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Gel polymer electrolytes for rechargeable batteries toward wide-temperature applications

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Rechargeable batteries, typically represented by lithium-ion batteries, have taken a huge leap in energy density over the last two decades. However, they still face material/chemical challenges in ensuring safety and long service life at temperatures beyond the optimum range, primarily due to the chemical/electrochemical instabilities of conventional liquid electrolytes against aggressive electrode reactions and temperature variation. In this regard, a gel polymer electrolyte (GPE) with its liquid components immobilized and stabilized by a solid matrix, capable of retaining almost all the advantageous natures of the liquid electrolytes and circumventing the interfacial issues that exist in the all-solid-state electrolytes, is of great significance to realize rechargeable batteries with extended working temperature range. We begin this review with the main challenges faced in the development of GPEs, based on extensive literature research and our practical experience. Then, a significant section is dedicated to the requirements and design principles of GPEs for wide-temperature applications, with special attention paid to the feasibility, cost, and environmental impact. Next, the research progress of GPEs is thoroughly reviewed according to the strategies applied. In the end, we outline some prospects of GPEs related to innovations in material sciences, advanced characterizations, artificial intelligence, and environmental impact analysis, hoping to spark new research activities that ultimately bring us a step closer to realizing wide-temperature rechargeable batteries.

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1. Introduction

Global energy consumption has risen greatly with the development of human society. To replace traditional fossil fuels, developing clean and renewable energy sources such as solar, wind and tide has become an urgent demand to an unprecedented level. However, these energy sources are intermittent



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and random, requiring reliable energy storage and conversion devices to balance the electricity supply and flexible deployment of the electrical grid system and to fill the gap between the installed capacities and the actual amount of electricity.^{1–3}

Among the various types of energy storage and conversion devices, rechargeable (or secondary) batteries are among the most successful proven technologies that can repeatedly convert chemical energy into clean electricity and have achieved a giant leap in energy density over the past two decades. Meanwhile, diversifying rechargeable batteries with different design configurations (*e.g.*, coin/cylindrical/prismatic shape, blade/magazine stack) and functionalities were developed, and novel battery chemistries were proposed to meet the requirements of extensive application scenarios including portable electronic consumer devices, electric vehicles (EVs) and large-scale electricity storage in smart or intelligent grids.^{4–7}



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Though energy/power density, safety, and stability of rechargeable batteries at room temperature have been greatly improved, the demand for batteries that can survive and operate under extreme temperatures is soaring.

In this century, severe summers and winters occurred frequently over the world. As displayed in Fig. 1(a), the record temperatures of capital cities around the world have been frequently broken: the high temperatures in some countries were over 40 °C, *e.g.*, India (45.6 °C in 2020), Egypt (45 °C in 2019), and Australia (46 °C in 2020), and low temperatures approximately approach –30 °C, *e.g.*, Russia (–33 °C in 2012), Canada (–36 °C in 2013), and Ukraine (–30 °C in 2012). It is worth noting that these are only the data for capital cities rather than the whole countries, taking the United States as an example, the extreme temperatures can hit down to –62.2 °C and up to 56.7 °C.⁸ Given the diversity and variability of climate on a global scale, EVs are expected to operate stably and safely under a wide temperature range. Extensive research has demonstrated that the driving range of EVs heavily depends on the environmental temperatures (Fig. 1(b)). According to the test results on thousands of EVs performed by Seattle-based Recurrent, some vehicles saw a 30% drop in their driving ranges at near subzero temperatures.^{9,10} Moreover, some special application scenarios, such as military, high-flying drones, and space exploration, urgently need batteries that can be operated under more challenging temperature conditions, for example, –40 to 70 °C for military applications.^{11–13}

Until now, however, the state-of-the-art rechargeable batteries face formidable challenges of serious electrochemical performance degradation and safety issues at both low and high temperatures due to the chemical/electrochemical limitations of conventional electrolytes, thus severely limiting their applications at harsh temperature conditions:^{14–16} (i) at low temperatures, the increased viscosity or even solidification of electrolytes causes sluggish ion-transfer kinetics both in bulk electrolytes and electrolyte/electrode interfaces as well as the



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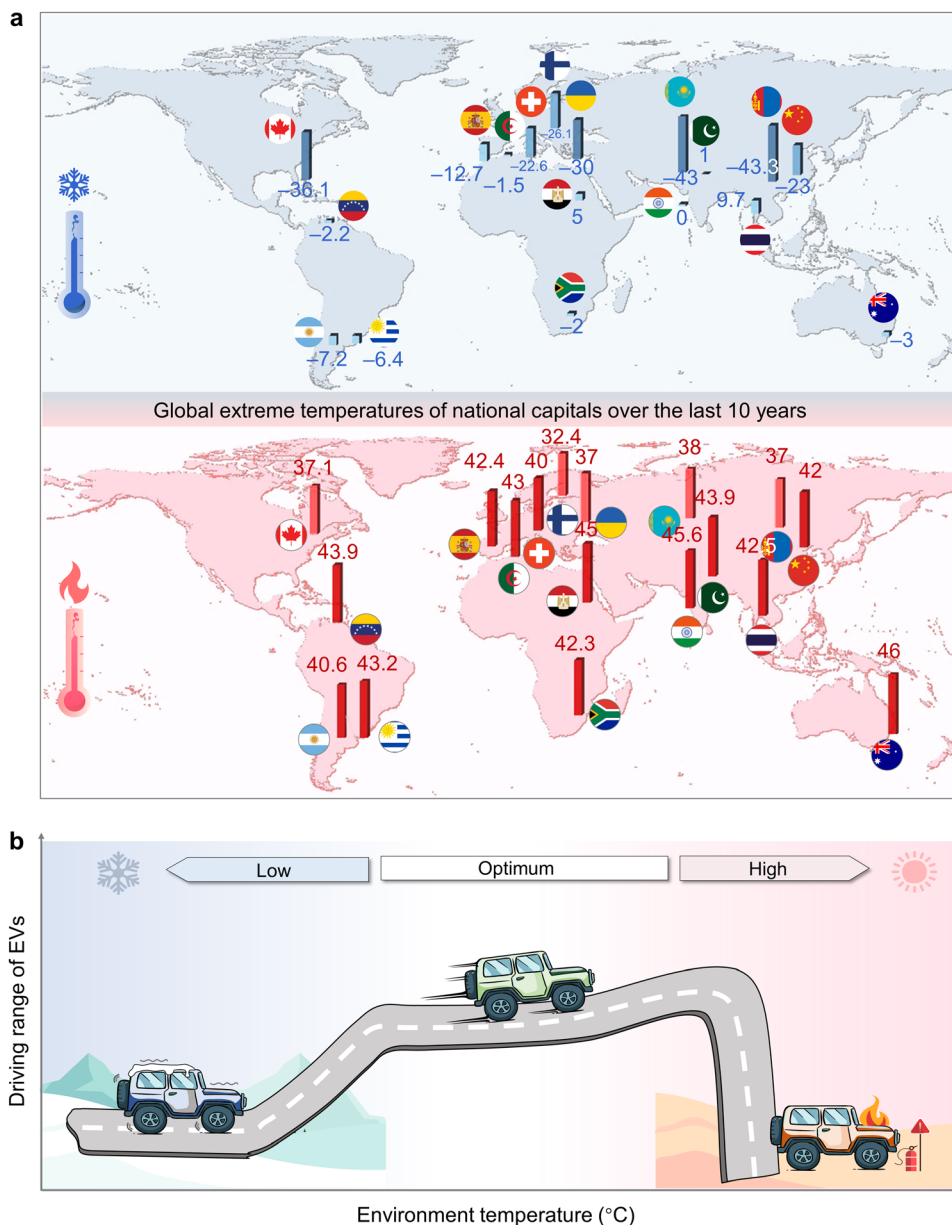


Fig. 1 Schematic showing the driving range anxiety and safety issues of EVs raised by global extreme temperatures. (a) Global extreme low (blue) and high (red) temperatures of capital cities of different countries in the last 10 years (from Jan. 2012 to July 2023), data collected from the Climate Change Institute, USA. (b) Change in driving range of EVs with the environmental temperature.

slow ion solid-state diffusion within the active electrode materials, lowering the practical energy densities; (ii) at high temperatures, the thermodynamic instability and increased reactivity of electrolytes toward the electrodes would lead to serious safety hazards of rechargeable batteries. Considering these demands, rechargeable batteries capable of operating at wide temperature ranges have attracted tremendous research attention in recent years.

To date, most research on wide-temperature rechargeable batteries has been largely devoted to the discovery of novel electrolytes due to their crucial bridge role in electrochemical battery systems (Fig. 2(a)). However, it remains challenging to explore a kind of electrolyte that can operate well in extremely

cold and hot environments. Standing out among various kinds of electrolyte systems, gel polymer electrolyte (GPE) combines the high ionic conductivity and excellent interfacial compatibility of liquid electrolytes as well as the high safety of all-solid-state electrolytes, promising to realize rechargeable batteries with wide-temperature operability (Fig. 2(b)) and has become a research hot spot in both academic and industrial world.^{17–19} Although great progress on GPEs has been made, the investigation on GPEs and batteries operating at extreme temperatures is still in their early stage.^{20–22} To date, reviews on GPEs have been mainly focused on the synthetic strategies, properties, and their battery performance at room temperature, with limited coverage of extreme temperature conditions.



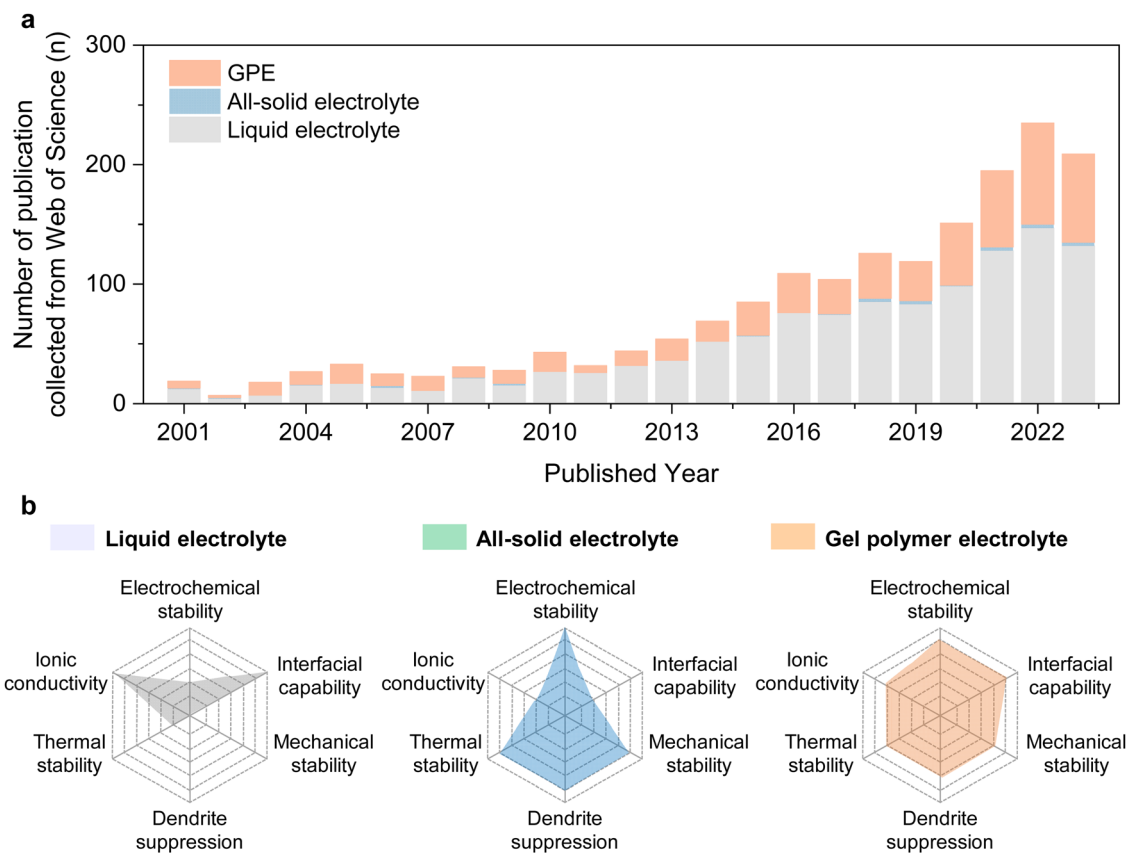


Fig. 2 Overview of the research trend of wide-temperature rechargeable batteries based on various electrolyte systems. (a) Number of publications on “wide-temperature/all-climate/high-temperature/elevated-temperature/low-temperature/extreme-temperature/extreme-condition/all-weather/extreme-weather/environment-adaptable/all-temperature”, up to December 2023. Database: Web of Science. (b) Comparison of wide-temperature performance between liquid, all-solid electrolytes and GPE.

To the best of our knowledge, there has been no comprehensive review discussing the design and engineering strategies of GPEs toward wide-temperature rechargeable batteries. Also, the structure–property relationships of GPEs and their effects on the wide-temperature performance of rechargeable batteries have not been comprehensively and thoroughly summarized. Given these facts, it is crucial and timely to review on the progress and advances and to discuss the design principles, engineering strategies, challenges, opportunities, and prospects of wide-temperature adaptable GPEs.

In this review, we first overview the major challenges encountered in the development of wide-temperature adaptable GPEs. Next, the key requirements and design principles of GPEs for wide-temperature applications are discussed. Then, we thoroughly discuss the engineering strategies of GPEs in terms of salts, polymer matrices, solvents, additives, and hosts to enable their wide-temperature operability. Finally, we offer our perspectives on the future research directions and opportunities of wide-temperature adaptable GPEs through innovations in material sciences, advanced characterizations, artificial intelligence, and environmental impact analysis for addressing the global energy challenges using gel polymer-based materials.

2. Main challenges of wide-temperature rechargeable batteries

The electrolyte, which physically interacts with all other components, stands out as the most distinctive component in a rechargeable battery.^{23,24} Temperature significantly affects the electrolyte and its interfaces with other components whether it is an organic nonaqueous or aqueous system. At both high and low temperatures, the electrolyte experiences substantial degradation, leading to increased interfacial polarization and a decline in the performance of rechargeable batteries. Given the wide range of physical and electrochemical events that occur in rechargeable batteries, particularly under extreme temperatures, there are numerous complex challenges to overcome in developing suitable electrolytes.

2.1 Challenges at low temperature

During the charging process of rechargeable batteries, the cations experience several steps: (i) solvation in bulk electrolyte, (ii) migration of solvated cations across the electrolyte to the surface of active material, (iii) desolvation of solvated cations at the electrolyte/electrode interphase, (iv) cations passing through the electrolyte/electrode interphase layer and intercalation into/plating onto the anode. At low temperatures,



however, the electrolyte freezing and the intrinsic kinetic constraints of charge carriers/mass-transfer can induce the reduction of ionic conductivity of GPE, and high desolvation barrier, SEI thickening and dendrite growth in the GPE/electrode interfaces, significantly deteriorating the discharge capacity and causing excessive overpotentials (Fig. 3(a)).

The conductivity of an electrolyte is a measure of its ability to transport ions. Researchers often use two empirical equations, namely, the Arrhenius equation (eqn (1)) and Vogel–Tammann–Fulcher (VTF) equation (eqn (2)), to describe the temperature dependence of ionic conductivity in the electrolyte (Fig. 3(b)).^{25,26}

$$\sigma(T) = \sigma_0 \exp\left(\frac{E_{a1}}{k_B T}\right) \quad (1)$$

where σ_0 , E_{a1} , and k_B are the pre-exponential factor, activation energy, and Boltzmann constant, respectively.

$$\sigma(T) = AT^{-1/2} \exp\left[\frac{E_{a2}}{k_B(T - T_0)}\right] \quad (2)$$

where A is the pre-exponential factor related to the number of charge ions, k_B is the Boltzmann constant, E_{a2} is the pseudo-activation energy associated with the polymer segmental motion and T_0 ($T_0 = T_g - 50$ K) is the temperature corresponding to zero configurational entropy. The linear and curvilinear behavior of Arrhenius and VTF plots can be respectively ascribed to the coupling and decoupling mechanisms between the ionic motion and long-range motion of the polymer branches and/or solvent molecules. It is widely accepted that the electrolyte should have a

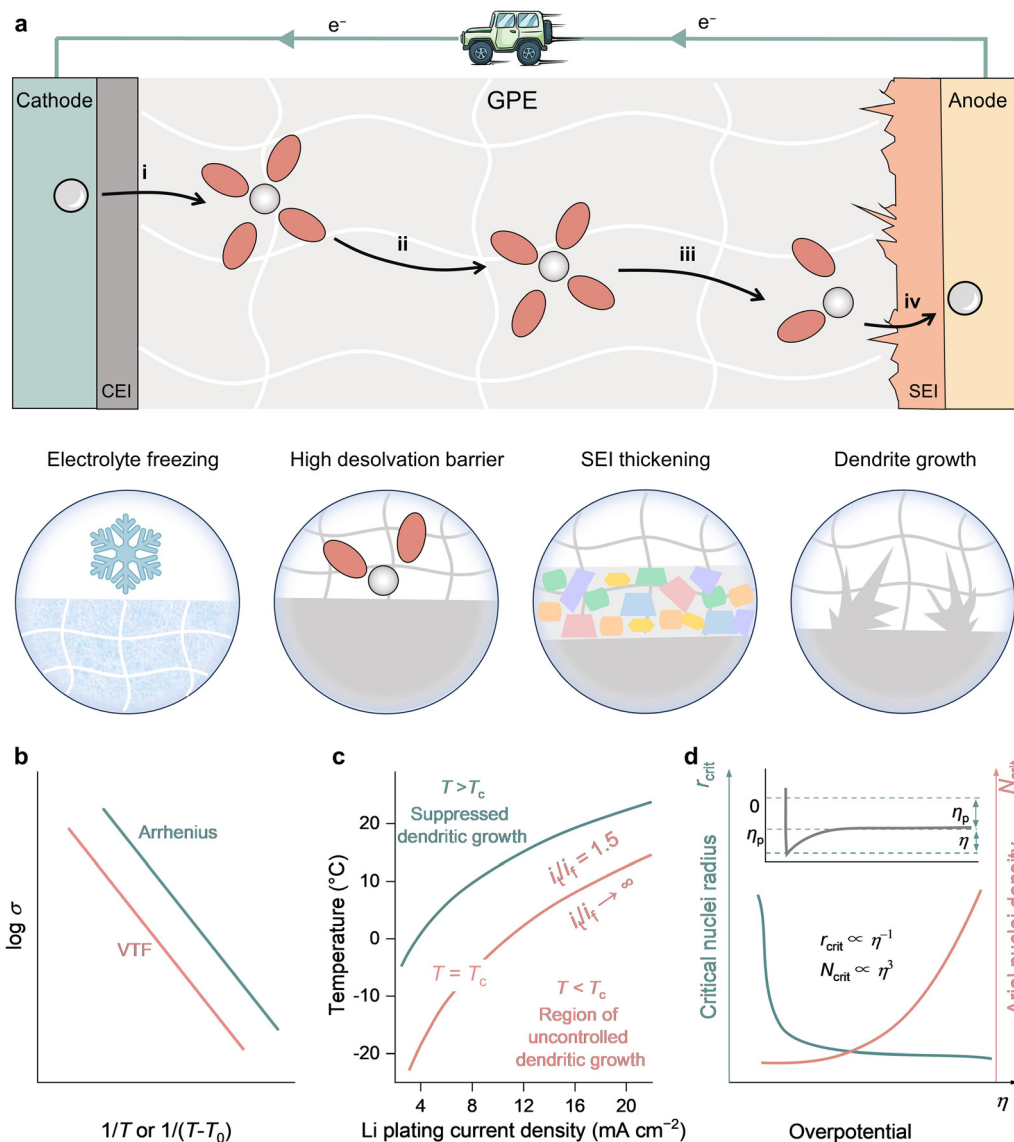


Fig. 3 Main challenges for rechargeable batteries at low temperatures. (a) Schematic representation of the main challenges for rechargeable batteries at low temperatures. The arrows represent the main ion transport processes during the charging process. (b) Schematic illustration of the temperature-dependent ionic conductivity based on the Arrhenius and VTF equations. (c) Variation of the dendrite growth rate ratio i_p/i_t with temperature and Li plating current density, data extracted from ref. 45 with permission from Elsevier, Copyright 2013. (d) Schematic plot of the dependence of the critical Li nuclei radius and areal nuclei density on the Li deposition overpotential, data extracted from ref. 51 with permission from American Chemical Society, Copyright 2017.



low activation energy (E_a) to maintain conductivity and ensure good electrochemical performance at low temperatures.

The ionic conductivity of conventional carbonate-based liquid electrolytes can be determined by eqn (3)

$$\sigma = \sum_i n_i \mu_i Z_i e \quad (3)$$

where n_i is the free-ion number, e is a unit charge, Z_i is the charge valence, and μ_i is the ion mobility of different ions, which could be calculated from eqn (4).

$$\mu_i = \frac{1}{6\pi\eta r_i} \quad (4)$$

where r_i represents the solvation radius and η is the viscosity. At low temperatures, the spiking viscosity (η) of the electrolyte increase while the solvated solvent forms a close ion pair with its counterions (causing decreased r_i),²⁷ leading to the retarded ionic mobility (μ_i) and the reduced ionic conductivity (σ). According to eqn (4), the lower viscosity of the electrolyte, the more readily the solvated ions will move in solvents. However, solvents with low viscosity usually have low dielectric constants, which hinders the dissociation of salts and the improvement of ionic conductivity. Therefore, it is necessary to combine the design of low viscosity electrolyte and the regulation of ion solvation structure to achieve optimal conductivity at low temperatures. Additionally, increased viscosity or even solidification/freezing at low temperatures reduces the wetting ability of the electrolyte-electrode interface, leading to interface separation and substantial increase in the interfacial electrochemical impedance.

Early research suggested that the ion diffusion within the anode material (such as graphite) was the main obstacle to battery performance.^{28,29} However, recent studies have recognized that the process of ion migration at the electrolyte/electrode interface, involving the desolvation of cations from the solvent sheath and subsequent ion diffusion across the solid interfacial interphase into the active electrode material, is the primary factor affecting battery performance, especially at low temperatures.^{30,31} The ion-transfer kinetics at low temperatures are reflected by charge transfer resistance (R_{ct} , the mid-frequency semicircle in EIS spectra). A high R_{ct} value indicates a slow charge transfer kinetics. Zhang *et al.* found that at temperatures below -20°C , the value of R_{ct} became dominant, almost equaling the total resistance (R_{cell}), whereas at 20°C , it only accounted for approximately 50% of the R_{cell} .³² According to the Arrhenius equation and the Butler-Volmer equation, R_{ct} is related to the activation energy (E_{a3}) of the charge transfer process.^{20,33}

$$\frac{1}{R_{ct}} = A \exp\left(-\frac{E_{a3}}{RT}\right) \quad (5)$$

where R is the gas constant, and A is a pre-exponential factor. It can be concluded from eqn (5) that the charge transfer impedance is

determined by the temperature and the E_{a3} of the charge transfer process. A lower E_{a3} results in faster charge-transfer kinetics. The E_{a3} was found to be dependent on the desolvation process of cations and the absorbing energy of cations on the surface of electrodes. Li *et al.* presented that desolvation was the main barrier for Li^+ migration in lithium-ion batteries under low temperatures rather than the diffusion in electrodes or across the SEI layer.³⁰ Therefore, optimizing the electrolytes to reduce the desolvation energy barrier at the electrode-electrolyte interface is crucial for enhancing the performance of rechargeable batteries in low-temperature conditions.

Besides the desolvation process, the ion migration across the SEI layer is also a key challenge at low temperatures, which depends on the nature and composition of the SEI layer. Thermodynamically, lowering the temperature not only slows down ion transport but also alters the decomposition reaction paths of electrolyte components. Wang *et al.* proposed that the SEI layer formed at low temperatures was mainly composed of intermediate products-rich organic species due to the incomplete decomposition of salts and solvents, which were metastable and easily dissolve in liquid electrolytes upon cell cycling, inhibiting the ion transport and inducing undesired side reactions.^{34,35} Furthermore, the lithium plating phenomenon of lithium-ion batteries was reported to occur at the anodic side at low temperature; it subsequently reacted with the solvent and lithium salt in electrolytes and the reaction products on the anodic side, increasing the thickness of the SEI layer. The increased SEI thickness lengthens the diffusion distance, causing a large interfacial impedance and polarization.^{36,37} It has been reported that an inorganic-rich SEI layer, particularly one containing fluorine, with a low thickness can lower the energy barrier for desolvation and facilitate rapid ion diffusion through the SEI layer.³⁸⁻⁴⁰ Given this understanding, it is crucial to develop suitable electrolyte compositions and SEI structures to enhance the performance of rechargeable batteries at low temperatures.

