⁺-poor phase runs out, the Li⁺-rich phase is the only present in the cathode, and its chemical potential will increase with more electrons and Li⁺ transported to the cathode. The cations Li⁺ move by the action of columbic forces as the electric field points to the cathode within the electrolyte (Fig. 1a).

While Gauss's law determines that the electric field outside an EDLC is null, as the cathode's chemical potential increases, the equilibrium between the electrolyte's electrochemical potentials and the cathode is dynamic. Fewer negative charges must accumulate at the electrolyte's surface on the cathode side. Therefore, the Li⁺-ion diffuses towards the cathode. In most electrolytes, the negative charges are negatively charged vacancies, Va_{Li}^- . Therefore, neutrality is restored when a Li⁺-ion occupies a Va_{Li}^- .

The potential difference between the two EDLCs in series sums and is equal to the battery's voltage, $\Delta V_{battery\ cell} = \Delta V_{AE} + \Delta V_{EC}$. As the chemical potential of the cathode increases due to electron transport from the anode, the voltage of the battery decreases, $\Delta V_{battery\ cell}$. When the cathode's chemical potential surpasses the electrolyte's intrinsic chemical potential, then the cell's potential

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difference is $\Delta V_{battery\ cell} = \Delta V_{AE} - \Delta V_{EC}$. The voltage will be zero when $\Delta V_{AE} = \Delta V_{EC}$. No EDLC forms the solution of the solution of

When the electrolyte is ferroelectric, the difference in chemical potentials at the interfaces is minimized [6], and the internal resistance is reduced to the activation energy associated with the mobility of the ions.

1.2 Solid electrolytes: Li/Na-SICON, argyrodite, garnet, perovskite; double perovskite; antiperovskites, halides, and oxyhalides

One key component of a battery is the electrolyte, making it one of the essential factors for battery performance. An ideal solid-state electrolyte is said to possess the following properties: (1) high ionic conductivity (Fig. 2); (2) low electrical conductivity; (3) a wide electrochemical window of stability to match with high-voltage cathodes (Fig. 3); (4) a high A⁺-ion transference number for inhibiting the formation of negative space charge near the anode and promoting uniform ion deposition; (5) sufficient mechanical strength to inhibit ion growth; (6) good electrode wettability and low interfacial resistance with the electrodes; (7) good flexibility to accommodate the change of the electrodes' volume during battery charge and discharge; (8) good thermal stability to ensure safe and stable operation of batteries at lower and higher temperatures; (9) easy fabrication and low cost.

It is possible to distinguish three types of solid electrolytes: Inorganic Electrolytes (ISEs) [7], polymer electrolytes (SPE) [8], and organic and inorganic composite electrolytes (O-ICSE). O-ICSE is the most recent class of electrolyte, as its composition combines ISE and SPE characteristics. However, studying these composites is a relatively recent subject, and much research still needs to be done. SPEs consist of a polymer matrix and Li salts without any liquid solvents. These electrolytes are lightweight and easy to shape, therefore capable of adapting to different battery structures. They also have good electrode wettability and flexibility that reduces the impact of electrode volume change during the charge-discharge process. However, they cannot always suppress the formation of Li dendritic structures, and the ionic conductivity of SPEs is very low, 10⁻⁴ - 0.01 mS.cm⁻¹, representing a significant drawback for application in all-solid-state batteries. On the other hand, inorganic solid electrolytes possess high ionic conductivities capable of competing even with liquid electrolytes > 10 mS.cm⁻¹ (Fig. 2). ISE is also associated with higher mechanical modules; they inhibit the growth of Li dendrites, are non-flammable, and are therefore associated with a higher safety level. Inorganic solid-state electrolytes can be divided into different types of structures or element-

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containing compounds that summarize the characteristics of the most studied families of the most studie

The general structure of LISICON (lithium superionic conductor) materials is based on the NASICON (sodium superionic conductor) crystal structure. LISICON belongs to a broader family of solid-state materials with the general chemical formula A₁₁B₂Ch₁₂, where A represents alkali metal cations (e.g., Li, Na, K), B represents metal cations (e.g., Ge, Ti, Zr), and Ch denotes chalcogen anions (e.g., O, S, Se). The crystal structure of LISICON can be described as a three-dimensional framework of corner-sharing polyhedral units. The B cations are surrounded by oxygen anions or other chalcogens, forming a B-centered polyhedron. These polyhedra are connected by shared oxygen (or chalcogen) ions to create a 3D network. Within this framework, the A cations (e.g., Li) occupy the interstitial spaces between the B-centered polyhedra. Lithium ions in these interstitial sites allow for efficient lithium-ion transport, making LISICON materials candidates for solid-state electrolytes.

The LISICON and argyrodite structures are usually found in sulfide-and-oxide-based electrolytes. The first LISICON electrolyte report can be traced back to 1980 when $Li_{3.6}Ge_{0.6}V_{0.4}O_4$ was successfully synthesized, resulting in an ionic conductivity of approximately 0.04 mS.cm⁻¹ at room temperature RT [9]. The crystalline structure of oxide-based LISICON electrolytes commonly resembles Li_4SiO_4 (monoclinic, P21/m space group) or γ -Li₃PO₄ (orthorhombic, Pnma space group) structures. These structures consist of tetrahedral units based on XO₄ (where X can be P, Al, Si, Ge, S, or Ti) and Li-O polyhedral units. For LISICON-oxide-based electrolytes, the highest reported value of ionic conductivity is approximately 0.9 mS.cm⁻¹ at 27 °C, for the $Li_4Al_{1/3}Si_{1/6}Ge_{1/6}P_{1/3}O_4$ electrolyte [10]. LISICON-oxide-based electrolytes are associated with relatively low ionic conductivity at RT, a significant limitation for employing these electrolytes in all-solid-state batteries.

Thio-LISICON solid-state electrolytes are derived from LISICON-oxide electrolytes by substituting the O^{2-} anion with the S^{2-} . This substitution improved the ionic conductivity of the LISICON electrolytes up to 12 mS.cm⁻¹ at RT for the Li₁₀GeP₂S₁₂ LGPS [11], which meets the demands of commercial all-solid-state-batteries (ASSBs). Albeit the thio-LISICON presents higher ionic conductivities due to the presence of sulfur in the backbone, this same element leads to a decrease in the overall stability of the electrolyte towards moisture.

Argyrodites are a group of compounds with the general chemical formula A_{7-x}BCh_{6-x}Hal_x, where A represents alkali metal cations (e.g., Li, Na, K), B is P or As, Ch is a chalcogen (e.g., S, Se) and Hal is Cl, Br, or I. In the argyrodite structure, the Ch chalcogens share the corners of the cube with the Hal halides; the P or As form a B-centered polyhedron with the Ch at the corners. The A cations are positioned at the cube's center, and the tetrahedral voids within the B-centered polyhedron. This crystal structure creates a three-dimensional network of corner-sharing polyhedrals, providing pathways for efficient ion transport, particularly for lithium ions. Argyrodites are known for their

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high lithium-ion conductivity, making them promising candidates for solid-state electrolytescinaline advanced lithium-ion batteries and other electrochemical devices.

Among the argyrodite-type sulfur electrolytes, Li_6PS_5Hal (Hal = Cl, Br, I) stand out as the most extensively studied and promising [12]. Substituting the halogen element between Cl, Br, and I leads to an evident variation in lattice site disorder, increasing the unit cell volume and the intra-cage and inter-cage jump distance [13].

In this perspective, our primary focus is on the chloride anion-based electrolytes. This choice is based on the following reasons: (1) fluorine is the most electronegative element, resulting in a strong bond with Li ions and further impeding fast Li-ion conduction; at the same time, as they have a small radius, the lattice volume is smaller, further hindering the Li⁺ conductivity; (2) Bromine is toxic, and as the world seeks safer and more environmentally friendly options, it has been disregarded; (3) iodine, while expected to yield better results, has limitations due to its photosensitivity, which poses challenges for mass-scale production applicability. Considering these factors, chloride anion-based electrolytes are a more favorable option, as they offer better Li-ion conduction while addressing the concerns associated with fluorine, bromine, and iodine.

Regarding $\text{Li}_6\text{PS}_5\text{Cl}$, the highest value of ionic conductivity reported for this electrolyte is 4.96 mS.cm⁻¹ at 26 °C when prepared via solid-state reaction [14]. The most common preparation method for this electrolyte is ball-milling [15]; however, several groups adopted a multiple-step synthesis that included both annealing and ball-milling to improve the crystallinity and ionic conductivity. The argyrodite-sulfur electrolytes display high ionic conductivity. However, the presence of sulfur in the backbone leads to decreased stability toward moisture [16]. In 2010, it was reported the first synthesis of oxide-based argyrodites $\text{Li}_6\text{PO}_5\text{Hal}$ [17]. The ionic conductivity of the oxygen-based $\text{Li}_6\text{PO}_5\text{Cl}$ electrolyte reported at RT was 10⁻⁶ mS.cm⁻¹, a value quite inferior to its sulfide-argyrodite counterpart.

The garnet structure is observed in oxide-based electrolytes, exhibiting exceptional stability when in contact with lithium metal and demonstrating satisfactory ionic conductivity. Garnet structures are commonly found in orthosilicates, characterized by the general formula, $A_3^{II}B_2^{III}(SiO_4)_3$ (A = Ca, Mg; B = Al, Cr, Fe) where A, B, and Si cations occupy the eight, six, and four coordination sites, respectively. These structures typically crystallize in a cubic arrangement with an Ia-3d space group. Among garnet-type electrolytes, there are four main subtypes: Li₃, Li₅, Li₆, and Li₇. Notably, the ionic conductivity of garnets containing lithium demonstrates a substantial exponential growth as the Li content expands throughout a wide range. Among all the famous Garnet-type electrolytes, the one that has emerged as the most promising is LLZO, Li₇La₃Zr₂O₁₂ [18]. This electrolyte has an ionic conductivity of 0.3 – 0.5 mS.cm⁻¹ at 25 °C [19], good thermal and chemical stability against reactions with prospective electrode materials, low cost of production, and environmental availability. On the

downside, the total ionic conductivity in this electrolyte is highly sensitive to grain boundariese These alive DOI: 10.1039/D3TA04228F structures tend to reduce the ionic conductivity of the electrolyte, introducing the need for careful control over a high-temperature sintering process to reduce grain boundaries and, therefore, achieve higher ionic densities and conductivities.

The general structure of NASICON is based on a three-dimensional framework of corner-sharing polyhedral units. NASICON is a family of solid-state materials with the general chemical formula $AM_2(PO_4)_3$, where A represents monovalent cations (e.g., Li, Na) and M represents a combination of divalent and trivalent cations (e.g., Sn, Ti, Zr).

In the NASICON crystal structure, the M cations are located at the corners of a distorted octahedron, forming an M-centered polyhedron. The PO_4 tetrahedral units (phosphate anions) are connected to these M-centered polyhedra, creating a three-dimensional network. The A cations are positioned in the interstitial sites between these polyhedral units.

The general structure of perovskites is based on a three-dimensional framework of corner-sharing polyhedral units. Perovskites are a family of materials with the general chemical formula ABX_3 , where A (e.g., Ba) and B (e.g., Ti) are cations, and X is an anion, for example, oxygen; the structure is cubic and Pm-3m. In the perovskite crystal structure, the X atoms (e.g., O) are located at the corners of an octahedron, forming a BO₆ polyhedral unit. The B cations are located at the center of the octahedron. The A cations are positioned in the center of the cube. The arrangement of these polyhedral units creates a three-dimensional network with a cubic symmetry. The general formula ABX_3 signifies that six X anions surround each B cation, and each A cation is coordinated to six X anions in an octahedral arrangement.

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Perovskites exhibit many properties and have applications in various fields, including solid-state materials, electronics, energy conversion and storage (e.g., photovoltaics, fuel cells), and catalysis. The versatility of perovskite materials stems from the ability to modify the A, B, and X elements, resulting in diverse electronic, magnetic, and optical properties.

The NASICON and the perovskite structures have lower ionic conductivity than previously mentioned electrolyte types and are still highly subjective to grain boundary issues. The NASICON electrolytes, discovered in 1976 by Goodenough et al. [20], have a general formula of $AM_2(PO_4)_3$, where A = Li, Na and M = Ge, Zr, Ti, and may be doped with Al. Generally, the ionic conductivity is higher when the A element is sodium Na rather than lithium Li. For instance, $LiGe_2(PO_4)_3 - LGP$ exhibits an ionic conductivity of approximately 10^{-4} mS.cm⁻¹. Adding aluminum as a dopant in $Li_{1.5}Al_{0.5}Ge_{1.5}(PO_4)_3 - LAGP$ significantly improves the ionic conductivity of this electrolyte, with a maximum achieved value of 0.58 mS.cm⁻¹ at RT [21]. Comparatively, the LAGP electrolyte showed a remarkable enhancement in ionic conductivity, surpassing LGP by approximately four orders of magnitude. Considering the sodium ion version, $Na_{1+x}Zr_2Si_xP_{3-x}O_{12}$, the highest ionic conductivity

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obtained is for x = 2, Na₃Zr₂PSi₂O₁₂ with an ionic conductivity of approximately 1.13 mS.cm^{4} at RT^{1} [22]. However, several doping strategies have been employed to achieve higher ionic conductivities: (1) doping with potassium ion, which demonstrated for Na_{2.303}K_{0.099}Zr₂Si_{1.880}P_{1.106}O₁₂ an ionic conductivity of 0.8 mS.cm⁻¹ at RT [23]; (2) doping with Ge, Na_{3.125}Zr_{1.75}Sc_{0.125}Ge_{0.125}Si₂PO₁₂, resulted in total ionic conductivity 4.6 mS.cm⁻¹ [24]; (3) doping with Sc, Na_{3.4}Zr_{1.6}Sc_{0.4}Si₂PO₁₂ renders an RT ionic conductivity of 4.0 mS.cm⁻¹ [22].

As for perovskite electrolytes, one notable example is $Li_{3x}La_{(2/3)-x}TiO_3$ LLTO. In 1987, Belous et al. [25] synthesized the LLTO perovskite for $0.04 \le x \le 0.17$ for the first time. In the LLTO solid electrolytes' structure, the Li-ion occupies the A-site, resulting in a lower activation energy. This electrolyte shows several advantages: (1) large ionic transference numbers, (2) superior chemical and (3) thermal stabilities in air, and (4) environmentally friendly character without any release of toxic gases during decomposition reactions. However, the significant grain boundary resistance reduces the total ionic conductivity of this electrolyte to $< 0.01 \text{ mS.cm}^{-1}$ at RT. LLTO is also very unstable when in the presence of Li metal. For potentials below -1.7 V vs. Li^o(-1.39 eV/e for E = 0 eV, electrons at rest in vacuum), LLTO is reduced, and Li can be intercalated in it, which causes the reduction of Ti^{4+} and enhances the electrical conductivity of $Li_{0.29}La_{0.57}TiO_3$ LLTO [26]. The latter observation indicates that the lowest energy of the conduction band is approximately -3.1 eV (E = 0 eV, electrons at rest in vacuum). The original calculations hereafter demonstrate that the lowest energy of the conduction band for LLTO is -3.6 eV, which is in good agreement with the experimental result, considering this value is the result of two different simulations: (1) the average potentials from a simulated surface; (2) the band structure from the bulk electrolyte [27]. Note that herein, the simulated LLTO was Li₂La₂Ti₃O₁₀.

