Review

A Review on Temperature-Dependent Electrochemical Properties, Aging, and Performance of Lithium-Ion Cells

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Abstract: Temperature heavily affects the behavior of any energy storage chemistries. In particular, lithium-ion batteries (LIBs) play a significant role in almost all storage application fields, including Electric Vehicles (EVs). Therefore, a full comprehension of the influence of the temperature on the key cell components and their governing equations is mandatory for the effective integration of LIBs into the application. If the battery is exposed to extreme thermal environments or the desired temperature cannot be maintained, the rates of chemical reactions and/or the mobility of the active species may change drastically. The alteration of properties of LIBs with temperature may create at best a performance problem and at worst a safety problem. Despite the presence of many reports on LIBs in the literature, their industrial realization has still been difficult, as the technologies developed in different labs have not been standardized yet. Thus, the field requires a systematic analysis of the effect of temperature on the critical properties of LIBs. In this paper, we report a comprehensive review of the effect of temperature on the properties of LIBs such as performance, cycle life, and safety. In addition, we focus on the alterations in resistances, energy losses, physicochemical properties, and aging mechanism when the temperature of LIBs are not under control.

Keywords: lithium-ion battery; temperature-dependent properties; battery resistances; aging mechanism; thermal parameters; physicochemical properties

1. Introduction

Lithium-ion batteries (LIBs) as rechargeable devices play a key role in electrochemical energy storage systems [1]. With the steady improvements in the performances in terms of energy and power densities, the application areas for LIB technology are increasing. Particularly in electrical vehicles (EVs), their widespread applications attract much attention [2,3]. Despite their advantages and widespread applications, the LIBs suffer from safety problems, high cost, and thermal issues [4–9].

Battery degradation rates, which affect lifetime, are influenced by many external factors such as battery operating temperature, charge/discharge current rate, depth of discharge (DOD), and time intervals between full charge cycles [10–14]. Among these factors, the temperature is the major parameter affecting the battery performance such as power output, capacity, safety, and cycle life.
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For example, at higher operating temperatures, active material and electrolyte decomposition, changes in the solid electrolyte interface and side reaction rates increase resulting in capacity fading, power fading, and thermal runaway [5,16,17]. Similarly, at lower operation temperatures, reduced ionic conductivity of electrolytes, increased charge transfer resistance, high polarization of graphite anode (lithium plating), and limited diffusivity of lithium ions in electrodes result in low energy and power output [17–19]. A significant example of the influence of temperature is, for instance, reported by Nagasubramanian [20] who showed that when the ambient temperature is changed from 25 into −40 °C, the power and energy density of the 18,650 Panasonic cells are dropped by 98.7% and 95%, respectively. Similarly, Gong and Mi [21] studied the collected data from 7375 distinctive individual trips of one major commercial EV national wide in the United States, and showed that the average EV available range decreased from 120 miles at 20 °C to 60 miles at −20 °C.

The physical laws and mathematical functions that link material parameters with temperature are essential to simplify the evaluation of new materials by providing, through an inductive process, the scale-up of performance at the cell level. Moreover, with these functions properly parameterized, the sizing and integration of the thermal conditioning system to the battery pack is greatly facilitated, as simulation tools based on these equations can be then adopted. This engineering process is key, as the parameters influencing the battery lifetime are highly temperature-dependent. For example, Motloch et al. [22] reported that each grade of increase in temperature in a range of 30–40 °C results in approximately two months decrease in the battery cycle-life. Therefore, in order to comply with the life and safety requirements, the cells of a pack must remain within a well-defined temperature range, this range for the most common type of LIBs is between 10–50 °C [10,13,23]. In this work, a comprehensive and systematic review of a wide range of relevant studies on the temperature-dependent properties of LIBs and their effects on battery performance and degradation are investigated and summarized. Therefore, this work provides a hands-on overview of all the most relevant phenomena affected by the temperature within a lithium–ion cell, so that material developers as well as engineers in the battery design could use it for simulations, cell performance estimations, battery pack integrations, and thermal management developments.

2. Energy Conservation and Energy Losses

There have been published numerous studies on the modeling of the thermal behavior of Li-ion cells. The general energy balance for LIBs is derived from principles of nonequilibrium thermodynamics and is given by Equation (1) [24]

\[
\rho C_p \frac{\partial T}{\partial t} = k \frac{\partial^2 T}{\partial x^2} + Q_v
\]

where the temperature T can change spatially and temporally, and \(Q_v\) is the volumetric heat generation source. The primary source of heat generation in the Li-ion cells are the charge transfer reactions, which result in an irreversible reaction heat and a reversible entropy heat component. The rate of the irreversible reaction heat is given as a product of the reaction flux, the electrode surface area per unit volume, and the overpotential as below [25,26]

\[
Q_{\text{irr},v} = F a_s \hbar_{\text{o,cat}} \eta_n = F a_s \hbar_{\text{o,cat}} (\Phi_n - \Phi_e - E_0); n = c, a
\]

where \(n\) represents cathode (c) or anode (a). Regardless of charging or discharging the cell, the irreversible heat generation is an exothermic quantity. On the other hand, the reversible heat, which is generated due to entropy changes during intercalation or deintercalation, is given as below [25,26]

\[
Q_{\text{rev},v} = F a_s \hbar_{\text{o,cat}} T \frac{\partial E_{\text{Ap}}}{\partial T}; n = c, a
\]

Based on the rate of the reaction, the reversible heat can be either exothermic or endothermic. The secondary source of volumetric heat generation in the battery cell is ohmic heat, which is due to electronic current conduction in the solid phase and ionic current conduction in the electrolyte phase. The ohmic heat generation is given as the product of current density in the solid or electrolyte phase and the corresponding potential gradient. Thus, in the electrode region, it is given as [25,26]
\[ Q_{\text{ohm,V}} = -i_e \left( \frac{\partial \phi_e}{\partial x} \right) - i_s \left( \frac{\partial \phi_s}{\partial x} \right) \] (4)

and in the electrolyte region, it is given by

\[ Q_{\text{ohm,V}} = -i_e \left( \frac{\partial \phi_e}{\partial x} \right) \] (5)

In addition to heat generation originating from charge transfer processes (Butler–Volmer reaction) and ohmic heat, in some cells, a heat source due to phase transitions in the electrodes is also considered. This quantity, the phase transition heat is given as a product of the difference in the enthalpies of two phases (\( \beta: \text{Li poor phase}, \gamma: \text{Li rich phase} \)), and the rate of phase transition at the \( \beta - \gamma \) interface [24].

\[ Q_{\text{pt,V}} = \Delta H_\beta \text{v}_\beta \phi_\gamma \] (6)

Therefore, the overall energy conservation for a Li-ion cell can be rewritten as

\[ \rho c_p \frac{\partial T}{\partial t} = k \frac{\partial^2 T}{\partial x^2} + Q_{\text{irr,V}} + Q_{\text{rev,V}} + Q_{\text{ohm,V}} + Q_{\text{pt,V}} \] (7)

The maximum amount of electric energy derived by the electrochemical oxidation and reduction in the electrodes is proportional to the change in the standard free energy \( \Delta G (x) \) of the electrochemical couples, as stated in Equation (8) [27].

\[ \Delta G = -nFE_0 \] (8)

However, chemical energy cannot be fully converted to useful electric energy because of the polarization loss and the cell internal impedance during discharge [28–30]. The sources of polarization during charge/discharge are activation polarization, concentration polarization, and ohmic polarization, which will be discussed in this section. Therefore, the cell potential \( E \) is derived as shown in Equation (9) [30]

\[ E = E_0 \pm (\eta_a + \eta_c + iR_i) \] (9)

where: \( \pm \) is + during charge and – during discharge.

Activation polarization (charge transfer polarization) is the voltage loss associated with overcoming the energy barrier to the electrochemical reactions. It usually happens when the rate of oxidation or reduction reactions on the electrodes are not enough to yield a desired current. To reduce the activation polarization, the electrode materials need to be highly porous to have maximum reaction rates. Activation polarization is a function of the operating temperature, of the state of charge (SOC), and of the electrode properties, and can be expressed as in Equation (10) [30]

\[ \eta_{\text{act}} = \frac{RT}{anF} \ln \left( \frac{i}{i_0} \right) \] (10)

The activation polarization can also be expressed as the Tafel Equation [31]

\[ \eta_{\text{act}} = a + b \ln i \] (11)

where \( a = -\frac{RT}{anF} \ln i_0 \) and \( b = \frac{RT}{anF} \) [32].