Kinetically, the decreased temperature reduces the mass transport limitation at the electrolyte/electrode interface and increases the reaction kinetic energy barrier of metal-ion deposition, leading to the slow and uneven plating of metallic species. This causes the formation and accumulation of dendrites near the anode.⁴¹⁻⁴³ Love *et al.* found that the temperature directly affected the dendrite morphology and the dendrite formation rate (v_d) and thus tuned the short-circuit time (t_{sc}),⁴⁴ as shown in eqn (6).

$$t_{sc} = t_f f(T) + \frac{1}{v_d f(i, T, \text{morphology})} + \text{morphology} f(T) \quad (6)$$

where l is the distance between electrodes, and i is the applied current. The analytical model proposed by Akolkar *et al.* demonstrated the critical temperature (T_c) of the dendrite formation rate (i_d/i_f) on the anodic side,^{45,46} as estimated by eqn (7).

$$\frac{i_t}{i_f} = \left\{ -\frac{1}{bC_0} \ln \left[\exp(bC_0) + \frac{i_f(1-t_+)b\delta}{nF\alpha_0 \exp\left\{A_1\left(\frac{1}{T_0} - \frac{1}{T}\right)\right\}} \right] \right\}^{-\left(\alpha_c^0/n\right) \exp\left\{A_2\left(\frac{1}{T_0} - \frac{1}{T}\right)\right\}} \quad (7)$$



where C_0 is the bulk concentration, α_c is the charge transfer coefficient, i_f represents the deposition current density on the flat electrode surface, α_0 is the temperature-independent pre-exponent, and T_0 is a reference temperature. It can be concluded that lithium battery systems can only operate at low current density (charge rate) at a low temperature to prevent uncontrolled dendrite formation (Fig. 3(c)). Temperature also affects the deposition process, including the formation of dendrites and the final surface morphology of deposits (e.g., moss-like, particulate, granular, or needle-like).^{47–50} In a typical galvanostatic electrodeposition process, there are two important characteristic overpotentials—the nucleation overpotential (η) and the mass-transfer controlled plateau overpotential (η_p) (Fig. 3(d)). According to the classical theory for homogeneous nucleation in the electrodeposition process, the ratio of nucleus (r_{crit}) is inversely proportional to the nucleation overpotential η while the areal nuclei density (N_{crit}) is proportional to η^3 eqn (8) and (9).^{51–53}

$$r_{\text{crit}} = \frac{2\gamma V_m}{F|\eta|} \quad (8)$$

$$N_{\text{crit}} = \frac{2z^3 F^3 \eta^3}{16\pi N_A V_m^2 \gamma_{\text{SL}}^3} \quad (9)$$

where γ , V_m , z , F , and N_A are the surface free energy between the deposits and the electrolyte, molar volume of the anode, number of electrons per monomer unit, Faraday constant, and Avogadro constant, respectively. Both the nucleation and plating overpotential increased with decreasing temperature, thus inducing smaller and dense nuclei and promoting the growth of high-aspect-ratio needle-like dendrites at high current densities.^{54,55}

2.2 Challenges at high temperature

Batteries often perform better at high temperatures due to increased ionic conductivity and speedier chemical reactions, this is only somewhat reliable within a -20 to 55 °C temperature range, based on numerous past studies. However, the performance of most documented rechargeable batteries at temperatures exceeding 55 °C remains questionable. This is mainly attributed to the electrolyte evaporation and decomposition, pronounced solubility and reconstruction of SEI, and uncontrollable dendrite growth. These factors trigger dynamic and intricate structural alterations during the mass and charge transfer process within the batteries, eventually leading to battery performance degradation and even causing safety hazards (Fig. 4(a)).⁵⁶ Physically, elevated temperatures would increase the vapor pressure of the liquid components in the electrolytes, raising the risk of evaporation or leakage, electrochemical reactions instability, and even safety hazards. Thermodynamically, the instability and decomposition of electrolytes at high temperatures can be attributed to the fact that these batteries operate at voltages beyond the thermodynamic stability range of the electrolytes, which is determined by the highest occupied molecular orbitals (HOMO) and lowest unoccupied molecular orbitals (LUMO). The chemical potential of cathode materials (μ_c) and anode materials (μ_a) was reported to shift with the increase in temperature. This process can be evaluated

according to the Nernst equation⁵⁷

$$V = V^0 \frac{RT}{nF} \ln \alpha \quad (10)$$

where V , R , and T signify the cell potential, the molar gas constant and the absolute temperature, respectively. n is the number of electrons transferred per mole, V^0 is the standard potential and F is the Faraday constant. As shown in Fig. 4(b), high temperature boosts the electron energy level and contributes to the positive shift of the equilibrium μ_c and negative shift of μ_a .¹⁶ In addition, the high temperature also shifts the HOMO/LUMO and narrow the gap between them. Based on the model proposed by Goodenough *et al.*, when the $\mu_a >$ LUMO, the electrons on the anode will preferentially transfer toward the unoccupied molecular orbital of the electrolyte and persuade the reduction reactions of electrolyte and the formation of SEI layer on the electrolyte/anode interface. Likewise, as the $\mu_c <$ HOMO, the redox reactions of electrolyte occur and facilitate the formation of the CEI layer on the cathode/electrolyte interface⁵⁸ (Fig. 4(c)). Kinetically, high temperature favors the chemical reaction rate because of the improved molecular thermal motion. According to Arrhenius equation, a 10 °C increase will double the reaction rate.⁵⁹ Therefore, higher temperature accelerates interfacial chemical reactions by enhancing interfacial mass and ionic transfer. As displayed in Fig. 4(d), the thickness of the anode turns thick with the temperature increasing from the low zone (<0 °C) to the high zone (>50 °C), and the thickness of the SEI on the single anode particles adds up to a macroscopic value.⁶⁰ In addition, the trace impurities contained in the electrolyte or dissolved matters such as organics, polysulfides, and transition metal ions from electrodes during electrochemical processes may participate in the interfacial chemical reactions at high temperature, exacerbating the destruction and reconstruction of the SEI/CEI while consuming the metal-ion salts to continuously form unwanted byproducts in the electrolyte/electrode interfaces.^{61–64} Furthermore, differing from metastable SEI/CEI layers in the electrolyte/electrode interfaces at lower temperatures, the electrochemical decomposition reactions of the electrolyte will be accelerated and completed to form the higher-modulus SEI/CEI composed of inorganic species at elevated temperatures (Fig. 4(e)).⁶⁵ The inorganic-rich SEI/CEI layers are considered to be relatively brittle, making it vulnerable to stress-induced cracking, which adversely deteriorates the battery performance. All of the above contributes to the thickening and breaking of the SEI/CEI layers, obstructing the migration of charge carriers, increasing the interfacial resistances, and leading to an excessive interfacial polarization and thus a dramatic drop in battery capacity.

In addition to unfavorable interfacial reactions, dendrite growth is also a main obstacle at high temperatures, especially in rechargeable metal batteries. The uncontrollable growth of these metallic protrusions can cause internal short-circuits. Theoretically, high temperatures have an inhibitory effect on the growth of dendrite.⁶⁶ As displayed in eqn (8), (9) and Fig. 3(d), a high temperature can facilitate ion diffusion, leading to the decrease of η and contributing to the metal deposits



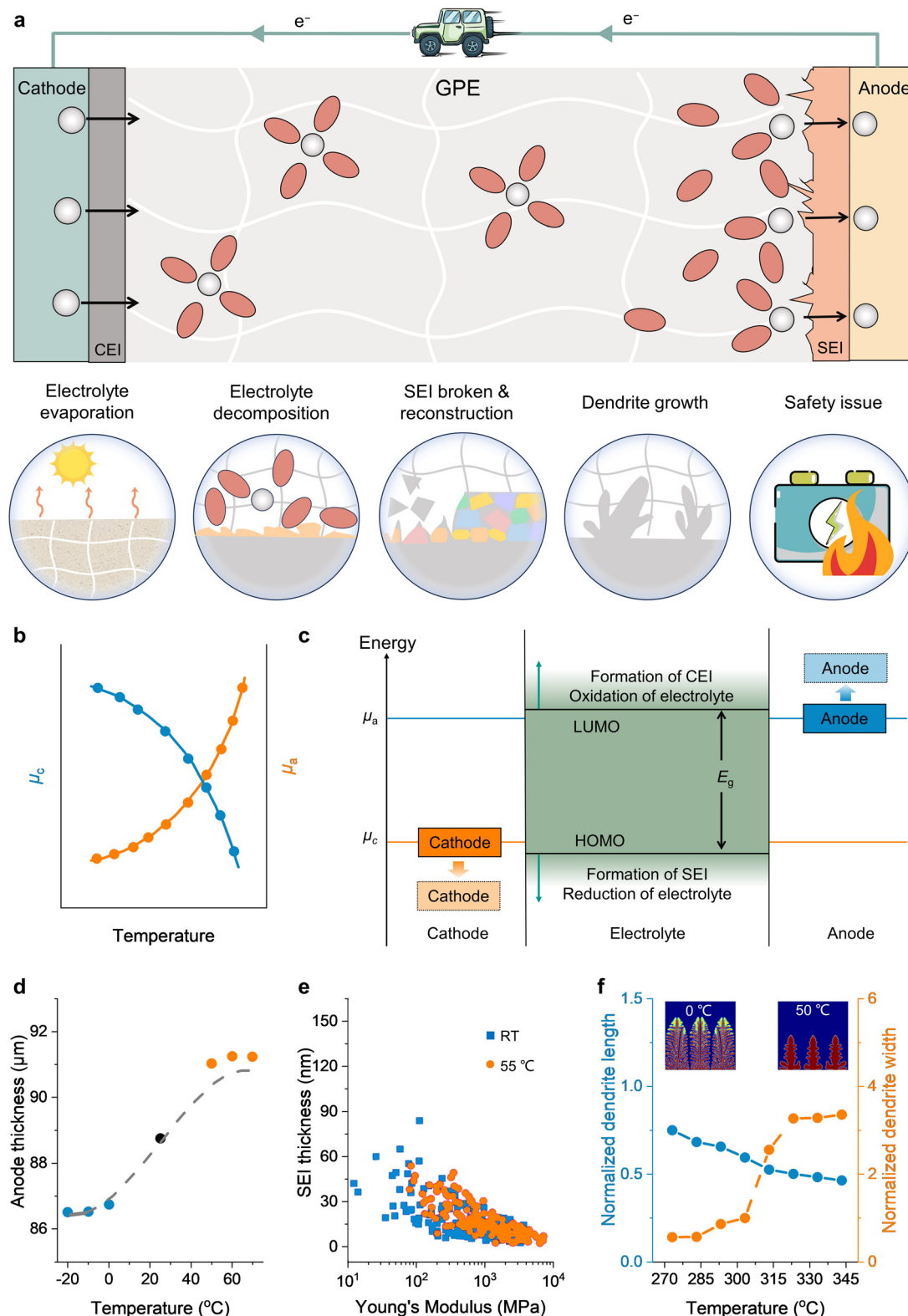


Fig. 4 Main challenges of rechargeable batteries at high temperatures. (a) Schematic illustrations of the main challenges of rechargeable batteries at high temperatures. (b) Change of chemical potential of cathode materials (μ_c) and anode materials (μ_a) with the increase in temperature, data extracted from ref. 16 with permission from RSC publishing, Copyright 2022. (c) Schematic illustration of the SEI/CEI energy level shift to reach a new steady state as the chemical potential of the cathode (anode) shifts with temperature. (d) Change of the thickness of the anode particles with temperature, data deduced from ref. 60 with permission from Elsevier, Copyright 2014. (e) Relationship between SEI thickness and Young's modulus at different temperatures, data extracted from ref. 65 with permission from RSC publishing, Copyright 2014. (f) Change in the dendrite length and width with the temperature, data extracted from ref. 67 with permission from Elsevier, Copyright 2018. Insets: The morphology of Li dendrites at the temperature of 0 and 50 $^{\circ}\text{C}$.



with larger nuclei size and low nucleation density. The integrated temperature-dependent phase-field models of Li-dendrite presented by Chen *et al.* and the Zn-dendrite model presented by Wang *et al.* both demonstrate that high temperature leads to a shorter dendrite length and larger width than those at low temperatures (Fig. 4(f)), thus weakening the dendrite growth.^{67,68}

However, in fact the experimental results of dendrite formation at high temperatures have been widely reported.^{69,70} In these cases, it is commonly assumed that the incompatible kinetics and uneven charge distribution at the electrolyte/electrode interfaces are responsible for dendritic growth at elevated temperatures. On the one hand, the ionic conductivity of the SEI is several orders of magnitude lower than that of the bulk electrolyte, causing faster ion diffusion in the electrolyte than in the SEI and leading to the formation of ionic depletion layer and strong concentration gradient, which aggravates dendrite growth. On the other hand, the unevenly distributed component and morphology of SEI accumulated by the degradation products induces an uneven impedance and current density distribution, promoting dendrite formation.^{70,71} Additionally, the irregular SEI breakdown (solubility or mechanical instability, voids and/or cracks formation, *etc.*) at the electrode vicinity due to the large volume change of the anode and the porous SEI augmentation by the electrolyte decomposition during repetitive plating/stripping at high temperature results in variations of ion concentration and mobility across the electrolyte/electrode interfaces, further exacerbating the dendrite formation.^{57,72}

3. Key requirements of wide-temperature rechargeable batteries

As discussed above, both high and low environmental temperatures would have an impact on the ion-transport kinetics and stabilities of electrolytes and electrolyte/electrode interfaces in rechargeable batteries. Rational electrolyte design is required for both aqueous and non-aqueous battery systems to enhance the electrochemical performance and avoid severe degradation and safety risks at extreme temperature conditions. In order to meet the requirements of rechargeable batteries to withstand wide-temperature applications, advanced electrolytes are required to overcome the above-mentioned drawbacks and achieve performance including rapid ionic charges and mass transport, excellent chemical/electrochemical stability against electrodes, and satisfactory thermal/mechanical tolerance. The detailed requirements are shown in Fig. 5 and comprehensively discussed in the following sections.

3.1 High ionic conductivity

Ionic conductivity is considered the most important parameter in all electrolyte research works as it primarily determines whether the electrolyte could be operated to deliver the theoretical energy density of the designed battery system. It is timely to establish a temperature-dependent threshold value of ionic conductivity, applicable to various electrolyte systems, to guide

the electrolyte optimization towards wide temperature operability.^{73–76} Due to that the ionic conductivity is not the limiting factor for high temperature battery operation, the ionic conductivity requirements in this section refer specifically to conditions of low temperature. In contrast to the relatively short research history of GPEs toward wide temperature applications, research progress on the more developed liquid electrolytes may provide some reference for evaluating the ionic conductivity requirements, it is demonstrated that the rationally designed electrolyte with an ionic conductivity of $\sim 2.7 \text{ mS cm}^{-1}$ at -20°C could allow a NCM811//Li battery with a high areal capacity of 3.5 mA h cm^{-2} to operate efficiently and stably at -20°C and even below.⁷⁷ Another groundbreaking example is the recent demonstration of a remarkable energy density of 110 W h kg^{-1} at extremely low temperature of -40°C for Ah-level sodium-ion batteries using a low-temperature specific liquid electrolyte.⁷⁸ Summarizing these achievements, it is found that at the very low environmental temperature of -20°C (approximately the lowest temperature that most regions of the earth could reach), an ionic conductive value of $3\text{--}10 \text{ mS cm}^{-1}$ is required for the efficient operation of these practical-level batteries. Considering that the ionic conductivities of liquid electrolytes would retain only one tenth to one sixth of their initial values when being impregnated in the commercial polyolefin separator, 1 mS cm^{-1} at -20°C could be preliminarily considered as the threshold requirement to guide the exploration of GPEs applicable at low temperatures.^{79–81} Furthermore, a low activation energy (*i.e.*, E_a) is desired for easy ion conduction and to minimize the impact of temperature on ionic conductivity.^{82,83} This can be achieved by improving cation concentration, enhancing cation migration rate and providing excess cation conduction paths. Since GPEs possesses cohesive properties of solids and diffusive properties of liquids simultaneously, proper tuning of GPE components, including polymer matrix and enclosed liquids, is necessary to achieve high ionic conductivity and low activation energy over a wide temperature range.^{84,85}

3.2 High target-ion transference number

Although necessary, high ionic conductivity alone does not guarantee excellent performance of wide-temperature rechargeable batteries, given the fact that ion conductivity is the result of the collective movement of all ionic species, whereas the faradaic current is predominantly balanced by the cations rather than their counterions.⁸⁶ In practical electrolyte systems, cations and anions move in opposite directions. However, interactions (*e.g.*, coordination bonds and hydrogen bonds) between cations and polar molecules such as solvents cause anions to move faster than cations, leading to the anion accumulation and the buildup of a concentration overpotential near the electrode, which can seriously lower the energy efficiency, particularly at extreme temperatures.^{87,88} In this case, the importance of target-ion transference number (t^+) is highly emphasized since it reflects the ionic conductivity contribution of specific ions ($t^+ = N_{v,s}D_s / \sum N_{v,s}D_s$, where $N_{v,s}$ and D_s are the number density and diffusion coefficient of individual ions,



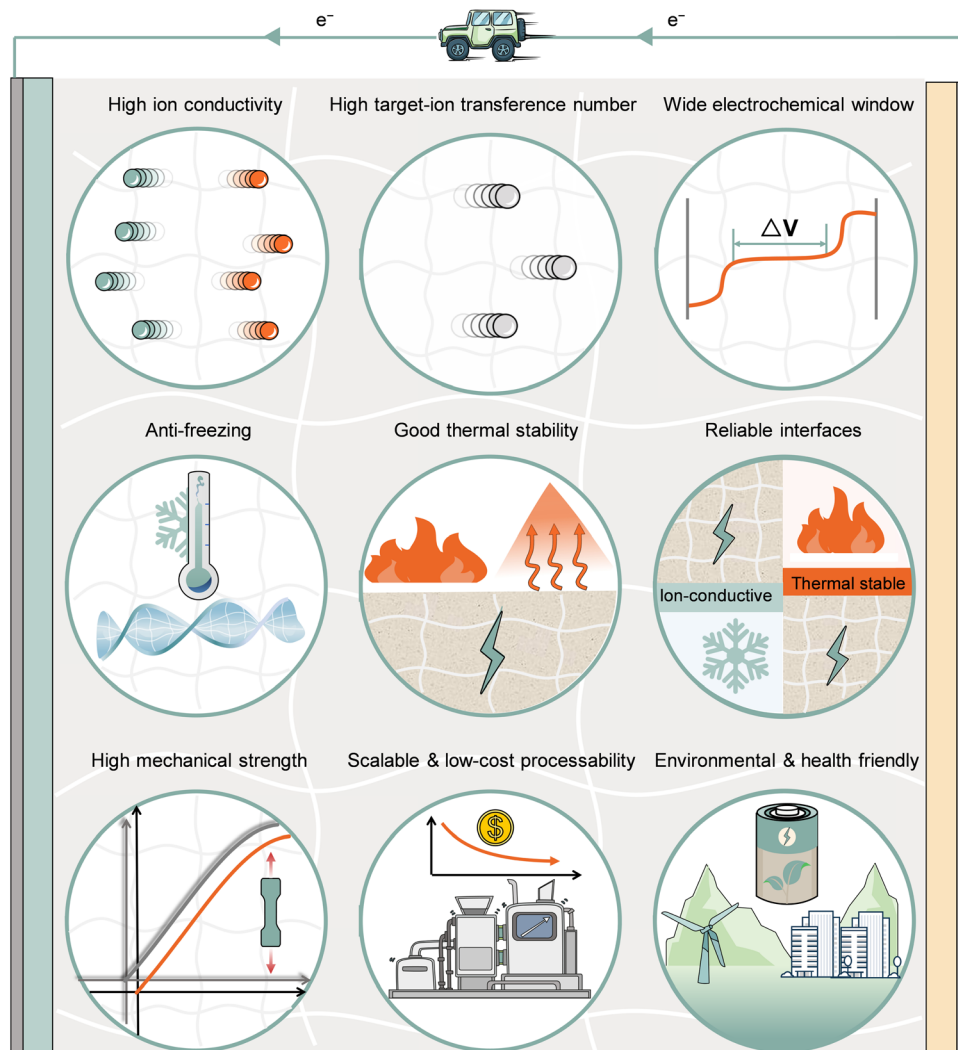


Fig. 5 Key requirements of wide-temperature rechargeable batteries. The temperature-dependent electrochemical/physicochemical/mechanical properties, cost, and environmental impact should be comprehensively considered.

respectively), a high t^+ would represent a relative higher target-ion mobility than other ionic species in electrolyte.^{89,90} According to the classical Newman model, an electrolyte with a lithium-ion transference number (t_{Li^+}) close to unity can warrant a considerably higher discharge capacity compared to the traditional liquid electrolytes with t_{Li^+} of approximately 0.2, primarily attributed to the negligible concentration polarization.^{91,92} Following the theoretical Chazalviel model and experimental measurements, it was found that the increase in dendrite growth is favored in electrolytes with low t^+ due to the cation depletion and formation of space charge layer.^{93,94} Conversely, improved t^+ promotes the cation distribution homogeneity and inhibits dendrite growth.^{95,96} It is suggested that modest improvements in t^+ , to 0.5 to 0.7, would be beneficial, particularly allowing more uniform electric field distribution on the electrode surfaces, which reduces the risk of hotspots and thus extends the life of the battery.⁸⁸ This suggests that optimizing the transference number along with the ionic conductivity could be a key strategy in the development of advanced electrolytes, especially for low temperature applications

where the total ionic conductivities of electrolytes have been significantly lowered.

3.3 Wide electrochemical stability window

For achieving stable and reversible redox reactions in batteries, a desirable electrolyte must have a sufficient electrochemical stability window against electroreduction and electrooxidation over a wide range of operating temperatures and working voltages without causing detrimental side reactions.^{97–99} The electrochemical stability window is typically defined as the gap between the HOMO and LUMO of the electrolyte. However, recent research suggests that the widening of electrochemical stability window requires the consideration of chemical/electrochemical interfacial compatibility beyond just the HOMO and LUMO of electrolyte components.^{100,101} The traditional methods of predicting electrochemical stability windows, such as HOMO/LUMO, overlook the electron transfer between reactants (neutral molecules) and products (charged molecules), as well as the structural evolution in the redox reaction, can lead



to large prediction errors of up to ~ 3.25 V.¹⁰⁰ It can be expected that the extreme temperatures can further increase the error due to the temperature-dependent nature of the electron transfer behaviors. Additionally, temperature can affect the thermodynamic reactivity of electrolyte components and the kinetic behavior of the SEI/CEI layer, which determines the electrochemical stability window. Higher temperatures can push the potential of the cathode or reduce the potential of the anode into the unstable zone of the electrolytes. Constructing electrochemically stable SEI/CEI layers can effectively broaden the electrochemical stability window, which can prevent direct contact between the electrodes and electrolytes consistently even when the electrolyte operates at extremely high temperatures.^{102–104} Therefore, choosing thermodynamically stable components for the electrolyte and constructing reliable SEI/CEI layers are two major pathways to widen the electrochemical stability window of the electrolyte over a wide temperature range.

Regulating the ionic solvation structure is a practical strategy to customize SEI/CEI and improve the electrochemical stability window of electrolytes, in which the size of ions, the coordination number of polar solvent/water molecules in the solvation sheath structure, and the interaction strength between ion and polar molecules affects the properties of SEI/CEI.^{105,106} For example, modifying the solvation structure using additives or cosolvents (*e.g.*, dimethyl sulfoxide (DMSO), dimethyl carbonate (DMC), and triethyl phosphate (TEP)) that are easier to coordinate with cations to reduce the solvent/water molecules in the primary solvation structure or by forming an anion-derived interphase has been reported to weaken the activity of the solvent and inhibit the active solvent-related decomposition and parasitic reactions and enlarge the electrochemical stability window.^{107–109} In some studied GPEs, the polar functional groups in their polymer chains have been reported to interact with the electroactive metal ions (such as Li⁺ in lithium-ion batteries) and thus participate in the regulation of the solvation structure.^{84,85} Although there is a preliminary understanding of some polymers, such as polyacrylonitrile, and their ability to extend the electrochemical stability window when they are used as a polymer matrix in GPEs, it remains unclear how and to what extent they affect the electrochemical stability windows, there is an urgent need for quantitative studies and internal guidelines.^{14,110} Of note, the influence of properties (*e.g.*, components, morphologies, and structures) and electrochemical reaction thermodynamics (*e.g.*, the formation mechanism and ion transport across SEI/CEI) of SEI/CEI on the electrochemical stability window of the electrolyte and the electrochemical performance of rechargeable batteries remains controversial, calling for deeper exploration in the future.

3.4 High thermal stability

High thermal stability is crucial for wide-temperature operability of GPEs as it encompasses thermodynamic, kinetic, and process stability related to heat generation and material stability upon exposure to extreme temperatures.

Several factors affect the thermostability of electrolytes. Firstly, the inherent electrolyte components affect the thermal behaviors of the overall bulk electrolyte system. The most used electrolytes for rechargeable batteries at large scale are carbonate solvents in non-aqueous electrolyte systems and water in aqueous electrolyte systems; they generally possess low boiling temperatures and are easy to volatilize or burn at elevated temperatures. Meanwhile, most salts with low electrostatic energy suffer from decomposition at elevated temperatures, indicating low thermal stability. Na salts showed better stability compared to the Li counterparts (*e.g.*, LiPF₆ (~ 175 °C) < NaPF₆ (~ 325 °C)).¹¹¹ In addition, the disproportionation and the reactions with charged electrodes of hexafluorophosphate are more likely to occur at high temperatures (above 50 °C), even in the presence of traces of protic impurities, to produce toxic and corrosive HF.^{112,113} Therefore, it is required to choose salts with resistance to heat for wide-temperature electrolytes.

Secondly, the interplays among the electrolyte components critically affect the freezing and boiling points of electrolytes and influence the resulting thermal stability. It was reported that in some electrolyte systems such as localized high concentration electrolyte systems, the solvation structure possessed enhanced anion–cation interactions to reduce the reduction of the un-coordinated “free” solvent molecules, resulting in reduced solvent evaporation and improved thermal stability of the electrolyte.^{114,115} According to Chen *et al.*, the enhancement of interactions between the polymer matrix and solvents can improve the thermal stability of GPEs by modifying the polymer matrix with similar functional groups to those found in the solvent based on the similarity-intermiscibility theory.¹¹⁶

Thirdly, when the electrolytes are used in rechargeable batteries, the thermal stability of electrolytes at the electrolyte/electrode interfaces must be taken into account. Side reactions between the electrolyte and the charged electrodes are one of the main factors contributing to the safety hazard of rechargeable batteries. In particular, the chemical transformations and parasitic reactions of electrolytes were susceptible to accelerate at elevated temperatures. Taking commercial lithium-ion battery as an example, the previous study presented that the SEI/CEI layers had the lowest decomposition onset temperature (60–120 °C) prior to the melting of the PE separator (~ 135 °C) and the decomposition of the cathode (~ 170 °C), electrolyte (> 200 °C) and binder (> 260 °C). Thus, the decomposition of the SEI/CEI layers determines the thermal stability of the overall rechargeable battery system.^{117,118} Due to the strong dependence of SEI/CEI stability on the solvent structure of the electrolyte, rational solvation structure engineering to construct thermally stable SEI/CEI is further highlighted.^{119,120} Recently, several methods have been successfully applied to facilitate the formation of a thermally stable SEI/CEI, including adjusting the participation and contribution of ion species, solvent/water molecules, and polymer dipoles in the coordination of metal-ion solvation structure by component regulation.^{116,121,122} Still, the effects of salts, solvents, polymers, additives, and concentrations of salt on the thermally stable SEI/CEI are unclear, and further research is needed.