Figure 2 shows the highest ionic conductivities in the literature, and Figure 3 shows our data on the simulated chemical potentials and band gaps for a selection of solid electrolytes.

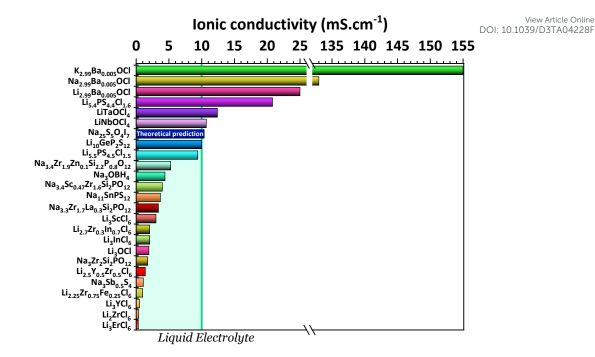


Figure 2. Ionic conductivities for the electrolytes: K_{2.99}Ba_{0.005}OCl and Na_{2.99}Ba_{0.005}OCl [28]; Li_{2.99}Ba_{0.005}OCl [29]; Li_{5.4}PS_{4.4}Cl_{1.6} [30]; LiTaOCl₄ and LiNbOCl₄[31]; Na₂₅S₅O₄I₇ (theoretical study) [32]; Li₁₀GeP₂S₁₂ [11]; Li_{5.5}PS_{4.5}Cl_{1.5} [33], $Na_{3.4}Zr_{1.9}Zn_{0.1}Si_{2.2}P_{0.8}O_{12} \ [34]; \ Na_{3}OBH_4 \ [35]; \ Na_{3.4}Sc_{0.47}Zr_{1.6}Si_{2}PO_{12} \ [36]; \ Na_{11}Sn_{2}PS_{12} \ [37]; \ Na_{3.3}Zr_{1.7}La_{0.3}Si_{2}PO_{12} \ [38]; \ Na_{3.4}Sc_{0.47}Zr_{1.6}Si_{2}PO_{12} \ [36]; \ Na_{11}Sn_{2}PS_{12} \ [37]; \ Na_{3.3}Zr_{1.7}La_{0.3}Si_{2}PO_{12} \ [38]; \ Na_{12}Si_{2}PO_{12} \ [38]; \ Na_{1$ Li₃ScCl₆ [39]; Li_{2.7}Zr_{0.3}In_{0.7}Cl₆ [40]; Li₃InCl₆ [41]; Li₃OCl[29]; Na₃Zr₂Si₂PO₁₂ [42]; Li_{2.5}Y_{0.5}Zr_{0.5}Cl₆ [43]; Na₃Sb_{0.5}S₄ [44]; $Li_{2.25}Zr_{0.75}Fe_{0.25}Cl_{6}[45]; Li_{3}YCl_{6}[46]; Li_{2}ZrCl_{6}[45]; and \ Li_{3}ErCl_{6}[47].$

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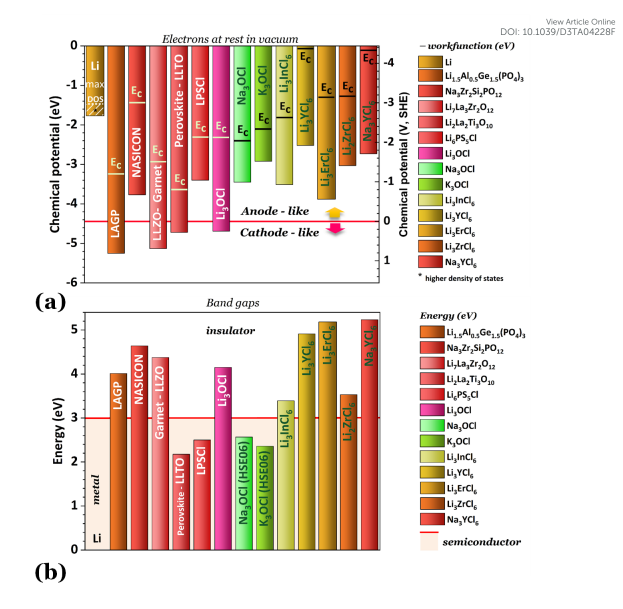


Figure 3. Original data: (a) chemical potentials for electrically insulated materials and the lowest energy in the conduction band; (b) band gap reflecting the insulator or semiconductor character of the electrolytes. Note: the energy of the band gaps in (a) is $E_g = 2 \times (E_c - \mu_{electrolyte}) eV$.

1.3 What makes a good solid electrolyte?

Our perspective showed that every electrolyte class has advantages and disadvantages. This realization highlights the necessity to identify a set of parameters to answer the question, *what truly defines a GOOD electrolyte?*

To address this question, we have conducted *ab initio* simulations using density functional DFT and exchange-correlation GGA-PBE or, when doubts arise concerning the electronic properties of the semiconductors, a hybrid functional HSo6 was employed. These simulations allowed us to determine band structures and electronic properties (Figs. 3b and 4) [48], [49] and the chemical potentials (Figs. 3a) [50], [51] and to compare them with their respective ionic conductivity and the performance of the electrolytes in general. The crystal structures used are described in Table 1. The

surfaces were simulated with a minimum of a 10 Å vacuum layer to break the periodicity Arthenime planewave cutoff was 500 eV, and the projection was in the reciprocal space. The spacing of k-points was 0.2-0.3 Å⁻¹. The cell contained the minimum number of atoms, allowing for a better approach to stoichiometry and experimental occupancy. The chemical potentials were obtained from the average potentials integrated over 10 Å.

Solely relying on ionic conductivity (Fig. 2) to enumerate the characteristics of a suitable electrolyte proves to be complex since it merely represents a property and is likely not a driving force of good performance. To progress with this discussion, the energy width of the band gaps has been calculated for various electrolytes representing the previously mentioned categories in Fig. 3b. It is anticipated that, in general, solid electrolytes exhibit insulator behavior, characterized by a band gap greater than 3 eV, indicating poor electrical conductivity. The band gap values may be distinguished into three different categories: (1) conductors or metals, $E_g = 0$ eV; (2) semiconductors, $E_g \le 3$ eV, and (3) insulators, $E_g > 3$ eV. In Fig. 3b, the semiconductor/insulator barrier is represented by a red horizontal line marking 3 eV.

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Based on this perspective, it can be inferred that LAGP, NASICON, Garnet, Li₃OCl, Li₃YCl₆, Li₃ErCl₆, and Na₃YCl₆ exhibit insulating behavior. However, Li₃YCl₆ and Na₃YCl₆ are not stable against Li^o and Na^o. The latter is understandable as, on the contrary to most insulator materials, there is a considerable density of states DOS at the lower energy of the conduction band corresponding to the orbitals *d* of Y³⁺ (Fig. 4). As this energy corresponds to an almost null electron affinity, however entirely above the chemical potential of the Lithium $\mu(Li^0) = -1.39 \text{ eV}$, the reduction of Y³⁺ may occur from the surface exposure directly. The Li⁺ required to form Y is $3Li^+ + 3\bar{e} + Li_3YCl_6 \rightarrow 6LiCl + Y$, may also be obtained within the electrolyte while charging the cell, as Li⁺ will move toward the negative electrode to accommodate the crescent bias between electrodes' voltages.

An equivalent process should drive the reactive character of Na_3YCl_6 and likely Li_3ErCl_6 , although the latter has a higher electron affinity but similar DOS at E_c (Fig. 4).

The cathode-like solid electrolytes (Fig. 3) have low DOS on the conduction band, which prevents the drift of electrons from the alkali metal to the conduction band of the electrolytes.

The ferroelectric character of the perovskites, antiperovskites, and glass electrolytes promotes stability against the alkali metals. In the electrolytes of the family $A_{3-2x}M_xOHal$, with A = Li, Na, K, M = Mg, Ca, Sr, Ba, Hal = Cl, Br, I, and $x \le 0.5$, a ferroelectric topologic insulator character was found to protect the alkali metals as self-charge happens with electrons tunneling from the electrolytes' surface to the negative electrode. Like the latter, other perovskites were found to manifest the phenomenon discussed hereafter. This phenomenon may also protect the perovskite Li₂La₂Ti₃O₁₀

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LLTO, also used as a cathode. Zhang *et al.* [52] reported $Li_2La_2Ti_3O_{10}$ electrolytes as displaying the policity of th

The stability of $\text{Li}_6\text{PS}_5\text{Cl}$ LPSCl toward Li^o is not expected because of its semiconductor character and the position of its lower conduction band energy level, allowing the drift of electrons from the alkali metal. We have detected a reversible reaction with Cu, $\mu(Cu^0) \approx -4.6 \text{ eV}$, which may indicate electronic surface conduction in the electrolyte and tunneling from its surface to the negative electrode. However, this result requires further confirmation.

From the analysis with different electrolytes, it is possible to establish a pattern for best-performing electrolytes (Figs. 2, 3, and 4): (1) display semiconductor behavior instead of insulator behavior (e.g., LPSCl, LLTO, Na₃OCl, K₃OCl, A_{2.99}Ba_{0.005}OCl – glass with A = Li, Na, K); (2) possess a low energy work function W, i.e., anode-like behavior; but not as small as to have a null, or almost null, electron affinity energy (Fig. 4); (3) show a ferroelectric behavior where the ionic movement establishes the polarization; (4) display topologic electronic emergent phenomena. These latter phenomena require further demonstrations in different electrolytes.

Details on the properties and behavior of all-solid-state electrolytes and all-solid devices, including synthesis methods, electrolytes showing ferroelectric and topologic insulator features, and quantum thermal batteries, will be given hereafter.

Solid electrolyte Crystal structure		Stable against the alkali metal?	
$\begin{array}{c} \text{Li}_{1.5}\text{Al}_{0.5}\text{Ge}_{1.5}(\text{PO}_4)_3\\ \text{LAGP} \end{array}$	Rhombohedral, R-3c, NASICON	~	
Na ₃ Zr ₂ Si ₂ PO ₁₂ NASICON	Rhombohedral, R-3c, NASICON Monoclinic, C2/c, disordered	~	
Li ₇ La ₃ Zr ₂ O ₁₂ LLZO	Cubic, Ia-3d	~	
Li ₂ La ₂ Ti ₃ O ₁₀ LLTO -also used as cathode- ferroelectric	Tetragonal layered perovskite P4/mmm	\checkmark	
Li ₆ PS₅Cl LPSCl	Trigonal, P-3m1, disordered	\checkmark	
glassy Li ₃ OCl ferroelectric	Cubic, Pm-3m, disordered obtained from antiperovskite	\checkmark	
glassy Na ₃ OCl ferroelectric	Cubic, Pm-3m, disordered obtained from antiperovskite	\checkmark	
glassy K ₃ OCl ferroelectric	Cubic, Pm-3m, disordered obtained from antiperovskite	\checkmark	
Li ₃ InCl ₆	Monoclinic, C2/m	×	
Li ₃ YCl ₆	Trigonal, P-3m1, disordered	×	
Li ₃ ErCl ₆	Trigonal, P-3m1, disordered	?	
Li ₂ ZrCl ₆	Monoclinic, C2/m	\checkmark	
Na ₃ YCl ₆	Monoclinic, P21/c	×	

Table 1. Solid electrolytes in Fig. 3; their crystal structure and their stability against the correspondent alkali metal.

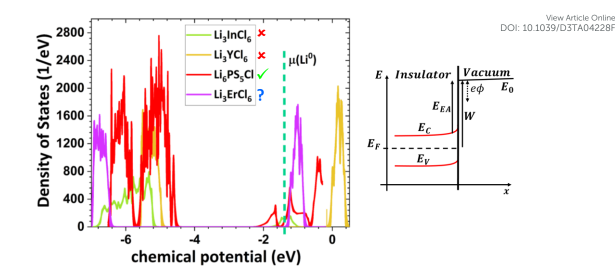


Figure 4. The density of states DOS compared with the chemical potential of $\mu(Li^0)$ for Li₃InCl₆, Li₃YCl₆, Li₆PS₅Cl, and Li₃ErCl₆. The first two are unstable against Lithium. $-W = E_F = \bar{\mu}_i = \mu_i + z_i e\phi$; where *W* is the workfunction; E_{FA} is the electron affinity; E_F the Fermi level or electrochemical potential $\bar{\mu}_i$; μ_i is the chemical potential; $z_i e\phi$ is the surface chemical potential; z_i is the valence of the mobile ion; *e* is the electron charge; ϕ is the surface potential that is majority produced by the materials in electrical contact with the material. Phenomena at the surface of the material may also create surface chemical potential. When the insulator is at equilibria with the materials in electrical contact $\bar{\mu}_i - \bar{\mu}_j = 0 = \mu_i - \mu_j + z_i e(\phi_i - \phi_j) \Leftrightarrow \mu_i - \mu_j = z_i e \Delta V$. If $\mu_j = E_0$ is the chemical potential or energy of the electrons at rest in a vacuum on the surface of the material *i*, μ_i is its chemical potential.

ВΥ (cc)

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Electrolyte	Fabrication Method	Parameters	Ionic conductivity (mS.cm ⁻¹)	Crystal Structure	Ref.
S ₅ Cl - LPSC	Ball Milling	10h - 600 rpm;	1.3 (25 °C)	S S S S S S S S S S S S S S	[67]
	Solid State Reaction	 (1) LiCl+Li₂S+P₂S₅ 100 rpm for 1h; (2) 10h of annealing; 	5.0 (26 °C)		[14]
	Wet Phase	Mixture with ethanol solution;	1.4 × 10 ⁻² (RT)		[68]
	Wet Phase (LiCl+Li ₂ S+ P_2S_5 +	 P₂S₅ + pyridine 400 rpm, RT add LiCl + Li₂S to 400 rpm at 70 °C removing pyridine in vacuum at 120 °C 	4.3 (RT)		[69]
TE Deposition Laser annealed Pulsed Lase Deposition	Reaction	(1) 200 rpm 12h in 2-propanol; (2) 900 °C 4h; (3) 200 rpm 12h;	0.2 (25 °C)		[19]
	Frequency magnetron	RF power of 40, 60, and 80 W at a working pressure of 7 MPa under Ar atmosphere;	4 × 10 ⁻⁴ (25 °C)		[70]
		$f_{laser \ pulse}$ = 10 Hz; 7.5 J.cm ⁻² ; 7700 shots; thickness ~1 µm; laser annealing;	7.4 × 10 ⁻⁴ (RT)		[71]
	Pulsed Laser Deposition – annealed at 800 °C	$f_{laser \ pulse}$ = 10 Hz; 7.5 J.cm ⁻² ; 7700 shots; thickness ~1 µm; annealing 800 °C;	1.8 × 10 ⁻⁴ (RT)		[71]
VOISU - Co- precipitation		 (1) Calcined at 1100 °C for 12 h; (2) ball milling at 150 rpm for 18 h; 	0.2 (50 °C)		[72]
			1.2 (RT)		[73]
	precipitation	diluted with 1-propanol; Solution 1 dipped on solution 2, forming a precipitate;	1.7 (25 °C)		[74]
	Spark Plasma Sintering	Powders were placed into a	1.8 (RT)		[75]

 Table 2. Synthesis methods for the electrolytes LPSCl, LLZO, and NASICON and their possible influence on their intermediate DOI: 10.1039/D3TA04228F

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 conductivity. CIF files: [53]-[66].