Concentration polarization is the mass transfer resistance associated with the concentration difference between the electrode surface and the bulk of the electrolyte. To have a minimum concentration polarization, electrolyte diffusivity should be maximum to lower the mass transport limitations. The concentration polarization is given in Equation (12) [30,33]

\[ \eta_{\text{co}} = \frac{RT}{nF} \left( \frac{C_B}{C_E} \right) \] (12)

The other source of energy loss is ohmic polarization that causes the voltage (IR) drop which is proportional to the current drawn from the cell. The total cell internal impedance is originated from the ionic electrolyte resistance and the electronic resistance of the electrode active material, current
collectors, positive and negative tabs, and contact resistance between the current collectors and electrodes. Therefore, ionic conductivity of the electrolyte in conjunction with the electronic conductivity of the electrodes and current collectors should be high enough to lower the ohmic drop [33].

\[ \eta_R = -iR_i \]  

Therefore, Equation (9) could be rewritten as

\[ E = E_0 \pm \left[ (\eta_{ct})_a + (\eta_{ce})_a \right] + \left[ (\eta_{ct})_c + (\eta_{ce})_c \right] + iR_i \]  

Equation (14) shows that part of the useful potential is lost as a result of the aforementioned polarizations (Figure 1). The open circuit potential (OCP) of the cathode and the anode can be approximated by Taylor’s expansion as a function of the temperature and concentrations as in Equation (15) [34]

\[ E_{o,n}(T, SOC) = E_{o,n}(T_{ref}, SOC) + \frac{\partial E_{o,n}}{\partial T}(-T_{ref}); \ n = c,a \]  

Figure 1. Typical polarization of a cell (re-elaborated from [30]).

3. Temperature Effects on Electrochemical Resistances

Battery resistances are strongly temperature-dependent parameters [35,36]. In the literature, three major resistances are considered for the LIBs, which are the bulk resistance, the surface film resistance, and the charge transfer resistance [37,38]. The temperature dependency of these resistances is reviewed in the following section.

3.1. Ohmic Resistance

The ohmic resistance of the cell includes ionic resistance of the electrolyte and electronic resistance of the other cell components such as the electrode materials, the separator, the current collectors, and the cell terminals. In general, it specifies the capability of the cell to handle a certain current [39]. The instantaneous voltage drop at the beginning of the discharge process is due to the ohmic resistance.

The operating temperature of an LIB has opposite effects on the electrolyte ionic resistance and the electronic resistance of other electronic conductors. The ionic resistance of the electrolyte increases as the operating temperature decreases. This can be explained by the fact that the increase in the number of free ions due to the dissociation of molecules exceeds the growth of ion scattering during collisions with solvent molecules. On the other hand, the electronic resistivity of the conductors increases with increasing temperature [40].

Andre et al. [41] used electrochemical impedance spectroscopy to investigate the effect of operating condition, such as temperature and SOC, on the electrochemical performance of commercial high-power batteries. They showed that the ohmic resistance is almost unaffected by the
SOC. However, it is strongly influenced by the operating temperature of the cell due to the high sensitivity of the electrolyte diffusion rate on the temperature. These results were confirmed by many researchers, such as Zhange et al. [42] (Figure 2) and other researchers [43,44], however, Yang et al. [45] showed that the ohmic resistance increases rapidly when the SOC is between 0 and 0.2, and it becomes stable when SOC is larger than 0.2.

![Figure 2. Temperature dependency of internal resistance [42].](image)

### 3.2. Surface Film Resistance

The common electrolytes used in LIBs are thermodynamically unstable at the LIBs’ operating potential range, and they are reduced/oxidized and get chemically decomposed at the electrode/electrolyte interface [46,47]. The decomposed products are not soluble in the electrolyte solution, and they precipitate and form a protective film on the surface of electrodes. This protective layer is called solid electrolyte interphase (SEI) which protects the electrolyte components from being further reduced on the anode surface [16,48]. In addition, the SEI layer is not thermally and mechanically stable and in the case of abuse conditions such as an internal short circuit, overvoltage, and high temperatures’ exposure, the cells can undergo failure or safety issues such as thermal runaway. It is well-known that the SEI decomposition is the first process that leads to exothermic reactions under thermal abuse conditions.

The resistance associated with the SEI layer is expressed as the surface film resistance ($R_{SEI}$). Mogi et al. [49] evaluated the effect of temperature on surface film formation on nickel-based electrode at a temperature range of from 40 to 80 °C. According to their study, even though at higher temperatures the film on the anode surface gets thicker, the resistance associated with the SEI layer decreases with the increasing temperature. Suresh et al. [40] reported that the $R_{SEI}$ is nearly unaltered with SOC, while there is a significant variance with the temperature (Figure 3). They showed that variation in the $R_{SEI}$ with the temperature obeys Arrhenius’ law, and its magnitude decreases as temperature increases.

\[
\frac{1}{R_{SEI}} = A_0 \exp \left( - \frac{E_a}{RT} \right)
\]
3.3. Charge Transfer Resistance

The charge transfer process is expressed as transforming a solvated Li+ in the electrolyte into Li in the positive or negative electrode by accepting an electron. Generally, it includes three main steps: de-solvation of Li+ at the SEI/electrolyte interface, Li+ transfer through SEI layer or bulk of active material, and intercalation in the electrode. Normally, the last step is fast and the other two steps can be rate-limiting processes depending on the operating temperature, and electrode or electrolyte material [50]. For example, Wang et al. [51], who studied the low-temperature performance of graphite and mesocarbon microbeads (MCMB) electrode, found that the stage transformation (diffusion in the bulk electrode) is a rate-limiting step at room temperature, while transfer through SEI layer is a rate-determining step at −30 °C.

The resistance associated with the charge transfer process on the electrode–electrolyte interface is defined as the charge transfer resistance ($R_{ct}$). It is identified with the reaction kinetics from the Butler–Volmer Equation for charge-transfer-controlled electrochemical reactions [52].

$$R_{ct} = \frac{RT}{F i_0}$$  \hspace{1cm} (17)

The exchange current density, $i_0$, has a nonlinear relationship with the Lithium ion concentration in the solid phase as given below

$$i_0 = F A k_s c_s^\alpha$$  \hspace{1cm} (18)

The solid phase concentration can be given as

$$c_s = c_{s,max} \cdot SOC$$  \hspace{1cm} (19)

Therefore, $c_s^\alpha$ in Equation (18) can be written as

$$c_s^\alpha = c_{s,max}^\alpha \cdot SOC^\alpha = k_1 \cdot SOC^\alpha$$  \hspace{1cm} (20)

Thus, combining Equations (18)–(20), Equation (17) can be written as

$$R_{ct} = \frac{RT}{F^2 A k_s k_1 (SOC)^\alpha}$$  \hspace{1cm} (21)

The reaction standard rate constant is obtained from the Arrhenius Equation as a function of temperature
then, Equation (21) can be rewritten as

\[ R_{ct} = \frac{RT \exp \left( \frac{E_a}{RT} \right)}{F^2 A_i A_0 (SOC)^{\alpha}} = \frac{k \exp \left( \frac{E_a}{RT} \right)}{(SOC)^{\alpha}} \]  

(23)

where \( k = \frac{R}{F^2 A_i A_0} \) is a constant. The charge transfer resistance \( (R_{ct}) \) dependence on SOC and temperature can be obtained separately. For this purpose, Equation (23) can be written as follows

\[ \ln \left( \frac{R_{ct}}{T} \right) = \ln k + \left( \frac{E_a}{RT} \right) - \alpha \ln SOC \]  

(24)

and then, on differentiation, we obtain

\[ \frac{d \ln \left( \frac{R_{ct}}{T} \right)}{d \ln T} = \frac{E_a}{R} \]  

(25)

\[ \frac{d \ln \left( \frac{R_{ct}}{T} \right)}{d \ln SOC} = -\alpha \]  

(26)

By differentiation and rewriting Equation (24) in the form of Equations (25) and (26), Suresh et al. [40] evaluated the activation energy \( (E_a) \) and the transfer coefficient \( (\alpha) \) by plotting the variation of \( \ln \left( \frac{R_{ct}}{T} \right) \) with \( (\frac{1}{T}) \), and variation in \( \ln \left( \frac{R_{ct}}{T} \right) \) with \( \ln SOC \).