3.5 High dimensional/mechanical stability

Existing GPE materials generally have limitations in weak mechanical properties at room temperature, which would be further magnified due to their vulnerability to freezing at low temperatures and matrix structure failure at high temperatures.^{123,124} The freezing of electrolytes at low temperatures could induce hardening issues such as salt precipitation, volume contraction of the GPEs. At high temperatures, structure failure induced by the evaporation and leakage of solvent, as well as the softening, shrinkage and even melting of the polymer matrix may lead to damage to the structural integrity and cause many dire consequences such as battery performance degradation, short-circuit between cathode and anode and even hazardous thermal runaway.^{125–127} Targeting high reliability in extreme temperature conditions, the ideal electrolyte should possess both high mechanical strength and good flexibility across a wide temperature range. Moreover, considering the external stress that the GPEs would bear during battery manufacturing, certain safety margins should be included in setting the mechanical performance parameters.

Due to the shortcomings discussed above, previously, one could only increase the thickness of the GPE or incorporate a reinforcing matrix, both of which inevitably come at the expense of ionic conductance. Therefore, what the high mechanical strength nature of a GPE can additionally promote is the dramatic reduction of the thickness towards high mass and volume energy density of the assembled batteries.^{128,129} According to the development trend of solid polymer electrolyte, if the tensile strength of GPEs could be improved to the level of 10 MPa, ultrathin GPEs with a thickness of 5–20 μm , even thinner than state-of-the-art separators, are expected without experiencing damage during the battery assembly and operation process.^{130,131}

3.6 Facile processability, low cost and low environmental/health impact

The most widely reported preparation methods of GPEs include *ex situ* solution impregnation/casting and *in situ* polymerization. The solution impregnation method is widely used to fabricate the GPE membranes with linear polymer matrices, such as polyethylene oxide (PEO), polydimethylsiloxane (PDMS), polyvinylidene fluoride (PVDF), and poly(acrylic acid) (PAA). It typically involves dissolving the polymer and forming dry membranes through casting and solvent evaporation, followed by a liquid swelling (soaking) step to produce GPEs. This approach is well-suited to current lithium-ion battery production if the dry membranes are used as the separator before solvent infiltration. In this case, these dry membranes need to meet similar general requirements for separators to overcome the damage caused by subsequent processing in battery production, such as curling, stacking, and packaging during, battery assembly.¹³² According to the USABC stipulations, the puncture strength of the separator needs to be greater than 100 kgf mil⁻¹ (1 kgf = 9.8 N, 1 mil = 25.4 μm) and the tensile strain should be less than 2% when an external force of 1000 psi (1 psi = 0.006895 MPa) is applied.¹³³ At the same time, the thickness of GPEs must be well-controlled to be comparable to commercial polyolefin separators (15 to 25 μm)

and be competitive. The solution casting method produces self-standing GPE membranes without requiring solvent evaporation, streamlining the manufacturing process. However, it presents challenges concerning the mechanical properties of the resultant GPE membrane. For instance, mechanical brittleness can be induced by the osmotic swelling of liquid solutions. Likewise, molecular treatments aimed at enhancing the wide temperature operability can result in weakened mechanical strength. All these issues should be considered in the context of the reel-to-reel production process in battery manufacturing. Furthermore, the inadequate interfacial contacts between the prepared GPEs and the porous electrodes with high mass loading could be a tricky question that needs to be addressed prior to practical applications.^{134,135} The *in situ* polymerization method offers an alternative by generating GPE directly within the battery assembly.^{136–138} The process involves injecting a homogeneous precursor solution containing low-viscosity monomers or oligomers, salts, solvents, and initiators into the battery's electrodes and inducing polymerization *via* UV or thermal energy to form the GPE, ensuring good electrolyte-to-electrode contact. This technique is particularly useful for fabricating cross-linked GPEs, such as polyacrylate, polycarbonate, and polyether. A noteworthy advancement in this field was made by Lee and colleagues, who achieved a uniform distribution of components in the through-thickness direction of an NCM811 cathode with high areal mass loading of 60 mg cm⁻² by the *in situ* introduction of a UV-cured GPE in the cathode.¹³⁹ Although this result offers a practical solution, its universality in other practical battery systems needs further exploration. Future investigations should also consider the limited penetration of UV light and the light-absorbing characteristics of carbon materials in electrodes. Furthermore, the enclosed environments of the battery system limit the use of UV-induced *in situ* polymerization, making thermally-induced free radical or ionic *in situ* polymerization a more practical approach. The use of appropriate monomers and initiators that avoid gas or by-product formation is essential for quality GPE production. Typically, a separator is included to prevent short-circuits during *in situ* polymerization, but it often obstructs the ion transport, inevitably adding unnecessary weight and reducing ion conductivity. Research is therefore aimed at developing thin or even separator-free electrolytes and designing ion-conducting separators or hosts for *in situ* polymerization to improve the overall battery performance.

The development and application of GPEs on a large scale necessitates a significant consideration of material and production costs. Economically viable alternatives or commercially proven materials can be used as polymer matrices and solvents within GPE systems, which are generally not prohibitive and can even be cost-effective. Certain additives, such as fluorine-containing species, despite being relatively expensive, can improve the performance of GPEs with a small dosage, thus minimizing the overall material cost. Therefore, material costs may not be the primary concern at this stage. The focus should rather be on production costs as the intricate preparation, production line, and sealing technologies associated with GPE represent significant challenges that need to be circumvented.



Innovative techniques, such as rapid room-temperature *in situ* polymerization, can help reduce production cycle time and enable continuous production. Additionally, the economic and environmental implications of GPE production, including the consumption and recycling of organic solvents such as NMP, AN, and acetone, should be considered. The feasible and scalable processability of GPEs is highly demanded in pursuing the practical applications of wide-temperature rechargeable batteries, especially when applied in large-scale energy storage scenarios.

The requirements for GPEs have extended beyond their physical and electrochemical properties, delving into the realm of environmental and health impacts. It is of paramount importance to ensure that the synthesis, use, and disposal of these materials do not pose possible threats to the environment and human health. A primary concern is the heavy use of organic solvents in the production of GPEs as these solvents generally present potential hazards to both the environment and human health due to their volatility and toxicity. Therefore, more benign alternatives need to be pursued. Some GPE systems based on green solvents, such as IL and water, have been proposed. Moreover, the incorporation of biomass materials or other biodegradable polymers, including polylactic acid, cellulose, chitosan, and xanthan gum, into the electrolyte matrix could offer a promising solution. In particular, cellulose and cellulose derivatives as ion-conductors have offered an interesting opportunity for replacing fossil oil-based polymers in GPEs.^{140,141} Not only would this approach reduce the reliance on non-renewable resources but it would also mitigate the environmental burden associated with polymer waste. Additionally, rigorous safety testing must be conducted to minimize any potential health risks associated with the use of GPEs. This includes examining the potential for harmful emissions during operation, assessing the risk of fire or explosion, and evaluating the potential toxicity of the materials to humans.

4. Design and engineering strategies of GPEs for wide-temperature rechargeable batteries

As can be expected from the above section, it is not easy to find an electrolyte that fulfills all the requirements for wide-temperature rechargeable batteries. To guarantee the workability of the rechargeable batteries over a wide temperature range, many strategies have shown the effectiveness of modifying the GPE chemistry by adjusting the relative composition of the electrolyte and species of salts, polymer matrices, solvents, additives, *etc.*

4.1 Salts

Salt is the key determinant of electrolyte performance. With respect to wide-temperature operability, the addition of salt reduces the freezing point of the electrolyte according to the colligative property, and the anionic species largely determine

the properties of the SEI/CEI layers.^{105,142} For achieving wide-temperature applications, an ideal salt is expected to have superior solubility to completely dissolve and dissociate in electrolyte and the solvated cations should have high mobility to achieve high ionic conductivity of the bulk electrolyte. Meanwhile, it should have high thermal and electro-/chemical stability.

4.1.1 Blended salts. Perchlorate salts (*e.g.*, LiClO₄, NaClO₄, and KClO₄) and hexafluorophosphate salts (*e.g.*, LiPF₆, NaPF₆, and KPF₆) are the most commonly used salts in GPEs. However, perchlorate salts have a strong oxidizing ability, which makes them prone to react easily with organic substances in the electrolyte, especially at high temperatures.^{143,144} The hexafluorophosphate salts are sensitive to water and high temperatures, leading to the formation of undesired reactive species like POF₃ and HF.¹⁴² Additionally, these salts have poor solubility at low temperatures, hindering their wide-temperature applications.

Over the past few decades, significant efforts have been dedicated to designing, synthesizing, and characterizing novel salts as potential alternatives both in liquid electrolytes and GPEs. The most intensively investigated salts are thermally and chemically stable borate salts (*e.g.*, lithium bis(oxalate)borate (LiBOB), sodium bis(oxalate)borate (NaBOB) and lithium difluoro(oxalate)borate (LiDFOB)) and imide salts (*e.g.*, lithium bis(fluorosulfonyl)imide (LiFSI), sodium bis(fluorosulfonyl)imide (NaFSI), lithium bis(trifluoromethanesulfonyl)imide (LiTFSI) and sodium bis(trifluoromethanesulfonyl)imide (NaTFSI)). However, borate salts have low solubility in carbonate solvents at room temperature, which worsens at low temperatures. While imide salts have shown good performance, especially at low temperatures, the significant corrosion toward aluminium current collector at relatively high voltages limits their application scenarios. Additionally, the high cost of these salts remains a major obstacle. Though most of the newly-developed salts demonstrated outstanding performance at room temperature, their use at low and high temperatures is still to be studied.^{145,146} Therefore, the strategy of using blended salts containing at least two main salts (usually binary/ternary main salts) offers a feasible and cost-effective approach to enhance the practical applications of salts and extend the operating temperature ranges of GPEs.

As shown in Table 1, the successful application of blended salts in electrolytes has been demonstrated at both low and high temperatures.^{147–151} The reasons for the substantial improvement in the low- and high-temperature performance using blended salts in electrolytes are summarized in Fig. 6. At low temperatures, the functions of blended salts are summarized as follows: (i) improving the ionic conductivity of the electrolytes. It was found that the dissociation ability of a single salt could be improved by introducing other salts, further influencing the dissociation balance reaction of a single salt, inducing the forward reactions, and improving the ionic conductivity.^{152,153} For example, the addition of LiDFOB in the LiBF₄-based electrolyte enhances its ionic conductivity by reducing salt association (see Fig. 6(a)) thanks to the larger size and electron-withdrawing nature of the ODFB[−] anions compared to BF₄[−] anions.¹⁵⁴



Table 1 Summary of blended-salt electrolytes and their wide-temperature battery performance

Electrolyte formulation	Cathode//anode	Working temperature (°C)	Battery performance (initial capacity/capacity retention/cycles)	Ref.
0.05 M LiPO ₂ F ₂ + 1 M LiTFSI _{0.6} -LiTPFPB _{0.4} in PC:EC:EMC	NMC532//Li	-40 to 90	154.4 mA h g ⁻¹ /69.6%/300 cycles@60 °C 108.1 mA h g ⁻¹ /93.7%/50 cycles@-20 °C	147
1 M LiTFSI _{0.6} -LiTPFPB _{0.4} in PC:EC:EMC			153 mA h g ⁻¹ /39.2%/300 cycles@60 °C 105.9 mA h g ⁻¹ /91.4%/50 cycles@-20 °C Failed after 300 cycles@60 °C	148
1 M LiPF ₆ in PC:EC:EMC	LiCoO ₂ //Li	RT to 80	2.4 mA h cm ⁻² /900%/100 cycles@80 °C -14%/100 cycles@80 °C	149
0.8 M LiTFSI-0.2 M LiDFOB-0.01 M LiPF ₆ in EC/PC	LFP//Li	RT to 60	Unstable at 60 °C	169
1 M LiTFSI _{0.5} -LiBOB _{0.5} in EC/EMC	PANI@V ₂ O ₅ //Zn	-20 to 60	Unstable at 60 °C	165
0.5 M LiPF ₆ -0.5 M LiBOB in EC/EMC	LFP//Li	25	216 mA h g ⁻¹ /250 cycles@60 °C	170
0.8 M Zn(OTf) ₂ -0.2 M Zn(ClO ₄) ₂ in 2-MeTHF/H ₂ O	LiCoO ₂ //Li	RT to 40	116.9 mA h g ⁻¹ /87.93%/300 cycles@25 °C	171
Cross-linked PEGDA-ETPTA + LiTFSI-LiPF ₆ in EC:DMC:DEC	LMO + LMFP//graphite	-18 to 45	138.3 mA h g ⁻¹ /~58%/60 cycles@40 °C 84%/100 cycle@45 °C	166
poly-DOL + 0.3 M LiDFOB-2 M LiTFSI			~63% of RT capacity/—/—@-18 °C	172
PVDF-HFP + 0.8 M LiBF ₄ -0.8 M LiTFSI in EC			51% over 100 cycles@45 °C	168
PVDF-HFP + 0.8 M LiBF ₄ -0.8 M LiTFSI in EC			~60% of RT capacity/—/—@-18 °C	166
PTFE + PVDF-HFP in FEC/adiponitrile	NMC811//Li	-10 to 80	143 mA h g ⁻¹ /78.3%/1000 cycles@RT	166
PVDF-1 M LiTFSI-1 M LiBOB in PC/EC/FEC	LFP//Li		165 mA h g ⁻¹ /92.9%/300 cycles@70 °C -1-(79 mA h g ⁻¹)/300 cycles@-10 °C	172
Poly-DOL-LiPF ₆ -LiDFOB in DOL	LiCoO ₂ //Li	5 to 100	118 mA h g ⁻¹ /87.2%/1200 cycles@15 °C -1/86.5%/520 cycles@80 °C	172
Polysaccharide κ-carrageenan/PAAM 1 M ZnCl ₂ -7 M LiCl in H ₂ O	LFP//Li	-40 to RT	-1/78.3%/1500 cycles@RT -1/80.5%/2000 cycles@-20 °C -1/95.2%/2000 cycles@-40 °C	168

(ii) Facilitating the desolvation process of target ions. Recent studies have shown that anionic coordination with target ions can regulate the solvation structure by reducing the interaction strength between the target ions and the solvents, which facilitates easier desolvation at the electrolyte/electrode interfaces. In a study by Hu *et al.*, both experimental results and DFT calculations showed that with the addition of LiFSI in the LiTFSI-based electrolyte, the inclusion of FSI⁻ in the Li⁺ solvation sheath decreases the Li⁺ desolvation energy from 63.9 kJ mol⁻¹ to 56.7 kJ mol⁻¹, thereby enhancing the Li⁺ desolvation kinetics at low temperatures (Fig. 6(b)).¹⁵⁵ (iii) Promoting the formation of SEI/CEI with low ion transport barrier. The salts with good film-formability can preferentially decompose to form low ion-resistive SEI/CEI layers on the surface of electrodes, allowing fast ion-migration across the interfaces.

At high temperatures, the blended salt strategy can improve the battery performance in two ways: (i) improving the thermal stability of electrolytes. The thermal stability of electrolyte systems can be improved by blending salts that are more thermally stable.^{156,157} For example, due to the higher thermal stability of LiBF₄, the exothermic reaction of the LiPF₆-LiBF₄ dual salt carbonate electrolyte system has a higher onset temperature compared to the pure LiPF₆ carbonate electrolyte system (Fig. 6(c)).¹⁵⁸ (ii) Forming thermostable SEI/CEI layers to prevent electrolyte decomposition.^{159,160} Some salts in blended salt systems can serve as sacrificial agents. When working at elevated temperatures, these salts can decompose to form thermally and chemically stable SEI/CEI layers on the surface of charged electrodes, protecting electrolytes from further decomposing and ensuring the stability of the interfaces (Fig. 6(d)).^{161,162} Thus, the effectiveness of these electrolytes is mainly attributed to the blended salts used, which synergistically modify the interfacial chemistry on the surface of the cathode and anode. As a typical example, Luo *et al.* reported a dual-salt electrolyte based on sulfite solvent (0.4 M LiDFOB + 0.6 M LiFSI in dimethyl sulfite (DMS)) for the wide-temperature operation of lithium-ion batteries. LiFSI has a strong dissociation ability in DMS, while LiDFOB has a strong affinity for Li⁺. At the low-temperature end, LiFSI provides ultrahigh ionic conductivity, and the weakly dissociated LiDFOB promotes Li⁺ desolvation by adjusting the solvation structure. At the high-temperature end, the interface stability of both the cathode and anode is improved by the preferential self-limiting decomposition of DFOB⁻ anions to generate inorganic-rich SEI/CEI layers, largely suppressing the decomposition of DMS solvent molecules. This synergistic effect of LiFSI and LiDFOB allows the LCO//graphite full cells to withstand a wide temperature range from -78 to 60 °C.¹⁶³

The above advantages make blended-salt electrolytes a shining star for wide-temperature GPEs.^{164,165} For example, Guo *et al.* designed a poly(ethylene glycol)diacrylate (PEGDA)-based GPE using a LiTFSI-LiPF₆ dual-salt. They demonstrated that the introduction of dual salts improved the ionic conductivity to 5.6 × 10⁻⁴ S cm⁻¹ @ 25 °C, which is higher than that for single LiTFSI and LiPF₆ salt in GPEs.¹⁶⁵ In a recent study, Cuicci and co-workers used a LiTFSI-LiBOB dual-salt in a PVDF-based GPE.



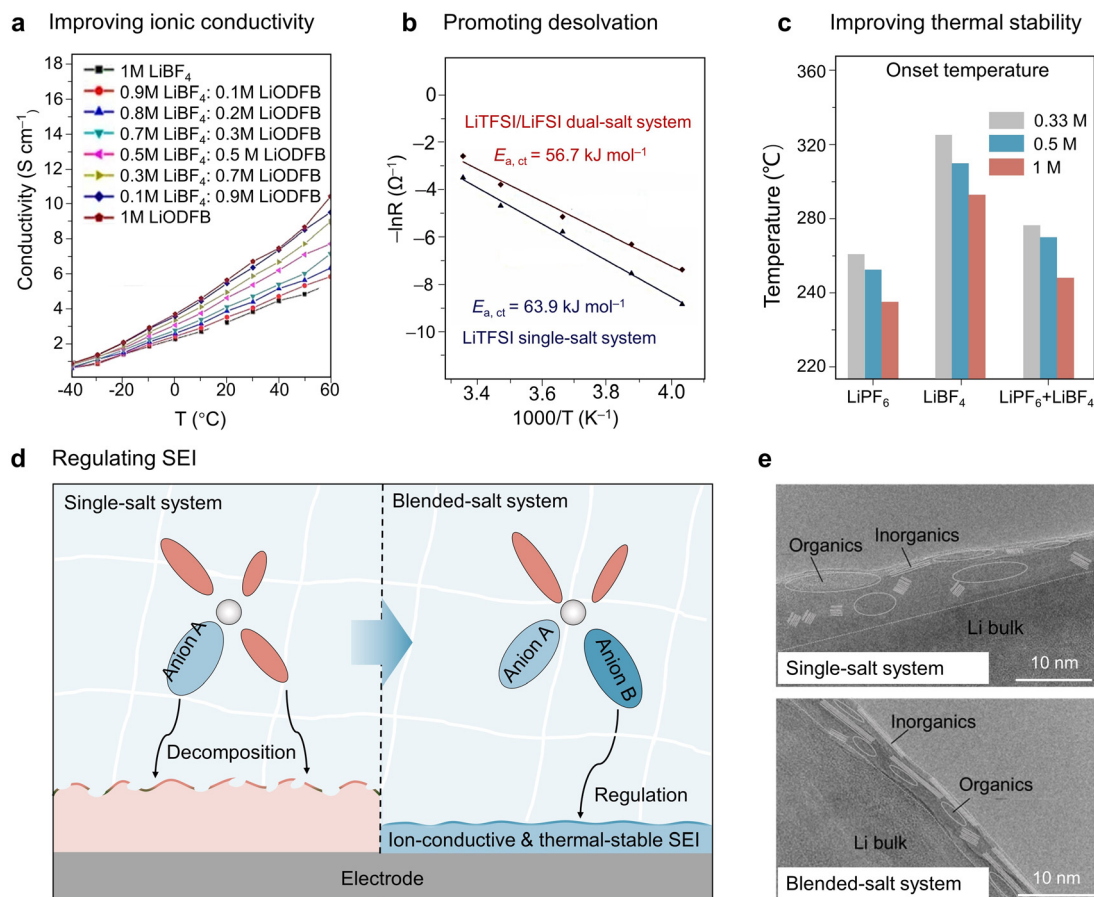


Fig. 6 Fundamental electrochemical aspects of the blended salt strategy towards wide-temperature batteries. Effects or proposed mechanisms of this strategy in regulating/improving (a) ionic conductivity, data extracted from ref. 154 with permission from Springer Nature, Copyright 2018; (b) desolvation kinetics, data extracted from ref. 155 with permission from Elsevier, Copyright 2023; (c) thermal stability, data extracted from ref. 158 with permission from IOP publishing, Copyright 2004; (d) formation of ion-conductive and thermostable SEI layer; (e) TEM images of cycled Li anode disassembled from single- and blended-salt systems. Reproduced from ref. 167 with permission from Springer Nature, Copyright 2023.

The synergistic effect of LiBOB and LiTFSI helped to achieve high ionic conductivities at both low and high temperatures (0.16 mS cm^{-1} @ $-20 \text{ }^\circ\text{C}$, 1.93 mS cm^{-1} @ $100 \text{ }^\circ\text{C}$). In addition, LiBOB has a lower LUMO energy level than LiTFSI, making it more likely to be preferentially reduced on the surface of the anode to form a stable SEI layer enriched with ion-conductive and thermostable LiF and Li-B-O-based species. Therefore, the LiTFSI-LiBOB dual-salt in GPEs enables fast ion migration at low temperatures and high interfacial stability at high temperatures, resulting in the extremely stable cycling performance of LFP//Li batteries over a wide operating temperature range (between -10 and $80 \text{ }^\circ\text{C}$).¹⁶⁶ Very recently, from the aspect of salt, Wagemaker *et al.* raised the entropy of electrolytes by introducing multiple salts (*e.g.*, LiFSI, LiTFSI, LiDFOB, and LiNO₃) in dimethoxyethane (DME). Due to the participation of more anionic groups in the solvation structures, the solvation strength of Li⁺-DME is largely weakened, and the formation of a thin ($\sim 6 \text{ nm}$) SEI layer enriched with ion-conductive inorganics is promoted. In contrast, the electrolyte with LiFSI as the single salt forms a thick organic-rich SEI layer (Fig. 6(e)).¹⁶⁷ Although this study does not specifically address the low- and

high-temperature performance of the electrolyte, we believe that this strategy has great potential for wide-temperature rechargeable batteries as the fast desolvation and migration of Li⁺ ions across the interfaces are widely recognized as necessary for improving the low-temperature performance, while a stable SEI layer holds promise for enhancing the high-temperature performance. However, although the positive effects of high-entropy salts on liquid electrolytes have been proven, there is a lack of knowledge regarding the properties of this class of liquid solutions in GPEs as well as their wide-temperature applications.

In addition to the abovementioned nonaqueous GPEs, the blended-salt GPEs with water as solvent have been also reported to achieve stable wide-temperature performance. For instance, Li *et al.* designed a polysaccharide κ -carrageenan/PAAM-based hydrogel electrolyte containing ZnCl₂-LiCl dual-salt. LiCl provides Li⁺ ions for combining H₂O molecules (competing against Zn²⁺ ions) and Cl⁻ ions for forming more Zn-Cl super halide anions, suppressing the formation of insoluble hydrolysis byproducts of Zn(OH)₂/ZnO and the freezing of the GPE. As a result, the LFP//Zn battery displays outstanding cycling stability at a low temperature of $-40 \text{ }^\circ\text{C}$.¹⁶⁸ In another study, Wang *et al.*



reported an aqueous electrolyte containing $\text{Zn}(\text{OTf})_2\text{-Zn}(\text{ClO}_4)_2$ dual-salt. The ClO_4^- anions in $\text{Zn}(\text{ClO}_4)_2$ act as hydrogen bonding acceptors to change the solvation structure and disturb the hydrogen bonding network, lowering the electrolyte solidification point and improving the Zn^{2+} transport kinetics. At the same time, $\text{Zn}(\text{OTf})_2$ facilitates the formation of ZnF_2 -rich SEI layer on the zinc anode surface, effectively inhibiting the growth of zinc dendrites and the parasitic reaction of H_2O molecules. As a result, the $\text{PANI@V}_2\text{O}_5//\text{Zn}$ cells delivered improved cycling stability across a wide temperature range from -20 to 60 °C.¹⁶⁹

Overall, the blended-salt strategy opens up a wide range of opportunities for designing GPEs with improved thermal stability, desirably high ionic conductivity and the ability to form more durable electrolyte/electrode interfaces in practical batteries. Due to the very limited research experience yet high chemical/structural complexity of blended salt systems, previous efforts in this direction have mainly focused on the characterizations of the GPEs' basic properties and demonstrations of battery performance improvements, lacking mechanistic investigation on how different salts can work collaboratively to realize these achievements. Continual research should focus on the precise detection of electrolyte/electrode interface evolution behaviors and the reconstruction of interactions among various ionic and molecular species within the blended-salt systems, this could involve the use of cutting-edge spectroscopic and imaging techniques to observe the dynamic processes at the molecular level, potentially leading to the discovery of novel blended-salt formulations for wide-temperature GPEs.

4.1.2 Concentration effects. Apart from the species of the added salts, the concentration of salts significantly affects the wide-temperature properties of the GPEs (as shown in Table 2) as it significantly influences the physicochemical properties of electrolytes, including ionic conductivity, anti-freezing performance and thermal stability.