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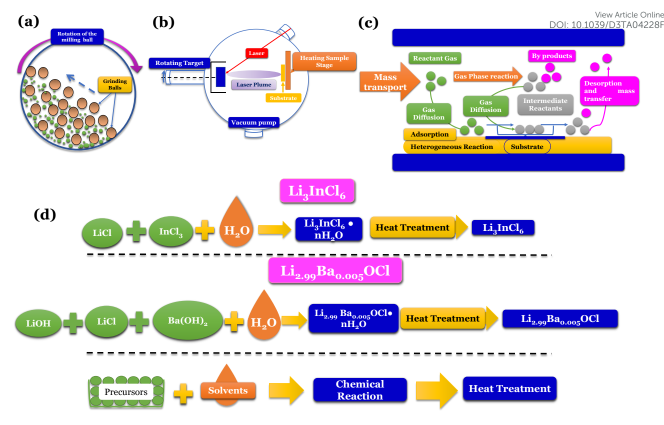


Figure 5. Synthesis methods: (a) high energy density ball milling [76]; (b) pulsed laser deposition PLD [77]; (c) chemical vapor deposition CVD [78]; (d) water mediated synthesis Li₃InCl₆ [41], Li_{2.99}Ba_{0.005}OCl [29], K_{2.99}Ba_{0.005}OCl and Na_{2.99}Ba_{0.005}OCl [28].

2.1. Synthesis Methods

When different methods prepare an electrolyte, the final characteristics of the electrolyte differ; this occurs because the preparation methods influence the crystal structure of the solid, the structural disorder, and mobile ion distributions, resulting in significantly different values of ionic conductivities in Table 2.

Mechanochemical methods, specifically ball-milling (Fig. 5a), are commonly linked to forming distinct phases, such as amorphous and/or nanocrystalline structures. On the other hand, when solid-state electrolyte SSEs are annealed at elevated temperatures, they tend to exhibit thermodynamically stable phases.

The vapor-phase deposition is widely used to prepare inorganic SSE membranes for thin-film batteries. These methods involve the formation of the vapor phases of the reactants, subsequent nucleation, and product growth on substrates, allowing the achievement of a homogeneous mix of the reactants on a molecular or atomic level. The gas-phase techniques involve deposition methods such as Pulse Laser Deposition PLD (Fig. 5b), Atomic Layer Deposition ALD, and Chemical Vapor Deposition CVD (Fig. 5c).

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To secure a dominant role in the industry, SSEs must be easily reproducible, and the veost confinine DOI: 10.1039/D31A04228F fabrication must be low. Techniques like ball milling don't allow reproducibility, and vapor-phase deposition techniques are usually expensive. Recently, a new technique only compatible with some types of electrolytes has been developed; the latter is a water-mediated synthesis (Fig. 5d) with great potential for mass production due to the relatively low cost of implementation.

Ball-Milling. High-energy methods are among the favorite synthesis methods to produce inorganic solid electrolytes. This technique generates chemical reactions through a force caused by the collision between the milling balls and the reactants, Figure 5a. During this process, the reagents are pulverized and mixed, with point and lattice defects forming. The mechanochemical method's main advantage is its ability to overcome the diffusion limitation in the solid-state reaction without the necessity to add bulk solvents or exposure to heat. This technique was introduced by Hayashi *et al.* [79] in 2001 for electrolytes and has since been a promising technique to achieve improved ionic conductivity; the strain during ball-milling increases the activation volume, allowing ion hopping.

Additionally, ball-milling can induce defects in inorganic solid electrolytes, enhancing their ionic transport. However, the results obtained by this technique don't show satisfactory reproducibility. Even using the same equipment, the ball milling process is influenced by chamber atmosphere, ball milling time, rotation speed, and ball size; this represents a significant setback for using this technique for mass-scale production.

Pulsed Laser Deposition. Pulse Laser deposition (PLD) is a process where a high-energy laser beam is used to ablate a target material onto a substrate inside a vacuum chamber. The target is heated and vaporized by the energy of the laser. The vapor is energized and is transformed into plasma with high energy. The atoms in the plasma collide and recondense on the substrate and form a thin film Fig. 5b. The main advantage of this technique is that the highly focused laser can effectively deliver energy to a localized area and ensure the evaporation of the target materials even with very low vapor pressure [71]. Additionally, the PLD technique exhibits a high deposition rate and accurate control of film growth precursor. In contrast to the conventional evaporation techniques, it is hard to control the chemical composition of the thin film because of possible precursor loss due to volatilization.

Chemical Vapor Deposition. Chemical vapor deposition CVD is a process that involves the reaction of a volatile precursor previously injected into a chamber, typically under a vacuum. The chamber is heated to a temperature that causes the precursor gas to react or break down into the desired coating and bond to the material surface. The process begins with the introduction of the reactants to the deposit region. After this, the reactants are transported by diffusion from the main gas stream through the boundary layer to the substrate surface, where the absorption of the reactants will occur. When this process starts, surface reactions and decomposition take place. The CVD process ends with the forced convection transport of the by-products away from the deposition nume DOI:10.1059/D31A04228F region, Fig. 5c. The CVD process involves highly complex flow dynamics since gases flow into the reactor, reacting, and then the by-products are exhausted.

The CVD technique is relatively new for fabricating thin film batteries [80]. The main advantage of CVD is the dependence on relatively low kinetic energy particles and the multitude of chemistries available to perform the deposition.

Water-Mediated Synthesis. To achieve mass-scale production of solid-state electrolytes, it is essential to utilize cost-effective synthesis methods. However, a few previously discussed methods are expensive and impractical for large-scale manufacturing. A recent liquid or wet-phase synthesis development provides a potential solution. When applied to sulfide-based electrolytes, this method, also known as wet chemical synthesis, can be executed in two ways: **(1)** generating a precipitate by mixing a precursor with a solvent or **(2)** dissolving the precursor to create a homogeneous solution [81]. Both approaches are followed by solvent evaporation to obtain the final solid-state electrolyte. However, these methods have limitations in terms of time consumption and solvent compatibility. Considering environmental concerns, water seems to be the ideal solvent due to its favorable properties. Unfortunately, it can only be employed in moisture-stable precursors, which excludes most sulfide electrolytes.

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Nevertheless, despite the incompatibility of sulfide-electrolytes with water-mediated synthesis, various oxyhalides, and halide electrolytes have demonstrated compatibility with this method, as shown in Fig. 5d. Moreover, these electrolytes exhibit exceptional ionic conductivities, further emphasizing their potential for practical applications. For example, Li₃InCl₆, when synthesized via solid-state synthesis, shows an ionic conductivity of 1.49 mS.cm⁻¹ [82]. When prepared via water-mediated synthesis, the ionic conductivity is 2.04 mS.cm⁻¹ [41]. However, the most significant advantage of this method is the increased humidity resistance when prepared by water-mediated synthesis.

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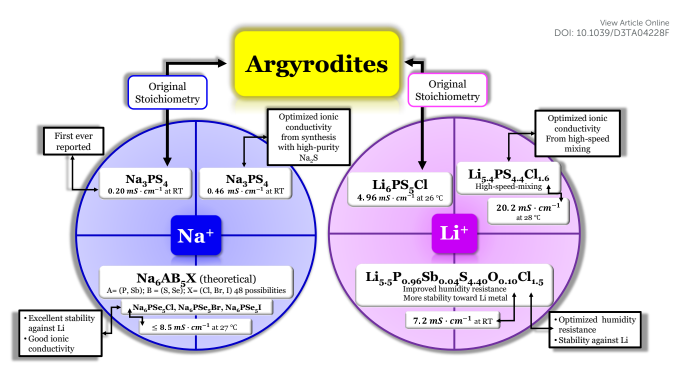


Figure 6. Argyrodites summary scheme, Li₆PS₅Cl [14]; Li_{5.4}PS_{4.4}Cl_{1.6} [30]; Li_{5.5}P_{0.96}Sb_{0.04}S_{4.40}O_{0.10}Cl_{1.5} [83]; Na₃PS₄[84], [85], and Na₆AB₅X, A = (P, Sb), B = (S, Se), X = (Cl, Br, I) [86].

2.2. Argyrodite electrolytes

Argyrodite electrolytes are derived from the Ag₈GeS₆ mineral and are reported in sulfide and oxidebased electrolytes. Considering the great interest in the mobility of Li⁺ ions in solids, it was surprising that the argyrodite structure had not been reported more in association with electrolytes until the beginning of the 21st century. In the Li-argyrodite family, compounds are described by the general formula, $\text{Li}_{7+x-y}M_x^{\text{IV}}M_{1-x}^{\text{V}}Ch_{6-y}\text{Hal}_y$ (M^{IV} = Si, Ge, Sn; M^V = P, Sb; Ch = O, S, Se; Hal = Cl, Br, I, BH₄; $0 \le x \le 1$; $0 \le y \le 2$) [87]. In 2008, Deiseroth *et al.* [88] reported the first Li₆PS₅Hal, Hal = Cl, Br, I, argyrodite electrolyte, showcasing its ionic conductivity in the 1 to 10 mS.cm⁻¹ range at RT. Since then, numerous reports have emerged regarding this stoichiometry, with the highest ionic conductivity obtained for Li₆PS₅Cl (LPSCl) being 4.96 mS.cm⁻¹ at 26.2 °C when prepared via solidstate reaction [14]. However, this is not the only stoichiometry configuration possible for a lithium argyrodite (Fig. 6).

In a recent study [30] the Li-argyrodite electrolyte Li_{5.4}PS_{4.4}Cl_{1.6} was synthesized using high-speed mixing. This electrolyte demonstrated an astonishingly high ionic conductivity of 20.2 mS.cm⁻¹ at 28 °C, Figs. 2 and 6, the highest reported value for a lithium argyrodite. The highest reported ionic conductivity for the standard high-energy ball milling Fig. 6 is 9.4 mS.cm⁻¹[33]. The study by Chen *et al.* [30] introduced a cost-effective and efficient method known as *high-speed mixing*, which utilizes an affordable consumer-grade commercial mixer. The high-speed mixing process enables

rapid material mixing, producing more than 500 g per batch of LSPCl solid electrolyte. In contrastontine DOI: 10.1039/D31A04228F the traditional high-energy ball-milling method is limited to 50 g per batch. Notably, the high-speed mixing technique is both energy-efficient and highly effective.

But how does this high-speed mixing technique affect the observed high ionic conductivity?

Using the ultra-high speed mixing preparation process, no obvious grain boundary was observed in the cross-sectional microstructure, which is one of the main factors for the reported high ionic conduction. This presents an effective mixing method for the mass production of $Li_{5.4}PS_{4.4}Cl_{1.6}$ solid electrolyte. The authors also noted that an all-solid-state electrolyte cell containing graphite/LPSC-mixing/NMC622 cell delivered an outstanding discharge capacity of 165 mAh.g⁻¹ at C/10 and a % coulombic efficiency of 82.4%.

Despite the numerous advantages offered by the conventional argyrodite electrolyte, concerns have been raised due to its vulnerability to atmospheric air and its limited structural and electrochemical stabilities. Nevertheless, completely replacing sulfur with oxygen in the argyrodite structure has significantly reduced ionic conductivity [17], indicating that this approach may not be the ideal solution. However, an alternative worth exploring involves doping the argyrodite structure with oxygen (Fig. 6). This approach combines the best attributes of sulfur and oxygen, leading to meaningful results [89].

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Hwang et al. [89] investigated the substitution of oxygen (O) by sulfur (S) in Li₆PO_xS_{5-x}Br_{0.5}Cl_{0.5} SEs to comprehend the underlying mechanisms and improve the stability of these materials. In his study, the author combined first-principles calculations and electrochemical experiments, spanning from bulk to surface models. Moreover, the author developed systematic experimental investigations to explore the impact of oxygen substitution on suppressing/reducing structural decomposition. The materials were synthesized and thoroughly characterized. The goal of doping the sulfide electrolyte with oxygen is to increase the resistance to moisture absorption, delaying phase decomposition. However, what is the mechanism underlying this effect? To answer this question, these electrolytes' atomic and electronic structures were calculated from bulk to surface. The theoretical analysis determined that the introduction of oxygen causes a more noticeable contraction of the lattice constants compared to the substitution of halogen elements. This suggests that replacing sulfur (S) with oxygen (O) significantly impacts the chemical bonding between isovalent chalcogens, leading to a stronger electrostatic bonding between O and Li ions. This strong bonding is closely associated with structural stability. Following this hypothesis, the authors studied the effect of the introduction of oxygen in the electrochemical environment of the electrolyte. The latter realization was made by considering that oxygen has a superior electronegativity than sulfur.

A new study by Wei et al. [83] described the synthesis of a chlorine-rich lithium argyrodite, $Li_{5.5}P_{0.96}Sb_{0.04}S_{4.40}O_{0.10}Cl_{1.5}$, which demonstrated enhanced stability in the presence of air and

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lithium-ion stability (Figs. 2 and 6). The reported ionic conductivity of this compound is 7.2 VinStemPnline ¹ at RT, slightly lower than that of the Li_{5.5}PS_{4.5}Cl_{1.5} electrolyte that exhibits 9.4 mS.cm⁻¹ at RT. However, it was observed that when both electrolytes were exposed to moisture, the ionic conductivity of $Li_{5.5}PS_{4.5}Cl_{1.5}$ decreased significantly to 1.1 mS.cm⁻¹, while Li_{5.5}P_{0.96}Sb_{0.04}S_{4.40}O_{0.10}Cl_{1.5} electrolyte showed a comparatively higher ionic conductivity of 3.6 mS.cm⁻¹. Furthermore, a comparison of the X-ray diffraction (XRD) spectra revealed that the Li_{5.5}PS_{4.5}Cl_{1.5} electrolyte exhibited weakened diffraction peaks and the presence of various impurity phases when exposed to open air for 20 minutes. In contrast, the Li_{5.5}P_{0.96}Sb_{0.04}S_{4.40}O_{0.10}Cl_{1.5} electrolyte displayed improved stability with fewer impurity phases, indicating its superior moisture resistance compared to the bare Li_{5.5}PS_{4.5}Cl_{1.5} electrolyte. The introduction of the Sb element helped to enhance the stability of the electrolyte towards lithium metal. The compound known as sulfur argyrodite is commonly found in Li⁺-ion conducting electrolytes.

However, there have been reports of Na+-ion argyrodite electrolytes as well (Fig. 6). In 2012, Hayashi et al. [84] reported the synthesis of Na₃PS₄ as a superionic glass-ceramic electrolyte for the first time, displaying an ionic conductivity of 0.2 mS.cm⁻¹ at RT. In 2014, the same author reported the synthesis of the same electrolyte with an improvement of the ionic conductivity of 0.46 mS.cm⁻¹ at RT [85]. The improvement was a result of an optimization during the synthesis mode. The author synthesized Na₃PS₄ in cubic phase directly via mechanical milling using a highly pure Na₂S reagent; with this, the authors could reduce the reaction time necessary to produce the glass ceramic effectively. This glass-ceramic electrolyte was synthesized by milling 75 mol% Na₂S and 25 mol% P_2S_5 for 1.5 hours and subsequent heat treatment at 270 °C for 1 hour, as reported in [84]. This method reduces the reaction time and facilitates the production of glass ceramics. Moreover, in this study, the author also synthesized an all-solid-state cell containing Na₁₅Sn₄/Na₃PS₄/NaCrO₂. The cell was tested electrochemically at RT and displayed an average capacity of 60 mAh.g⁻¹ of NaCrO₂ for 15 cycles.