Zheng et al. [53] and Kumar et al. [54] found that the charge transfer resistance at the electrolyte/electrode interface is the most temperature-dependent resistance source of Li-ion cells. This is also confirmed by Liao et al.[55], who studied the low-temperature performance of a LiFePO₄ cathode and showed that, compared to \( R_b \) and \( R_{SEI} \), the \( R_{ct} \) is most significantly increased resistance, especially at subzero temperatures. This is because the \( R_{ct} \) is controlled by the kinetics of the reactions, while the \( R_b \) and \( R_{SEI} \) are mainly controlled by the electrolyte ionic conductivity, which is less temperature-dependent than the reaction kinetics. Kumar et al. [54] showed that the \( R_{ct} \) of a LiCoO₂ cell drops from 4 to nearly 0 Ω in a temperature range of -20 to 60 °C.

4. Temperature-Dependent Chemical–Thermal–Physical Properties of LIBs

The parameters used in the electrochemical–thermal models such as the kinetic reaction parameters, electrolyte conductivity, electrolyte and solid phase diffusivities, and open circuit voltage of the anode and cathode are mainly functions of the material type, the cell operating temperature, and the concentration. The most temperature-dependent physiochemical properties of the Li ion cell are the electrolyte density, the dynamic viscosity of the electrolyte, the solid-phase diffusion coefficient, the exchange current density, the electrolyte diffusion coefficient, the electrode electronic conductivity, the electrolyte ionic conductivity, and the chemical reaction rate, which will be discussed separately in the following sections.

4.1. Electrolytes

Generally, four types of electrolytes are used in LIBs: liquid electrolytes (solutions of lithium salt in aprotic solvents), solid/gel systems, solid-state electrolytes and ionic liquids [56]. In the following section, the effects of the operating temperature on the thermal, physical, and chemical properties of these electrolytes are reviewed.

4.1.1. Liquid Electrolytes (Standard)

The selection of solvents for liquid electrolytes is constrained since it must be mechanically, thermally, and chemically stable at the low potential of the anode and the high potential of the cathode [57]. Furthermore, the ideal liquid electrolyte should benefit from a wide operating
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temperature window, meaning that it should have a low melting point and a high boiling point. Xu et al. [58] reported the operating temperature range for a list of organic carbonates, esters, and ethers as an electrolyte solution. The cyclic organic carbonates are found to be the most desired solvents because of their high dielectric constants and formation of low energy complexes with lithium cations. However, these solvents suffer from the high viscosity and a high melting point. For instance, the ethylene carbonate (EC) is solid at room temperature. In the commercial cells, the acyclic alkyl carbonates are commonly added to benefit from their low viscosity and low melting point [56,59]. For example, Logan et al. [60] studied the effect of adding methyl acetate (MA) as an ester co-solvent on the low-temperature performance and rate capability of cells. They found that adding MA significantly decreases the viscosity of EC, while its effect on overall electrolyte conductivity is negligible. Generally, the LiPF6 as salt and EC as solvent are two indispensable components used in the present liquid electrolyte. Despite their many advantages for use as a salt and solvent, both are highly sensitive to the operating temperature. Simply, the LiPF6 contributes to a higher temperature failure due to its reactivity with solvents, and the EC contributes to a lower temperature instability due to its higher melting point and lower conductivity, especially at subzero temperatures [58].

The operating temperature range for the liquid electrolytes is defined based on a liquidus temperature, at which point one of the electrolyte components begins to crystallize and a bubble temperature, at which point one of its components begins to vaporize. Tarascon et al. [61] and Ding et al. [62] studied the temperature range for nonaqueous electrolytes used in Li-ion cells and found that the upper temperature limit (bubble temperature) in a binary or ternary electrolyte system is dominated by the lower boiling temperature. This is different from the operating temperature range defined for the Li-ion cell, since other factors are playing a role when an overall cell system is considered. For instance, the upper temperature limit is approximately 90 °C for dimethyl carbonate (DMC-), 110 °C for ethyl methyl carbonate (EMC-), and 120 °C for diethyl carbonate (DEC-) based electrolytes, while it is 70 °C for the LiPF6 salt. On the other hand, a lower boundary of the electrolyte range contributes to the low-temperature instability of a liquid electrolyte. Tarascon and Guyomard showed that the low-temperature limit for a binary EC/DMC (3:7) electrolyte is −25 °C [61]. Nonflammability of the electrolytes is also important if safety issues are considered [63]. Hu et al. [64] reported a dual-anion deep eutectic solution (combination of nitrile and lithium salts) as a nonflammable electrolyte with superior interphase stability and ionic conductivity.

Ion conductivity is one of the main characteristic properties of the electrolyte, which determines how fast the energy stored in the electrodes can be delivered. It mainly consists of two steps: (1) the solvation/dissociation of ionic compounds and (2) the migration of the solvated ions through the solvent. The ionic conductivity of liquid electrolytes is highly temperature-dependent and normally it increases monotonically with temperature. However, at very high temperatures, the ionic conductivity is more dominantly affected by the dielectric constants of the solvent mixture. One of the most common equations used for fitting the temperature dependence of conductivity data is the Vogel–Tamman–Fulcher (VTF) Equation [65,66]:

\[
\sigma = AT^{-0.5}\exp \left( \frac{-B}{RT - T_b} \right) \quad (27)
\]

The potential drop in the electrolyte and the absolute ionic current, without considering diffusion and convection, is directly proportional to the ionic conductivity. However, the binary diffusion coefficient is a proportionality factor between the ion flux and the concentration gradient [67]. In other words, the driving force for migration is the electric potential gradient, while the diffusion is due to the concentration gradient. Landesfeind and Gasteiger studied the binary diffusion coefficient over a large concentration and temperature range and for three different electrolyte mixtures. They reported that at 0.1 M LiPF6, the diffusion coefficients range from \( \approx 2.3-7.4 \cdot 10^{-6} \text{ cm}^2/\text{s} \) for EC:DMC (1:1 w:w) and EC:EMC (3:7 w:w) as well as from \( \approx 2.7-8.0 \cdot 10^{-6} \text{ cm}^2/\text{s} \) for EMC:FEC (19:1 w:w), as the temperature increases from −7.5 °C to +50 °C, respectively; at 3.0 M LiPF6, the diffusion coefficients values range from \( \approx 0.2-2.4 \cdot 10^{-6} \text{ cm}^2/\text{s} \) (EC:DMC (1:1 w:w)), \( \approx 0.2-1.6 \cdot 10^{-6} \text{ cm}^2/\text{s} \) (EC:EMC (3:7 w:w)), and \( \approx 0.2-2.0 \cdot 10^{-6} \text{ cm}^2/\text{s} \) (EMC:FEC (19:1 w:w)) for the same temperature increase [67]. Xu
et al. [26] reported the temperature-dependence of the Li ion diffusion coefficient in the electrolyte phase ($D_e$) as
\[
D_e(C_e, T) = 10^{\frac{4.43 + 54}{T - (229 - 5 \times 10^{-3} C_e)^4 - 0.22 \times 10^{-3} C_e}}^4
\] (28)