In conventional liquid electrolyte recipes, the ionic conductivity increases drastically as the salt concentrates until it peaks at an optimum value, in which the optimum ionic conductivity can be achieved by the trade-off between the ionic carrier number proportional to the dissolution and dissociation of salts and the ionic mobility that is inversely proportional to the viscosity of the solution. After that, the increase in the concentration will lead to a reduction in the ionic mobility due to an increase in viscosity (Fig. 7(a)–(c)).^{173–175} Another critical feature brought by concentration alternation involves ion-solvation structure and the interfacial chemistry (Fig. 7(d)). In low-concentration electrolyte (typically, <1 M), the number of solvent molecules is much greater than that of salts, in which metal ions are completely coordinated with solvent molecules in the solvation structure, and free solvent molecules are the dominant species (Fig. 7(d₁)).^{106,176} Conventional medium-concentration electrolytes (usually 1 to 2 M) exhibit a balanced number of anions and solvents; the solvent separated ion pairs (SSIPs) and free solvent molecules are the main species in solvation structure (Fig. 7(d₂)). Such an electrolyte is desired because only a small amount of salt is required,

which reduces the cost of electrolyte. However, the free solvent molecules are prone to suffer from freezing, and the strong binding of solvent molecules to the ions in SSIPs may limit the desolvation dynamics at low temperatures. Moreover, free solvent molecules and loosely-coordinated anions in SSIP can intensely decompose near the electrolyte/electrode interface at high temperatures. All these effects will limit the wide-temperature use of low- and medium-concentration electrolytes. With ascending salt concentration to above a threshold (usually >2 M, depending on the salt-solvent combinations), the interactions between cations and anions are enhanced, and the solvation sheath of the cations is increasingly dominated by salt anions, leading to the structural transition from SSIPs to contact ion pairs (CIPs) and ionic aggregates (AGGs) (Fig. 7(d₃)). The new solution structure makes the location of the LUMO shift from the solvent towards the salt, resulting in the reductive predominant decomposition of the anions before the solvent to form an anion-derived SEI at low potential. This anion-derived passivation was revealed to be inorganic-rich, providing low interfacial charge-transfer resistance and high stability, which is beneficial for wide-temperature performance.^{167,177} Moreover, the dramatical drop of free solvent molecules in the liquid solution shows less decomposition and passivation in the electrolyte–electrode interfaces, improving the thermodynamic stability and electrochemical stability of the electrolyte.¹⁷⁸ In addition, as anions are confined kinetically to CIPs and AGGs, the cationic transference number will increase in high-concentration electrolytes, which favors the improvement of rate capability. Therefore, increasing the salt concentration of the electrolyte provides a potential solution for widening the operating temperature window.

High-concentration salt electrolytes have been demonstrated to boost the cycling stability of batteries at high temperatures. For example, Sun *et al.* developed an electrolyte for Li-S battery with 5 M LiFSI dissolved in tetraethylene glycol dimethyl ether (G4). They suggested that the SEI layer formed in this high-concentration electrolyte consisted mainly of LiF, which was derived from the LiTFSI salt rather than the G4 solvent. The LiF-rich SEI layer helps to suppress the unwanted side reactions and dendrite growth on the Li anode, enabling the safe and stable operation of Li-S batteries at 60 °C.¹⁷⁹ Similarly, a concentrated electrolyte of 2.0 M LiDFOB in EC/DMC was proved to boost the high-temperature cycling stability of a graphite//LCO cell at 90 °C. They attributed this to the formation of a compact LiF-rich CEI layer in the high-concentration electrolyte, which could effectively inhibit the solvent decomposition and the dissolution of Co^{2+} in the LCO cathode.¹⁸⁰

In spite of the improved high-temperature electrochemical performance enabled by high-concentration electrolytes, their applications below room temperature are rarely reported. The main reason is their dramatically increased viscosity at low temperatures, which leads to low ionic conductivity and inferior wetting property.¹⁷⁴ The novel concept of localized high concentration electrolytes (LHCE), as proposed in recent studies, is particularly interesting and is attracting increasing



Table 2 Summary of high-concentration GPEs and battery performance under a wide-temperature range

Electrolyte	σ	Cathode/anode	Working temperature (°C)	Battery performance	Ref.
—/5 M LiTFSI in G4	$9 \times 10^{-5} \text{ S cm}^{-1}$ @25 °C	S//Li	25 to 60	666 mA h g ⁻¹ over 100 cycles@60 °C	179
—/2.0 M LiDFOB in EC/DMC	—	LCO/graphite	25 to 90	99.6% after 160 cycles@90 °C	180
—/1M LiTFSI MDFA/MDFA-TTE	0.6 mS cm ⁻¹ @20 °C	NMC811/graphite	—60 to 60	160.2 mA h g ⁻¹ /80% after 360 cycles@-30 °C	108
PVDF-HFP/6 M LiTFSI/DMSO	0.89 mS cm ⁻¹ @50 °C; 0.031 mS cm ⁻¹ @0 °C	LFP//Li	—10 to 100	158.7 mA h g ⁻¹ /50C@100 °C; 158.7 mA h g ⁻¹ /2C@-10 °C	188
—/19 M CH ₃ COONH ₄ /H ₂ O	—	NiHCF@CNTs/ poly(1,5-NAPD)	—40 to 80	0.05 A g ⁻¹ , 130 mA h g ⁻¹ @-40 °C; 4 A g ⁻¹ , 84 mA h g ⁻¹ @25 °C 4 A g ⁻¹ , 102 mA h g ⁻¹ @80 °C	197
Sodium polyacrylate (PVNa)/6 M KOH in H ₂ O	5.7 S m ⁻¹ @-20 °C; 12.6 S m ⁻¹ @24 °C; 16.3 S m ⁻¹ @50 °C	NiCo//Zn	—20 to 25	87%/10 000 cycles@RT	198
Polyelectrolyte poly(2-acrylamido-2-methylpropanesulfonic acid potassium salt) and methyl cellulose/5 M KOH in H ₂ O	0.18 S cm ⁻¹ @-20 °C; 1.05 S cm ⁻¹ @25 °C	Co ₃ O ₄ /carbon black/ air	—20 to 50	764.7 mA h g ⁻¹ @25 °C; 758.5 mA h g ⁻¹ @0 °C; 754.2 mA h g ⁻¹ @-20 °C	199
PVA/33 M LiNO ₃ in H ₂ O	$2.51 \times 10^{-2} \text{ S cm}^{-1}$ at RT	LiMn ₂ O ₄ /V ₂ O ₅ cathode//Zn	RT to 80	64.7 mA h g ⁻¹ —/@RT	192
PAM/3 m Zn(BF ₄) ₂ in H ₂ O	2.38 mS cm ⁻¹ @-70 °C	PANI@SWCNTs	—70 to 25	54.5 mA h g ⁻¹ 76.6%/176 cycles@60 °C —/≈100%/100 cycles@-70 °C	189
Polyacrylamide (PAM)/7.5 M ZnCl ₂ in H ₂ O	6 mS cm ⁻¹ @RT	WO ₃ //Zn	—30 to 30	~106 mA h g ⁻¹ /94%/1000 cycles@25 °C —/—(278.3 mA h m ⁻²)/9200 cycles@RT	200
Xanthan gum/12 M ZnCl ₂ in H ₂ O	1.28 mS cm ⁻¹ @-30 °C 13.8 mS cm ⁻¹ @20 °C 5.0 mS cm ⁻¹ @-20 °C	NH ₄ V ₃ O ₈ ·1.9H ₂ O//Zn	—20 to 20	At 1.5 A g ⁻¹ , 166 mA h g ⁻¹ @20 °C; 97 mA h g ⁻¹ @0 °C 185 mA h g ⁻¹ /90.3%/800 cycles@20 °C; 129 mA h g ⁻¹ /90.7%/800 cycles	201
PAM/3 M Zn(OTf) ₂ in H ₂ O	1.9 mS cm ⁻¹ @-30 °C	MgVO//Zn	—30 to 80	484 mA h g ⁻¹ /without any capacity sacrifice/100 cycles@60 °C	191
Poly(2-acrylamide-2-methyl-1-propanesulfonic acid co-acrylamide)/3 M Zn(ClO ₄) ₂ in H ₂ O	27.1 mS cm ⁻¹ @30 °C 59.9 mS cm ⁻¹ @25 °C	PANI//Zn	—35 to 25	—/almost unchanged/100 cycles@-30 °C —/great stability for > 30 000 cycles@25 °C	202
PVDF-HFP/6 M LiTFSI in DMSO	14.2 mS cm ⁻¹ @-35 °C 0.27 mS m ⁻¹ @20 °C 0.89 mS m ⁻¹ @50 °C	LFP//Li	—10 to 100	—/great stability for > 70 000 cycles@-35 °C 163.5 mA h g ⁻¹ /89.6%/800 cycles@25 °C —/stable for over 100 cycles@100 °C	188
PVA/33 M LiNO ₃ in H ₂ O	$2.51 \times 10^{-2} \text{ S cm}^{-1}$ @RT	LiMn ₂ O ₄ /VO ₂	RT to 80	2C, 133.5 mA h g ⁻¹ @-10 °C 73.0 mA h g ⁻¹ —/@80 °C	192
PAA/1 M Zn(OAc) ₂ and 40 M KOAc in H ₂ O	$3.74 \times 10^{-3} \text{ S cm}^{-1}$ @RT	MnO ₂ //Zn	25 to 80	71.1 mA h g ⁻¹ /76.6%/176 cycles@60 °C 299.0 mA h g ⁻¹ /90.6%/300 cycles@60 °C	194



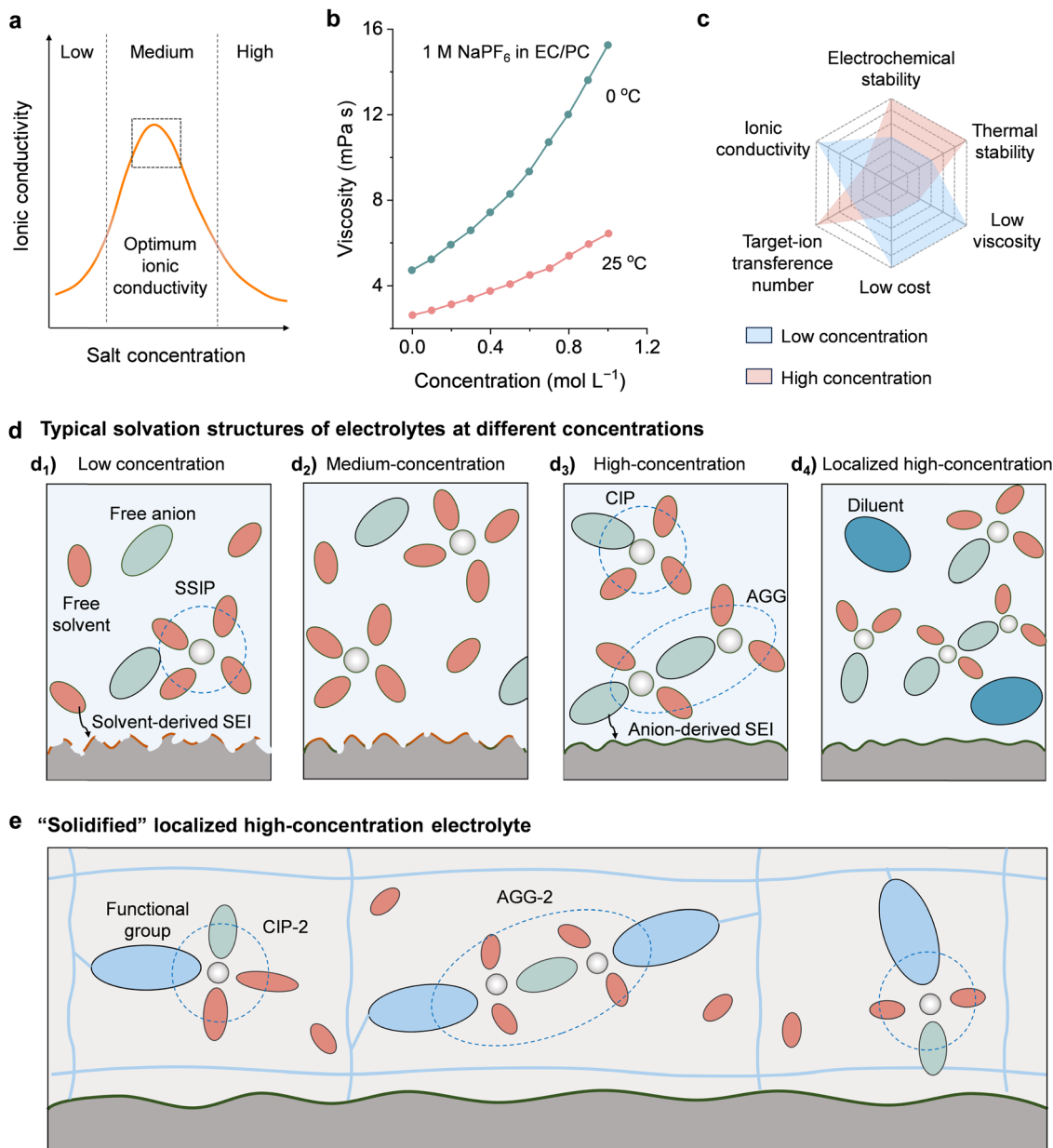


Fig. 7 Extending the operating temperature of GPEs *via* salt concentration adjustment. (a) Dependence of ionic conductivity on the salt concentration; (b) Variations of viscosity on the salt concentration at different temperatures, data extracted from ref. 174. Reproduced from ref. 174 with permission from American Chemical Society, Copyright 2020; (c) the comparison of high- and low-concentration electrolytes.¹⁹⁶ Reproduced from ref. 196 with permission from CC BY 4.0; (d) the solvation structure and SEI evolution of (d₁) low, (d₂) medium-concentration, (d₃) high-concentration and (d₄) localized high-concentration electrolytes; the schematic illustrations of (e) the intermolecular interactions and solvation structures in a solidified localized high-concentration electrolyte.

attention.^{181–183} By adding a small amount of non-polar or low-polar solvents such as 1,1,2,2-tetrafluoroethyl-2,2,3,3-tetrafluoropropyl ether (TTE), bis(2,2,2-trifluoroethyl)ether (BTFE), and methyl difluoroacetate (MDFA), the concentrated electrolyte solution can be diluted while maintaining the originally favorable solvation structures (highly concentrated coordination clusters) (Fig. 7(d₄)).^{184–187} For example, Huang *et al.* revealed that the fluorinated ether cosolvent with low dielectricity and weak binding strength with Li⁺ intrinsically facilitated the anion–cation interactions by affording a specific low-dielectric

environment, thus promoting the generation of AGGs and favoring the formation of a dense and smooth SEI layer on the Li metal anode.¹⁸⁴ In a more recent study, Wang *et al.* developed an LiFSI-based LHCE diluted with TTE and MDFA. They revealed that the addition of TTE and MDFA minimized the Li⁺–solvent binding energy and promoted the formation of CIPs and AGGs, enabling the improvement of ionic conductivity and the formation of TFSI⁻ anion-derived LiF-rich SEI/CEI layers. As a result, the NMC811/graphite full cells achieved stable cycling performance over a wide temperature range from



–60 to 60 °C.¹⁰⁸ However, developing solvent systems that can concurrently show good salt solubility and high compatibility with polymer matrices remains challenging. Very recently, a “solidified localized high-concentration electrolyte” was proposed through the gelation of a concentrated solution containing 11.92 M LiTFSI/DMSO within the PVDF-HFP membrane, in which the non-solvating PVDF-HFP framework participated in the regulation of the solvation structure. By interacting with Li⁺ ions and DMSO molecules, the PVDF-HFP framework can confine solvent molecules and promote the formation of ion aggregates (donated as CIP-2 and AGG-2) in an ultrahigh salt concentration regime (Fig. 7(e)). The unique solvation structure enables a high ionic conductivity (0.031 mS cm⁻¹ at 0 °C) as well as the formation of an LiF-rich SEI layer, resulting in the stable operation of LMBs over a wide temperature range from –10 to 100 °C.¹⁸⁸ Although only very limited progress has been achieved, we anticipate that studies on GPEs with (localized) high salt concentration will increase in the years to come.

In aqueous electrolyte systems, high-concentration salts are thought to help reduce hydrogen bonding density between free water molecules, inhibiting the formation of ice crystals and lower the freezing point of aqueous electrolytes.¹⁸⁹ The wide-temperature effectiveness of high-concentration (“salt-in-water”) electrolytes has been demonstrated for both liquid electrolytes and GPEs.^{157,190} A concentrated Zn(OTf)₂-based polyacrylamide (PAM) GPE, for example, was reported to show satisfactory low temperature ionic conductivity of 1.9 mS cm⁻¹ at –30 °C and thus boosting the wide-temperature performance of the Zn–Mg_xV₂O₅·*n*H₂O (MgVO) battery from –30 to 80 °C. This is because the strong coordination between water and ions in the concentrated electrolyte reduces the intermolecular hydrogen bonds, thus preventing the electrolyte from freezing at low temperatures and suppressing the electrochemical activity of water from side reactions on both the surfaces of the MgVO cathode and Zn anode at high temperatures.¹⁹¹ Even when increasing the salt concentration to an oversaturated state, GPEs still showed an expanded operating temperature range.^{192–194} A recent work by Tao *et al.* demonstrated that a poly(acrylic acid) (PAA)-based GPE with oversaturated salts (1 M Zn(OAc)₂ + 40 M KOAc) and configuration merely composed of inactive “frozen-like” solvent and contacted cation solvation sheath helped to improve the electrochemical stability and suppress Zn dendrites formation at an elevated temperature of 60 °C.¹⁹⁴

Therefore, constructing salt-concentrated GPEs may be a promising direction in developing wide-temperature adaptable batteries. However, the lack of a general principle for guiding the construction of salt-concentrated GPEs poses many uncertainties to researchers in this field. Laborious trial and error experiments are needed for the optimization of salt type and concentration. In addition, developing low-cost salt alternatives that do not compromise the electrochemical performance of the GPEs is a critical yet thorny issue. It should also be noted that increasing the salt concentration may negatively affect the GPEs’ mechanical properties.¹⁹⁵ Therefore, the rational design of polymer matrices becomes crucial in balancing their electrochemical performance and mechanical stability, especially under extreme temperature conditions.

Table 3 Summary of polymer matrix features in GPEs and their wide-temperature battery performance

Polymer matrices	Functional groups	Liquid solution	Ionic conductivity	Working temperature (°C)	Cathode//anode	Battery performance	Ref.
PAM-SC	–COO [–]	6 M KOH in H ₂ O	324.68 mS cm ⁻¹ @20 °C 73.96 mS cm ⁻¹ @–40 °C	–40 to 40	Air//Zn	–/–/700 cycles@–40 °C –/–/400 cycles@0 °C	250
G-CyBA/PAAm double network	–NH ₂ –OH	6 M KOH in H ₂ O	300.1 mS cm ⁻¹ @–80 °C 431.7 mS cm ⁻¹ @100 °C	–50 to 100	Air//Zn	–/–/170 cycles@40 °C –/–/42 cycles@100 °C –/–/192 cycles@70 °C	225
Poly(TMPTA)	–C–O–C– –C=O	1 M NaPF ₆ in EC:DEC:FEC	1.13 × 10 ⁻⁴ S cm ⁻¹ @0 °C 7.16 × 10 ⁻⁴ S cm ⁻¹ @30 °C 1.13 × 10 ⁻³ S cm ⁻¹ @60 °C	0 to 60	NMT//Zn	–/–/240 cycles@–50 °C 71.6 mA h g ⁻¹ /98%/100 cycles@0 °C 83.8 mA h g ⁻¹ /80%/1000 cycles@60 °C	233
Polymeric acid PAAm	–SO ₃ O=C–	3 M Zn(ClO ₄) ₂ in H ₂ O Zn(CF ₃ SO ₃) ₂ in DMSO/H ₂ O	14.2 mS cm ⁻¹ @–35 °C	–35 to 25 –40 to 60	PANI//Zn Zn ₃ V ₂ O ₈ //Zn	79.6 mA h g ⁻¹ /–/70 000 cycles@–35 °C –/95.27%/3000 cycles@RT; ~175 mA h g ⁻¹ /–/75 000 cycles@–40 °C	202 249



4.2 Polymer matrices

In addition to conditioning salts, the modification of polymer matrices in GPEs is another commendable option to achieve stable performance under wide-temperature conditions (Table 3). The most frequently used homopolymer and copolymer matrix materials in GPEs include synthetic and natural polymers. Different polymers possess different polymer chain structures, functional groups, properties and costs, as presented in Fig. 8. Synthetic polymers, such as PEO, PVDF, polyacrylates (polymethyl methacrylate) (PMMA), poly(acrylamide) (PAM), polyacrylonitrile (PAN), and PAA, possessing exceptional ionic conduction properties and excellent thermal stability, are widely used as polymer matrices of wide-temperature adaptable GPEs. However, these synthetic

polymer matrices generally come with high costs due to their complex synthesis processes and raw material costs. This can significantly impact the overall cost of the GPEs and, consequently, the final battery products. Therefore, natural polymers derived from biomass, such as cellulose, chitin, agar, sodium alginate, starch, and silk fibroin, are emerging as sustainable and cost-effective alternatives. In addition, these renewable materials offer decent mechanical properties for GPEs.^{141,203,204} However, their performance, particularly their ionic conduction properties, might not match those of synthetic polymers to satisfy the demand for long cyclability over a wide temperature range. Therefore, engineering the polymers to balance these factors—ionic conduction, thermal stability, and cost—is crucial for developing wide-temperature adaptable GPEs.

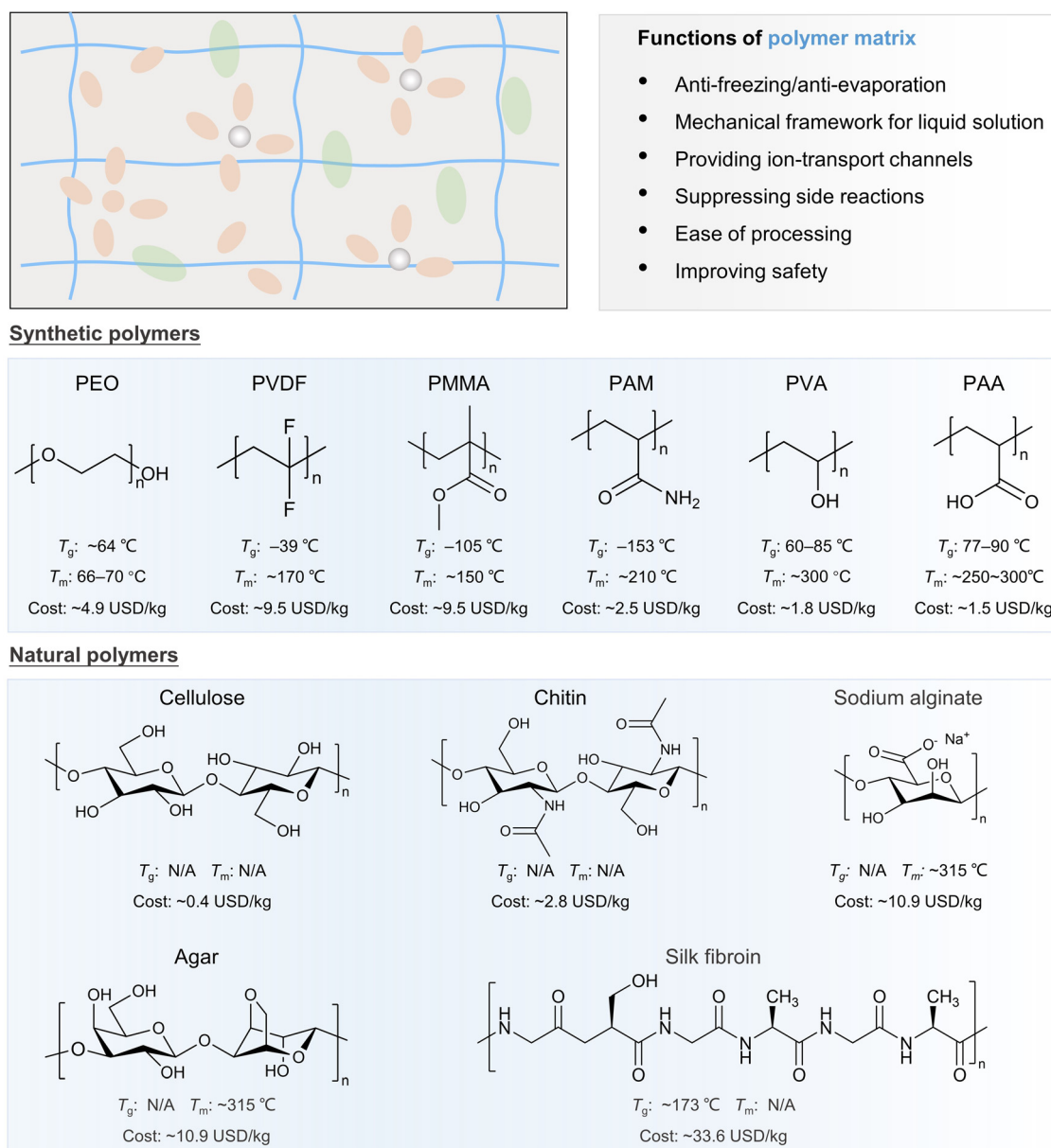


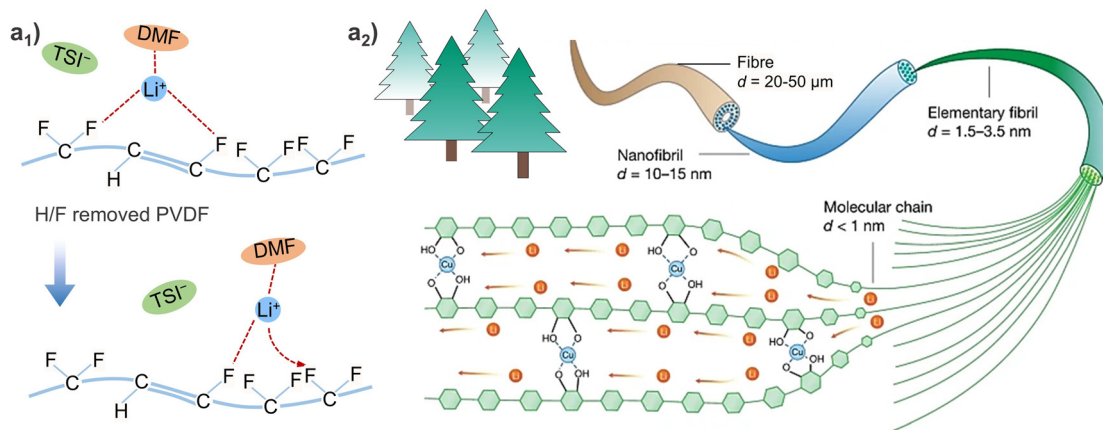
Fig. 8 Functions of polymer matrices in extending the operating temperature. Molecular structures, thermal properties and estimated costs of various synthetic and natural polymers having been widely used or showing great application potential.