Following the previous reports, in 2016 de Klerk and Wagemaker [90] conducted a study to comprehend the diffusion mechanism in Na₃PS₄ and explore potential ways to enhance Na⁺-ion conductivity. Na₃PS₄ is known to exist in two distinct phases, cubic and tetragonal. However, the differences between these phases are small; the tetragonal phase is tetragonally distorted from the cubic phase [90]. The main difference between these two structures lies in the positioning of sodium within the crystal. Regardless if both phases are synthesized similarly, they display similar ionic conductivity. A significant boost in ionic conductivity is observed when transitioning from the crystalline to the glass-ceramic form of the cubic phase, increasing 30 times the ionic conductivity. The introduction of halogen dopants plays a crucial role in enhancing ion conductivity. This effect is attributed to a significant increase in the concentration of Na⁺ vacancies, which is vital to microscopic diffusion. The role of temperature is also an essential factor in Na+-ion diffusion. In this study, the

the

authors [90] conducted molecular dynamics simulations to investigate the effects of relative entire parameter increase on the conductivity of cubic $Na_{2.50}PS_4$ and $Na_{2.94}PS_4$ at 252 °C. Considering a 1% expansion in lattice parameters, corresponding to a 3 % volume expansion, the conductivity increased by 10 % for $Na_{2.94}PS_4$ and 17 % for $Na_{2.50}PS_4$, highlighting the significant impact of lattice expansion on conductivity when the temperature increases.

In 2020, Ouyang et al. [86], conducted research utilizing computational simulations to explore the discovery of halogen-substituted Na⁺-argyrodites. In this study, the author gained insights into the necessary conditions to achieve high ionic conductivity in Na⁺ argyrodites, Fig. 6. This investigation focused on the phase stability, electrochemical stability, and ionic conductivity of 48 potential Nabased argyrodites. In the Na⁺-based argyrodite Na₆ACh₅Hal, A represents a pnictogen atom P or Sb, Ch is typically a chalcogenide atom such as S or Se, and Hal is a halogen element, similar to its Li+ counterpart (Fig. 6). Regarding the phase stability, the authors concluded that the formation of the SbS₄ polyanion in Na⁺ Argyrodites performs more favorably compared to the PS₄ polyanion. The halogen element Hal had minimal impact on the electrochemical stability or the decomposition phases, considering the established limits imposed by the polyanion and chalcogen elements. The Na⁺ argyrodites are generally electrochemically stable within the potential range of 1 - 2 V vs Na/Na⁺, which means that the lower energy of the conduction band of the Na⁺ argyrodites varies from $-1.73 \le E_c(eV) \le -2.73$ where $E_0 = 0$ eV is the energy of the electrons at rest in a vacuum. The latter value for the lowest energy of the conduction band is shown to agree with the equivalent value for the Li⁺ argyrodite, LPSCl calculated herein and shown in Fig. 3a. The previous limits, as already mentioned, are imposed by the pnictogen element, which reduces from +5 to +3 for potentials above the stability window for $E > E_c$ and by the chalcogen element since it oxidizes for values of potentials lower than the argyrodite stability window $E < E_V$ [86] (Fig. 4). Furthermore, a comparison of stability windows revealed that the Na-Sb-S(Se)-Hal systems have a narrow electrochemical window, whereas the Na-P-S(Se)-Hal systems generally possess a wider electrochemical window [86]. When considering the ionic conductivity, the halogen element has a dominant role since the occupancy of the halogen, especially in a sulfide-based system, deeply affects the ionic conductivity. In contrast, the ionic conductivity in selenide-based systems remains similar, regardless of the halogen occupancy. The compression effect generated by halogen substitution is less significant in the selenide systems, which generally leads to good ionic conductivity [86]. Ouvang et al. [86] concluded that among the predicted Na $^+$ argyrodites, the compositions within the Na-P-Se-Hal (Hal = Cl, Br, or I) system demonstrated excellent stability and ionic conductivity.



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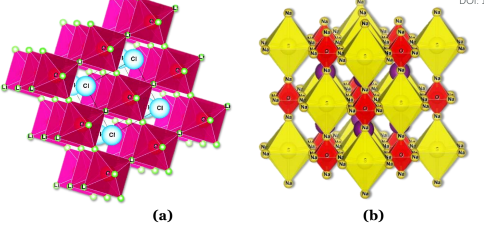


Figure 7. Antiperovskite electrolytes structures: (a) Li₃OCl [29]; (b) Li₆SO(BH₄)₂[91]; CIF files: [53]–[66].

2.3. Antiperovskites

Recently, a new type of structure called antiperovskites has been receiving increased attention in inorganic material research. The antiperovskites have the same structure as the perovskites. However, they adopt a reversed ionic arrangement, which means unique anion-centered cubic units in antiperovskites compared to the conventional cation-centered ones in perovskites. The cation position in the antiperovskite causes significant differences in these structures, providing a distinct local environment for mobile ions and altering their mobility. The first time ionic conductive antiperovskites with X_3BA structures were reported dates to 1960 when Reuter *et al.* [92] reported the Ag₃SI compound with Ag⁺ as a mobile cation. In 2012, Zhao *et al.* synthesized and designed the first Li⁺ conducting X_3BA shape antiperovskites [93]. In the antiperovskite formula X_3BA , X is a cation, and A and B are different-sized anions. The high-temperature superionic conductivity of $NaMgF_3$ perovskites inspired the discovery of the ionic conductivity of Li-rich antiperovskites (LIRAP). The electronically inverted antiperovskite Li_3OHal (Hal = Cl, Br, I) allowed Li^+ ionic conductivity like the F-superionic conductivity in NaMgF₃. However, the perfect X₃BA antiperovskite structures are considered a disadvantage for ionic conduction due to the lack of intrinsic defects that would work as charge carriers in ionic conduction [94]. A variety of methods for defect engineering, such as mixing, doping, and depletion by doping, for example, with divalent ions such as Ba²⁺, was proposed theoretically in [94]. Following this study, Braga et al. [29] also reported a series of divalent cation-doped Li₃OCl glass electrolytes, $Li_{3-2x}M_xOCl$ with M = Mg, Ca, Sr, Ba, synthesized by watermediated synthesis followed by a heat treatment to remove the solvent. These latter electrolytes are not antiperovskites anymore as they are glassy-amorphous, disordered materials.

In solids, as explained before, ion conduction occurs by ion hopping from one lattice site to another by vacancy mechanisms. The rate of ion transport in a lattice is dictated by diffusivity and

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concentration of vacancies mediating transport. Glassy materials are usually associated with high nine DOI: 10.1039/D31A04228F ionic conductivity because their open structures facilitate the hoping process.

In the Li₃OCl crystalline allotrope, the antiperovskite structure, the pattern followed is the following: (1) the Cl⁻ occupies the center atom of the cube⁻; (2) the O²⁻ occupies the center of the octahedral; (3) the Li⁺ occupies the vertices of the octahedral, making it a Li⁺-rich structure, Fig. 7. Doping this structure with higher valence cations such as Ba²⁺ and Ca²⁺ creates vacancies in the cation sublattice, one per each Ba²⁺ that replaces two Li⁺. This substitution lowers the activation energy for Li⁺-ion diffusion as the formation of vacancies is unnecessary or assured at synthesis.

Of all the reported electrolytes [29], the one that displayed better results was the one doped with Ba^{2+} cation; Barium ion has a larger doping ion radius, resulting in a higher anisotropy of the lattice, which favors a higher ionic conductivity. The glassy electrolyte $Li_{2.99}Ba_{0.005}OCl$ obtained a ground-breaking ionic conductivity of 25 mS.cm⁻¹ at 25 °C. After this study, Braga *et al.* synthesized the analogous Na⁺ and K⁺ electrolytes [28]. The Na_{2.99}Ba_{0.005}OCl and K_{2.99}Ba_{0.005}OCl electrolytes, synthesized as glass materials, displayed an astonishing ionic conductivity of 133 mS.cm⁻¹ and 155 mS.cm⁻¹ at RT. The high ionic conductivity reported in these electrolytes is attributed to their ferroelectricity driven by the mobility of the ions and the electrolytes' semiconductor nature with surface-protected states as in higher-order topological insulators that contribute to the fast polarization of the electrolytes. These concepts will be further discussed in detail in Section 3.1.

Recently, a structural modification on Li⁺/Na⁺ antiperovskite electrolytes has been proposed with the introduction of large polyatomic units to replace single ions on crystallographic sites. In 2017, Fang et al. [95] developed a simulation that allowed them to propose a new Li⁺ antiperovskite structure composed of BH₄- polyatomic ions in A-sites (i.e., Li₃O(BH₄)). BH₄- is commonly known as a super-halogen, which has been suggested to replace halogen ions due to their similar chemistry but stronger electron affinity when compared to halogens. In 2019, Sun et al. [35] synthesized a sodium electrolyte containing BH₄- polyatomic ions into X₃BA-type antiperovskite structure for the first time, Na₃O(BH₄) from a solid reaction between Na₂O and NaBH₄ mixtures. This material was identified as having an antiperovskite structure and exhibiting an ionic conductivity of 4.4 mS.cm⁻¹ at RT and additionally displayed a low activation energy of 0.25 eV. Since the first synthesis reported for this electrolyte, other attempts to synthesize this double antiperovskite have been performed. In 2020, Ahiavi et al. [96] reported the mechanochemical synthesis of the $Na_3O(BH_4)$ electrolyte; however, the reported ionic conductivity was seven orders of magnitude inferior to the one proposed by Sun et al.[35]. The author explained that the hot-pressing pellets method used in [35] could alleviate the effects of blocking grain boundary, improving the ionic conductivity. Further research is still necessary to determine which synthesis method enhances the characteristics and nature of this electrolyte.

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Recently Tsai *et al.* reported the synthesis of double paddle-wheel sodium amide boroindridenine Na_{3-x}O_{1-x}(NH₂)_x(BH₄) [97]. The electrolyte contains two clusters of anions, the super halogen $BH_4^$ and the amide cluster anion NH_2^- . They can incorporate various ionic substitutions, including multiple cluster anions. This characteristic makes them highly versatile for investigating and understanding the paddle-wheel phenomenon. The *paddle-wheel* effect refers to the correlated rotations of cluster anions, enabling more accessible cation translation with reduced energy barriers and thereby enhancing cation mobility. Antiperovskites are appealing for studying paddle-wheel effects due to their ability to accommodate a wide range of ionic substitutions, including multiple cluster anions. The Na₃O(BH₄) electrolyte already discussed corresponds to a single paddle wheel system, whereas the Na_{3-x}O_{1-x}(NH₂)_x(BH₄) corresponds to a double paddle wheel system. The authors verified [97] that for x=1 Na₂(NH₂)BH₄ has a low activation energy of 0.268 eV, and when x = 0.5, the ionic conductivity is a hundred times greater than the extrapolated conductivity at x = 1.

Another common structural modification performed on X₃BA antiperovskite is the formation of double antiperovskites. Islam *et al.* [91] reported the first synthesis of a super-halogen Li-rich double antiperovskite Li₆OS(BH₄)₂ as a solid electrolyte. The synthesis consisted of a 3-step process starting with ball milling and liquid phase process, followed by heat treatment at 380 – 400 °C and water removal under a high vacuum to promote the creation of the Li₆OS(BH₄)₂ product, $Li_2O + Li_2S + 2Li(BH_4) \rightarrow Li_6OS(BH_4)_2$ [91], [98]. In the study [91], the band gap of the electrolyte was determined to be 4.03 eV. Furthermore, a symmetric cell was fabricated as Li/Li₆OS(BH₄)₂/Li; this cell is reported to have a long cyclic stability of about 3000 h at 60 μ A.cm⁻² current density, even though the increment of the current density seems to reduce the cyclic stability. The crystal structure of Li₆SO(BH₄) has been found to stabilize in the cubic Fm-3m phase [91] which is consistent with the antiperovskite structures of Li₃O(BH₄) and Li₃S(BH₄) but different from Li₃OCl whose cubic crystalline structure belongs to the perovskite conventional Pm-3m space group (Table 1). Moreover, this electrolyte exhibits exceptional mechanical stability, with a bulk modulus of 128.5 GPa. The combination of these properties renders this double perovskite suitable for ensuring a safe operation in all-solid-state ASS technologies.

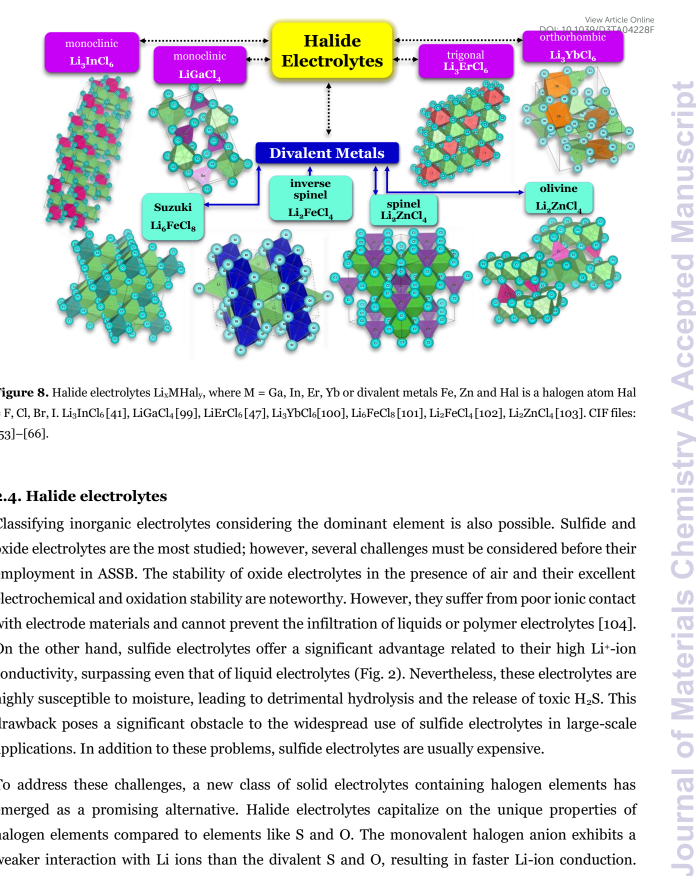


Figure 8. Halide electrolytes Li_xMHal_y, where M = Ga, In, Er, Yb or divalent metals Fe, Zn and Hal is a halogen atom Hal = F, Cl, Br, I. Li₃InCl₆[41], LiGaCl₄[99], LiErCl₆[47], Li₃YbCl₆[100], Li₆FeCl₈[101], Li₂FeCl₄[102], Li₂ZnCl₄[103]. CIF files: [53]-[66].

2.4. Halide electrolytes

Classifying inorganic electrolytes considering the dominant element is also possible. Sulfide and oxide electrolytes are the most studied; however, several challenges must be considered before their employment in ASSB. The stability of oxide electrolytes in the presence of air and their excellent electrochemical and oxidation stability are noteworthy. However, they suffer from poor ionic contact with electrode materials and cannot prevent the infiltration of liquids or polymer electrolytes [104]. On the other hand, sulfide electrolytes offer a significant advantage related to their high Li⁺-ion conductivity, surpassing even that of liquid electrolytes (Fig. 2). Nevertheless, these electrolytes are highly susceptible to moisture, leading to detrimental hydrolysis and the release of toxic H₂S. This drawback poses a significant obstacle to the widespread use of sulfide electrolytes in large-scale applications. In addition to these problems, sulfide electrolytes are usually expensive.