### 4.1.2. Solid/Gel Polymer Electrolytes

Polymer electrolytes (PEs) are generally a membrane that consists of a polymer host and a lithium salt. They possess good transport properties and compared to liquid electrolytes have some advantages such as non-flammability, no-internal shorting, non-volatility, no electrolyte leakage, improved safety, and lower decomposition at the electrodes’ surface [68–70]. The most commonly studied PEs are based on poly(oxa alkanes), poly(aza alkanes) or poly(thia alkanes) [71]. The materials used in polymer electrolytes (PEs) are required to be mechanically and thermally stable over a wide range of temperatures, to be electrochemically stable at high and low electrode potentials, to have low activation energy for conduction, and to be not reactive at the electrolyte-electrode interface. More importantly, they are required to have a low glass transition temperature ($T_g$) since the ionic conductivity becomes lower at the highly polymer crystalline structure [72]. Mindemark et al. [73] synthesized an electrolyte based on the copolymers of trimethylene carbonate and ε-caprolactone and reported that increasing the ester content improves the ionic conductivity and lowers the $T_g$. The polyether–lithium salt complexes are the most studied PEs for which the ionic conductivity is in the order of $10^{-7}$ to $10^{-8}$ S.cm$^{-1}$. Their low ionic conductivity is due to their highly crystalline structure at ambient temperatures which limits their applications [56]. Sun et al. [74] reported that plots of the ionic conductivity versus temperature for poly(trimethylene carbonate)-based polymer electrolytes demonstrate Vogel–Tammann–Fulcher (VTF) behavior in a temperature range of 294 to 357 K. Aside from the restricted ionic conductivity, there is additionally a high interfacial obstruction brought about by the poor contact and wetting of polymer electrolytes on regular composite cathodes/anodes. Sun et al. [75] reported improved electrode/electrolyte interphase properties by adding poly(trimethylene carbonate) oligomer as an interfacial mediator. The PEs also suffer from the low thermal conductivity together with the low lithium transference number, which varies between 0.1–0.2 [56]. For better mechanical and thermal stability, the PEs are crosslinked using a crosslinking agent such as isocyanates. Sukeshini et al. [76] studied the temperature dependence of the ionic conductivity for a ternary polymer electrolyte containing poly(vinyl chloride) as a solvent, lithium bis{(trimethyl sulfonyl)imide as a salt, and dibutyl phthalate as a plasticizer. Nest et al. [77] developed a relationship between the glass transition temperature ($T_g$) and the crosslinking agent molarity ($C$) as below
\[
\frac{1}{T_g} = \frac{1}{T_{g0}} - 7.6 \times 10^{-4} C
\] (29)
where $T_{g0}$ is the glass transition temperature at $C = 0$.

### 4.1.3. Solid-State Electrolytes

Solid-state electrolytes are thermally more stable and non-flammable with higher melting points compared to the commercially used liquid- and polymer-based electrolytes [78–80]. The functional solid-state electrolytes are required to have high room-temperature ionic conductivity and fast interfacial charge transport to be applicable in solid-state batteries [81–83]. They can be classified into three main types: garnet-type, sulfide-based, and phosphate-based electrolytes.

Garnet-type Li$_7$La$_3$Zr$_2$Ta$_2$O$_{12}$ (LLZTO) is considered as a good alternative for solid electrolytes because of its' sensibly high conductivity around 1 mS cm$^{-1}$ at room temperature and stability against lithium metal [78]. Its conductivity follows an Arrhenius-like behavior and reaches 16 mS cm$^{-1}$ at 100 °C [84].

Sulfide-based solid electrolytes (SE) are very alluring for application in all-solid-state batteries (ASSB) due to their high ionic conductivities and low grain boundary resistance. However, restricted chemical and electrochemical stability necessitates protection on both the cathode and anode side.
The initial challenge of poor ionic conductivity at ambient temperature has been met with sulfide-based materials like Li\textsubscript{10}GeP\textsubscript{2}S\textsubscript{12} (LGPS) or Li\textsubscript{2}S·P\textsubscript{2}S\textsubscript{5}. Their conductivities is as high as 25 mS cm\textsuperscript{-1}, surpassing that of current liquid electrolytes. [86,87]. In addition, their softness allows for good contacting, hence giving lower grain-boundary and interfacial resistance contrasted with oxide-based solid-state electrolytes [85]. A significant issue for sulfide-based SEs is their chemical instability in contact with air and moisture, as well as with Li metal and the cathode materials [88].

Solid-state electrolytes made of phosphate-based ion conducting glass-ceramics with a NASICON structure have rather lower conductivity and poor plasticity compared with sulfide-based materials. However, they have different focal points, for example, their chemical stability and ease of handling [89]. They additionally have great mechanical stability, which could stop the dendrite growth, which causes significant issues in some liquid electrolyte-based batteries [90]. A promising material system is LAGP (Li\textsubscript{1+x}Al\textsubscript{x}Ge\textsubscript{2−x}(PO\textsubscript{4})\textsubscript{3} with x = 0.3–0.7. The measured values of the ionic conductivities were in the range of 0.1 to 1 mS cm\textsuperscript{-1} at room temperature, but exhibited an increasing behavior as a function of temperature, reaching a level of the order of 10 mS cm\textsuperscript{-1} above 200 °C [91]. In addition, it has been shown by Cui et al. and Mahmoud et al. that the ionic conductivity can be influenced by the microstructure using different sintering methods [92,93]. Figure 4 shows, e.g., that microwave-assisted sintering can enhance the ionic conductivity of LAGP in comparison to melt-quenching [92]. Only recently, Rohde et al. have reported that the thermal conductivity of LAGP varies only slowly as a function of temperature, but its level depends on the composition. The apparent specific heat relies likewise upon the composition and exhibits enthalpy changes because of phase transitions at higher temperatures for LAGP samples with x > 0.5 [61].

![Figure 4. Ionic conductivity of LAGP glass-ceramics samples heat-treated at 800 °C for 6 h using 30 GHz microwave and conventional heating [92].](image)

4.1.4. Ionic Liquids

Unlike the standard electrolytes that are obtained by dissolution of salts in molecular solvents, the ionic liquids (ILs) are obtained by the melting down or liquefaction of a salt by providing a heat to counterbalance the salt lattice energy. The ionic liquids benefit from excellent properties, for example, non-volatility, non-combustibility, thermal and chemical stability, high ionic conductivity, and wide potential range, which makes them an ideal alternative to be used as electrolyte in batteries [94].

Zheng et al. [53] reported the effect of temperature on the physiochemical properties of an ionic liquid comprising of Trimethylhexylammonium (TMHA) cation and bis (trifluoromethane) sulfonylimide (TFSI) anion containing various concentrations of LiTFSI salt. They fitted the experimental density and viscosity data of the TMHA-TFSI ionic liquid to the linear model presented in Equation (30) and to the Arrhenius model in Equation (31), respectively. Figure 5a shows a linear
decrease in the solution density with the temperature, and Figure 5b shows the decrease in the solution viscosity with the temperature for the TMHA-TFSI ionic liquid containing different concentrations of LiTFSI salt.

\[ \rho = a + bT \]  \hspace{1cm} (30)

\[ \mu = A \exp \left( \frac{E_\mu}{RT} \right) \]  \hspace{1cm} (31)

Figure 5. (a) Variation in solution density, and (b) solution viscosity with temperature for TMHA-TFSI ionic liquid containing different concentrations of LiTFSI salt [53].

Shobukawa et al. [95] studied the temperature-dependent molar conductivity and ionic conductivity for novel lithium salts of borates having two electron-withdrawing groups (either 1,1,1,3,3,3-hexafluoro-2-propoxy or pentafluorophenoxy group) and two methoxy-oligo (ethylene oxide) groups (number of repeating unit: \( n = 3, 4, 7.2 \)). They used the Nernst–Einstein Equation to correlate the molar conductivity of the ionic liquids with the diffusion coefficients of the cation and anion as below

\[ \Lambda_{\text{diff}} = \frac{F^2(D_+ + D_-)}{RT} \]  \hspace{1cm} (32)

They also exhibited the Arrhenius plots of the ionic conductivity for the ionic liquids as in Equation (33).

\[ \sigma_\text{e}(T) = A_0 \exp \left( -\frac{E_{\sigma}}{RT} \right) \]  \hspace{1cm} (33)

They measured the ionic conductivity of the developed ionic liquids as \( 10^{-5} \) to \( 10^{-4} \) S cm\(^{-1} \) at 30 °C, which is lower than that of typical ionic liquids by two orders of magnitude. They explained the increase in the ionic conductivity of the solution with increasing temperatures by the inverse proportion of the solution viscosity with temperature based on the Vogel–Fulcher–Tammann–Hesse law

\[ \log_{10} \mu = A + \frac{B}{T - T_0} \]  \hspace{1cm} (34)

where \( A \), \( B \) and \( T_0 \) are constants, and \( T \) is the absolute temperature.

4.2. Electrode Materials

Temperature can strongly affect the mass transfer, reaction kinetics, and charge transfer rates in the electrodes. The most temperature-dependent parameters in the solid phases are the current density, the diffusion rates, the conductivity, and the reaction rate constant. The local current density
due the electrochemical reactions per active material area is governed by the Butler–Volmer Equation and is given by

$$i_{oc}(T) = a_i a_0(T) \exp \left( -\frac{\alpha \eta F}{RT} \right) - \exp \left( -\frac{\alpha \eta F}{RT} \right)$$

(35)

where $i_0(T)$ is the temperature-dependent exchange current density, and it is expressed as [96]

$$i_0(T) = A_i \exp \left( -\frac{\Delta H_0}{RT} \right)$$

(36)

Kumar et al. [54] calculated the value of $i_0$ as a function of the temperature and showed that it changes from 5.5 to 730 mA/cm$^2$ when the temperature changes from $-20$ to $60$ °C. Therefore, they reported that the low discharge capacity of the cell can be related to very low values of the $i_0$ at subzero temperatures.