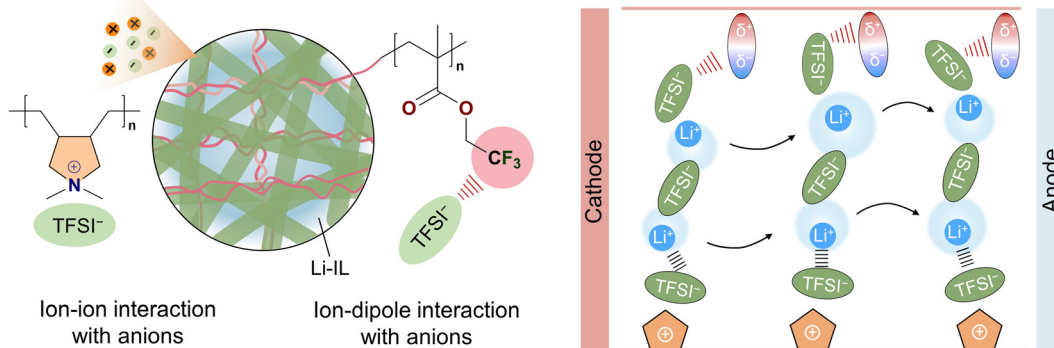
4.2.1 Ion conduction. For GPEs, their polymer matrices are generally considered to be a physical obstruction to the ion conduction process, leading to the classical trade-off issue between ionic conductive ability and mechanical strength. To address this, regulating the solvation structure and thereby the ion transport behavior by structuring/functionalizing the polymer matrix is attracting soaring interest in the GPE research community.^{205,206} Chen's group suggested that the interaction between DMF-dehydrofluorinated PVDF and the [DMF-Li⁺]

complex promotes the charge carrier along PVDF segmental chains (Fig. 9(a1)).²⁰⁷ Similar ion conduction mechanisms have also been observed in GPE systems such as PVDF-HFP/LiTFSI/tetraglyme and PVDF-HFP/LiTFSI/DMSO.^{116,188} The polymer-facilitated approach to ion conduction has also been observed in electrolytes based on natural polymers, such as cellulose, the most abundant natural polymer on earth, with a wealth of polar chemical groups (such as -OH and -O-) in its molecular chains.²⁰⁸⁻²¹⁰ Engineered into an ion-conductive material,

a Polymer matrices promoting the movement of target ions



b Polymer matrices inhibiting the movement of non-target ion species



c Polymer matrices with combined effects

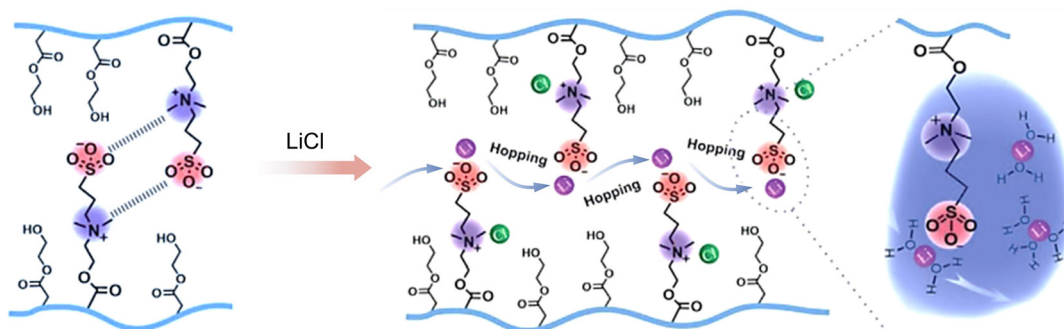


Fig. 9 Three proposed mechanisms possibly to facilitate the ion transport in GPEs. (a₁) Schematic illustration for the interactions and lithium transport in the "dried" PVDF-DMF-LiTFSI electrolyte. Reproduced from ref. 207 with permission from Elsevier, Copyright 2022. (a₂) Schematic illustration of the structure and ion-transport channels in cellulose-based GPE. Reproduced from ref. 209 with permission from Springer Nature, Copyright 2021. (b) Schematic illustration of the Li⁺ transport mechanism in IGEM. Reproduced from ref. 122 with permission from John Wiley & Sons, Copyright 2021. (c) Proposed Li⁺ migration mechanism in polySH electrolyte. Reproduced from ref. 215 with permission from John Wiley & Sons, Copyright 2021.



cellulose has shown impressive ionic conductivity. In a pioneering study, Hu's group successfully developed an Li^+ -conducting polymer electrolyte using cellulose nanofibrils, with a surprisingly high room-temperature ionic conductivity of 1.5 mS cm^{-1} along the molecular chain direction.²⁰⁹ The possible fundamental behind fast ion-transport dynamics is described in Fig. 9(a₂). The coordination of Cu^{2+} ions with the oxygen-containing groups helps to expand the spacing between the cellulose polymer chains into molecular channels and Li^+ transport through the molecular channels by hopping between the anionic sites of oxygen-containing groups with the assistance of bound water molecules. In a more recent study, Wang and colleagues further enhanced ion transport by esterifying cellulose acetate, weakening hydrogen bonds between the cellulose chains, facilitating the coupling/decoupling of Li^+ ions and oxygen atoms, and thus speeding up Li^+ ion transport.²¹⁰

Apart from interaction with cations, some polar functional groups can also interact with anions to affect ion conduction.^{211,212} Yu *et al.* reported an ionogel membrane (IGEM) composing polymerized ionic liquid (PIL) of poly(diallyldimethylammonium) bis(trifluoromethanesulfonyl)imide (PDAD-MATFSI) nanofibers and cross-linked poly(2,2,2-trifluoroethyl methacrylate) (PTFEMA), in which the positively charged PIL and functional groups ($-\text{CH}_2\text{CF}_3$) with strong electronegativity on PTFEMA were able to balance the ion-dipole interactions and selectively anchor TFSI⁻.¹²² By regulating the environment surrounding the Li^+ ions, the ion transport within the IGEM shifts from a sluggish vehicular Li^+ transport, where Li^+ migrates together with its coordination shell, to fast structural Li^+ diffusion, where Li^+ migrates by anion exchange in the first coordination shell, thus boosting the Li^+ mobility (Fig. 9(b)). Benefiting from the enhanced Li^+ conductivity, the LFP//Li cells with the IGEM allows low interfacial polarization, reduces the side reactions and achieves reasonably high rate capability and stable cycling stability in a wide temperature range from 0 to 90 °C. In another study, Coskun's group introduced a Lewis-acidic fluorinated alkyl side chain to functionalize the GPE matrix. The Lewis acidic sites immobilize TFSI⁻ anions through noncovalent interactions, decreasing the affinity of Li^+ towards oxygen atoms of glycol chains and promoting the Li^+ mobility, thus realizing a high room-temperature Li^+ conductivity (9.16 mS cm^{-1}) and high t^+ value of 0.69 simultaneously.²¹³

Beyond the individual interactions with either cations or anions, some polymer matrices can interact with both anions and cations in GPEs. One such example is the zwitterionic polymer, in which the cation and anion are covalently bonded to form the side chain. This unique structure allows the dissociation of salts and increases the abundance of free ions through electrostatic interactions. Additionally, the zwitterionic polymer can also serve as migration pathways for both cations and anions under an applied electric field. These properties have made zwitterionic polymers highly attractive as matrices for constructing high-performance GPEs.^{214–216} Liu's group designed a zwitterionic polymer by randomly copolymerizing zwitterionic monomer (SBMA) and 2-hydroxyethyl acrylate (HEA) in the presence of LiCl salt. They proposed a direct

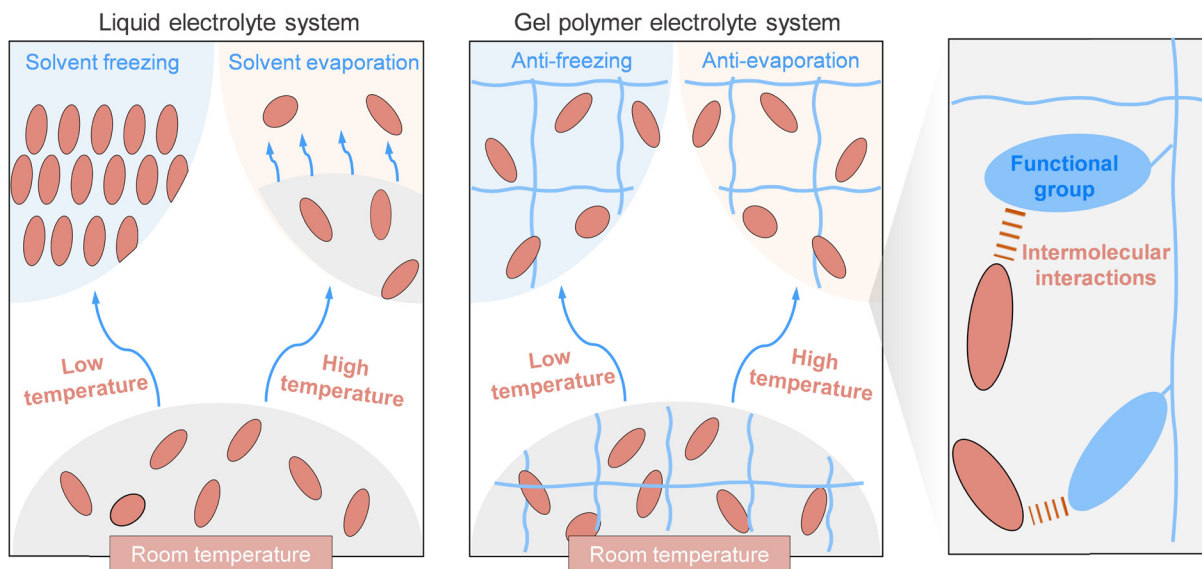
hopping migration mechanism, wherein the interactions between anionic and cationic counterions on zwitterionic chains and Li^+ ions facilitates the dissociation of lithium salt, allowing the formed hydrated lithium-ion ($\text{Li}^+(\text{H}_2\text{O})_n$) to hop and migrate through the channel of zwitterionic groups, as depicted in Fig. 9(c). This unique migration mechanism greatly improves the ionic conductivity of the GPE. As a result, the GPE exhibited an impressive ionic conductivity of 146 mS cm^{-1} at room temperature and 12.6 mS cm^{-1} at $-40 \text{ }^\circ\text{C}$, which is much higher than that of the conventional PVA-based GPEs.²¹⁵

In these cases, apart from providing mechanically stable networks for the liquid solution, the polymers serve as the ion transport medium. GPE with rationally tailored polymer chain chemistry is therefore promising to deliver desirable ionic conductive ability. This would be further highlighted under the development trend that liquid-lean GPEs are nowadays actively pursued.^{217,218} as the liquid fraction is reduced, the ion transport may not depend largely on the liquid solvent molecules, and the role of the polymer matrix in ion conduction can become very prominent. This leads to a new sub-class of GPEs with improved stability and safety and may imply new thoughts for designing batteries with an extended operating temperature range.

4.2.2 Anti-freezing and anti-evaporation abilities. In addition to being involved in ionic conduction, polymer-solvent interaction (*e.g.*, electrostatic interactions, van der Waals interaction and hydrogen bonding interactions) are thought to be responsible for the anti-freezing ability at low temperature and anti-evaporation ability at high temperature, thus extending the operating temperature range of GPEs (Fig. 10(a)). This strategy has been demonstrated in a large number of GPEs to extend the operating temperature range where the polymers contain solvent-philic polar groups.^{219,220} For example, the abundant hydrophilic functional groups of polymer matrices of hydrogel electrolytes, such as carboxyl ($-\text{COO}^-$), hydroxyl ($-\text{OH}$), amide groups ($-\text{CONH}_2$), amino, and sulfonic acid, promote salt dissolution and enhance the interactions between polar water molecules and hydrogel networks through strong hydrogen bonds, thus reducing the fraction of free water. A recent study demonstrates that the carboxyl alkylated-PAA hydrogel exhibits anti-freezing property at $-20 \text{ }^\circ\text{C}$ comparable to ILs, higher than that of the bare PVA hydrogel electrolyte ($-13 \text{ }^\circ\text{C}$) when accommodated with the same aqueous solution (10 wt % KOH).²²¹ Liu *et al.* confined DMSO within a 4,4'-bis(stearoylamino)diphenyl ether-based polymer matrix to fabricate GPE. The strong interactions between the N-H bonds in BSDE and the S=O bonds in DMSO ($\text{N-H} \cdots \text{O}=\text{S}$) broke the original weak hydrogen bonds and ordered molecular arrangement of DMSO, damaging the frozen structure of DMSO solvent at sub-zero temperatures.²²² Wu's group reported that the abundant functional groups in modified polysaccharides carboxymethyl chitosan-PAM hydrogel could form ternary and weak hydrogen bonding with chaotropic ClO_4^- anions and water molecules, which enabled the polymer chains to break the hydrogen bonds between water molecules to remarkably decrease the electrolyte freezing point.²²³

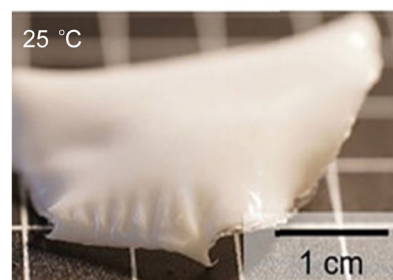
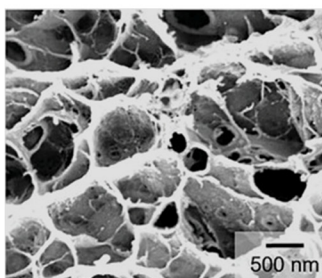


a Regulating polymer matrix-solvent interactions to achieve anti-freezing/anti-evaporation



b Modifying polymer matrix structures to achieve anti-freezing/anti-evaporation

Semi-closed-cell hydrogel electrolyte



Open-cell hydrogel electrolyte

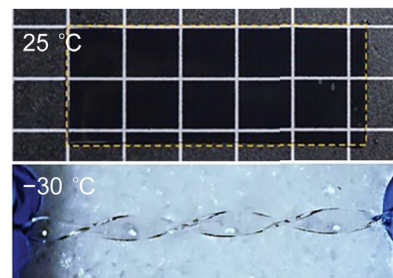
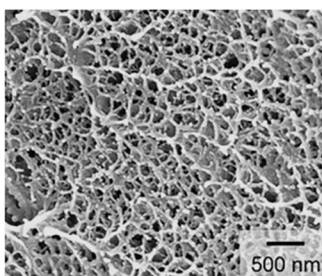
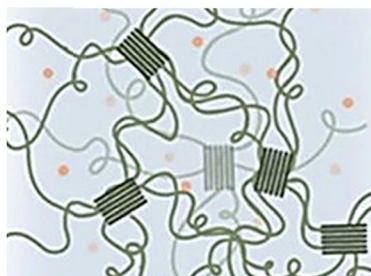


Fig. 10 Beneficial effects of polymer matrices on suppressing the freezing and evaporation of confined solvents. (a) Illustrations of the mechanisms involved in the suppression of freezing and evaporation. (b) Schematics, SEM images and optical photos of GPEs featuring polymer matrices with open-cell and semi-close-cell structures. Reproduced from ref. 224 with permission from John Wiley & Sons, Copyright 2023.

In addition, the structure of polymer matrices influences the temperature tolerance, mechanical and ion transporting properties of GPEs. As shown in Fig. 10(b), He's group fabricated a PVA-based GPEs with unique open-cell porous structures *via* a strategy combining co-nonsolvency and "salting-out". They used "salting-out" K^+ and acetate to toughen hydrogels *via* ion-promoted chain aggregation and added DMSO cosolvent to the precursor, synergistically promoting the formation of open-cell porous structures with a densified polymer network. The open-cell porous structure allows the fast diffusion of the ions

and achieves enhanced mass transport ($10\times$ lower overpotential) compared to the PVA hydrogel with a semi-closed-cell structure. Moreover, the hydrogel electrolyte contains strongly aggregated polymer chains and disrupted hydrogen bonds among free water molecules, contributing to a high mechanical strength (tensile strength of 15.6 MPa) and excellent freezing tolerance ($< -77^\circ\text{C}$).²²⁴ Gu *et al.* designed a supramolecular double-network hydrogel electrolyte with a hierarchical porous structure by interpenetrating guanosine-cyclohexylboronic acid (G-CyBA) with a polyacrylamide (PAAm) network in the



presence of concentrated aqueous KOH. The O atom in the B–O bonds of G-CyBA nanowires intensively interact with -NH_2 groups in PAAM *via* multiple $\text{N-H}\cdots\text{O}$ hydrogen bonds. Also, the additional hydrogen bonding sites in the -OH groups of the G-quadruplex nanowires and -NH_2 groups of the PAAM chain were supposed to further interact with water molecules to immobilize the solvents. The obtained hydrogels exhibit high mechanical robustness and stretchability, ultra-high conductivity, as well as superior anti-freezing ($-196\text{ }^\circ\text{C}$) and anti-dehydration ($100\text{ }^\circ\text{C}$) properties.²²⁵

Of note, the use of nonflammable polymer matrices is effective to improve the safety of GPE at high temperature. For example, Liu *et al.* fabricated the succinonitrile-based PIL GPE. With the thermally stable PIL as the polymer matrix, the thermal stability of GPE was improved up to $300\text{ }^\circ\text{C}$, $70\text{ }^\circ\text{C}$ higher than that of GPE without the PIL matrix.²²⁶ In another study, Zhang *et al.* synthesized nonflammable or less-flammable poly(phosphonate/phosphates)-containing polymer matrices. These GPEs show good thermal stability and efficient flame retardance even filled with 90 wt% of traditional liquid electrolytes.^{227,228}

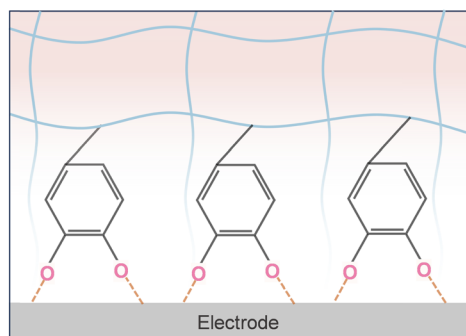
Overall, by fine-tuning the polymer–solvent interactions and the structure of the polymer matrix, it is possible to extend the operating temperature range of GPEs, making them more versatile for a wide range of applications.

4.2.3 Improvement of interfacial compatibility. In addition to lowering the freezing point and suppressing the evaporation of GPEs, the polymer matrices can also regulate the electrolyte/electrode interface chemistry through the chemical/physical interactions between polymeric scaffolds and liquid species.^{229–232}

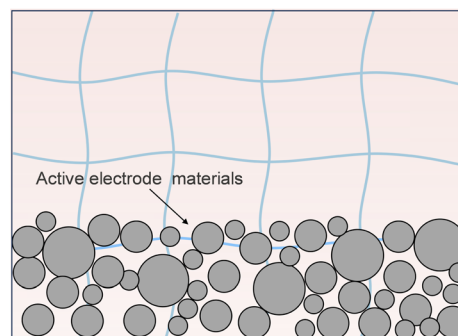
First, the optimization of polymer matrices of GPEs can enhance the interfacial compatibility by introducing adhesive functional groups or creating “conformal” interfacial contacts (Fig. 11(a)).^{233,234} For instance, Li *et al.* reported a adhesive hydrogel electrolyte prepared through a mild free radical copolymerization with sodium lignosulfonate and acrylamide as the monomers. The catechol groups from sodium lignosulfonate strongly interact and adhere well with both the cathode and anode materials, which significantly reduces the contact resistances and improves the affinity between the electrolyte and electrode (Fig. 11(a₁)).²³⁵ In another study, Cao *et al.* demonstrated a cellulose nanofibril-based morphing GPE to improve the interfacial contact, it is mechano-adaptive and beneficial to the formation of interlocked structure on the surface of electrodes, which largely reduces the interfacial contact resistance and promotes uniform nucleation.²³⁶ In addition, chemical bonds form between polar groups in polymer matrices of GPE and electrodes (especially when employing organic electrodes), which enhances the affinity of electrolyte and electrodes. Recently, Shim *et al.* fabricated a GPE for PANI//Zn

a Enhancing interfacial compatibility

a₁) Introducing adhesive functional groups

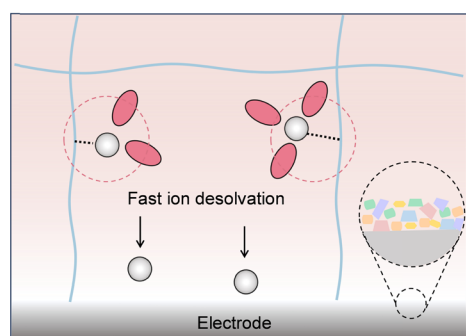


a₂) Creating “conformal” contact



b Accelerating interfacial ion transport

b₁) Regulating ion solvation structure



b₂) Guiding uniform ion transport

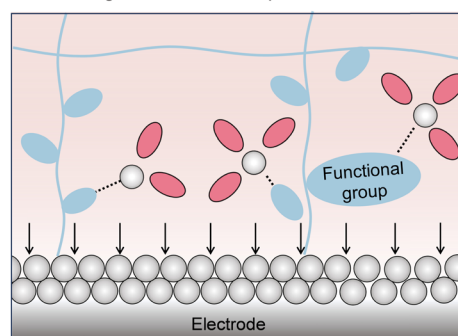


Fig. 11 Influence of polymer matrix on the interfacial chemistry. (a) Enhancing interfacial compatibility by: (a₁) introducing adhesive functional groups; (a₂) creating “conformal” contact. (b) Accelerating interfacial ion transport by: (b₁) regulating the ion solvation structure; (b₂) guiding uniform ion transport.



batteries using methanesulfonic acid and PVA. The intermolecular interactions between the electrolyte and the PANI cathode enable the close contact between the electrolyte and the polyaniline cathode, thus increasing the adhesion of the electrolyte and the electrode. The increased adhesion not only reduces the interfacial impedance but also enhances the interfacial charge transfer performance by increasing the localized sites for ion hopping.²³⁷ Additionally, the enhancement of the interfacial compatibility can be achieved by *in situ* synthesis methods such as UV/thermal polymerization (Fig. 11(a₂)).^{238–240} The polymer electrolyte with compact interfacial contact developed by the *in situ* polymerization of trihydroxymethylpropyl triacrylate (TMPTA) not only provides an integrated 3D network throughout the whole interior of battery with well-connected Na⁺ ion migration pathways but also forms a favorable NaF-rich SEI layer to prevent the sodium metal anode from forming dendrites, endowing the Na//Na_{2/3}Ni_{1/3}Mn_{1/3}Ti_{1/3}O₂ (NMT) batteries with excellent cycling stability in the temperature range from 0 to 60 °C.²³³

Second, the molecularly designed polar groups in the polymer chain were reported to coordinate with target-ions and delicately modulate the solvation structure, thus accelerating the target-ion desolvation and regulating the SEI layer on the surface of the electrode (see Fig. 11(b₁)).^{84,206,229,241} For example, Luo and co-workers developed a GPE composed of 0.5 M NaPF₆/diglyme in polymerized 1,3-dioxolane (poly-DOL). They indicated that the poly-DOL segments in the GPE help to weaken the interaction between Na⁺ ions and diglyme solvents and confine the PF₆⁻ anions. This leads to a local high concentration of Na⁺ ions and significantly speeds up the Na⁺ desolvation process, promoting the formation of a thin inorganic SEI like Na₂O and NaF. These inorganic components in SEI restrain side reactions and provide faster channels for interfacial Na⁺ migration. As a result, the NVP//Na cells presented excellent performance after 1000 cycles with a capacity retention of ~99% at -20 °C, better than the regular liquid electrolyte.²⁴¹ The participation of the polymer matrix in the solvation structure can expel partial active solvent molecules from the solvent sheath, thus reducing the formation of undesirable substances in the SEI layer.^{84,242} In a recent work, Zhou *et al.* found that the abundant functional carbonyl groups (O=C) groups in the polymer chains of the PEGDA-based GPE weakened the interactions between Na⁺ and propylene carbonate (PC) solvent molecules, greatly decreasing the activity of organic PC and inhibiting the parasitic reactions.⁸⁴

Additionally, the abundant charged groups and the unique backbone structure of polymer matrices can provide additional driving force and specific channels to accelerate ion transport (Fig. 11(b₂)).^{202,243,244} Recent studies have demonstrated that some polymer networks possess metal-ionophilic properties to uniformize metal ion flux distributions and constrain two-dimensional diffusion over the surface of anode, thus achieving highly efficient ion migration.^{245–247} For instance, Hu's group developed a negatively charged carboxylate hydrogel electrolyte (gelatin/sodium alginate-acetate), in which the zincophilic carboxyl functional groups and acetate ions guided the directional

migration of hydrated Zn²⁺ ions along polymer chains, stabilizing the interfacial electrochemistry.²⁴⁸ A recent study showed that the abundant negatively charged carbonyl groups in the PAAm chain of a cross-linked PAAm-based GPE could coordinate with Zn ions to improve their distribution, which could restrict the cusp effect and attract more accumulation of Zn salts. As a result, the dendrite growth was suppressed and the operation temperature of the Zn₃V₂O₈//Zn full-cell was extended to -40 to 60 °C.²⁴⁹

As seen in this research direction, previous interest has mainly been focused on the role of polymer matrices on the electrolyte/electrode interfacial compatibility. Future efforts such as molecular and structural engineering of polymer matrices to tailor the solvation/desolvation dynamics of electroactive ions towards better electrochemical performance of batteries are of great significance. It is expected that a thorough understanding and thus control of the interactions between the polymer matrices and ionic/molecular species of GPEs could lead to further breakthroughs in all-temperature batteries.