To address these challenges, a new class of solid electrolytes containing halogen elements has emerged as a promising alternative. Halide electrolytes capitalize on the unique properties of halogen elements compared to elements like S and O. The monovalent halogen anion exhibits a weaker interaction with Li ions than the divalent S and O, resulting in faster Li-ion conduction. Moreover, the larger radius of halogen anions leads to longer ionic bonds and increased polarizability in the electrolytes, facilitating the migration and plasticity of Li⁺-ions [91], [105].

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Selecting the proper central element M is fundamental to design halide SSEs successfully. Halide electrolytes can be divided in three classes, considering the central element: (1) Group 3, Li_3YCl_6 [46], Li_3ErCl_6 [47], and Li_3YbCl_6 [100]; (2) Group 13, $LiAlCl_4$ [106], $LiGaCl_4$ [99], and Li_3InCl_6 [41]; (3) Divalent metals: (1) Li_2ZnCl_4 standard spinel structure at RT [103]; (2) Li_2ZnCl_4 olivine structures [103]; (3) Li_2FeCl_4 inverse spinel structure [102] and (4) Li_6FeCl_8 Suzuki structure [101], Fig. 8. Considering the group 3 and 13 of halogen electrolytes, it is possible to highlight three structures: (1) orthorhombic ($Pna2_1$), (2) monoclinic ($P12_1/c1$), and (3) trigonal (P321). The trigonal and orthorhombic structure is based on the hexagonal closed-packed (hcp) anion arrangement, and the monoclinic structures are usually found in relatively large anionic compounds, such as Cl- and Br-, or relatively smaller metal cations. Regarding the divalent metal electrolytes, the reported structures are (1) olivine, (2) spinel, (3) spinel inverse, and (4) Suzuki; however, these structures are usually associated with lower ionic conductivity electrolytes.

The first ionic conduction was detected on lithium halide electrolytes in 1930 with the synthesis LiHal (Hal = F, Cl, Br, I) [107]. However, the electrolytes had low ionic conductivities, with the maximum being 0.001 mS.cm⁻¹ at RT [107]. Only in 2018, this class of electrolytes gained the interest of the scientific community when, for the first time, Tetsuya Asano et al. [46] mechanochemically synthesized Li₃YCl₆ LYC using a stochiometric mixture of LiCl and YCl₃. LYC electrolyte consists of a hexagonal close-packed (hcp)-like anion arrangement structure, where yttrium resides at the octahedral sites, and yttrium halides display high stability octahedrons YX_6^{3-} . The ionic conductivity of the LYC electrolyte is 0.51 mS.cm⁻¹ at RT. The previous conductivity was found to be a revelation since it was observed in a close-packed anion sublattice material; the latter varies from the source of high Li+-ion conductivity found for sulfide or oxide lithium-ion conductors. The electronic conductivity of LYC is 2.8×10^{-6} mS.cm⁻¹ corresponds to the upper characteristic values of electronic conductivities of an insulator [46] and agrees with our data for the band gap and DOS in Figs. 3 and 4. The activation energy was determined to be 0.40 eV [46]. In 2021, Schlem Romam et al. [108] developed a study to assess the lithium sub-structure in lithium conducting halides via temperaturedependent neutron diffraction of the LYC electrolyte. In this study, the author considered two different synthesis methods: ball milling and ampule synthesis. It was verified that for the LYC, the mechanochemical synthesis led to a local distortion of the structure, therefore improving the ionic conductivity. Large disorders motivated by a frustrated energy landscape enhance ionic transport. The disorder in the LYC electrolyte affects the polyhedral volumes of Li⁺ and their directional thermal

amplitude. Sendek *et al.* [109] performed a DFT simulation guided by a machine learning-based-nine approach to find new fast ionic conductors. From this study, it was concluded that Li_3ErCl_6 was an acceptable candidate to achieve good ionic conductivity and was expected to demonstrate an ionic conductivity of 3 mS.cm⁻¹ at 25 °C. However, when synthesized via ball milling and annealing, this electrolyte only displayed an RT ionic conductivity of 0.3 and 0.02 mS.cm⁻¹, respectively [47]. This electrolyte shows a trigonal structure $P\overline{3}m1$ [109], Table 1, Figs. 2-4. Similarly to what was reported for the LYC electrolyte, the ionic conductivity is reduced with increasing crystallinity during subsequent annealing processes [109].

Of all the halide electrolytes, the one that showed a higher conductivity at RT was Li_3InCl_6 . This electrolyte was first synthesized by a mechanochemical method in 2019 by Li *et al.* [82], which obtained an ionic conductivity of 1.49 mS.cm⁻¹, surpassing all its predecessors. Additionally, this electrolyte exhibited good stability toward conventional oxide cathode materials and good stability in air. The breakthrough of this electrolyte study arose when Li *et al.* [82] proposed a water-mediated synthesis technique to prepare Li_3InCl_6 [41] (Fig. 5d). The obtained RT ionic conductivity was 2.04 mS.cm⁻¹ (Fig. 2).

All the previously reported halide electrolytes have a setback to mass production; the middle element is usually a rare earth metal, leading to high production costs and compromising the commercialization of halide-based solid-state batteries.

In 2021, Kwak *et al.* reported the first cost-effective halide SSE, Li_2ZrCl_6 [45]. This electrolyte was prepared via ball milling and heat treatment (Fig. 5a). The use of different synthesis methods introduced considerable variations in the crystal structure and ionic conductivity. When prepared by heat treatment, the electrolyte presented a monoclinic structure with poor Li+ conduction 0.006 mS.cm⁻¹ at 30 °C. When it was synthesized by ball milling, it showed a hcp structure with a Liion conduction of 0.4 mS.cm⁻¹ at 30 °C. These results are atypical in the halide electrolyte world; the monoclinic structure is usually associated with fast ionic conduction, for example, in the Li₃InCl₆ electrolyte. Zirconium is an abundant metal on Earth, approximately an order of magnitude greater than yttrium or scandium and more than three orders of magnitude than indium [110]. The cost to produce most no-Li-containing chlorides needed to synthesize Li₃MCl₆ is usually above \$1000/kg, completely offsetting the cost-effectiveness of the LiCl precursor rounding (\$5.88/kg) [110]. Wang Kai et al. [110] reported the synthesis of Li₂ZrCl₆ electrolyte, using LiCl and ZrCl₄ as precursors. The cost of raw material for Li₂ZrCl₆ at a thickness of 50 μm is approximately \$1.38/m², which is significantly lower than the cost of Li_3YCl_6 (\$23.05/m²) [110]. Li_2ZrCl_6 is considered the most economical chloride system based on existing literature. Moreover, its cost is well below the \$10/m² threshold required for cost-competitive all-solid-state batteries. Additionally, the electrolyte was reported to be moisture-resistant at 5% relative humidity and displayed an ionic conductivity at room temperature of 0.81 mS.cm⁻¹.

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Until now, solely lithium halide electrolytes have been introduced. Nevertheless, some sodium halidenine DOI: 10.1039/D31A04228F analogues have already been synthesized. The primary challenge encountered in synthesizing sodium-based halides lies in achieving a significant ionic conductivity while preserving the thermal stability of the electrolyte.

In 2020, Schelm *et al.* [111] reported successfully synthesizing Na_{2.4}Er_{0.4}Zr_{0.6}Cl₆ with an ionic conductivity of ~0.035 mS.cm⁻¹ at RT. The research centered around synthesizing the Na_{3-x}Er_{1-x}Zr_xCl₆ compound series, exploring different values of $0 \le x \le 1$ to identify the optimal stoichiometry that maximizes the electrolyte properties. The study investigated the case where x = 0 corresponds to Na₃ErCl₆. The analysis revealed that Na₃ErCl₆ crystallizes with a monoclinic structure containing ErCl₆³⁻ octahedra frameworks at the corners of the unit cell. Each NaCl₆⁵⁻ prism is connected by an edge or corner sharing arrangement with another NaCl₆⁵⁻ octahedron. The obtained ionic conductivity for the Na₃ErCl₆ electrolyte at approximately 25 °C was 10⁻⁶ mS.cm⁻¹. In the case where x = 1 corresponds to the electrolyte Na₂ZrCl₆ displaying a trigonal crystal structure, the maximum reported ionic conductivity is 3.5×10^{-7} mS.cm⁻¹. When the incorporation of Zr⁴⁺ occurs, the structure adopted by the Na_{3-x}Er_{1-x}Zr_xCl₆ is monoclinic, as already discussed. Introducing smaller Zr⁴⁺ ions reduces lattice parameters. Additionally, the substitution of Er³⁺ by Zr⁴⁺ introduces sodium vacancies, leading to a tuneable ionic conductivity in the series of solid solutions Na_{3-x}Er_{1-x}Zr_xCl₆, which in turn provides a promising step toward halide-based sodium-ion conductors.

In the realm of Yttrium metal sodium halide electrolytes, a recent contribution by Xu *et al.* [112] presented a new electrolyte, Na₃Y₂Cl₉, designed using a building block method. A 3-step structure was proposed to create new structures incorporating Na⁺ions and (Y₂Cl₉)³⁻ units. Theoretical analysis conducted in the study demonstrates that this electrolyte exhibits a low energy barrier for ionic diffusion, leading to an ionic conductivity of 0.1 mS.cm⁻¹ at RT. Although the ionic conductivity may not be considered exceptional, the electrolyte demonstrates excellent chemical and electrochemical stability. Additionally, the calculated band gap for this electrolyte was determined to be 7.34 eV. Both agree with the wide herein simulated band gap of 5.2 eV and lower energy value for the conduction band $E_C = -0.11 \text{ eV}$ (for E = 0, the electrons at rest in a vacuum) for Na₃YCl₆ (Figs. 3a and 3b). The lowest energy for the E_C of Na₃YCl₆ is 1.62 eV higher than Na^o; therefore, Na₃YCl₆ is supposedly very stable against Na-metal, although with such a reduced electron affinity energy that it may decompose by being reduced from the surface in contact with a reductant.

Using DFT simulations, Liu *et al.* [113] proposed a new spinel chloride sodium electrolyte, $Na_2Y_{2/3}Cl_4$. This electrolyte is expected to exhibit a high ionic conductivity of 0.94 mS.cm⁻¹ at RT with a low activation energy barrier for Na^+ ion diffusion of 0.29 eV while possessing a semiconductor energy gap [113] from -3.85 to -0.68 eV, $E_0 = 0$ electrons at rest in vacuum, $E_g = 3.17$ eV; Fig. 4 shows the energy concepts. Changing to Indium, in April 2023, Zhao *et al.* [114] reported the controlled synthesis of NasAdZr Prime DOI: 10.1039/D51A04228F $_x$ In_xCl₆ electrolyte. The study was motivated by the limited number of reports on sodium halides, none discussing sodium halides containing indium. The authors investigated a series of solid solutions of Na_{2+x}Zr_{1-x}In_xCl₆ (0 ≤ x ≤ 1). The most promising candidate from this study was the electrolyte for which x = 0.5, N_{2.5}Zr_{0.5}In_{0.5}Cl₆, with an ionic conductivity of 0.066 mS.cm⁻¹ at RT.

Even though some promising improvements have been made in the production of halide electrolytes, the ionic conductivities remain considerably low when compared to other electrolytes, such as sulfide electrolytes. The advantages of doping existing electrolytes with halide elements such as chlorine are clear. A new class of electrolytes called oxyhalide seems to be the most promising. The previously discussed electrolytes and two recently reported ones, LiTaOCl₄ and LiNbOCl₄ [31], are plotted considering their optimized stoichiometry and reported ionic conductivities (Fig. 2).

Both oxide and halide electrolytes show considerably smaller ionic conductivities than sulfide-based electrolytes. Overall, oxyhalides display groundbreaking ionic conductivities, highlighting the Na⁺ and K⁺-glass electrolytes that have been reported to display self-charging behavior due to their ferroelectric nature [28], [115], [116] (Fig. 9).

3. Towards the future: Quantum Power Harvesting and Storage

Although perhaps the future will bring diversity into energy harvesting and storage-tailored devices, expensive solutions, such as the current rule in the switching (transistor) world, must be much superior to others to compensate for the extra cost. In the battery field, this means being safe, having high energy and power densities, and cycle and shelf lives. Solid electrolytes potentially better cover the first two, but being sustainable, permitting high cycle and shelf life, and being affordable are attributes just for a few chosen electrolytes.

Ferroelectrics are complex systems based on quantum mechanics principles; they may be protagonists of a disruption in the energy storage realm. Notably, halide-perovskites are being successfully used as photoelectric semiconductors in solar cells and may achieve ~35 % cell efficiency today [117], some with ferroelectric heterojunctions with $BaTiO_3$ [118]. Ferroelectrics, for example, HfO₂, may also recently be found in field effect transistors FET for a more efficient switch at RT [119].

Until now, from the best of our knowledge, the only electrolytes displaying a ferroelectric character, having a Li⁺, Na⁺, or K⁺ mobile ion, are LLTO, Na₃OCl, K₃OCl, A_{2.99}Ba_{0.005}OCl – glass with A = Li, Na, K; with LLTO sub-performing when compared to the others. These ferroelectric electrolytes are glassy-oxyhalides, uniting the best properties previously discussed. They are also inexpensive, especially the sodium-based versions, and are eco-friendly. However, they are hygroscopic, but from our experience, promising sulfides such as LPSCl, compared with A_{2.99}Ba_{0.005}OCl – electrolytes, are slightly more hygroscopic but, worse, release hazardous vapors (e.g., H₂S).

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As reported hereafter, not only are $A_{2.99}Ba_{0.005}OCl - glass$ with A = Li, Na, K ferroelectric, but the online properties such as self-charging and self-cycling based on coherent complex dynamics [28], [115], [116] sustained by internal feedback, picturing Otto cycles in quantum thermal machines such as described hereafter. Therefore, quantum battery solutions are a reality, and they are based on spontaneous polarization of the material driven by the difference in Fermi levels of the electrodes; polarization that, in traditional batteries, was seen as an obstacle.

The caveat of these neophytes is the difficulty in taming their properties driven by what they have best, their complex behavior, and their vast possibility of increasing their *energy information* even while set to discharge with a load.

Hereafter, the working principles of quantum batteries in their panoply of possibilities will be discussed.

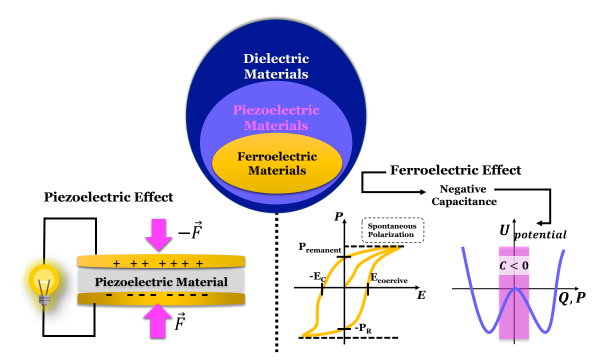


Figure 9. Dielectric, piezoelectric, and ferroelectric materials and devices. Force \vec{F} , polarization *P*, capacity *Q*, electric field *E*, potential energy *U*, and capacitance *C*, remanent polarization *P*_{remanent}, coercive electric field *E*_{coercive}.