The temperature-dependent behavior of the Li ion diffusion coefficient in the solid phase ($D_s$) [34,97–99], the solid phase electronic conductivity ($\sigma_s$) [100], and the reaction constant ($k_0$)[26], follows an Arrhenius law as stated in Equations (37)–(40). The constant diffusion coefficient $D_{s,ref}$ at the reference temperature is a function of SOC as presented in Equation (38).

$$D_s(T) = D_{s,ref} \exp \left[ \frac{E_{Ad}}{R} \left( \frac{1}{T} - \frac{1}{T_{ref}} \right) \right]$$

(37)

Xu et al. [26] reported the $E_{Ad}$ values to be 20 kJ/mol for the LFP (LiFePO$_4$) cathode and 4 kJ/mol for the anode. They also reported the $D_{s,ref}$ as below

$$D_{s,ref} = 3.9 \times 10^{-14}(1.5 - \text{SOC})^{3.5}$$

(38)

$$\sigma_s = \frac{\sigma_{s,0}}{T} \exp \left( -\frac{E_s}{k_BT} \right)$$

(39)

The coefficients for different Lithium metal oxide cathodes in the temperature range $-198$ to $27$ °C are reported by Heikes and Johnston [100].

$$k_0(T) = k_{0,ref} \exp \left[ \frac{E_a}{R} \left( \frac{1}{T_{ref}} - \frac{1}{T} \right) \right]$$

(40)

The values for $k_{0,ref}$ and $E_a$ are reported as $3 \times 10^{-11}$ m$^{2.5}$ mol$^{-0.5}$/s and 20 kJ/mol for the LFP cathode, and similarly, they are reported as $1.2 \times 10^{-14}$ m$^{2.5}$ mol$^{-0.5}$/s and 30 kJ/mol for the anode [99].

5. Temperature Effects on LIB’s Aging Mechanisms

5.1. Temperature-Dependent Aging Mechanisms

Identifying the exact aging mechanisms of LIBs has been one of the main challenges for the researchers. There are numerous factors that influence battery capacity loss and resistance growth which makes understanding of the aging mechanisms more complicated [101–105]. The main factors known to influence the degradation rate of the LIBs are the temperature [16,106], the low or high SOC [107,108], and the current [109]. The degradation process mainly takes place in the electrodes, and there are significant differences between the aging mechanisms of positive and negative electrodes. The main aging mechanisms have been known for a long time [16] and can be summarized as: [110]

- Structural changes in the insertion electrode;
- Electrolyte decomposition;
- Active material dissolution;
- Phase change in the insertion electrode;
- Passive film formation over electrodes and current collector surface.

By developing an electrochemical based electrical (ECBE) model and linking the model parameters to specific ageing mechanisms, Leng et al. [111] have shown recently on a graphite/LCO coin cell that the LIB degradation is a hierarchical process that returns, as indicated by a progression of associated forms. These procedures are: (i) solid electrolyte interphase (SEI) growth on the anode
surfaces; (ii) cathode phase change from active hexagonal to less active spinel phases; (iii) Joule heating because of the raised cell resistance resulting in the cracking and re-growth of SEI along with surface spallation at the cathode, as well as degradation of the separator; (iv) active Li losses as a result of metal plating on the electrodes during overcharge and over discharge; (v) exfoliation of the negative electrode and the degradation of positive electrode; and (vi) intergranular contact detachment inside electrodes. These procedures results in overall decay in the capacity limit and the LIB working potential in an unexpected way.

The operating temperature of the LIBs greatly influences the electrochemical performance, the cycle life, and the safety of the batteries [5,7,110–112]. It is also one of the main factors affecting the aging rate of the batteries. In recent years, many researchers have studied the effects of operating temperature on the aging mechanisms. Markevich et al. [113] studied the behavior of graphite electrode at elevated temperatures (up to 80 °C); Ramadass et al. [114] studied the capacity fade of Sony 18,650 Li-ion cells at room temperature, 45, 50, and 55 °C; Handel et al. [115] studied the thermal decomposition of the electrolyte, and Bodenes et al. [116] investigated the high-temperature influences on the degradation process of positive electrode’s binder, SEI, and positive active material. Uddin et al. [117] summarized the effects of high operating temperature on negative and positive electrodes, respectively. The effects of temperature on the degradation rate of electrodes will be discussed separately in the following sections. The electrolyte aging mechanisms can be regarded as a part of electrode aging and will not be discussed separately.

5.1.1. Temperature Effect on Aging of the Negative Electrode

The most common materials chosen as the negative electrode for LIBs are graphite, carbon, silicone or Lithium titanate Li4Ti5O12 (LTO) [101,118]. Graphite anodes have some advantages over the other anode electrodes such as high capacity, low cost, long cycle life, low volume expansion, and safety [101,119,120]. Despite these advantages, as an anode material, its aging is significantly depending on the operating temperature, which limits the LIBs’ application on subzero and high ambient temperatures. The temperature dependent aging mechanisms of the anode material are reviewed below, and they are summarized in Figure 6.

![Figure 6. Schematic of the temperature effect on aging of the negative electrode (re-elaborated from Uddin et al. [117]).](image-url)
5.1.1.1 High Temperature

The common electrolytes used in Li-ion cells are not electrochemically stable at the voltage range that battery anodes are operating. Therefore, the main aging mechanism of a graphite electrode is believed to be the formation and growth of the solid electrolyte interphase (SEI) on the electrode surface with time [105]. The SEI layer is developed through the reduction in electrolyte components and the formation of a passive film at zero potential on the graphite surface [16,102,121]. Since the SEI layer is not fully permeable for the lithium ions, the amount of recyclable lithium ions decreases as the thickness of the SEI layer increases with the number of cycles, which leads to capacity fading [122]. Another possible graphite aging mechanism can be solvent co-intercalation into the carbon and exfoliation of the carbon particles. This process results in the formation of carbon intercalation compounds (CIC), which in turn lead to irreversible capacity fading [12,123,124].

The developed SEI layer is not thermally stable, and increased temperature results in an increased degradation rate of the SEI layer [125]. The collapsed SEI layer must be replaced by a new one. Thus, more Lithium ions are consumed in the new SEI layer replacement process. Moreover, the old broken layer fills the empty volumes, resulting in decreased available surface area on the active material [126]. A high temperature also accelerates the formation of CIC. Leng et al. [110] used an electrochemistry-based model to investigate the effect of temperature on the aging behavior of a cycled LIB within the temperature range from 25 to 55 °C. They have studied the impact of temperature on the aging rate of LCO electrode (cathode), graphite electrode (anode), and electrolyte separately. Based on their study, both of the aging mechanisms of graphite electrode (SEI and CIC growth) lead to more degradation at higher operating temperatures. Another negative impact is that at high temperatures, metastable organic SEI components are converted into more stable inorganic components. Since these inorganic components are more stable, their growth may decrease the conductivity of the SEI layer [127].

5.1.1.2 Low Temperature

On the other hand, the cold operating temperature, in conjunction with the high charging rate and non-uniform current distribution result in lithium plating on the negative electrode [128]. Lithium plating is defined as precipitation of the lithium ions on the electrode surface when the potential of the electrode is very close to that of the lithium deposition potential. The main reason is the reduced rate of Li ion diffusion into the anode particles and accumulation of the ions as inactive metallic Li on the particles surface at the low potential of the anode. Therefore, lithium plating at lower temperatures decreases the amount of cyclable lithium ions and results in capacity fading [129,130]. Uddin et al. [117] summarized the effects of low operating temperature on a negative electrode.

Another effect of low temperature on the anode is the intercalation gradient with cycling which accelerates the electrode aging. At low temperatures, when the cell is charged at higher current rates, a mechanical strain is applied on the anode due to a sharp ion gradient which results in crack formation and splits in the graphite particles [127,131,132].