4.3 (Co-)solvents

For GPEs, the solvent, both water and organic ones, primarily functioned as the liquid plasticizer to reduce the polymer matrix crystallinity, thus enabling good mechanical reliance and significantly improved ionic conductivity. Additionally, it also plays an important role in determining wide-temperature operability, mainly referring to high-temperature thermal/electrochemical stability, freezing tolerance, and SEI stability at both high and low temperatures, by regulating the microstructure (especially solvation structure of target ion) of liquid components confined in the matrix.

For achieving wide-temperature applications, the ideal solvent should satisfy the requirements as follows: (i) good solvation ability to salts; (ii) chemical inertness toward battery components; (iii) wide electrochemical window; and (iv) low viscosity and freezing point to favor ion mobility at low temperature, high boiling point to improve the safety at high temperature. However, as displayed in Fig. 12, the properties of single-solvent electrolytes are limited and do not meet the requirements of wide-temperature or even room-temperature electrolytes. In addition, the complex interfacial chemistry at extreme temperatures further complicates the solvent selection. A common strategy to overcome these challenges and extend the operating temperature range is the addition of cosolvents.

4.3.1 (Co-)solvents toward low-temperature applications. To improve the low-temperature operability, the first thought that comes to the mind of battery researchers is to ensure low freezing point and low viscosity of liquid solution within the polymer matrix. As a very representative example, in early research efforts of lithium battery electrolyte, chain carbonates, such as dimethyl carbonate (DMC), diethyl carbonate (DEC) and ethyl methyl carbonate (EMC), were introduced to efficiently reduce the freezing point and viscosity of cyclic ethylene carbonate (EC)-based and PC-based electrolytes.²⁵¹ Jow *et al.* investigated the thermal behaviors of five liquid/solid binary carbonate mixtures (PC/DMC, PC/DEC, DEC/EC, DEC/DMC, DEC/EMC) and observed that an increasing amount of cosolvent



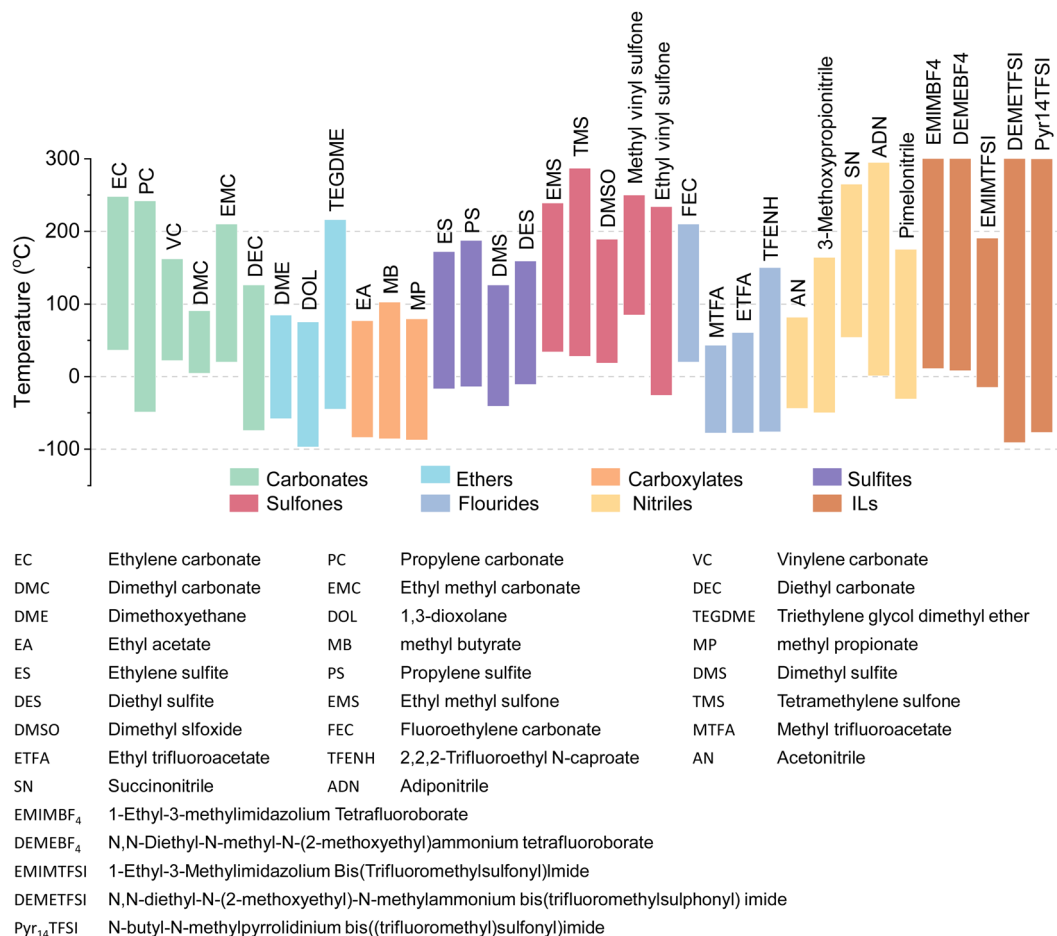


Fig. 12 Summary of freezing points and boiling points of popular solvents used to prepare GPEs. This data could roughly guide the design of GPEs with the desired working temperature range. For example, sulfones and ILs may be applied to stabilize the GPEs toward high temperature applications.

could result in a significant decrease in the freezing point of binary mixtures, expanding the liquid region of binary systems toward low temperature.^{252,253} Smart *et al.* used the binary (EC/DMC, EC/DEC) and ternary cosolvent of EC/DMC/DEC in LiPF₆-based electrolytes to achieve high low-temperature ionic conductivity ($\sim 1 \text{ mS cm}^{-1}$ at -20°C).²⁵⁴ According to the same mechanism, other solvents with low freezing point were also added to the non-aqueous electrolytes as cosolvents for achieving excellent low-temperature performance, including carboxylates, esters, and sulfites.^{255–259} For example, the addition of methyl butyrate (MB) with a low freezing point of -84°C into LiPF₆-based carbonate electrolyte (1.2 M LiPF₆ in EC/EMC) extends the operating temperature of lithium-ion batteries down to -60°C .²⁶⁰ By sharp contrast, traditional electrolytes tend to solidify at temperatures below -20°C . The sulfolane is another effective co-solvent that is able to achieve high ionic conductivities at wide range of temperatures varying from -60 to 55°C . Similarly, the addition of methyl propionate (MP), which freezes at -88°C , greatly enhances low-temperature ion transport.²⁶¹ In a recent work, the poly-DOL-based GPE with MP as the cosolvent presented outstanding ionic conductivity of 1.0 mS cm^{-1} at -30°C , enabling the LFP//Li battery to operate effectively at a low temperature of -20°C .²⁶² It is important to highlight that in

mixed solvent-based electrolyte systems, the electrolyte can attain its minimum freezing point only when a specific ratio of each solvent is attained. Additionally, the interactions between polymers and solvents may influence the interactions among solvents, thus potentially influencing the low-temperature performance of GPEs. Further comprehensive research is warranted to delve into these aspects.

Different from reducing the freezing point of the electrolyte directly, some cosolvents were found to participate in the solvation structure, helping to improve the ion transport kinetics of batteries at low temperatures.^{256,263} In commercial electrolytes, EC has been established as the main constituent in the primary solvation sheath of Li⁺, making EC the major chemical source for the desolvation and formation of SEI. However, the strong affinity toward Li⁺, together with the organic-rich EC-derived SEI, unavoidably hinders Li⁺ desolvation kinetics and slows the migration rate of Li⁺ across the SEI, especially at low temperatures.²⁶⁴ It was reported that the addition of linear DMS or DES as the cosolvent is capable of simultaneously decreasing the freezing point of electrolytes and forming a robust SEI layer on the anode, thus keeping the low irreversible charge losses at low temperatures. Guo *et al.* very recently found that the single-electron reduction of DMS in



baseline electrolyte (1 M LiPF₆ in EC/EMC) facilitated the formation of ionic conductive SEI layer, leading to a low R_{ct} and R_{SEI} in graphite//Li half-cell and retarding the capacity decay of LiNi_{0.5}Co_{0.2}Mn_{0.3}O₂//graphite batteries at $-20\text{ }^{\circ}\text{C}$.²⁶³ Fluorinated solvents, such as fluoroethylene carbonate (FEC), 2,2,2-trifluoro-*N,N*-dimethylacetamide (FDMA) and methyl-3,3,3-trifluoropropanoate (MTFP), have been widely used as low-temperature cosolvents due to their low freezing points and favorable effects of electron-withdrawing fluorine on desolvation and interfacial chemistry.^{182,265,266} A recent study by Xiong *et al.* showed that the Li⁺-dipole interaction strength decreased from -1.90 to -1.66 and -1.44 eV with the increased fluorination degree from EC to FEC and difluoroethylene carbonate (DFEC), respectively. Owing to the lower interaction strength, FEC and DFEC can desolvate from the Li⁺ solvation sheath more easily than EC molecules to produce a LiF-rich SEI layer. As a result, the NCM811//Li batteries with FEC- and DFEC-based electrolytes retain 45% and 51% of room-temperature capacity at $-30\text{ }^{\circ}\text{C}$, respectively, superior to the nearly zero capacity in the EC-based electrolyte.²⁶⁷ He *et al.* fabricated a PEGDA-based GPE containing 1 M LiPF₆ in ethyl propanoate (EP)/EMC/FEC cosolvent. Benefiting from the high ionic conductivity of 1.2 mS cm^{-1} at $-40\text{ }^{\circ}\text{C}$ and LiF-rich CEI and SEI, the LFP//Li batteries show a high discharge capacity (72.1% of the room temperature capacity) even at a low temperature of $-40\text{ }^{\circ}\text{C}$.^{268,269} Guo *et al.* demonstrated a NCM811//Li battery with a low-temperature adaptable GPE consisting of FDMA and FEC as the cosolvent, LiDFOB as the salt, and poly(1,3,5-trioxane) as the polymer matrix.¹⁷ The implementation of this design allows the stable cycling of NCM811//Li batteries at $-30\text{ }^{\circ}\text{C}$. This impressive behavior is mainly attributed to the formation of a Li_xBO_yF_z/LiF-rich and Li₂CO₃-poor SEI with a low energy barrier for Li⁺ diffusion. At the same time, the formed amorphous CEI helps to stabilize the NCM811 cathode from the dissolution of transition metal and structural cracking.

The cosolvent strategy is also found useful in improving the freezing tolerance and ionic conductivity of hydrogel electrolytes. Organic cosolvents commonly used in hydrogel electrolytes include nitriles, esters, and amides.^{270,271} For example, using the low-freezing point DMF ($-60.5\text{ }^{\circ}\text{C}$) as a cosolvent, the freezing point of the ZnSO₄/DMF/H₂O electrolyte drops down to $-32\text{ }^{\circ}\text{C}$, lower than that of the ZnSO₄/H₂O electrolyte ($-11\text{ }^{\circ}\text{C}$).²⁷² Similarly, Chen *et al.* prepared a PVA-based hydrogel using an IL/water (EMImAc/H₂O) binary solvent. The freezing points of the prepared hydrogels ($-65\text{ }^{\circ}\text{C}$) were much lower than those of pure water and EMImAc ($-20\text{ }^{\circ}\text{C}$) due to the strong interactions between EMImAc and H₂O.²⁷³ Unlike aprotic organic solvents, water molecules can be used as both electron acceptors and electron donors. Therefore, another strategy to weaken the water-water interaction and improve the low-temperature performance of aqueous electrolytes is to add cosolvents such as dimethyl sulfoxide (DMSO) and polyol solvents.²⁷⁴⁻²⁷⁶ When polyol cosolvents are added to the electrolyte, the strong interactions of -OH groups of polyol solvents with water molecules through hydrogen bonding severely impair the interactions among water molecules, thus inhibiting

the ice crystallization and lowering the freezing temperature. At the same time, the attraction of polyol molecules to H₂O could reduce the solvation degree of Zn²⁺-H₂O, which is beneficial to ion migration in the electrolytes. For example, Li *et al.* added a low freezing point EG cosolvent to water (EG:H₂O = 3:2 by volume), reducing the freezing point of the electrolyte to $-33\text{ }^{\circ}\text{C}$.²⁷⁷ DFT calculation results showed stronger hydrogen bonding interactions of EG-H₂O ($10.40\text{ kcal mol}^{-1}$) than H₂O-H₂O ($5.78\text{ kcal mol}^{-1}$), indicating the capability of EG in locking free water and inhibiting freezing.²⁷⁸ In GPE systems, the polyol cosolvents can assist the PAM polymer network to break the hydrogen bond networks of water, endowing PAM-based hydrogel electrolyte with excellent anti-freezing ability down to $-40\text{ }^{\circ}\text{C}$.²⁷⁹ In addition to polyols, DMSO as a cosolvent has also been extensively evaluated in hydrogel electrolytes to improve their low-temperature operability. For example, Li *et al.* designed a DMSO-containing PAAM/agar gel electrolyte that could operate at $-30\text{ }^{\circ}\text{C}$.²⁸⁰ A recent study by Liu *et al.* demonstrated that DMSO in GPE could stabilize the Zn²⁺ solvation structure and reduce the proportion of networked water, inhibiting the freezing of the hydrogel electrolyte and lowering the operating temperature of the Zn battery to $-20\text{ }^{\circ}\text{C}$.²⁸¹

Recently, liquid electrolytes using high-entropy solvents have been recognized as a potential solution for low-temperature electrolytes. By increasing the solvent molecular diversity to modulate the solvation entropy and tune the mesoscopic solvation structure, the operating temperature of the liquid electrolyte could be extended to extremely low temperatures.²⁸² For example, Chen *et al.* configured a series of high-entropy solvent-based electrolytes. They indicated that the freezing point of electrolytes decreased greatly with an increase in the number of solvents from binary to decimal. The high entropy of solvents endows the high disorder degree of mixtures, making the EC solvent molecules to nearly completely separate from each other and largely reducing the viscosity of the electrolyte. As a result, the decimal high-entropy EC-based electrolyte (EC/DEC/PC/EMC/EP/EA/MB/BA/MP/PB) showed an ultralow freezing temperature of $-130\text{ }^{\circ}\text{C}$. Moreover, the electrolyte achieved a remarkably high ionic conductivity of 0.62 mS cm^{-1} at $-60\text{ }^{\circ}\text{C}$ and excellent electrochemical performance of the LMO//LTO batteries at $-40\text{ }^{\circ}\text{C}$, which was far superior to that of commercial counterparts.²⁸³

Obviously, the development of high-entropy electrolytes with more than five organic solvents is of great research value for both low-temperature liquid electrolytes and GPEs. However, the precise selection and optimization of solvent combinations from the vast number of possible solvents remains challenging. Moreover, the complexity of high-entropy solvents increases the difficulty of understanding the physical and chemical interactions within the electrolyte system, not to mention at extreme temperatures. In addition, the introduction of multiple solvents into the GPE systems may lead to potential compatibility issues or unwanted side reactions. It is expected that these issues will be well addressed prior to the low temperature applications of GPEs with high-entropy solvents.



4.3.2 (Co-)solvents toward high-temperature applications.

Compared to achieving low-temperature performance, for nonaqueous battery systems, improving their upper operating temperature limit is to some extent easier due to the wide choice of thermally/electrochemically stable electrolyte components, including but not limited to sulfites, dinitriles, ILs, and their mixtures. Among them, ILs, characterized by a unique combination of near-zero vapor pressure, excellent thermal/electrochemical stability, and non-flammability, are found in a majority of gel polymer studies that are devoted to realizing the high temperature applications of rechargeable batteries. In the exploration of battery systems capable of being operated at temperatures above 100 °C, GPEs immobilized by pure IL electrolytes (usually referred to as ionogels or ion gel electrolytes) are almost the only choice.^{284–287} For example, 1-butyl-1-methyl-pyrrolidinium bis(trifluoromethyl-sulfonyl)imide (P14TFSI) was adopted to prepare a PVDF-HFP-based ionogel electrolyte. The resulting ionogel electrolyte maintains stable up to 350 °C, allowing the LFP//LTO batteries to stably operate at a high temperature of 100 °C.²⁸⁸ Despite these achievements, the high viscosity nature of IL electrolytes compared to conventional organic ones makes it challenging to prepare ionogel electrolytes having a room-temperature ionic conductivity of the 10^{-3} mS cm⁻¹ level. Hence, to date, batteries using ionogel electrolytes with satisfactory performance at moderate and low temperatures are rarely reported. The addition of small-molecule cosolvents to IL electrolytes was proposed for combining the advantages of small-molecule organics to improve the transport properties at low temperatures and the thermal stability of ILs at elevated temperatures so as to realize wide-temperature applications. It was reported that the electrolyte obtained by combining PC with PYR₁₄TFSI cosolvent has a lower viscosity and higher ionic conductivity than that of pure PYR₁₄TFSI-based electrolyte in the temperature range between -40 °C and 100 °C.²⁸⁹ However, this approach deviates from the original intention of using ILs for security design.

Despite their less pronounced stability, molecular-type solvents of sulfites and dinitriles have also been widely applied.²⁹⁰ For example, sulfolane, one of the most common sulfites, was successfully applied to prepare a PVDF/PEO-based GPE, which remarkably enhanced the cycling and safety performances of LMBs. When used in the Li//LCO battery, the sulfolane-containing electrolyte enables long-term cycling stability at high temperatures of up to 90 °C. Moreover, the full-charged Li//LCO pouch cells exhibited a higher onset temperature of thermal runaway at 190 °C, 60 °C higher than their commercial counterpart of LiPF₆//EC/DMC/EMC.²⁹¹ Succinonitrile (boiling point: 266 °C), a representative dinitrile, has a similar function to sulfolane. It has been reported that the PEO-based GPE plasticized with succinonitrile achieved a superior high-temperature-tolerance characteristic up to 170 °C.²⁹² At a high temperature of 55 °C, the LCO//Li batteries using succinonitrile/LiTFSI-plasticized polyurethane acrylate GPE demonstrated improved cycling stability, compared to the one plasticized with liquid electrolyte (LiPF₆ in EC/DMC).²⁹³

Despite the high boiling points of these solvents, their limited salt solubility and low reductive stability pose

challenges in exceeding the upper temperature limits for batteries using them as plasticizers in GPEs. The addition of cosolvents, such as carbonates, and carboxylates, can help to solve these issues. Laskovic reported a PVDF-HFP-based GPEs gelled by liquids of LiTFSI in PC/isobutyronitrile/trimethyl acetonitrile/ethylene sulfate (ES)/vinylene carbonate (VC). Using ES and VC as SEI forming agents and employing EC and PC to dissolve LiTFSI, along with isobutyronitrile and trimethyl acetonitrile to enhance the thermal stability at high temperatures, PVDF-HFP-based GPEs demonstrated high thermal stability and improved cycling performance in NMC622//graphite full cells, even at 60 °C.²⁹⁴ In particular, fluorinated solvents have been extensively utilized as high-temperature cosolvents due to their ability to promote the formation of thermally stable LiF-rich SEI/CEI layers. Deng *et al.* introduced a fluorinated ether co-solvent (1,1,1-trifluoro-2-[(2,2,2-trifluoroethoxy) methoxy]ethane (TTME)) to the conventional ether electrolyte system comprising DME and LiFSI to form a fluorinated ether-based electrolyte. TTME not only helps to generate a stable SEI on the surface of graphite and to inhibit the electrolyte decomposition but also improves the overall oxidation stability of the electrolyte to prevent the degradation of the electrolyte on the surface of the LCO cathode. As a result, the LCO//graphite batteries performed much better in terms of cycling performance than their counterparts using carbonate electrolyte at elevated temperatures.²⁹⁵

Compared to non-aqueous electrolytes, achieving high-temperature applications with aqueous electrolytes poses more challenges. The lower boiling point of water, in comparison to most non-aqueous solvents, leads to the evaporation of water solvent and increased water activity in aqueous electrolyte systems operating at elevated temperatures. This can result in significant electrolyte loss and decomposition at the electrolyte/electrode interfaces. The addition of high boiling point cosolvents in the plasticizer of hydrogel electrolytes can improve their thermal stability and inhibit their interfacial side reactions. The main cosolvents of high-temperature aqueous electrolytes include amides, sulphones, and polyols. These cosolvents not only possess high boiling point values but can also manipulate the solvation structure and interfacial chemistry, inhibiting the evaporation of water inside the electrolyte and suppressing the interfacial side reactions, thus improving the high-temperature performance of electrolytes and batteries.^{296–298} Xiong *et al.* modified the ZnSO₄ aqueous electrolyte with DMF (boiling point: 153 °C). They demonstrated that the DMF can intensify the hydrogen bonds of DMF-H₂O and weaken the solvation interaction of Zn²⁺ with H₂O, facilitating the *in situ* formation of a Zn²⁺-conducting SEI layer (Zn₅(CO₃)₂(OH)₆) on the electrode surface during the cycling process. The formed SEI layer, featuring a high thermal decomposition temperature of 250 °C, enables the electrolyte to achieve outstanding performances and stabilities at high temperatures of 70 °C.²⁷² The sulphones (*e.g.*, DMSO) and polyols (*e.g.*, glycol, glycerol, *etc.*) have been proven to be able to form hydrogen bonding with water molecules and suppress the water activity, mitigating the electrolyte decomposition at the electrolyte/electrode interfaces at high



temperatures.²⁴⁹ It is foreseeable that the high temperature performance of aqueous electrolytes can be improved by increasing the content of organic co-solvents to water. However, this approach would defeat the original purpose of aqueous batteries as low cost, safe and environmentally friendly; thus, the ratio of organic co-solvents to water should be carefully adjusted.

For this consideration, the incorporation of ILs into hydrogel electrolytes appears to be a promising approach for high-temperature aqueous batteries as it combines the benefits of the high-temperature stability of ILs while preserving the advantages of aqueous electrolytes. However, the use of ILs in aqueous battery systems has been limited due to their thermodynamic immiscibility with water and poor salt solubility. Nevertheless, a recent study by Yu *et al.* has demonstrated a breakthrough in this regard. They developed an electrolyte consisting of a water-immiscible ionic liquid diluent, specifically 1-ethyl-3-methylimidazolium bis(fluorosulfonyl)amide (EmimFSI), in water. Remarkably, this electrolyte enabled the operation of $\text{Zn}_{0.25}\text{V}_2\text{O}_5 \cdot n\text{H}_2\text{O} // \text{Zn}$ cells at a temperature as high as 100 °C.²⁹⁹ This work opens new possibilities for the high-temperature utilization of ILs in aqueous battery systems regardless of liquid electrolytes or GPEs.

4.4 Additives

Unlike the salts and co-solvents, which are the dominant constituents of the electrolytes, additives are generally characterized by their small dosage but “quick-effect” nature. Due to their unique roles in regulating the ion transport behaviors and in improving the stability of electrolytes and the electrolyte/electrode interfaces, numerous types of additives are employed to improve the low- and/or high-temperature performance of GPEs and thus the batteries. In accordance with their respective

roles under different temperature ranges, the functions of the additives are summarized in Fig. 13. In the case of low-temperature operation scenario, additives are strategically incorporated to (i) increase ionic conductivity as solubilizers; (ii) accelerate the rate-determining desolvation process as solvation tailors; and (iii) assist in the formation of thin and ion-conducting SEI as interface modifiers. Under high-temperature conditions, the additive engineering of GPEs shifts mainly to (i) improving the thermal stability as a thermal tolerance enhancer; (ii) protecting the GPE from flame as fire preventers; and (iii) helping to form thermostable SEI at the GPE/electrode interfaces as interfacial protectors.

4.4.1 Organic additives. Carbonates, sulfides, fluorides, sulfones, ILs, *etc.*, are the most-widely used organic additives to improve the low-temperature and/or high-temperature performance.^{260,301–303}

At the low-temperature end, although some organic additives with low freezing point and low viscosity can inhibit the solvent solidification or freezing of liquid solution and improve the ionic conductivity of GPE to a certain extent, considering their small dosages in the whole GPE systems, researchers are currently focusing more on their functions in regulating the interfacial chemistry of batteries.³⁰⁴ The organic additive can participate in the regulation of the ion solvation structure and formation of interfaces and interphases on the surface of electrodes, reducing the barrier of the desolvation process and across the SEI/CEI layers and eventually facilitating the ion-transport kinetics.^{305–307} Some organic solvents, such as VC, ES, AN, EC, and FEC, can function as film-forming additives to create interfacial layers with low ion transport barriers between the electrolyte and electrode.^{302,308,309} For example, the co-addition of 2 wt% ES and 2 wt% VC in a PVDF-HFP-based

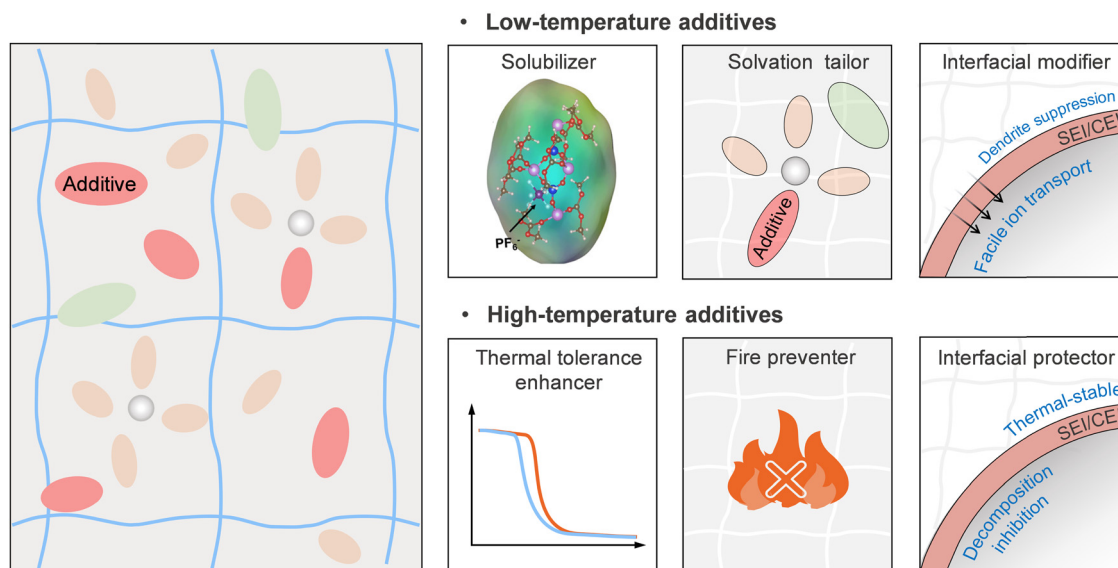


Fig. 13 Effects of additives on extending the operating temperature of GPEs. Additives can function as solubilizer, solvation tailor, and interfacial modifier to ensure the low-temperature battery performance, while in the cases aiming at high-temperature operability, they usually act to improve the thermal stability, fire resistance and structural stability of the solid electrolyte interphase. Solubilizer image adapted from ref. 300 with permission from John Wiley & Sons, Copyright 2022.