3.1. Ferroelectric devices for harvesting and storing energy.

The development of self-charging systems is fundamental to elevating the next generation of electronic devices. These systems will simultaneously display the ability to storage energy as well as the ability to harvest energy. Piezoelectric materials are an option to transition the concept of self-charging to reality, specifically when these piezoelectric materials are also ferroelectric materials. The piezoelectric phenomenon occurs in dielectric materials that, when subjected to mechanical stress, can generate charge carriers and polarize (Fig. 9). Under an electrical field, the piezoelectric material undergoes a mechanical deformation or strain.

Ferroelectric materials have awakened the scientific community's interest due to their ability: to the display negative capacitance and, therefore, are expected to overcome fundamental limits of power dissipation in integrated circuits [120]. Ferroelectric materials can polarize spontaneously below the Curie temperature, forming permanent dipoles, even when no electric field is applied. However, above the Curie temperature, the ferroelectric materials transition into a paraelectric phase with no spontaneous polarization. The negative capacitance NC was predicted by the Landau model for ferroelectricity [120]. From this model, it is possible to define a capacitance as the reciprocal of the second derivate of the potential energy *U* with respect to the electric polarization charge, Q_F : C_F =

 $\left(\frac{d^2U}{dQ_F^2}\right)^{-1}$ [121]. These phenomena have been studied and observed in different harvesting and storage devices. In 2014, Salvatore *et al.* [122] verified experimentally that the negative capacitance phase stability in a field transistor (FET) varies with temperature. The NC region narrows down at high temperatures when approaching the Curie temperature of the ferroelectric material, validating experimentally Landau's theory. In 2015, Khan *et al.* [123] first synthesized a thin epitaxial ferroelectric film displaying negative capacitance. When the applied potential difference is inferior to the coercive voltage (Fig. 9), the capacity is positive, and therefore, the ferroelectric transistor behaves as a simple transistor. However, the authors verified that the ferroelectric switches to negative capacitance when the applied potential difference was superior to the coercive voltage (Fig. 9).

The negative capacitance phenomenon is not limited to transistor technology alone; it has also been observed in the field of batteries. In 2020, Braga *et al.* [116] reported the discovery of negative capacitance in an electrochemical cell that demonstrated the ability to self-charge. The researchers conducted the study using a cell containing two electrodes separated by a ferroelectric oxyhalide glass-electrolyte, previously investigated by the author in [28], [29]. The study established a clear linear relationship between negative capacitance double-layer capacitors and the system's remarkable capacity for self-charging, even when initially configured to discharge with a material load; the cells were set to discharge while maintaining a fixed external resistance. Although the current was set to be negative (discharge direction), the recorded voltage increased, and the current consistently remained above zero, indicating the presence of negative resistance alongside a negative capacitance regime. During this period of negative resistance, the battery exhibited self-oscillations, entering into a self-cycling regime.

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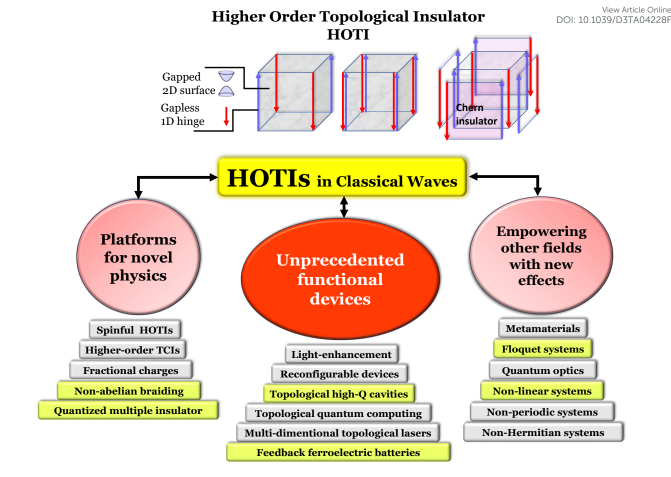


Figure 10. Higher order topologic insulators, HOTI. From left to right, the arrows in HOTI indicate chiral hinge currents, pairs of hinge modes running along the corners, and a Chern insulator with Hall conductivity. Below possible future applications of HOTIs [124]–[126] highlighting possible harvesting and storage coherent systems

3.2. Topologic insulators for harvesting and storing energy.

The theoretical underpinnings of solid-state physics rest upon two fundamental pillars: electromagnetism and quantum mechanics. One fascinating concept built on these two pillars is the quantum topological materials, highlighting topological insulators (TI). Studying topological insulators without referring to the quantum anomalous Hall effect is impossible.

What are, in fact, topological insulators? A topological insulator is a material whose bulk behaves as an electrical insulator. In contrast, its surface or edges behaves as an electrical conductor due to protected surface states, Fig. 10. The scientific community interest and potential applications of TIs in electronics, spintronics, and quantum information have driven tremendous efforts to find robust material realizations of TIs. The TI state has already been reported and observed in some materials. Some examples of topological insulator behavior have already been reported: halide perovskite displaying a non-trivial topological insulator phase [127], oxide topological insulator based on the superconductor BaBiO₃ [128], and the existence of a topological phase in the non-centrosymmetric NaSnBi material [129].

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A prominent area of research involves the search for materials displaying ferroelectricity-and automotion of the consequences of the coexistence of ferroelectric structural phase transition under strain. Pressure can also induce a topological phase transition from a standard insulator to a topological insulator in CsPbI₃. One of the clear advantages of FETI over the ferroelectric normal insulators FENI is the guaranteed presence of metallic surface states, which can serve as metallic electrodes to create an intrinsic short circuit condition, allowing automatic surface charge dissipation and internal device self-charging feedback. FETI may, therefore, work as a dual tandem dielectric-electrode with the advantage of reducing to the minimum the interfacial resistance and configuring one-electrode battery or capacitor where the electrode performs to maximize the potential bias in the cell.

Another interesting study was in 2017 by Monserrat *et al.* in 2017 [131]. They reported antiferroelectric topological insulators (AFTI) in orthorhombic AMgBi compound where A = Li, Na, K. From this study, the authors started to distinguish three types of AFTIs, depending on which states are topological insulators: (1) type-I AFTI with an antipolar normal state and polar topological state, (2) type-II AFTI with an antipolar topological state and polar normal states, and (3) type-III AFTI with antipolar and polar topological states. From the first principles calculations, the authors determined the conditions where each material behaves as the most promising candidate. The LiMgBi performs better under epitaxial strain, whereas KMgBi under hydrostatic pressure. In the LiMgBi and KMgBi AFTI phases of type I, an electric field could induce a topological phase transition by switching between the antipolar and polar states. The author concluded the crucial necessity of undertaking experimental investigations to explore these systems, particularly studying KMgBi under standard or moderate pressure and examining LiMgBi under epitaxial strain. Such experimental exploration may contribute valuable insights to our understanding of these materials.

High-order topological insulators (HOTIs) (Fig. 10) are a type of topological insulator that exhibit topologically protected states on their surfaces and on their boundaries of higher dimensions. This means that in addition to surface states, HOTIs possess robust and protected states along their hinges, corners, or other high-dimensional boundaries.

The unique properties of HOTIs make them attractive for potential applications in energy harvesting and storage devices. Here are a few ways HOTIs can be related to such devices [121]: **(1)** HOTIs can be utilized for efficient energy harvesting due to their topologically protected edge or boundary states. For example, the hinge states in 3D HOTIs can carry current and be engineered to exhibit specific energy-momentum dispersion relations. By properly designing the device geometry and

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connecting appropriate electrodes, energy can be extracted from external sources, such as vheat opinine mechanical vibrations, by converting these topologically protected states; **(2)** energy storage, HOTIs can also be relevant for energy storage applications. Their topologically protected states can facilitate the efficient transport and storage of charge carriers, enhancing the performance of energy storage devices such as batteries or supercapacitors. By leveraging the unique electronic properties of HOTIs, it may be possible to design more efficient and stable energy storage systems.

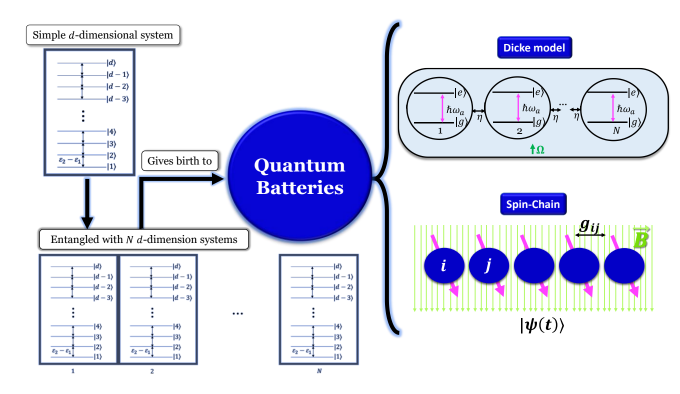


Figure 11. Quantum batteries sum up schematics.

3.3 Quantum or condensed matter batteries

The primary function of a battery is to be a reservoir of energy. Currently, technologies are evolving towards smaller devices, and batteries power these devices and, therefore, are needed to follow this tendency. However, as the unit cell of a battery approaches the size of a molecule or atom, quantum effects must be considered. The study of quantum batteries was first introduced by [129] in 2013 as a small quantum mechanism that temporarily stores energy. These batteries are described as *d*-dimensional quantum systems without degenerated energy levels, as demonstrated in Fig. 11. Work can be reversely extracted and deposited at the energy levels. As a result, the time evolution of the battery will be described by unitary operations.

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3.3.1 General Modulation

The Hilbert space, \mathbb{H} , of the wave function of the battery can be chosen to be *d*-dimensional [132], with a Hamiltonian, \mathcal{H}_0 , written in its base $|j\rangle$ as

$$\mathcal{H}_0 = \sum_{j=1}^d \epsilon_j |j\rangle\langle j| \tag{2}$$

where $\epsilon_j < \epsilon_{j+1}$ are the eigenvalues of $|j\rangle$ and $|j+1\rangle$, respectively. This condition assumes no degeneracy.

Considering a Hermitian operator field $\mathcal{V}(t) = \mathcal{V}^{\dagger}(t)$, which is used to extract energy and is only different from zero for time t, so that $0 < t < \tau_c$, where τ_c is the charging time. This field allows for a temporal evolution of the system, which is described by the Schrödinger equation, where the Hamiltonian, \mathcal{H} , will be $\mathcal{H} = \mathcal{H}_0 + \mathcal{V}(t)$. The general solution for this equation can be given by the unitary operator $\mathcal{U}(t_1, t_2)$, where a given state $|\psi(t_1)\rangle$ evolves to the state $|\psi(t_2)\rangle$, mathematically represented as $|\psi(t_2)\rangle = \mathcal{U}(t_1, t_2)|\psi(t_1)\rangle$. This operator is the solution for the Schrödinger equation when defined as the time-ordered exponential, with \hbar being the reduced Plank constant,

$$\mathcal{U}(t_1, t_2) = \mathcal{T}\left\{ e^{-\frac{l}{\hbar} \int_{t_1}^{t_2} \mathcal{H}(t') dt'} \right\}$$
(3)

The state $|\psi(\tau_c)\rangle$ is then written as,

$$|\psi(\tau_c)\rangle = \mathcal{T}\left\{e^{-\frac{i}{\hbar}\int_{t_1}^{t_2}\mathcal{H}(t')dt'}\right\}|\psi(0)\rangle.$$
(4)

It is possible to describe a state as its density matrix $\rho(t) = |\psi(t)\rangle\langle\psi(t)|$, becoming clear that the value of $\rho(\tau_c)$, is given by,

$$\rho(\tau_c) = \mathcal{U}(0, \tau_c) |\psi(0)\rangle \langle \psi(0) | \mathcal{U}^{\dagger}(0, \tau_c)$$

$$= \mathcal{U}(0, \tau_c) \rho(0) \mathcal{U}^{\dagger}(0, \tau_c).$$
(5)

The work extracted from the state is W,

$$W = \operatorname{Tr}\{\rho(0)\mathcal{H}_0\} - \operatorname{Tr}\{\mathcal{U}(0,\tau_c)\rho(0)\mathcal{U}^{\dagger}(0,\tau_c)\mathcal{H}_0\}$$
(6)

where Tr is the trace of the matrix. As \mathcal{U} depends on \mathcal{V} , a controllable variable, it is possible to control \mathcal{U} by choosing \mathcal{V} , allowing for the work extracted to be maximized (*ergotropy*) and defined as

$$W_{\max} = \operatorname{Tr}\{\rho(0)\mathcal{H}_0\} - \min\left\{\operatorname{Tr}\{\mathcal{U}(0,\tau_c)\rho(0)\mathcal{U}^{\dagger}(0,\tau_c)\mathcal{H}_0\}\right\}$$
(7)

where the minimum is taken over all unitary transformations.

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Charging the quantum battery. Charging a battery consists of varying its state $\rho(t_0)$ to a state $\rho(\tau_c)$, which gives $W = \text{Tr}\{\rho(\tau_c)\mathcal{H}_0\} - \text{Tr}\{\rho(t_0)\mathcal{H}_0\} \ge 0$. Conversely, discharging will take the state $\rho(t_0)$ to a lower energy state $\rho(\tau_d)$, where τ_d is the discharging time, $W = \text{Tr}\{\rho(t_0 + \tau_d)\mathcal{H}_0\} - \text{Tr}\{\rho(t_0)\mathcal{H}_0\} \le 0$. As shown in the previous section, this can be achieved by applying a $\mathcal{V}(t)$ field.

In practice, it is generally preferable to charge a battery quickly, which mathematically means maximizing the power $P \propto 1/t$. However, the average work must be considered, as it also needs to be maximized. The way to ensure both *P* and *W* are maximized without one being a detriment to the other is by maximizing the function \mathcal{F} , $\mathcal{F} = P^{\alpha}W^{1-\alpha}$, where α is a free parameter $0 \leq \alpha \leq 1$. Note that for $\alpha = 0$, $\mathcal{F} = W$ and for $\alpha = 1$, $\mathcal{F} = P$.

In [133] some remarks are made: (1) the cyclicity of the process is an important requirement since we are interested in the power; (2) if $t \to 0$, the quantum speed limits, which bond the time evolution by the inverse of the mean energy, would be violated; (3) there are natural constraints to the Hamiltonian that must be incorporated into the analysis. In particular, [133] restricts the trace norm of the Hamiltonian $Tr{H} \leq E_{max}$ with the gauge convention that states that the lowest eigenvalue of the Hamiltonian is zero.

3.3.2 The Dicke Quantum-Battery

The Dicke model in Fig. 11 consists of a collection of *N* two-level systems that interact with a singlephoton cavity model with an external driving field. Ferraro *et al.* [134] propose a model with a Hamiltonian $\mathcal{H} = \hbar \left(\omega_a J_z + \omega_c a^{\dagger} a + \lambda(t) \omega_c J_x (a^{\dagger} + a) \right)$, where parameter $\hbar \omega_a$ is the energy between the ground state $|g\rangle$ and the excited state $|e\rangle$, *a* and a^{\dagger} annihilates and creates a cavity photon with frequency ω_c , $J_{\alpha} = \sum_{i=1}^N \sigma_i^{\alpha}$, for $\alpha = \{x, y, z\}$, and $\sigma_i^{\alpha} = \mathbb{I}^{\otimes i-1} \sigma_a \mathbb{I}^{\otimes N-i}$ is the Pauli matrix of the i-th term, and \mathbb{I} is the identity matrix; $\lambda(t)$ is the step function, $\lambda(t) = \begin{cases} 1, \text{ for } 0 \leq t \leq \tau_c \\ 0, \text{ for all others} \end{cases}$, where τ_c is the charging time.