5.1.2 Temperature Effect on Aging of the Positive Electrode

A large number of previous studies have been focused on the degradation mechanisms of the lithium metal oxide cathodes [104,133–135]. Briefly, the changes that have been observed on the lithium oxide metals are: [16,103,105,136,137]

- Degradation of active material;
- Degradation of cathode components such as conductive agents, binder, corrosion of current collectors;
- Electrolyte oxidation and SEI formation;
- Interaction between the aging products of a positive electrode (dissolved within the electrolyte) with a negative electrode.

The thermal stability of a cathode material has been one of the most important safety-related concerns of cell manufacturers and researchers [138]. The LiMPO4 cathodes, where M stands for
transition metals such as Fe, Mn and Co, have attracted much attention due to their better stability at elevated temperatures and good safety features [139]. However, the transition metal oxide cathodes suffer from the dissolution of the transition metals into the electrolyte at high temperatures and high SOC. This process not only affects the cathode morphology, but also changes the SEI layer composition on the anode surface. After the dissolution in the electrolyte, the transition metals migrate through the electrolyte and subsequently deposit on the anode side where they can form dendrites on the anode surface or react with the SEI layer to form new compositions [140]. According to many researchers [141–143], Ni is the most stable transition metal in terms of dissolution in electrolytes and, in comparison with the other common transition metals, the following decreasing order has been observed: Mn > Co > Ni. The low Ni dissolution can be explained by the stable NiO-like rocksalt phase generated at high potential via loss of oxygen [140].

The other high-temperature degradation mechanism at the cathode side is related to the oxidation of the electrolyte at the cathode electrolyte interface (CEI) and the instability of the CEI itself [144]. These processes are irreversible and result in a passivation film formation on the cathode surface, which is called a surface layer or passivation film. The electrolyte oxidation occurs when the Fermi level of the cathode material is located below the electrolyte HOMO (highest occupied molecular orbital) level [145]. Edstrom et al. [146] studied the temperature effect on the cathode-electrolyte interface for different cathode chemistries: LiMn2O4, LiCoO2/LiNi0.8Co0.2O2, and carbon-coated LiFePO4. They reported that the composition of the SEI layer on the cathode surface is dependent on the chemistry of the cathode material. However, in all of the cases, the thickness of the SEI layer on the cathode side is increased with the increasing temperature indicating the higher electrolyte oxidation at higher temperatures. Leng et al. [110] studied the effect of temperature on the degradation of an LCO (LiCoO2) cathode. They found that degradation of an LCO electrode could be related to two main mechanisms: formation of SEI layer and structural/phase change of the electrode. Both of these two mechanisms reduce the reaction rate of the Li ion insertion/de-intercalation and the charge transfer rates. Furthermore, the influence of these mechanisms on degradation of the LCO electrode increases with temperature. This impact is more evident for a larger number of cycles [147,148]. The temperature dependence of the cathode material is summarized in Figure 7.

**Figure 7.** Schematic of the temperature effect on aging of the positive electrode (re-elaborated from Uddin et al. [117]).

Unlike the high-temperature effects on the cathode aging, there are limited studies on the low-temperature effects. Many researchers such as Gandoman et al. [149] and Uddin et al. [117] have neglected the low-temperature effects on the cathode aging. However, Wu et al. [150], who applied non-destructive aging detection methods on LIBs, reported that the vulnerability to low-temperature (10 °C) primarily originates from cathode. They explained this phenomena by the increased charge-
transfer resistance in the electrode-electrolyte interface due to the passive layer growth and phase transitions on the cathode surface.

5.1.3. Temperature Effect on the Electrolyte Decomposition

The concerns over aging of electrolytes necessitates to study the thermal stability of the electrolyte at elevated and low operating temperatures. Contrary to electrode material, thermal stability of the electrolytes has not been studied so much. The electrolytes which tolerates both elevated and very low temperatures are especially important in military and space-related applications [151]. The main reason of electrolyte decomposition at elevated temperature is reported to be the decomposition or hydrolysis of LiPF$_6$ by trace moisture which results in the formation of Lewis acid, HF, PF$_5$, or PO$_4$F$_3$. These formed components later interact with the cyclic or linear carbonates of the solvent and causes the electrolyte to be decomposed [58]. Botte et al. [152] used a differential scanning calorimeter to study thermal stability of the LiPF$_6$–EC:EMC electrolyte, and they reported that it is highly dependent on the salt concentration, the solvent concentration, and the heating rates. Eshetu et al. [153] reported that cyclic solvents such as EC and PC, are more thermally stable than linear solvents such as DMC and DEC. Especially for LiNi$_{1-x-y}$Mn$_x$Co$_y$O$_2$ (NMC) cathodes with high Ni content, such as LiNi$_{0.8}$Mn$_{0.1}$Co$_{0.1}$O$_2$ (NMC811), even at room temperature, the electrolyte decomposition plays a major role at elevated operating voltage [154].

5.2. Modeling of Temperature-Dependent Aging Mechanisms

Modeling of the aging mechanisms and degradation rate is necessary for lifetime prognosis of LIBs [155] and reducing the time and the cost needed for the experimental aging tests [23,156–158]. There are a number of aging models in the literature such as models based on neural networks (machine learning) [159–164], physiochemical models [165–167], and empirical models [108,168–170]. The temperature effect on the aging of LIBs has been well studied and formulated. The proposed models for both calendar and cycle aging are reviewed below.

5.2.1. Calendar Aging Model

Calendar aging is described as the capacity loss due to progress in time that occurs independent of charge/discharge cycling. The calendar capacity loss is described with a square-root dependence on time, which accounts for the diffusion-limited capacity loss. Most studies done on calendar aging mechanism suggest that the rate of calendar aging of the battery strongly depends on the temperature and the SOC (potential that the battery is kept) [171–174]. A high temperature accelerates the rate of chemical side reactions, which in turn results in increasing the SEI layer thickness and irreversible Li ion loss [175–177].

Schimpe et al. [178] reported a model predicting the cyclic aging with time in which the capacity loss is proportional to the square root of time.

$$Q_{Cal} = K_{Cal}(T, SOC) \sqrt{t} \tag{41}$$

where $K_{Cal}$ is a stress factor and is a function of time and SOC, as shown in Equation (42)

$$K_{Cal}(T, SOC) = K_{Cal.ref} \exp \left[ \frac{-E_{a,Cal}}{R \left( \frac{1}{T} - \frac{1}{T_{ref}} \right)} \right] \cdot \exp \left[ \frac{\alpha F}{R \left( E_{0.ref} - E_0(SOC) \right)} \frac{T_{ref}}{T} \right] + k_0 \tag{42}$$

where $K_{Cal.ref}$ is defined as the calendar reference stress factor at $T = 298.15$ K and SOC = 50%, $E_{0.ref}$ is the open circuit potential at SOC = 50%, and $k_0$ is a constant.

Xu et al. [179] also modeled the calendar aging as time, temperature, and SOC dependent stress factors as the following equations. However, they proposed that the time stress factor changes linearly with time.

$$Q_{Cal} = S_t(t) S_{SOC}(SOC) S_T(T) \tag{43}$$

where the stress model for temperature and SOC is derived from Arrhenius law, and the stress model for time is obtained as:
\[ S_t(t) = k_t t \] (44)

Wang et al. [175] proposed that calendar aging is mostly affected by temperature and time, but they ignored the SOC effect.