GPE with PC as the solvent contributes to the formation of a stable SEI, making the charge transfer and SEI impedances independent of the cycling process. This formulation achieved a remarkable cycle life for the NMC532/graphite cells.³¹⁰ At low temperatures, however, VC leads to a thicker SEI with a low lithium ion conductivity, decreasing the discharge capacity.³¹¹ FEC, as an alternative, has been investigated in detail as a film-forming additive to create a LiF-rich SEI with superior low-temperature performance and improve the ionic conductivity of the electrolytes at low temperatures.^{307,312,313} In a recent study, Liao *et al.* found that the addition of 2 wt% FEC in the liquid electrolyte could increase the discharge capacity and CE by forming SEI layers on the graphite anode and LFP cathode at $-40\text{ }^{\circ}\text{C}$.^{314,315} Moreover, Ciucci *et al.* demonstrated that FEC additive in a poly-DOL-based GPE participated in the first Li^+ solvation sheath and reduced the binding energy of MP solvents with Li^+ , resulting in easier Li^+ transport. At the same time, the presence of FEC induced the formation of fluorine-rich interphases, contributing to enhanced interfacial stability. Li//LFP and high-voltage Li//NCM811 batteries demonstrate excellent long-term stability (>2000 cycles at room temperature) and can operate effectively at a low temperature of $-30\text{ }^{\circ}\text{C}$.³¹⁶ Interestingly, the introduction of FEC is reported to be able to regulate the Li deposition morphologies on the Li anode from porous loosened deposits with dendritic structure into large particles with a dense structure at a low temperature of $-40\text{ }^{\circ}\text{C}$,³¹⁷ which provides a valuable solution for dendrite suppression of low-temperature adaptable alkali-metal batteries. These findings certainly motivate the exploration of more novel F-containing compounds.

At the high-temperature end, organic additives can also enhance the battery's thermal tolerance by forming a protective yet ion-conductive layer on the surfaces of both the cathode and anode. For example, the unsaturated organic compound of VC can be readily reduced to form a SEI layer rich in polymeric components *via* a ring-opening reaction at high temperatures. Its effectiveness has been proven in several battery systems including lithium-ion, lithium-sulfur, sodium-ion, and potassium-ion batteries.^{318,319} In addition, fluorides, such as FEC, can form LiF components on the anode and polyether species on the cathode, protecting the electrolyte from degradation, thus improving the reversibility of the battery at high temperature.³²⁰ Although some electrolytes discussed above could be operated at a high temperature, most of them contain flammable components, which may induce safety issues at abnormal temperatures. Flame retardants, such as organic phosphorus (*e.g.*, triethyl phosphate (TEP), triphenyl phosphate (TPP), and trimethyl phosphate (TMP)) and fluorinated compounds (*e.g.*, tris(pentafluorophenyl)borane, TB), can reduce the flammability of the electrolyte and passivate the electrodes by forming thermally stable SEI/CEI layers simultaneously, preventing possible hazards resulting from the thermal runaway of batteries.^{121,262,321–323} For example, TMP can release phosphorus-containing free radicals PO^{\bullet} and PO_2^{\bullet} to prevent the chain reaction in batteries by capturing the H^{\bullet} and HO^{\bullet} radicals released from the burning composition.^{318,324} In an attempt to develop a safe and wide-temperature operable quasi-solid-state Li-S battery, TB

additive was introduced to improve the flame retardance *via* decomposing and generating fluorine radicals to trap the highly reactive radical released by the GPE. At the same time, the TB with high fluoride content contributes to the formation of a highly stable and resilient LiF-rich SEI layer. With these merits, Li-S batteries assembled with this electrolyte demonstrated stable cycling over a wide temperature range from -20 to $50\text{ }^{\circ}\text{C}$.³²³

Similarly, for hydrogel electrolytes, the low-temperature performance was reported to be improved by adding organic additives such as polyols, sorbitol, and ILs.^{325,326} These additives feature high solubility in water, and the ability to strongly interact with water molecules *via* hydrogen bonding can improve the low-temperature ionic conductivities of hydrogel electrolytes. For instance, Chen *et al.* reported that the DMSO additive in PAAM-based GPEs could form abundant hydrogen bonds with both the functional groups of the polymer matrix and water molecules, enabling the formation of optimized GPEs with a high ionic conductivity of 0.82 S m^{-1} at $-20\text{ }^{\circ}\text{C}$, comparable to that at room temperature.³²⁷ Moreover, at the electrolyte/electrode interfaces, the organic additive molecules are able to participate in the solvation structure and affect the interfacial chemistry through the interactions with target ions.^{281,304} A study by Hou *et al.* demonstrated that the DMSO additive in a PAM-based hydrogel electrolyte could coordinate with Zn^{2+} in the solvation structure to reduce the $\text{Zn}^{2+}\text{-O}(\text{H}_2\text{O})$ pair, lowering the energy barrier for Zn^{2+} desolvation and improving the reversibility of Zn plating/stripping at low temperatures.^{281,319,328} At elevated temperatures, the organic additives were demonstrated to be able to inhibit the water decomposition *via* hydrogen evolution reaction and oxygen evolution reaction at the surface of discharged electrodes.^{329,330} A study by Hu *et al.* proposed that the addition of glycol and AN additive could adjust the affinity between H_2O and oxygen-containing groups in the PAM-based GPE, enabling high ionic conductivity at low temperatures and reduced H_2O reactivity at high temperatures. The GPE exhibited good anti-freezing properties at $-20\text{ }^{\circ}\text{C}$ and a slow water evaporating rate at $60\text{ }^{\circ}\text{C}$, enabling a high-performance Zn// V_2O_5 full cell that can cycle stably in a wide temperature range from -20 to $60\text{ }^{\circ}\text{C}$.²⁷⁴

Compared to the relatively limited choice of solvents/liquid plasticizers, it remains highly challenging to select suitable additives from the vast chemical space of possible compounds. Despite many successful additives for various battery chemistries, traditional trial and error experiments for additive searching is still far from efficient. Fortunately, the fast development of data-driven quantitative structure–property relationship models can substantially accelerate the findings of additives, promoting more advancements in the development of GPE-based batteries with wide temperature operability.

4.4.2 Auxiliary salt additives. In addition to organic solvents, organic or inorganic metal-ion salts are utilized as functional additives in GPEs for wide-temperature applications, which have a significant impact on the low-temperature and high-temperature performance of electrolytes by altering the degree of dissociation of anions and the ability of SEI formation.^{302,331,332}



Many salts, such as LiBOB, LiDFOB, and LiPO_2F_2 , have been investigated as auxiliary additives due to their film-forming ability.^{145,333,334} These salts can be sacrificially reduced or oxidized at the anode or cathode surface prior to solvent decomposition, thereby forming low-resistive and stable SEI/CEI layers composed of inorganic-rich components. For example, LiBOB can be electrochemically reduced to form a low-resistive SEI film at low temperatures. When cooperated with 4-vinyl-1,3-dioxolan-2-one (VEC) additive in a PVDF-HFP-based GPE, a robust and highly ionic-conductive SEI film can be built up to enhance the electrolyte/electrode interfacial stability, enabling the superior cycling performances of LMO+ $\text{LiMn}_{0.7}\text{Fe}_{0.3}\text{PO}_4$ (LMFP)//graphite battery at temperatures ranging from -18 to 45 °C.¹⁷¹ In addition to LiBOB, LiDFOB was also recognized as an effective auxiliary additive. With the contribution of LiDFOB decomposition to form low-resistance and stable F-rich and B-containing SEI film, the PVA-based GPE with LiDFOB presented lower capacity fading at low temperatures.³³⁵ F-containing metal-ion salt additives are especially attractive for constructing stable electrolyte/electrode interfaces/interphases. Their decomposition products, such as LiF and NaF, are believed to be able to reduce the interfacial resistance and improve the toughness of the SEI/CEI layers, therefore facilitating the interfacial ion transport and stability and boosting the wide-temperature performance.^{336–338} In a recent work by Cui and co-workers, 0.05 M LiPO_2F_2 was added to LiTFSI/lithium trifluoro(perfluoro-*tert*-butyloxy)borate (LiTFPB) dual salt-based GPE, dramatically improving the cyclability and rate capability of an $\text{LiNi}_{0.5}\text{Mn}_{0.3}\text{Co}_{0.2}\text{O}_2$ //Li battery over an impressively wide temperature range from -40 to 90 °C. LiPO_2F_2 plays a dominant role in forming an F-rich SEI with both low impedance and high-temperature durability. In particular, the inorganic Li-containing species (especially LiF and Li_2O , ionic conductive) and P–O species at the outer shell of the SEI layer can suppress Li dendrite growth and their reactions with carbonate solvents.¹⁴⁷ These mechanisms have also been proved *via* the construction of N-rich or S-rich or B-containing SEI/CEI layers.^{335,339–341}

Recently, nitrate salts (*e.g.*, LiNO_3 and NaNO_3) have also been used as auxiliary additives to improve the wide-temperature tolerance of GPEs. Archer *et al.*, for example, reported that the addition of LiNO_3 in PEG-based GPEs showed a high ionic conductivity of >1 mS cm^{-1} even at -30 °C, making it suitable for low-temperature batteries without any compromises in power density.³⁴² Furthermore, they presented that the physical interactions between the electron-lacking N-atom in LiNO_3 additive and DOL solvents could distort bonds in DOL and coupled motions of individual molecules, thus arresting the liquid to crystalline solid thermal transition and making the electrolyte preserve abnormally high bulk and interfacial ionic conductivities at temperatures down to -50 °C.³⁴³ Furthermore, the LiNO_3 -containing electrolyte can facilitate the formation of dendrite-free Li deposition.³⁴⁴ Additionally, it was reported that the addition of a small dosage of Lewis acidic salts (*e.g.*, $\text{Al}(\text{OTf})_3$, $\text{Mg}(\text{OTf})_2$, LiPF_6 , and LiDFOB) or halides (*e.g.*, AlI_3 or AlF_3) to a baseline electrolyte can initiate

the *in situ* polymerization of monomers inside a battery cell without the addition of initiators, enabling the facile preparation of GPEs with high thermal, mechanical, and electrochemical stability. Because the formed GPEs originated from liquids are able to wet all components of the cell, “conformal” interfacial contact could be realized and thus the low interfacial resistance and homogeneous interfacial charge transfer, which represents a promising approach for constructing GPEs and battery systems that are adaptable over a wide range of temperatures.^{164,238,345–347}

For aqueous GPEs, the introduction of soluble ions into water can lower the freezing point of electrolytes according to the colligative properties observed in dilute solutions. For example, the freezing point of water can be significantly lowered by the dissolution of inorganic salts such as CaCl_2 , LiCl, ZnSO_4 , or organic zwitterions.²⁵⁰ Chen’s group introduced sodium citrate (SC) into PAM-based hydrogel electrolyte. The intensive interactions between the hydrophilic $-\text{COO}^-$ functional groups of SC molecules with H_2O molecules facilitate the localization of water molecules in hydrogels, preventing water from freezing and evaporation. The electrolyte delivers a high ionic conductivity of 73.96 mS cm^{-1} even at a low temperature of -40 °C as well as high water retention of 96.85% after 96 h exposure, both of which are substantially higher than those of the control samples. As a result, high-performance zinc–air batteries with PAM-SC hydrogel electrolyte can be achieved over a wide temperature range from -40 to 40 °C.²⁵⁰

4.4.3 Inorganic fillers. Recently, the integration of inorganic fillers within GPEs is adopted as an effective approach to enhance ionic conduction and improve thermal stability in a wide temperature range but also maintain good electrochemical/chemical stability and compatibility with electrodes. These inorganic fillers, once integrated into the GPEs, can interact with polar species in electrolytes, providing additional sites or pathways for ion migration. This leads to improved ionic conductivity, accelerated charge transfer kinetics, and enhanced interfacial stability.^{348,349}

At low temperatures, inorganic fillers are reported to interact with ion species and ease the diffusion barrier.^{350,351} Zhang *et al.* incorporated an $-\text{NH}_2$ -functionalized vermiculite (Vr- NH_2) as fillers into GPE, in which IL was confined in the regular interlayer nanochannels of the lamellar Vr- NH_2 framework. The fine-tuning of the microstructure can trigger the rearrangement and crystallinity of ILs in nanochannels, leading to the formation of continuous and densely-aligned nanocrystalline IL layers and fast lithium-ion hopping transport along the 2D channels (Fig. 14(a)–(c)). With such unique structural merits, the developed electrolyte delivered a high ionic conductivity of 0.09 to 1.35 mS cm^{-1} over a wide temperature range from -40 to 100 °C. As a result, the LiFePO_4 //Li pouch cells presented an excellent wide-temperature electrochemical performance (93.8 and 45.0 mA h g^{-1} at 60 and -20 °C with capacity retention of 97% and 98% after 50 cycles, respectively).³⁵² Recently, Ren and coworkers utilized the graphene oxide quantum dots (GOQDs) as an additive of aqueous electrolyte. They found that GOQDs strongly adsorb on the ice crystal surface to inhibit their growth, thus effectively inhibiting the growth of ice



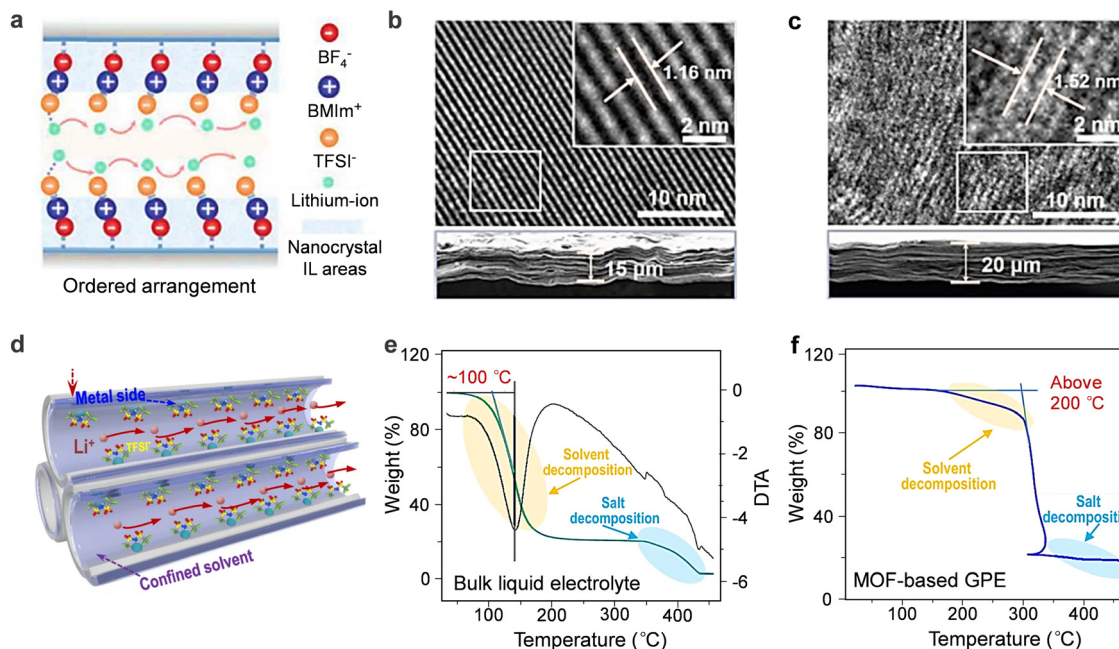


Fig. 14 Compositing GPEs with inorganic additives to extend the operating temperature limits. (a) Schematic illustration of the Li⁺ transport mechanism along the 2D interlayer channel of the ionogel electrolyte with Vr-NH₂ as the host; cross-sectional SEM and HR-TEM images (insets) of (b) the Vr-NH₂ framework and (c) L-ILCE-NH₂.³⁵² Reproduced from ref. 352 with permission from John Wiley & Sons, Copyright 2023. (d) Schematic illustration of the function of MOF filler at high temperature; TGA curves of (e) the typical liquid electrolyte (1 M LiTFSI-PC) and (f) GPE with MOF as the filler.³⁵³ Reproduced from ref. 353 with permission from Springer Nature, Copyright 2022.

crystals and expanding the interconnected liquid regions to offer more spacious ion-transport channels, resulting in improved cell capacity and cycling stability at low temperatures.

At high temperatures, the addition of inorganic fillers can improve the thermal stability of GPEs by confining the liquid components. For instance, Zhou and coworkers used MOF (CuBTC-PSS, 6.5 Å) that possesses a nanoporous (sub-nanoporous) structure as the host to fabricate the gel electrolyte. Due to the physical sub-nanoconfinement and chemical coordination with the MOF host (Fig. 14(d)), the solvents in GPE are unlikely to evaporate and decompose at the electrolyte/electrode interfaces. As a result, the decomposition temperature (highlighted in yellow) of the liquid solvent within the GPE began to decompose at nearly 200 °C, over 100 °C higher than that of the LE (Fig. 14(e) and (f)). Furthermore, the high-voltage NCM811/Li pouch cells using this GPE with a high NCM811 mass loading of 20 mg cm⁻² delivered highly stable electrochemical performances with a high capacity of 171 mA h g⁻¹ and 89% capacity retention after 300 cycles, even at a high working temperature of 90 °C.³⁵³

4.5 Hosts

Usually, the pursuit of mechanical strength and structural/dimensional stability of GPE is realized by increasing the mass ratio of the polymer matrix, which would unavoidably sacrifice the ion transport properties. Moreover, even with this strategy, the mechanical performance cannot always satisfy the requirements of GPE towards practical applications due to the theoretically limited strength and thermal stability of the molecularly dispersed polymer network, especially at extreme temperature

conditions. With this consideration, the incorporation of host materials that are swelling-resistant against water/organic solvent has been considered an effective approach of counter-balancing the plasticizing effect. Familiar hosts include cellulose membrane, glass fiber filter, silica aerogel, and polymer nonwovens, which are typically in the form of interwoven structured or 3D porous membranes.^{354,355} In addition to good swelling resistance, they share similar advantageous properties of high mechanical strength reaching the 10² MPa level and excellent thermal stability higher than 300 °C. Therefore, their use can dramatically improve the mechanical performance of GPE mainly in terms of tensile strength, Young's modulus and thermal/dimensional stability, preventing short-circuit caused by dendrite growth and thermal shrinkage/cold contraction.^{356–358}

4.5.1 Improvement of thermal/mechanical stability. From the above discussion, it is clear that improving the mechanical strength and thermal/dimensional stability is the intrinsic benefit of incorporating hosts. In addition, its structural flexibility warrants very good electrolyte/electrode contact of the resulting GPE like free-standing ones.^{359–361} The utilization of an alien matrix to reinforce GPE could be traced back to as early as 1985, in which Abraham *et al.* encapsulated plasticized polymer electrolytes into the pores of commercial Celgard membrane to prepare ultrathin and mechanically strong GPEs. However, it could be anticipated that the thermal and dimensional stability would not be improved due to the intrinsically poor structural stability against thermal degradation. A wide variety of thermally stable hosts were developed soon afterwards, specifically including various porous polymer membranes prepared through a phase inversion method,



electrospun polymer membranes (PVDF, PAN, PVDF-HFP, *etc.*), bio-based nanofibrous membranes (cellulose and their derivatives), and inorganic ones represented by glass fiber. In the quasi-solid/solid electrolyte research community, the preparation of mechanically strong GPE by incorporating a porous membrane is also widely recognized as a “rigid-flexible” strategy, which shares the same wisdoms with “Tai-Chi”. In such GPE systems, the “rigid” supporting membrane provides high mechanical strength and thermal/dimensional stability, while the ion transport properties are closely related to the “soft” swollen gel phase. Despite being studied for more than two decades, this strategy currently arouses popular interest from academic studies to industrial applications.

4.5.2 Enhancement in electro-/chemical stability. Apart from providing a mechanical supporting skeleton to enhance the physical stability of the GPEs, the electro-/chemical properties can also be affected once the polar species is immobilized in the host matrix since the intermolecular interactions occur between the hosts and the hybrid or/and polymers adapt to them.³⁶² Typically, the inorganic materials have received much attention because of their strong chemical adsorption of negative species in the electrolyte, such as anions, and some organic species dissolution from cathodes.^{213,363–367} The passive (such as SiO₂, Al₂O₃, ZrO₂, *etc.*) and active oxides (such as LISICON, NASICON, garnet, *etc.*) inorganic hosts can attract the anions of metal salts and elevate the t^+ value by Lewis acid-basic interactions, which plays a vital role in optimizing the diffusion,

deposition, and reaction behaviors of metal ions in rechargeable batteries in a wide temperature range.^{368,369} For example, Li *et al.* demonstrated that the GPE prepared by immobilizing IL ([EMIM][TFSI]) in the nanopores of HKUST-1 MOF, presenting ion conductivities and Li⁺ transference numbers of 0.68×10^{-4} S cm⁻¹ and 0.46 at 25 °C, and 6.85×10^{-4} S cm⁻¹ and 0.68 at 100 °C, respectively. The improved Li⁺ transference number has long been demonstrated to help to reduce the concentration polymerization; when exposed long-term to extremely thermal environments, the GPE with MOF hosts demonstrated high CE over 92% and long cycling stability.³⁷⁰ In another example, the oxygen-containing functional groups (*e.g.*, -COOH and -OH) of GO could form F–H hydrogen bonds with fluorine in PF₆⁻ ions, attracting the PF₆⁻ ions and accelerating Li⁺ transport at low temperatures, allowing the application of GPE in the LCO//graphite cell at a low temperature of -15 °C (with 92.7% of its room-temperature capacity).³⁷¹

Thanks to the integrated 3D hosts, which offer continuous paths for ion transport, the GPEs can deliver enhanced ionic conductivity across a broad temperature spectrum.^{372–374} Lu's group fabricated a PVDF-HFP-based GPE with a uniformly cross-linked β''-Al₂O₃ nanowire as the host (ANs-GPE) (Fig. 15(a)). They revealed that the electrolyte can suppress the side reactions between the sodium metal and liquid electrolyte, as well as sodium dendrite formation even at an elevated temperature of 60 °C. They attributed this to the effective immobilization of the liquid electrolyte by the

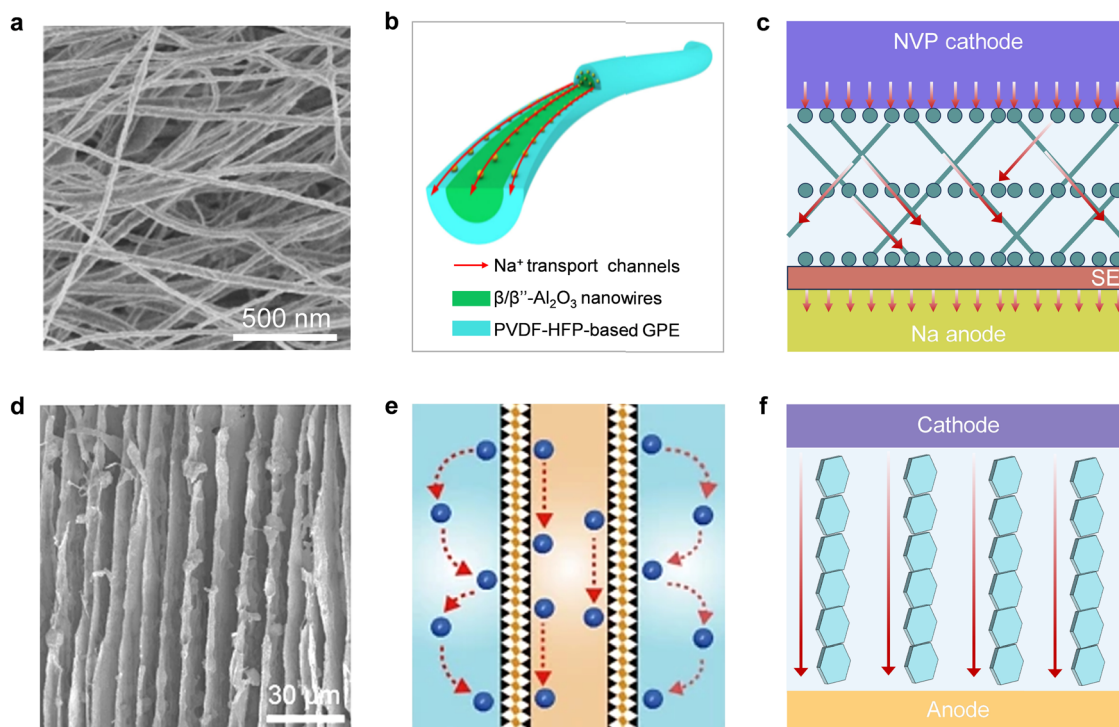


Fig. 15 Regulating ion transport behaviors in GPEs by incorporating functional hosts. (a) Planar SEM image of GPE with β''-Al₂O₃ nanowires as the host; (b) schematic diagram of the Na⁺ transportation path along the nanowire; (c) working mechanism of NVP//Na batteries with the ANs-GPE.³⁷⁵ Reproduced from ref. 375, Springer Nature, Copyright 2019. (d) Cross-sectional SEM image of the VTMMT membrane; illustrations of the Li⁺ diffusion pathways (e) along the VTMMT and (f) within the GPE.³⁷⁶ Reproduced from ref. 376 with permission from Springer Nature, Copyright 2022.