However, this model does not take into account the interatomic interactions and the driving field. To include these aspects, [135] proposes a model with new terms in the Hamiltonian that considers those interactions. The two-level systems will be regarded as having infinite-range interactions and are coupled to the single cavity mode via single-photon coupling. The Hamiltonian can be divided into two parts: (1) \mathcal{H}_0 the Hamiltonian of the battery, and (2) \mathcal{H}_1 the charging Hamiltonian $\mathcal{H}(t) = \mathcal{H}_0 + \lambda(t)\mathcal{H}_1$. The two Hamiltonian parts are given by,

$$\mathcal{H}_0 = \hbar \omega_a J_z; \tag{8}$$

$$\mathcal{H}_{c} + \mathcal{H}_{a-c} + \mathcal{H}_{a-a} + \mathcal{H}_{\text{field}} =$$

 $\mathcal{H}_1 =$

$$=\hbar\left(\omega_{c}a^{\dagger}a+2\omega_{c}\gamma J_{x}\left(a^{\dagger}+a\right)+\frac{\eta}{N}J_{z}^{2}+\Omega J_{x}\right)$$
(9)

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In Eq. (9) $\mathcal{H}_c = \hbar \omega_c a^{\dagger} a$ is the Hamiltonian of the cavity, $\mathcal{H}_{a-c} = 2\hbar \omega_c \gamma J_x (a^{\dagger} + a)_{DOT 101039/DSTA04228F}$ interaction between the atom and the cavity, $\mathcal{H}_{a-a} = \hbar \frac{\eta}{N} J_z^2$ is the interaction between atoms and $\mathcal{H}_{\text{field}} = \hbar \Omega J_x$ is the Hamiltonian due to the applied magnetic field along the *x* axis. The parameter $\hbar \omega_a$ is the energy difference between the ground state $|g\rangle$ and the excited state $|e\rangle$, γ is the two-level system cavity coupling strength, η is the atomic interaction strength, and Ω is the driving field strength.

Considering a close quantum system where the *N* two-level systems are in the ground state and coupled to the single-mode cavity in the *N* photon Folk state. The initial state of the total system is $|\psi(0)\rangle = |N\rangle \otimes |g, g, ..., g\rangle$.

The battery will start to charge when $\lambda(t) \neq 0$. By defining the density matrix $\rho(t) = |\psi(t)\rangle\langle\psi(t)|$, the total energy storage at a time *t* is obtained by, $W(t) = \text{Tr}\{\rho(t)\mathcal{H}_0\} - \text{Tr}\{\rho(0)\mathcal{H}_0\}$. In these operators' basis, the initial state $|\psi(0)\rangle$, can be written as $|\psi(0)\rangle = |N, \frac{N}{2}, -\frac{N}{2}\rangle$, giving all the two-level systems in the ground state (state of minimal energy).

The matrix element of equation (9) can be obtained by proceeding step by step and fixing $j = \frac{N}{2}$, so $m = -\frac{N}{2}, ..., \frac{N}{2}$. Summing all components, it is possible to obtain the Hamiltonian matrix. It is important to note that the number of photons is not conserved nor bounded by the Dicke Hamiltonian; thus, it can take an arbitrarily large integer value. In practice, it is necessary to design a cut-off so that $N_{\rm ph} > N$ on the maximum number of photons $N_{\rm ph}$. In studies [136], [134] the authors show that excellent numerical convergence can be achieved by selecting the maximum number of photons as $N_{\rm ph} = 4N$.

Charging Properties. Firstly, considering charging without the driving field, the Hamiltonian is $\mathcal{H} = \omega_c a^{\dagger} a + \omega_a J_z + 2\omega_c \gamma J_x (a^{\dagger} + a) + \frac{\eta}{N} J_z^2$. If $\eta < 0$, attractive interactions are being considered, whereas if $\eta > 0$, these interactions are repulsive. Additionally, in [135] $\eta = 0$ means no interactions.

When the battery system is in a weak ultra-strong coupling regime, the light-matter interaction energy is a comparable fraction of the bare frequencies of the uncoupled systems, and the atomic interaction, whether repulsive or attractive, always has a negative impact on the energy stored by the quantum battery. However, the repulsive interaction increases the charging power while attractive interaction decreases it. It is also possible to notice that the atomic interaction grows weaker as the coupling strength increases. When the coupling strength increases to the deep-strong coupling regime, characterized by interaction strengths surpassing the bare frequencies, the energy and charging power of the battery remain largely unaffected [135]. The latter shows an interesting feature from the atom-atom and atom-cavity field interactions. It is also shown that the maximum energy stored always has a critical behavior. The critical atomic interaction η_{crit} , was calculated in [135] as

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 $\eta_{\text{crit}} = \omega_a - \frac{4\gamma^2 N}{\omega_c}$, and at the critical point $\gamma = \frac{\sqrt{\omega_a \omega_c}}{2}$, where the system is in its normal-super-space phase.

Advantage of collective charging. Ferraro *et al.* [134] have shown that the energy in a singlephoton Dicke model scales with *N* and the charging power with $N^{\frac{3}{2}}$ for large *N*. Therefore, [135] assumes it reasonable to consider $P_{\max} \propto \beta N^{\alpha}$, for this extensive Dicke model. Then, $\log(P_{\max}) \propto \alpha \log(N) + \log(\beta)$, which by linear fitting allows us to obtain α .

Numerical simulations. To verify the benefits of collective charging, we generated a plot highlighting the relation between the maximum power and different numbers of two-level states. These simulations were conducted in the subspace of *j*, Fig. 12, and were developed using the programming language *Python* [137] through *Jupyter Notebook* [138], [139]. As anticipated, the maximum power and energy of the battery increases with the number of two-level systems, N. Furthermore, our simulation data exhibited a linear relationship when plotting the logarithm of the maximum power against the logarithm of N. The linear fit yielded an R-squared value of 0.9987, indicating a strong correlation consistent with the findings reported in the previous paper [135].

In the ultra-strong coupling regime, the scaling exponent can be approximately $\alpha = 1.8 > 3/2$. On the other hand, in the deep-strong coupling regime, the scaling exponent is approximately $\alpha = 1.5 = 3/2$ [135]. Regardless of the coupling regime, the external driving field strength increases the maximum charging power of the quantum battery. However, as the driving field strength increases, the advantage of the system's maximum charging power capabilities weakens.

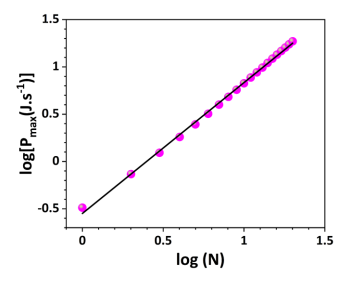


Figure 12. Maximum power vs. N two-level systems. By linear fitting, we have obtained α = 1.4.

Application of the Dicke quantum-battery. This model can be demonstrated experimentally by choosing an experimental set-up using a Bose-Einstein condensate, BEC, of ⁸⁷Rb atoms coupled

to a quantum electrodynamical QED cavity [140], [141]. The BEC with two levels [1] and 22 tigenine prepared in a time-averaged, orbiting potential magnetic trap [135]. After moving the BEC into an ultra-high-finesse optical cavity, an external controllable classical laser is applied to produce various transitions of the atoms between [1] and [2] states, thus charging the battery [135].

3.3.3 Spin Chain Quantum Battery

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Lee *et al.* [142] proposed another type of quantum battery, considering a one-dimensional XXZ Heisenberg spin-chain. This type of spin-chain has a coupling constant different in the *z* direction than the other directions, Figs. 11 and 13. In the absence of charging operations, the Hamiltonian is assumed to be static and represented by \mathcal{H}_0 , so that, $\mathcal{H}_0 = \mathcal{H}_B + \mathcal{H}_g$. \mathcal{H}_B is the Hamiltonian due to the magnetic field \vec{B} , and is defined as $\mathcal{H}_B = -\vec{B} \cdot \sum_{l=1}^{N} \vec{\mu_l}$, where $\vec{\mu_l}$ is the magnetic momentum. Assuming \vec{B} in the *-z* direction (Fig. 13) and assuming that the magnetic moment of the *i*-th member of the chain is given by $\vec{\mu_l} = \frac{1}{2}g_s\mu_B\vec{\sigma_l}$, where $g_s \approx 2$, μ_B is the Bohr magneton and $\vec{\sigma}$ is the vector of the Pauli matrices,

$$\mathcal{H}_B = B\mu_B \sum_{i=1}^N \sigma_i^Z,\tag{10}$$

 σ_i^a for $a = \{x, y, z\}$, is the Pauli matrix for the *a* coordinate and the *i*-th member of the spin-chain, and it is obtained by $\sigma_i^a = \mathbb{I}^{\otimes i-1} \sigma_a \mathbb{I}^{\otimes N-i}$.

The second term of the Hamiltonian \mathcal{H}_a , defines the interaction between different spins,

$$\mathcal{H}_{g} = -\frac{\hbar^{2}}{4} \sum_{i < j} g_{ij} \left[\sigma_{i}^{z} \cdot \sigma_{j}^{z} + \alpha \left(\sigma_{i}^{x} \cdot \sigma_{j}^{x} + \sigma_{i}^{y} \cdot \sigma_{j}^{y} \right) \right]$$
(11)

where g_{ij} is the interaction between spins *i* and *j* and α ensures that the interaction between the *z* direction and the other two directions is different, $|\alpha| \leq 1$. Lee *et al.* [142] consider two types of interactions: (1) the nearest neighbors g_{ij}^{NN} , and (2) the long-range interaction g_{ij}^{LR} given by $g_{ij}^{NN} = g\delta_{ij}$ and $g_{ij}^{LR} = \frac{g}{|i-j|^p}$, respectively, where *g* is the interaction strength and assuming that the spinchain is ferromagnetic $g \geq 0$, *p* is the decaying power of the long-range interaction.

Considering a ferromagnetic spin-chain, the ground state $|GS\rangle$, can be written as $|GS\rangle = |\downarrow\rangle \otimes |\downarrow\rangle \otimes$... $\otimes |\downarrow\rangle$ the tensorial product is obtained *N* times. All spins are, then, aligned with the \vec{B} field this way, the density matrix of the ground state ρ_{\downarrow} , is $\rho_{\downarrow} = |GS\rangle\langle GS|$.

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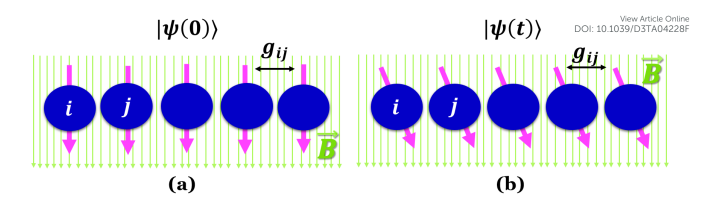


Figure 13. Initial and *t* state for the spin-chain battery. (a) $|\psi(0)\rangle$; (b) $|\psi(t)\rangle$.

Therefore, the initial energy will be given by $\text{Tr}\{\mathcal{H}_0\rho_{\downarrow}\} = \text{Tr}\{\mathcal{H}_B\rho_{\downarrow}\} + \text{Tr}\{\mathcal{H}_g\rho_{\downarrow}\}$ and using $\sigma^a|\downarrow\rangle = -\delta_{za}|\downarrow\rangle$, having into account that σ^a is the Pauli matrix for the *a* coordinate, and δ_{za} , is only one when a = z and zero in all the other cases,

$$\operatorname{Tr}\{\mathcal{H}_{0}\rho_{\downarrow}\} = -NB\mu_{B} - \frac{\hbar^{2}}{4}\sum_{i< j}g_{ij}$$
(12)

is the energy of the ground state (Fig. 13).

To import energy into the system [142] a charging Hamiltonian \mathcal{V} was considered, Eq (13). This Hamiltonian is obtained by varying \mathcal{H}_B the energy splitting Hamiltonian, Eq (10),

$$\mathcal{V} = \hbar\omega \sum_{i=1}^{N} \sigma_i^x \tag{13}$$

which is perpendicular to the original Zeeman splitting and uniform in space, Fig. 13. This can be obtained by applying another external magnetic field or simply rotating the system relative to the existing field.

Charging can be time-dependent; despite that, it is possible to consider the simpler case where \mathcal{V} is constant, $0 < t < \tau_c$. From Eq. (3) yet considering that $\mathcal{H} = \mathcal{V} + \mathcal{H}_g$ is constant with time, it is only necessary to use the exponential definition.

$$\mathcal{U}(0,t) = e^{\left\{-\frac{l}{\hbar}(\mathcal{H}_g + \mathcal{V})t\right\}}$$
(14)

where for *t*, the battery will be in the $\rho(t)$ state, obtained by, $\rho(t) = \mathcal{U}(0, t)\rho_{\downarrow}\mathcal{U}^{\dagger}(0, t)$. The work is then obtained by $W(t) = \text{Tr}\{\mathcal{H}_0\rho(t)\} - \text{Tr}\{\mathcal{H}_0\rho_{\downarrow}\}$ and the charging power by $P(t) = \frac{W(t)}{t}$.

Numerical simulations. To better understand how this model works and how its parameters^{nume} affect the performance of the spin-chain battery, a series of numerical simulations were developed. The simulations were written *in the coding language* Python [137] *through* Jupyter Notebook [138], [139] using the Quspin library recently developed by two faculty members of the Department of Physics at Boston University, Phillip Weinbe and Marin Bukov [143]. The battery's power was calculated and normalized with the power associated with a battery with isotropic interactions $\alpha = 1$ corresponding to that of four independent pins. In Fig. 14, it is demonstrated that independently of the type of interaction used, the power obtained will always be smaller when $\alpha = 1$, proving that a spin-chain without anisotropy provides less power. Furthermore, as the power increases with the anisotropy, the rotation or translation symmetries must be broken for a more powerful battery.

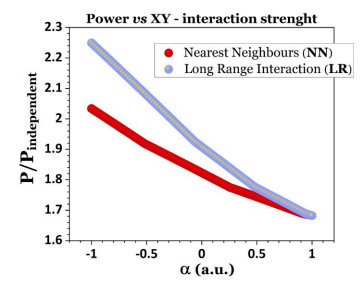


Figure 14. Charging power of a spin-chain quantum battery as a function of the anisotropy factor: N = 4, B = 3 *a.u.*, $\omega = 4B$, g = B and p = 1.

3.4 Quantum thermal machines

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Quantum thermal machines QTMs are advanced devices that operate on nanoscale systems, utilizing small two-level system quasi-particles known as qubits. These machines perform thermodynamic cycles and have various applications (Fig. 15). For example, they can function as nano-heat engines [144], leveraging the AC Josephson effect, or serve as quantum thermal thermometers as shown by Hofer *et al.* [145] and transistors in Joulain *et al.* [146].