5.2.2. Cyclic Aging Model

Cyclic aging includes all the degradation processes when the battery is cycled using any kind of load profile [180–182]. Cyclic aging mechanisms are different and depend on several factors. Schimpe et al. [178] proposed a model for the temperature and SOC-dependent cyclic aging and investigated the dominant cyclic aging mechanism separately. The main cyclic aging mechanisms are:

- High Temperature \( Q_{\text{Cyc,High}} T \)
- Low Temperature \( Q_{\text{Cyc,Low}} T \)
- Low Temperature, High SOC \( Q_{\text{Cyc,Low}} T \text{ High SOC} \)

Thus, the temperature-dependent cycling aging models are different at low and high temperatures. At high temperature levels, the cyclic aging is proportional to the total charge throughput during both charging and discharging (\( Q_{\text{Tot}} \)), as shown in Equation (45).

\[ Q_{\text{Cyc,High}} T = K_{\text{Cyc,High}} T(T) \cdot \sqrt{Q_{\text{Tot}}} \] (45)

The cyclic aging stress factor is a function of the temperature, which is expressed as

\[ K_{\text{Cyc,High}} T(T) = K_{\text{Cyc,High}} T, \text{ref} \cdot \exp \left( \frac{-E_{a,\text{Cyc,High}} T}{R} \left( \frac{1}{T} - \frac{1}{T_{\text{ref}}} \right) \right) \] (46)

At low temperatures, the cyclic aging depends on both temperature and current rate, and it is proportional to the square root of charging aging, \( Q_{\text{Ch}} \).

\[ Q_{\text{Cyc,Low}} T = K_{\text{Cyc,Low}} T(T, i_{\text{Ch}}) \cdot \sqrt{Q_{\text{Ch}}} \] (47)

The cyclic stress factor at low temperature as a function of the temperature and charging current is given as follows

\[ K_{\text{Cyc,Low}} T(T, i_{\text{Ch}}) = K_{\text{Cyc,Low}} T, i_{\text{Ch}}, \text{ref} \cdot \exp \left( \frac{-E_{a,\text{Cyc,Low}} T}{R} \left( \frac{1}{T} - \frac{1}{T_{\text{ref}}} \right) \right) \cdot \exp \left( \beta_{\text{Low}} T \cdot \frac{i_{\text{Ch}} - i_{\text{Ch},\text{ref}}}{C_0} \right) \] (48)

where \( \beta_{\text{Low}} T \) is a constant.

The last cyclic aging mechanism is associated with the mechanism at low temperature and high SOC which is expressed as

\[ Q_{\text{Cyc,Low}} T \text{ High SOC} = K_{\text{Cyc,Low}} T \text{ High SOC}(T, i_{\text{Ch}}, \text{SOC}) \cdot \sqrt{Q_{\text{Ch}}} \] (49)

For the mentioned mechanism, the cyclic aging stress factor is a function of the temperature, the charging current, and the SOC, and is expressed as follows

\[ K_{\text{Cyc,Low}} T \text{ High SOC}(T, i_{\text{Ch}}, \text{SOC}) = K_{\text{Cyc,Low}} T \text{ High SOC}, \text{ref} \cdot \exp \left( \frac{-E_{a,\text{Cyc,Low}} T \text{ High SOC}}{R} \left( \frac{1}{T} - \frac{1}{T_{\text{ref}}} \right) \right) \cdot \exp \left( \beta_{\text{Low}} T \text{ High SOC} \cdot \frac{i_{\text{Ch}} - i_{\text{Ch},\text{ref}}}{C_0} \cdot \left( \text{sgn}(\text{SOC} - \text{SOC}_{\text{ref}}) + 1 \right) \right) / 2 \] (50)

The parameters obtained from fitting the data for cyclic aging can be found in the work of Schimpe et al [178]. Overall, the combined calendar and cyclic aging can be expressed as follows: [178]

\[ Q_{\text{Loss}} = Q_{\text{Cal}} + Q_{\text{Cyc,High}} T + Q_{\text{Cyc,Low}} T + Q_{\text{Cyc,Low}} T \text{ High SOC} \] (51)
Wang et al. proposed two main aging mechanisms for the LIB with composite cathode material which are high current rates at low temperature (cycle aging) and Li ion loss at high temperature due to increased side reactions and SEI layer thickness increase (cycle and calendar aging) [138]. Their proposed model for total aging (cycle and calendar) is as below

\[
Q_{\text{loss}} = (a \cdot T^2 + b \cdot T + c) \exp[(d \cdot T + e) \cdot I_{\text{rate}}] \cdot \text{Ah}_{\text{throughput}} + f \cdot \sqrt{t} \cdot \exp[-E_a/RT] \tag{52}
\]

Xu et al. [179] proposed a semi-empirical aging model based on the stress models as in Equation 53. They ignored the C-rate effects in their model and linked it to the battery operating temperature.

\[
Q_{\text{Cat}} = S_{\text{DOD}}(\text{DOD})S_{\text{SOC}}(\text{SOC})S_{\gamma}(T) \tag{53}
\]

6. Temperature Effects on LIB’s Performance and Safety

The temperature effect on battery resistances, physiochemical-thermal properties, and aging mechanisms have been reviewed in the previous sections. Variations in the aforementioned properties with temperature affect LIB’s electrochemical performance, safety, and the cycle life of the battery. Tables 1 and 2 are summarizing the thermal effects on LIB’s performance and safety, respectively.

| Thermal Effect | Author | Material | Remark |
|----------------|--------|----------|--------|
| Performance    |        |          |        |
| Capacity/Power fade | Ramadass et al. [185] | LiCoO2/C | The main parameters that significantly attribute to capacity fade were loss of primary (Li+) and secondary (LiCoO2/carbon) active material together with the rate capability losses. |
|                 | Belt et al. [186] | SAFT America proprietary G4 lithium ion chemistry | Capacity and power fade showed a weak inverse temperature relationship. The expected lifetime of the system decreases drastically when a 10 °C increase in temperature (i.e., from 25 to 35 °C) was considered, from 102 months to only 58 months. |
|                 | Stroe et al. [187] | LiFePO4 | |
|                 | Cordoba-Arenas [188] | Li(Ni0.33Mn0.33Co0.33)O2/LiMn2O4 blend | |
| Self-discharge  | Utsunomiya et al. [189] | Hard carbon (HC), synthetic flake graphite (FG), and spherical natural graphite (SG) | Higher specific surface area of the electrode and higher storage temperature significantly increases the rate of self-discharging. |
| Discover **Kurzweil et al.** [190] | Li(Ni_{0.33}Mn_{0.33}Co_{0.33})O_2 | For lithium-ion batteries, capacitance obtained by the electrochemical impedance spectra dependably showed the accessible electric charge in the working extent between full charge (without overload) and cut-off voltage (without deep discharge). Present a pulse-measurement technique as a novel method for characterizing the self-discharge behavior as a function of temperature. |
| **Schmidt et al.** [191] | LiNi_{0.8}Co_{0.15}Al_{0.05}O_2/LiCoO_2 blend | The reported that the local electrochemical impedance changing with temperature in different regions of the jelly roll results in non-uniform current density. Temperature differences among the cells cause unbalanced discharging and aging. They reported that even low currents, for example, 0.1 C, can prompt critical inhomogeneities, while a higher cell temperature for the most part leads to more pronounced inhomogeneities. They attributed the poor low temperature performance of Li-ion battery to the significantly high R_{ct} of the electrodes. Based on these studies it seemed likely that Li-ion battery electrodes composed of nanoscopic particles of the electrode material could mitigate this low-temperature performance problem. |
| **Electrical balance** Fleckenstein et al. [106] | LiFePO_4 | |
| **Yang et al.** [192] | | |
| **Oswald et al.** [193] | LiFePO_4 | |
| **Low temperature performance** Zhang et al. [18] | lithium nickel-based mixed oxide/graphite | |
| **Sides & Martin** [194] | Li_{0.2}V_2O_5 | |
Huang et al. [195] LiCo0.2Ni0.8O2/mesocarbon microbead graphite

They reported that the main reasons for the poor performance in the graphite electrodes are (i) the low value and concentration dependence of the Li diffusivity and (ii) limited Li capacity.