β'' -Al₂O₃ host to construct highly efficient, uniform, and continuous solid–liquid hybrid Na-ion transportation channels through and along the β'' -Al₂O₃ nanowires (Fig. 15(b)), promoting uniform sodium deposition and formation of a stable and flat solid electrolyte interface on the sodium metal anode (Fig. 15(c)). As a result, the NVP//Na batteries using this electrolyte presented excellent cycling performance in a wide temperature range from 25 to 60 °C, with 95.3% and 78.8% capacity retentions after 1000 cycles at 1C at 25 and 60 °C, respectively.³⁷⁵

To further improve the ion transportation efficiency in GPE, aligned networks have been proposed.^{376,377} Ding *et al.* incorporated poly(ethylene glycol) diacrylate-based precursor into the vertically-aligned MMT scaffold (VTMMT, as shown in Fig. 15(d)) to form a GPE *via in situ* polymerization. The uniform and fast Li⁺ diffusion pathways along the continuous interface/interlayer of the low-tortuosity VTMMT provided benefits to the GPE by improving the Li⁺ transport capacity and reducing the occurrence of side reactions (Fig. 1(5e) and (f)). As a result, the LFP//Li full cell using this GPE exhibited outstanding rate performance and cycling stability with high CE and high-capacity retention at a wide temperature range from 0 to 60 °C.³⁷⁶ The development of GPEs with aligned host provides a promising approach for improving the ion conduction *via* novel nano-architecture designs.

Despite these advantages, there are still challenges associated with the use of inorganic hosts in wide-temperature GPEs. One challenge is the limited compatibility between the inorganic host and the polymer matrix. The incorporation of inorganic particles into the polymer matrix can disrupt the molecular order and hinder ion mobility, resulting in decreased ionic conductivity and deteriorated electrochemical performance. Therefore, the careful selection and optimization of both the inorganic host and the polymer matrix are essential to achieve optimal compatibility and maintain good ion transport properties.

4.6 Multiple combined strategies

The multiple combined strategies that can integrate the advantageous effects of various strategies are proposed to circumvent the limitation that most of the “single” strategies discussed above can only prioritize either low- or high-temperature performance. For instance, some solvents having low melting points can favor the low-temperature operability of batteries but are susceptible to volatilization and decomposition at slightly higher temperatures; IL plasticizers are particularly advantageous in ensuring excellent high-temperature oxidation stability, but the resulting demerit of low target-ion transference number would severely limit the rate performance of batteries. Obviously, to battery chemists, electrolyte design/engineering has always been the game of overcoming trade-offs between various property/performance indicators, while a single tuning strategy often appears to be less competent. In fact, the multiple combined strategies have been widely adopted by many works discussed in previous sections: strategies of co-solvent incorporation and salt concentration adjustment are simultaneously involved in the design of the organohydrogel with enhanced freezing tolerance and thermal

stability; the ionogel electrolyte with its polymer matrix rationally designed to assist the ion transport is exactly a multiple strategy combining both solvent optimization and matrix design. Nevertheless, this part is written separately to highlight the great significance of the multiple combined strategies.

In the effort to address the issue of low Li⁺ transference number of ionogel electrolytes for Li battery use, Song *et al.* designed a composite ionogel electrolyte showing superior comprehensive performance in terms of concurrent high ionic conductivity, oxidation stability up to 5.5 V, and, most importantly, a high t_{Li^+} of 0.89, by taking the advantages of the IL solvent of EmimTFSI, high-concentration Li salt of 3 M LiTFSI, and inorganic filler (host) of LLZO, in which the cooperation of high concentration Li salt and inorganic filler with t_{Li^+} approaching unity is believed to mainly responsible for the desired high t_{Li^+} value. With these advantages, LMBs using the as-obtained ionogels as GPE and high-voltage NCM materials as the cathode deliver a high discharge capacity, which is hardly achievable for conventional ionogel electrolytes.³⁷⁸

Additionally, from our perspective, the multiple combined strategies may also refer to the preparation of hybrid electrolytes that feature an A/B double-layer structure (or A/B/A and A/B/C three-layer structures in some cases), of which process A and B layers are prepared based on different strategies. For example, one can construct an ultrathin inorganic layer with high Young's modulus on the side of a conventional GPE interfaced with the metallic anode to physically inhibit the dendrite growth, without significantly sacrificing the ion transport properties; distinct chemical/electrochemical requirements of anodes and cathodes could be elegantly addressed by designing hybrid GPEs consisting of two thermodynamically immiscible functional layers.

As a notable example, Lv *et al.* designed a hierarchically-structured polymerized tetrahydrofuran (poly-THF) GPE with one layer doped with the Li_xGa₈₆In₁₄ alloy, in which the THF solvent with low viscosity and melting point ensured high ion conductivity at low temperatures, while the Li_xGa₈₆In₁₄-doped layer helped to homogenize the Li⁺ flux and formed a stable SEI layer, thus enabling high discharge areal capacity and good cycling stability of LMBs over a wide temperature range of –20 to 55 °C.³⁷⁹ Despite being in an early stage, these results suggest the great potential of the multiple combined strategies, namely, rationally integrating two or more strategies, in sparking unprecedented breakthroughs in the design and synthesis of advanced GPEs toward wide temperature applications.

5. Summary and perspectives

Advancing the electrolytes play a critical role in enabling rechargeable batteries to operate reliably in extreme conditions. The emergence of high- and low-temperature batteries highlights the potential of GPEs in facilitating wide-temperature adaptable batteries. This review focuses on the GPEs and their associated wide-temperature applications, emphasizes the interest of the scientific community in both (i) the factors leading to capacity loss of batteries at low and high temperatures, and (ii) the latest



strategies in GPE engineering and related interface chemistry to improve low-temperature kinetics and high-temperature stability. On this basis, new insights into viable solutions for wide-temperature GPEs are proposed, as displayed in Fig. 16, and discussed in more detail below.

5.1 In-depth study of battery performance degradation mechanisms at extreme temperatures

The practical application of GPE with wide-temperature operability hinges on continued research of the deterioration mechanism of the electrolyte itself and battery performance

under thermal extremes. Such investigations must be devoted to the deep understanding of the intermolecular interactions between the polymer matrices and liquid plasticizers in response to temperature variation in a wide range. Then, the cause-to-effect relationship between the nanoscale interaction, the macroscopic electrochemical properties of GPEs, and the battery working and failure mechanisms should be studied. Hence, from our perspective, the leading issue is to quantify the contribution of temperature-dependent polymer-ion interaction to the ionic conductivity of GPE with different mass ratios of the liquid plasticizer. Previously, most reports on GPEs

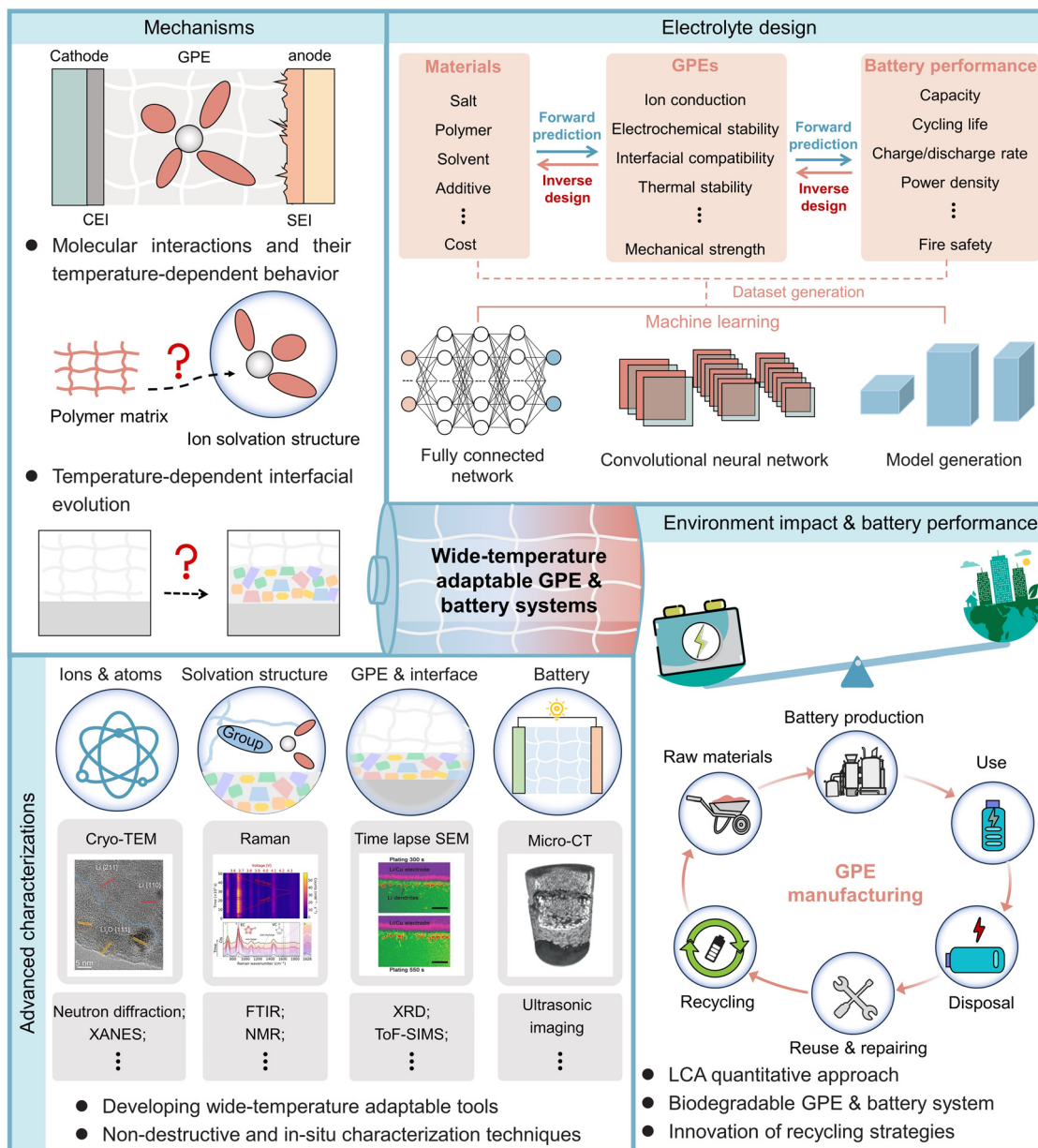


Fig. 16 Research and industrialization opportunities of wide-temperature GPEs. Opportunities stemming from the gain of deeper mechanism insights, AI-assisted GPE design, advanced characterizations, and balancing the environmental impact and battery performance are discussed. Adapted with permission from: ref. 104 (cryo-TEM image), John Wiley & Sons, Copyright 2022; ref. 380 (Raman image), Springer Nature, Copyright 2022; ref. 381, (time lapse SEM), John Wiley & Sons, Copyright 2017; ref. 382 (Micro-CT), Springer Nature, Copyright 2022.



with high ionic conductivity are realized by increasing the proportion of liquid plasticizers, the role of polymer matrices in ion transport is barely illustrated. Considering the fact that the mobility of chain segments increases with temperature, it is crucial not to disregard the influence of polymer matrices on ion transport within GPEs, especially at high temperatures. This effect has not been thoroughly elucidated thus far and warrants further investigation.

The in-depth investigations of the electrolyte/electrode interface chemistry of GPEs at extreme temperatures should be more underscored. It is widely accepted that the retarded ion kinetics and accelerated electrolyte decomposition at the electrolyte/electrode interfaces are the main factors causing battery performance degradation at low and high temperatures, respectively. However, the most fundamental questions such as the effect of operating temperature on the GPE/electrode interfacial chemistry and charge carrier/mass transport kinetics, remain unanswered. Special attention should be given to the impact of temperature on the interactions among various components and ion solvation structure. Though it has been reported that the ion-dipole interactions between solvent and salt increase with temperature,^{383,384} these are only hypothesis, and concrete characterizations are lacking. Recent studies have suggested that the interactions between polymer matrices and ionic species or solvents in GPE greatly affect the ion solvation structure and interfacial stability of batteries; yet, their dynamic changes and the effects on electrochemical kinetics under different temperatures should be thoroughly investigated.

At the battery level, further in-depth research on the influence of temperature on the electrochemical performance remains indispensable and urgent. Some research directions can be highlighted in future endeavors, such as the interfacial stress evolution induced by the low-temperature freezing and high-temperature melting of GPEs. In addition, the structural reversibility of GPE and GPE/electrode interfaces with the temperature changes, if any, remains ambiguous, which still requires in-depth research both theoretically and experimentally.

A thorough understanding of these mechanisms can guide the design of GPEs with efficient charge transfer, minimized side reactions, and enhanced overall battery performance across a wide temperature range.

5.2 Developing novel electrolyte systems with wide-temperature operability

Precisely controlled components with tailorable structures are also desired for wide-temperature GPEs and rechargeable batteries. As a prerequisite, highly thermally stable electrolyte components should be designed to accommodate temperature variations. On this basis, the high ionic conductivity and the component formation of the ion-conductive and thermostable SEI layer should be highlighted when exploiting novel electrolyte materials. Recent advancements in GPEs with wide temperature adaptability have shown that the introduction of ionophilic/solvent-philic polymer matrices can enhance ion diffusion in the bulk electrolyte and improve the interfacial stability, thereby enhancing the performance of GPEs at high

and low temperatures. However, further research is needed to systematically evaluate the impact of these advancements on the wide-temperature tolerance of GPE and battery systems.

It is likewise important to focus more on the self-protection design of novel GPE to prevent the hazards of thermal runaway of batteries when operating in extreme temperature conditions. Thermoresponsive materials show promise in this regard through several mechanisms: (i) inhibiting ion transport at high temperature to reduce the likelihood of thermal runaway; (ii) absorbing or dispersing heat generated to prevent the temperature from escalating to dangerous levels during thermal runaway events; and (iii) acting as thermal insulators, reduce the heat transfer from the surroundings to the battery or between different components within the battery, preventing overheating or excessive cooling in extreme temperature conditions. For instance, Qian *et al.* integrated a thermoresponsive phase-change material, polyethylene glycol (PEG), into agarose-based GPEs.³⁸⁵ Once the operating temperature reaches the phase-change temperature, the PEG can absorb heat and inhibit ion transport by undergoing a phase change from crystalline to amorphous, alleviating heat generation inside the batteries under thermal shock. Zhang *et al.* introduced thermal-polymerization VC as a thermoresponsive solvent that could boost the thermal stability of both the SEI and electrolyte, raising the critical temperature for thermal safety (the beginning temperature of obvious self-heating) from 71.5 to 137.4 °C.³⁸⁶ Leveraging the unique properties of the thermoresponsive materials, the safety and performance of GPEs and batteries could be collectively enhanced in extreme temperature conditions, making them potential candidates for novel wide-temperature GPE design as polymer matrices or functional components.^{386–388}

Further work is needed to develop unscalable manufacturing methods and achieve mass production and industrial scale-up of laboratory GPEs. This work should focus on using safe and cost-effective reactants and synthetic processes. The manufacturing of ionogels based on the biomass polymer matrices and ILs solvents shows promise and should therefore be given a higher priority. Considering the possible interfacial compatibility of *ex situ* production processes of GPEs, the facile synthesis technology based on the *in situ* thermo-polymerization seems like a prompt pathway as it enables the creation of an integrated battery structure with seamless interfacial contact, satisfactory electrochemical performance, as well as the good compatibility with existing lithium-ion battery production processes.^{239,389,390} Additionally, the balance between electrochemical properties and mechanical properties, such as strength, flexibility, and stretchability, should be carefully regulated through polymer engineering and optimizing the polymer-to-liquid plasticizer ratio. Furthermore, as the electrolyte thickness decreases, the importance of ionic conductance ($G = \sigma A/l$) in evaluating ionic conduction should be emphasized.^{391,392}

Overall, the physical, chemical and electrochemical properties of GPEs in a wide temperature range should be focused on for meeting the criteria for designing usable and safe rechargeable batteries. Since the wide-temperature battery is a complicated topic, in addition to the material challenges and



solutions presented in this review, the technical innovations in battery configuration and the full operational simulation in different temperature environments before stepping into the practical manufacturing market are also important research directions and should therefore be paid attention to. All of these will help to ensure the reliability and functionality of wide-temperature GPE-based rechargeable batteries.

5.3 Wide-temperature adaptable non-destructive and *in-operando* tools

The complex nature of GPEs makes it difficult to obtain detailed information about their bulk and molecular structures as well as their evolution upon the battery life cycle and under different temperature conditions. This complexity spans multiple scales, including atomic and ionic chemistry, microscopic molecular structures, mesoscopic electrolyte materials and interfaces, and macroscopic battery systems. Conventional characterization techniques, such as X-ray diffraction (XRD), SEM and X-ray photoelectron spectroscopy (XPS), are usually performed *ex situ* and require battery disassembly, hampering the mechanistic examination and continuous research of GPEs and rechargeable batteries, thereby obscuring some of the fundamental principles essential for the design of wide-temperature batteries.³⁹³

Recent emphasis has been applied to *in situ/operando* techniques to study batteries in operation. Such tools can avoid the negative effects and uncertainty from the post-treatment of samples and thus provide an accurate information of the electrochemical processes.^{394–396} Examples include the *in situ/operando* X-ray computed tomography (CT) for non-invasive information about a battery's morphology and internal structure, the non-destructive ultrasonic scanning and imaging technique for examining the battery state of charge and health, and time-lapse SEM for observing the detailed lithium deposition processes.^{381,397,398} Advanced techniques like time-of-flight secondary ion mass spectroscopy (TOF-SIMS) are suitable for interfacial composition characterization because of their high mass sensitivity simultaneously and ultrahigh spatial resolution. The *in situ* XRD techniques has been utilized for investigating the structure and interfacial evolution of graphite anode in Al-ion batteries assembled with GPE.³⁹⁹ At the microscale, Raman and FTIR spectra are important techniques to study the presence of intermolecular interactions between the constituents of polymer electrolytes, such as ion–ion, ion–solvent, ion–nanofiller, polymer–ion, polymer–solvent, and polymer–ion–solvent. It provides in-hand information about the complex formation and examines by identifying the shift in the peak position of the characteristic functional groups. Mao and coworkers elucidated the SEI formation and evolution pathways during the operation of LMBS using a depth-sensitive plasmon-enhanced Raman spectroscopy for the monitoring of dynamic nature of SEI formation and evolution.⁴⁰⁰ A deeper insight into the atomic/ionic structure can be provided by aberration-corrected high angle annular dark field scanning TEM (HAADF-STEM), annular bright field STEM, neutron diffraction, X-ray absorption near edge structure (XANES), *etc.*

Building upon the progress in *in situ/operando* characterizations, there are several considerations worth investigating.

Firstly, the liquid–solid duality of GPEs suggest that these *in situ* characterizations and analyses of both liquid and solid electrolyte-based batteries can be extended to GPEs-based battery systems. Secondly, combining various *in situ/operando* characterization techniques can provide diverse information at different scales for optimizing wide-temperature adaptable GPEs and rechargeable batteries. Thirdly, *in situ* sensing techniques show great promise for the *in situ/operando* evaluation of battery electrochemical behaviour by correlating them with parameters (*e.g.*, stress, temperature, and gas) of interest.^{401,402} As a notable example, Tarascon *et al.* monitored electrolyte evolution in 18 650 Na-ion and Li-ion cells under real working conditions by *in operando* IR fiber evanescent wave spectroscopy (FBWS). The FBWS helps to reveal changes in the dynamics of the solvents and additives in electrolytes and identify the nature of the electrolyte decomposition species without any latency as well as changes in the ion solvation dynamics as a function of voltage and current.⁴⁰³ These efforts can contribute to a deeper understanding of the reaction processes, degradation mechanisms, and thermal decomposition mechanisms of GPEs under various temperature conditions, guiding rational multiscale materials design for optimized properties and performance.

5.4 Artificial intelligence (AI)-assisted GPE design

The rechargeable battery with GPE is a complex electrochemical system governed by electro-/chemical reactions, charge, and ion transport in solid and liquid phases, making it challenging to describe with a simple physics formula, particularly under extreme temperature conditions. Additionally, the properties of GPEs, including ionic conductivity, cationic transference number and temperature tolerance, significantly impact the battery's performance in terms of charging and discharging rates, cycle life, and energy density at extreme temperatures. Although researchers have made significant progress in the experimental modification of electrolyte formulations and theoretical mole including DFT modeling and molecular dynamics, accurate and versatile prediction tools are urgently needed to explore optimal battery performance and corresponding gel electrolyte formulations under various temperature scenarios, crossing large time scale and under practical application conditions. Machine learning and deep learning algorithms offer a widely applicable and automated tool that have been carried out to assist the electrode and electrolyte design.⁴⁰⁴ For example, deep learning algorithms can be used to improve the accuracy and efficiency of numerical simulation. By implementing the neural network to learn the atomic polarizabilities and charges, the molecular dynamics prediction on the polarization terms in the liquid electrolyte can be effectively accelerated, and dependencies on the 3D conformation can be avoided.⁴⁰⁵ Moreover, by informing the neural networks about the surrounding environment of the atoms or molecules, the accuracy and speed of the prediction could be further improved.^{406,407} In addition, machine learning has been demonstrated to facilitate the guidance of empirical data on electrolyte synthesis. Cui's group used the elemental composition of electrolytes as model features of a machine learning model to identify key features for predicting CE and created fluorine-free



solvent-based electrolyte formulations, helping to achieve a remarkable CE of 99.70% according to the as-trained machine learning model.⁴⁰⁸ These advancements have demonstrated the potential of AI-assisted models and algorithms in efficiently designing GPE and rechargeable batteries that are capable of operating in a wide temperature range. By harnessing AI-assisted models and algorithms, researchers are expected to efficiently analyze large amounts of data, identify patterns, and establish correlations between composition and electrolyte and electrolyte/electrode interfacial evolution, as well as the corresponding battery performance at different temperatures, which can help to screen and optimize gel-polymer electrolyte candidates. Additionally, AI is able to predict material properties and behaviors, which can lead to a reduction in time-consuming and expensive experimental tests. As AI algorithms continue to evolve and improve, their integration into the design of wide-temperature GPEs is expected to revolutionize the development of advanced energy storage systems, leading to more efficient and durable devices for a sustainable future.

5.5 Balancing the environmental sustainability and battery performance

Balancing environmental sustainability and battery performance is an increasingly important consideration in the future design of GPEs and rechargeable batteries. A forward-looking approach involves the development of quantitative approaches, such as life cycle assessment (LCA), to provide a comprehensive analysis of the entire life cycle of the batteries, from raw material extraction through manufacturing, use, and disposal, and to evaluate the environmental and economic impacts and resource risks of GPEs and rechargeable batteries in a dispersive temperature regime. In recent years, there have been several LCA studies on lithium-ion and lithium-beyond batteries with liquid electrolytes based on different life cycle impact assessment methodologies, life cycle inventory databases, and different assumptions about the energy intensity of battery cell production.^{409,410} However, there are very few studies available on the environmental assessment of GPEs and corresponding rechargeable batteries. Moreover, most LCA studies focus on the carbon emissions and environmental indicators of the battery manufacturing phase rather than on the whole battery cycling lifetime, and, in particular, neglect the stages of material recycling and remanufacturing phases of spent batteries.⁴¹¹ In addition, there is still a lack of a consistent definition, and guidelines for functional units and sufficient data (original and current) for LCA assessment are still lacking.^{412,413} For example, several functional units such as “battery pack”, “1 kg of battery”, “1 kW h of storage capacity”, and “1 km (the distance traveled by the electric vehicle during its lifetime)” have been proposed in previous works, making it difficult to compare different LCA studies.^{383,414} Therefore, there is a need to promote reproducible, comparable, and verifiable LCA studies on different GPEs and rechargeable systems.

Exploring more biodegradable GPEs and battery systems is a promising approach to address the environmental concerns associated with traditional GPEs made from fossil oil-based

polymers, such as PEO, PMMA, and PVDF and their derivatives. These species are non-biodegradable and easily result in energy waste and environmental “white pollution”, making it hard to meet the ever-growing demands for both comprehensive performance and environmental protection of rechargeable batteries. In this respect, biomass materials, including polylactic acid, cellulose, chitosan, and xanthan gum, can offer an interesting opportunity for replacing synthetic polymers and developing GPEs with reduced environmental impact.^{415–417} A wide variety of cellulose and derivatives have gained considerable attention due to their abundant biomass sources (*e.g.*, bacteria, wood, bamboo, and grass), high ionic conductivity, satisfactory mechanical strength and good freezing tolerance, creating a range of possibilities for developing functional, inexpensive, safe, and environment-friendly GPEs.^{230,418–420} We expect that their applications could be extended to wide-temperature scenarios and more rechargeable battery systems.

In addition, the innovation of recycling strategies that align with the principles of green chemistry is also an integral part of the future development of wide-temperature GPEs and rechargeable batteries. Efficient and sustainable recycling practices will minimize the waste and environmental pollution associated with the disposal of GPEs and rechargeable batteries.⁴²¹ The traditional linear model of sending raw materials to landfills or partial recycling causes significant pollution and waste.^{422,423} Existing battery recycling strategies, such as classical pyrometallurgy, hydrometallurgy and emerging bio-metallurgy, and direct regeneration techniques focus on the extraction of valuable elements or materials from the cathode, which typically requires expensive reagents or complex processes and often neglects the recovery of low-value components such as electrolytes and graphite.^{424–426} In the future, researchers should focus on exploring more environment-friendly recycling processes, including the separation of GPEs from batteries, and the recovery and reuse of GPEs and other valuable materials from rechargeable batteries. All the above efforts will contribute to the creation of more environment-friendly and sustainable battery technologies, promoting a greener future for energy storage.

Author contributions

Conceptualization, X. Zhou, Y. Zhou, L. Yu, P. Hu, and C. Chen; data curation, X. Zhou and Y. Zhou; visualization, X. Zhou, L. Yu, L. Qi, and Y. Zhou; writing – original draft, X. Zhou and L. Yu; writing – review & editing, X. Zhou, L. Yu, Y. Zhou, K. Oh, S.-Y. Lee, P. Hu, and C. Chen; resources, C. Chen; supervision, P. Hu, S.-Y. Lee, C. Chen.

Conflicts of interest

There are no conflicts to declare.

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