One of the most significant challenges in quantum thermal machines is achieving optimal control of open quantum systems; therefore, improving this aspect has a crucial role in developing quantum information technologies. Erdman *et al.* [147] proposed an innovative approach that utilizes reinforcement learning (RL) to optimize the performance of QTMs. RL is a framework in which a computer agent learns and masters tasks through repeated interactions with an environment. The

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developed study focused on finite-time thermodynamics of QTMs within the Markovian regime Doi:10.1039/031A04228F employing a commonly used master equation. The case studies considered were **(1)** a heat engine based on a two-level system, **(2)** a refrigerator based on a superconductor, and **(3)** a heat engine based on a quantum harmonic oscillator.

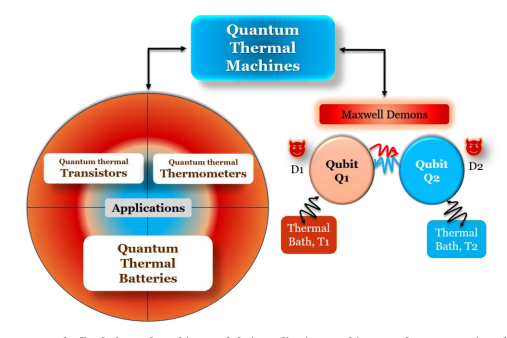


Figure 15. Quantum feedback thermal machines and their applications. Qubits Q1 and Q2 are monitored by D1 and D2, connected to thermal baths T1 and T2. In the case of Maxwell's demons, T1 = T2 = T. The two baths can also have different temperatures [148] and work as a refrigerator.

In the first case, the authors observed that initially, when the agent had no prior experience, it performed random actions, dissipating work into the heat baths instead of harvesting. However, the agent learned to control the heat engine over time, resulting in the average power saturating at a maximum positive value. Moving forward, the author considered a second system, which involved a refrigerator based on a superconducting qubit coupled to two resonant circuits acting as heat baths. In this case, unlike the previous scenario, the existence of coherence between instantaneous eigenstates needed to be considered. Similarly to the first case, in the beginning, the agent performs random actions; however, as time moves forward, the agent gains experience saturating the value of the average cooling power in a positive value. The study of the qubit-based refrigerator ended by evaluating the coefficient performance at maximum power COP. From this, it was possible to conclude that the COP at maximum power is 6 %; even though this value seems low, on-chip cooling applications typically aim to maximize the cooling power regardless of the efficiency of such a process. To end the study, the authors [147] considered a heat engine based on a collection of noninteracting particles confined in a harmonic potential. As in the first case, the agent makes the decision of which bath to couple with the oscillator. As it was observed in the first case, the power is negative for small steps, while it converges to a positive value as the agent gains experience. In this

study, the cycle discovered by the agent is similar to the Otto cycle (Fig. 16); however, it possessestime some additional features. The crucial difference regarding the Otto cycle is the ramping up of (t) while in contact with the baths, benefiting simultaneously from exchanging heat and modulating instead of achieving this in two separate systems.

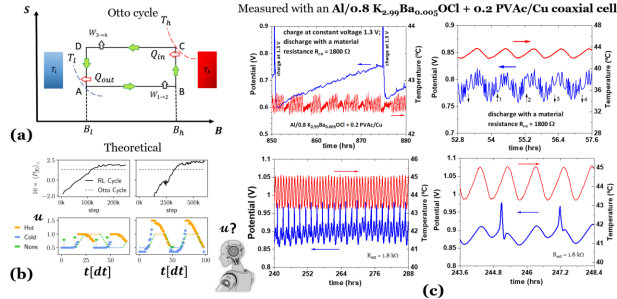


Figure 16. Otto cycle. **(a)** Entropy *S* vs applied magnetic field *B*. T_l – low reservoir temperature; T_h –high reservoir temperature; W – work; Q – heat. The system contacts the baths in the isomagnetic portions of the cycle (i.e., A \rightarrow D and A \rightarrow C). At A, $T = T_l$, and at C, the system's temperature reaches $T = T_h$. For the quantum version, the entropy S_B and S_D are calculated using the same thermal probabilities in A and C to ensure the quantum adiabatic portions A \rightarrow B and C \rightarrow D two-level quantum system. Reproduced from reference [149] CC BY 4.0; **(b)** Results of training the agent on the quantum harmonic oscillator heat engine model; the dashed lines represent the Otto cycle; Actions chosen by the agent, represented by the value of u, as a function of time. Reproduced from reference [147] CC BY 4.0; **(c)** the quantum thermal battery is a reality and shows spontaneous coherence obtained with a FETI in a double layer or, in this case, in a Al/0.8K_{2.99}Ba_{0.005}OCl + 0.2 PVAc/Cu coaxial cell when connected to a material resistor of 1.8 kΩ; the Otto field is an internal spontaneous electric field established to equilibrate de Fermi levels between the materials in electrical contact; the self-cycling drives the oscillation of the temperature of the cell [28].

Manzano [150] conducted a study to comprehend the influence of quantum effects on the operation and characteristics of thermal machines. Specifically, they focused on analyzing the entropy production and fluctuations in Maxwell's refrigerator with squeezing (Fig. 15). This study holds significant importance as it addresses a key challenge in quantum thermodynamics, which is to determine the impact of genuine quantum effects on the thermodynamic behavior and properties of small systems subjected to fluctuations. The study involved a thermal device between two resonant bosonic reservoirs at different inverse temperatures and an external memory system capable of erasing or storing information when in contact with the device. From this study, the authors verified ournal of Materials Chemistry A Accepted Manuscri

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that in scenarios of squeezed thermal reservoirs, the changes in entropy within the environmentation polition of squeezed thermal reservoirs, the changes in entropy within the environmentation result from the exchange of energy and coherence between the nonequilibrium reservoirs. This exchange mechanism also led to the induction of squeezing in the memory system at the steady state. While the fluctuation theorem for the total entropy production remained valid in this case, the fundamental processes associated with its increase were no longer straightforward jumps linked to the exchange of energy quanta between reservoirs.

How are quantum thermal machines related to quantum batteries? A recent study conducted by Tacchino *et al.* [151] aimed to prove that a battery can be charged when traversed by an incoherent heat flow. This heat flow arises from the weak interaction between a quantum system and two thermal baths at different temperatures. The process of charging the battery is initiated by applying a thermal gradient, which drives the transfer of energy and results in the battery being charged. The authors studied this subject by analyzing a model of two coupled qubits as the working fluid. The most important concepts to be taken into account were the **(1)** Hamiltonian of the two coupled qubits, Eq. (15), and **(2)** the heat absorbed from the hot bath.

$$H = \hbar\omega_0(\sigma_+^{(1)}\sigma_-^{(1)} + \sigma_+^{(2)}\sigma_-^{(2)}) - \lambda(\sigma_+^{(1)}\sigma_-^{(2)} + h.c.)$$
(15)

where $2\sigma_{\pm} = \sigma_x \pm i\sigma_y$ and $\{\sigma_j\}$ for j = x, y, z are the Pauli matrices, $\hbar\omega_0$ corresponds to the energy gap, λ is the coupling strength, and *h. c.* is the Hermitian conjugation. With this study, the authors verified that efficiency and output power are maximized in the short-limit cycle.

A study from October 2022 by Santos et al. [152] aimed to investigate the thermodynamic costs and limitations in the simplest case of optically pumping energy into an effective two-level quantum battery, as analyzed herein with the Dicke model (Figs. 11 and 12). The author started the study by analyzing a single quantum battery, highlighting the relationship between the pump system and efficiency and the dependence of the charging efficiency and the frequency of the pumping field. From this, it was possible to conclude that there is no optimal set of parameters to charge the battery. The process is more efficient for shorter times. For this time scale, the power storage in the battery is maximized; however, the efficiency decreases due to heat loss. The charging process is not affected by temperature if adiabatic elimination is verified. This condition is verified unless the dissipative thermal reservoir temperature is high enough compared to the energy gap of the eliminated level. Charging is most efficient at resonance, making this the optimal charging scenario.

Following, the author studied a second situation considering a quantum engine. It was verified that the best performance is achieved at low temperatures and short cycles. The maximum efficiency obtained is equivalent to a machine operating on an Otto cycle between two different reservoirs (Fig. 16). However, the efficiency of the Otto cycle is reduced due to energy dissipation in creating an effective high-temperature reservoir through optical pumping. This, however, can be minimized if the energy difference between the adiabatically eliminated level and the optically pumped level is not significant (Fig. 16). The highest efficiency in a short cycle situation is achieved when the pumping the provide the provided the provi

Bhandari *et al.* [148] conducted a study in May 2023 to explore using quantum measurements and feedback in coupled-qubit-based thermal machines. The researchers examined two configurations of these devices: (1) Maxwell's demon and (2) measurement-assisted refrigerator (Fig. 15). Before reviewing the results reported by the authors, it's fundamental to understand the model underlying both experiments. In their investigation, the authors employed a setup consisting of two coupled qubits, denoted as Q_1 and Q_2 . These qubits were connected to two thermal baths at temperatures T_1 and T_2 , respectively. Additionally, two measurement apparatuses, D1 and D2, enabled local quantum measurements on the respective qubits. The total Hamiltonian for the system was the sum of three components: (1) \mathcal{H}_Q describing the coupled-qubit system, (2) \mathcal{H}_b representing the Hamiltonian of the heat baths, (3) and \mathcal{H}_c , characterizing the tunnel-like coupling between the baths and the system. Hence, the total Hamiltonian can be expressed as $\mathcal{H} = \mathcal{H}_Q + \mathcal{H}_b + \mathcal{H}_c$.

In the case of Maxwell's demons, the environment has a uniform temperature, denoted as $T_1 = T_2 =$ T. This configuration aimed to extract heat from a single-temperature bath by leveraging invasive quantum measurements and feedback. On the other hand, the measurement-assisted refrigerator involved two distinct temperatures, T₁ and T₂, to maximize the heat extracted from the cold bath. The authors studied continuous and discrete measurements for Maxwell's demons scenario. In the case of one qubit measurement and considering a feedback-favorable situation, it was observed that the heat extraction from the thermal bath increases monotonously as a function of the coupling strength between the two qubits $\sigma_Z - \sigma_Z$. By comparing the heat extracted from a single setup when both qubits were measured together to the heat extracted from two parallel setups where only individual qubits were measured, the authors concluded that the former outperforms the latter due to a collective effect. The distribution of heat extraction for both one and two-qubit measurements was considered for the continuous measurement. The combined measurement of two qubits has a superior average heat extraction compared to individually measuring each qubit in two parallel setups. However, it is essential to note that higher fluctuations consistently accompanied this enhanced average heat extraction. In the study's second phase, the researchers examined the application of measurement-assisted refrigeration in a system consisting of coupled qubits connected to two thermal baths with different temperatures. It was demonstrated that while refrigeration can be achieved by using either measurement or swap operations independently, combining them leads to even more effective refrigeration.

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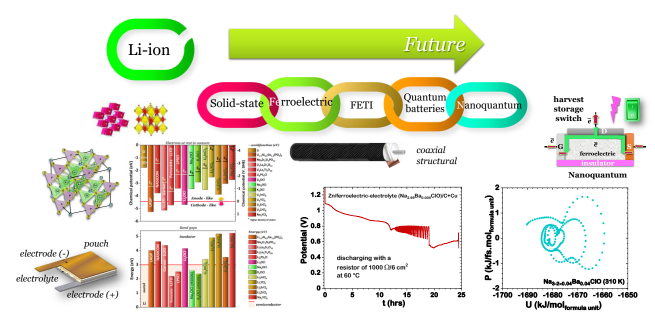


Figure 17. A perspective journey into the present and future of all-solid-state batteries. Note: for more information on the graphs on the right [115] reproduced under CC BY 4.0.

4. Conclusion

All-solid-state batteries are expected to be widely adopted once they become cost-effective or significantly more efficient than their liquid counterparts. Sustainability is a driving force for their adoption, as it requires promoting renewable energy sources, implementing energy-efficient practices, reducing greenhouse gas emissions, and promoting waste reduction and recycling. Consequently, the materials used must be non-hazardous. An exemplary achievement would be converting a portion of the wasted heat and thermal energy into electrical energy.

The best-performing solid electrolytes currently being adopted by the industry include argyrodites, oxyhalides, and combinations of argyrodites with oxides and halides and oxyhalides with sulfur. These solid electrolytes can be enhanced by doping them with higher valence cations to create vacancies that decrease the activation energy.

The synthesis method is also crucial as it relates to crystallinity and defects. Wet chemistry synthesis has proven to be the best method for obtaining higher ionic conductivity and even lower hygroscopicity. Being nanocrystalline or glass-amorphous has also been shown to improve cationic mobility within the solid electrolyte.

In addition to being an electrolyte, being a ferroelectric (e.g., perovskite LLTO and $A_{3-2x}M_x$ OCl where A = Li, Na, K and M = Mg, Ca, Sr, Ba) enhances ionic performance due to the spontaneous polarization of the ferroelectric driving ionic mobility, and vice versa.

Another intriguing characteristic, second only to the ferroelectricity dynamics described above clishine DOI: 10.1039/D31A04228F the semiconductor anode-like behavior of the best-performing electrolytes, including Li₆PS₅Cl, Na₃OCl, K₃OCl, and Li₃InCl₆. These electrolytes are semiconductors or have a band gap energy slightly above 3 eV, but their conduction band is placed at a similar energy level to the wellperforming electrolytes. This characteristic suggests that electrons likely play a role in the excellent performance of an electrolyte, even though an SEI layer forms as the Li⁺-ion, Li-batteries, and Na⁺ion traditional anodes have chemical potentials well above the conduction band of these electrolytes, facilitating the tunneling and drift of the electrons to the electrolyte's surface.

The ferroelectric electrolytes of the $A_{3-2x}M_x$ OCl family, where A = Li, Na, K, and M = Mg, Ca, Sr, Ba, are FETI (ferroelectric topological insulators) and, as previously mentioned, they have the guaranteed presence of metallic protected surface states that can serve as metallic electrodes. FETI can, therefore, function as a tandem dielectric-electrode, reducing interfacial resistance and configuring a one-electrode-battery or capacitor, where the electrode raises the bias between the chemical potential of the ferroelectrics' surfaces. Other ferroelectrics that have been proven to be topological insulators, although not electrolytes, are BaBiO₃, NaSnBi, and the cubic halide perovskite CsPbI₃, which has applications in photovoltaics and optoelectronics.

FETI electrolytes used in energy storage devices exhibit features of quantum thermal batteries, such as the possibility of establishing positive feedback from the positive to the negative electrode through their surface, leading to stability and self-charge even when the battery is set to discharge with a resistive material load.

Whether FETI is used in batteries or not, the future lies in carefully tailoring electrolytes and devices, utilizing quantum mechanics to our advantage. The future may be centered on quantum batteries with harvesting capabilities, which may exhibit coherent states even at macroscopic dimensions and room temperature. Combining energy harvesting and storage in one cell is environmentally advisable, especially if no additional efforts are required for their fabrication and use; the materials used are non-hazardous and are recyclable, safe, and readily available. These efforts and tendencies are not limited to batteries and energy storage devices but extend to photovoltaics, memories, optoelectronics, and transistors (Fig. 17).

Contributions

Original draft, visualization, quantum batteries simulations analysis: BMG; Original draft on quantum batteries and simulations' code: JFM; original draft on main perspectives, conceptualization, visualization, *ab initio* simulations, and supervision: MHB; review and editing: All. All authors have read and agreed to the published version of the manuscript.

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