Table 2. Review of thermal effects on LIB’s safety.

| (Thermal Abuse) | GP | A123 Lithium Ion Cell | Thermal Runaway Happened Due to Local Overheat Originated from Increased Contact Resistance |
|-----------------|----|-----------------------|----------------------------------------------------------------------------------|
| (Thermal abuse) | Beauregard [196] | LiFePO4 | They exposed the cells to high temperatures and found that unlike the oven test, the accelerated rate calorimetry (ARC) tests do not fully capture the self-heating and thermal runaway safety hazard of a cell. |
| (Electrolyte decomposition) | Bugryniec et al. [197] | LiCoO2 | The amount of released gases increased with the increase in the cell temperature. |
| (Review on thermal issues) | Ohsaki et al. [198] | LiNi0.8CO0.15Al0.05O2 | Described thermal runaway in 3 steps; Slow anodic reactions starting at 90 °C, exothermic reactions on cathode starting at 140 °C, O2 release from cathode materials and interfacial oxidation of electrolyte starting above 180 °C |
| (onset-of-thermal-runaway (OTR) temperatures) | Wen et al. [7] | LiNi0.33Mn0.33Co0.33O2 | Showed that the temperature at which thermal runaway occurs changes with the SOC. |
| (Gas formation & pressure rise under thermal abuse) | Lei et al. [200] | Li(Ni0.33Mn0.33Co0.33)O2, LiMn2O4, LiFePO4 | Measured the internal pressure increase in 18,650 cells due to gas formation by exothermal reactions, showed that onset temperatures, maximum temperatures and temperature rates during thermal runaway as well as activation energies depend on the cathode materials |
| (Lithium plating at low temperature) | Wang et al. [201] | LiFePO4 | The thermal stability of SEI layer is deteriorated with growth of dendrite reducing the thermal runaway temperature of the battery. |
| (Improved thermal stability of an electrolyte) | Hofmann et al. [202] | Li(Ni0.33Mn0.33Co0.33)O2 | Showed that electrolyte mixtures based on ethylene carbonate and dimethyl sulfone with higher flashpoints of 140 °C substantially improve the cell safety. |
| Topic                                                                 | Authors                  | Key Components                                                                 | Notes                                                                                                                                                                                                 |
|----------------------------------------------------------------------|--------------------------|--------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Preventing thermal runaway by a pressure reduction                    | Hofmann et al. [203]     | Li(Ni0.33Mn0.33Co0.33)O2                                                        | Showed that fire and cell explosion during thermal runaway could be prevented by a pressure reduction.                                                                                                   |
| Computational study of the safety regime                             | Zhang et al. [204]       | LCO (LiCoO2), NMC(LiNiMnCoO2), LFP(LiFePO4)                                     | Safety regime and thermal runaway zone for LIBs are computationally studied and the effects of cathode material are investigated.                                                                        |
| SOC influence on thermal reactions                                    | Perea et al. [205]       | LiFePO4, Li(Ni0.80Co0.15Al0.05)O2                                              | Effects of SOC on thermal stability and thermal runaway characteristics of cells are studied using ARC.                                                                                                |
| Review on safety issues under mechanical abuse loading               | Liu et al. [206]         | Lithium-ion batteries                                                           | The safety aspects in conjunction with the coupled mechanical–electrochemical–thermal behavior of LIBs under mechanical abuse are reviewed.                                                             |

Besides the absolute temperature of a battery, non-uniform temperature distribution between the cells inside a battery pack and within each cell causes electric unbalances, poor battery performance, and capacity and power fading [207–211]. Temperature non-uniformity along single cell layers causes non-uniform current density along the cell and, consequently, localized aging [15,193,212,213]. Chiu et al. [214] developed an aging model to simulate and analyze the capacity fade of ten series connected cells operated at different temperatures. According to their study, by setting each cell with a different temperature, the whole battery pack’s capacity is limited by the highest temperature cell. Troxler et al. [215] applied a controlled temperature gradient between the cells of a battery pack with one hundred cells and proposed a model for the effect of the temperature gradient on the cell resistances. In this approach, they computed the equivalent resistance of sets of resistors that are connected in parallel, and each resistor is a function of temperature

$$R_{\text{eff}} = \left[ \frac{1}{\sum_{j=0}^{N} 1/R(T(x_j))} \right]^{-1}$$

(54)

where $R_{\text{eff}}$ is the effective resistor of a pack with a temperature gradient between the cells, and $R$ is a temperature-dependent resistance parameter. They used an Arrhenius Equation to describe the SEI resistance and the charge transfer resistance as a function of temperature, as in Equation (55)

$$R_j(T) = R_{\text{ref}} \exp \left[ \frac{-E_a}{R \left( \frac{1}{T_{\text{ref}}} - \frac{1}{T} \right) } \right]$$

(55)

Klein et al. [216] studied the effects of a temperature gradient on the nominal capacity of a 10Ah Li ion pouch cell. They used a PID temperature controller to apply controlled non-uniform temperature conditions and performed hybrid pulse power characterization (HPPC) tests to figure out the instantaneous power capability effects. They found that the overpotential values increase as the temperature gradient ($\Delta T$) increases. Moreover, they reported that the overpotential sensitivity to $\Delta T$ is more distinguished at lower temperature levels, and the temperature gradient influences the maximum capacity as the SOC decreases with increasing $\Delta T$.

### 7. Conclusions

After a comprehensive analysis of the literature, this paper reports a consensus on the massive influence of the temperature on the key cell components and parameters. Among these, the open-
Circuit voltage, solid-state kinetics in the electrodes, electrolyte conductivity, and growth phenomena of the SEI layer play the main role.

Consequently, the temperature also affects the macroscopic behaviour of the cell, thus the available capacity and power capability are enhanced or massively reduced by temperatures outside of the favourable operating range between 10 and 35 °C for almost any lithium-ion cell. Outside of this range, the cycle or calendar life suffers from reduction. The predominant ageing mechanism caused by the low temperature is the plating of metallic lithium on the anode, while the elevated temperatures accelerate the degradation of the cathode and SEI layer growth on the anode, leading to capacity fade and internal resistance increase. In the worst case, the temperature triggers reactions which could give rise to critical hazardous conditions like a thermal runaway. In this work, the main thermal–chemical–physical governing equations and their dependency from temperature have been summarized. These are essential for overcoming the thermal issues of the LIBs. The work proposes an organically organized collection of the most relevant temperature-dependant cell-governing equations. Moreover, for these relationships, initial parameter sets have been provided which help the reader to select and implement the specific equation in his own investigations. Examples of typical application problems are the physicochemical simulation of new cells as well as the development of the battery management system, which usually aims to maintain the battery within the optimum temperature and voltage windows for maximal performance and cycle life achievement.

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**Nomenclature**

| Symbol | Description |
|--------|-------------|
| \(A_0\) | Preexponential factor |
| \(A_{io}\) | Cell exchange current density multiplier |
| \(a_s\) | Electrode specific surface area (m\(^2\)) |
| \(C_0\) | Nominal capacity (mAh) |
| \(C_s\) | Bulk concentration of electroactive species (mol/m\(^3\)) |
| \(c_e\) | Lithium concentration in electrolyte (mol/m\(^3\)) |
| \(C_t\) | Concentration in an electrode (mol/m\(^3\)) |
| \(C_p\) | Specific heat (J/kg.K) |
| \(c_s\) | Solid phase concentration (mol/m\(^3\)) |
| \(c_{s,max}\) | Maximum concentration that can be taken from the electrode (mol/m\(^3\)) |
| \(D_e\) | Diffusion coefficient of Li ion in electrolyte (m\(^2\)/s) |
| \(D_s\) | Diffusion coefficient of Li ion in solid (m\(^2\)/s) |
| \(E_0\) | Open circuit potential (V) |
| \(E_a\) | Reaction activation energy (J/mol) |
| \(E_{aD}\) | Diffusion activation energy (J/mol) |
| \(E_r\) | Energy barrier against conductivity (J/mol) |
| \(E_{aR}\) | Energy barrier hindering the ions to move (J/mol) |
| \(F\) | Faraday’s constant (96,487 C/mol) |
| \(i\) | Current (A) |
| \(i_{0}\) | Exchange current density (A/m\(^2\)) |
| \(i_{loc}\) | Local current density (A/m\(^2\)) |
| \(S_{DD}\) | DOD dependent stress factor |
| \(S_{SOC}\) | SOC dependent stress factor |
| \(S_t(t)\) | Time dependent stress factor |
| \(S_{T(T)}\) | Temperature dependent stress factor |
| \(t\) | Time (s) |
| \(T\) | Absolute temperature (K) |
| \(T_g\) | Glass transition temperature (K) |
| \(T_{ref}\) | Reference temperature (K) |
| \(x\) | Coordinate |
| \(t^+\) | Transferring number of Li\(^+\) |

**Greek Letters**

| Symbol | Description |
|--------|-------------|
| \(\alpha\) | Transfer coefficient |
| \(\beta\) | Li poor phase |
| \(\gamma\) | Li rich phase |
| \(\Delta G\) | The change in the cell’s standard free energy (J/mol) |
| \(\Delta H\) | Enthalpy change (J/mol) |
| \(\Delta H_0\) | Enthalpy change of activation for the Li-\(\text{Li}^+\) reaction (J/mol) |
| \(\Delta S\) | Entropy change (J/mol) |
| \(\eta\) | Overpotential (V) |
| \(\eta_{0}\) | Concentration overpotential (V) |
| \(\eta_{ct}\) | Charge transfer overpotential (V) |
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